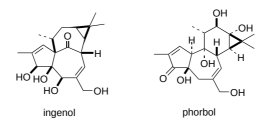


Tetrahedron Vol. 62, No. 7, 2006

Contents

REPORT

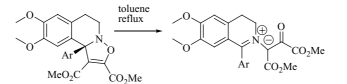
Synthetic approaches to ingenol Jin Kun Cha* and Oleg L. Epstein



ARTICLES

Synthesis of stable azomethine ylides by the rearrangement of 1,3-dipolar cycloadducts of 3,4-dihydroisoquinoline-2-oxides with DMAD

Necdet Coşkun* and Selen Tunçman

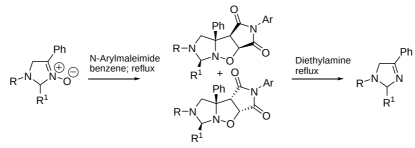


pp 1351-1359

and diethylamine induced ring-opening of *exo* and *endo* hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3-diones

Dipolar cycloadditions of imidazoline 3-oxides with N-arylmaleimides. Synthesis

Necdet Coşkun,* Habibe Mert and Nevin Arıkan

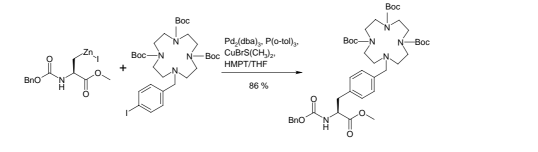


pp 1329-1343

pp 1345-1350

Synthesis of chiral amino acids with metal ion chelating side chains from L-serine using Negishi cross-coupling reaction

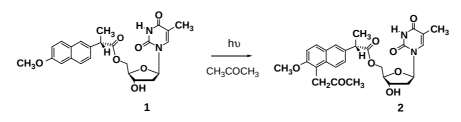
Michael Kruppa, Giovanni Imperato and Burkhard König*



A straightforward synthesis and structure of unprecedented iminium salts of dihydropyrido [3,2-*e*][1,3]thiazines

Margarita Suárez,* Hector Novoa,* Yamila Verdecia, Estael Ochoa, Amaury Alvarez, Rolando Pérez, Roberto Martínez-Alvarez, Dolores Molero, Carlos Seoane, Norbert M. Blaton, Oswald M. Peeters and Nazario Martín*

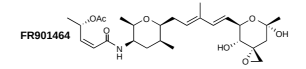
©□ Photochemistry of a naphthalene–thymine dyad in the presence of acetone Noureddine Belmadoui, María José Climent and Miguel A. Miranda*



The photobehavior of 1 is dominated by the naphthalene-derived chromophore. In the presence of acetone, this was found to be the photoreactive site, and only 2 was obtained. Thymine dimers and acetone photoadducts involving the thymine ring, which would arise from the thymine triplet excited state, were not detected.

Total synthesis of FR901464: second generation

Hajime Motoyoshi, Masato Horigome, Hidenori Watanabe and Takeshi Kitahara*



pp 1360-1364

pp 1365–1371



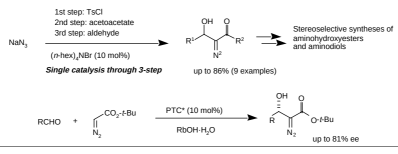
pp 1378-1389

pp 1390-1401

pp 1402-1409

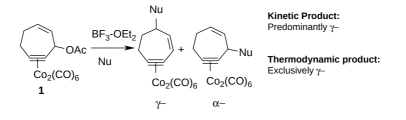
Synthesis of α -diazo- β -hydroxyesters through a one-pot protocol by phase-transfer catalysis: application to enantioselective aldol-type reaction and diastereoselective synthesis of α -amino- β -hydroxyester derivatives

Kazuya Hasegawa, Shigeru Arai* and Atsushi Nishida*



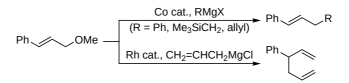
Lewis and protic acid mediated Nicholas reactions of 3-acetoxycyclohept-1-en-4-ynedicobalt hexacarbonyl: site selectivity of nucleophile incorporation

Joseph DiMartino and James R. Green*



Cobalt- and rhodium-catalyzed cross-coupling reaction of allylic ethers and halides with organometallic reagents

Hiroto Yasui, Keiya Mizutani, Hideki Yorimitsu* and Koichiro Oshima*



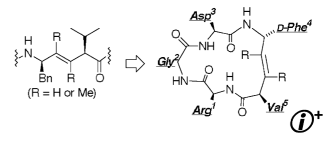
Application of tri- and tetrasubstituted alkene dipeptide mimetics to conformational studies of cyclic RGD peptides

pp 1416-1424

pp 1410-1415

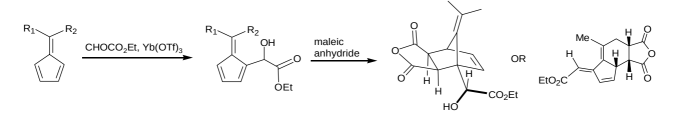
Shinya Oishi, Kazuhide Miyamoto, Ayumu Niida, Mikio Yamamoto, Keiichi Ajito, Hirokazu Tamamura, Akira Otaka, Yoshihiro Kuroda, Akira Asai and Nobutaka Fujii*

 $\psi[(E)$ -CMe=CH]- and $\psi[(E)$ -CMe=CMe]-type alkene isosteres are potential –CO–NH- and –CO–NMe-type peptide bond mimetics, respectively. The application of these chiral isosteres to conformational studies on cyclic RGD peptides having potent integrin antagonistic activities is reported.



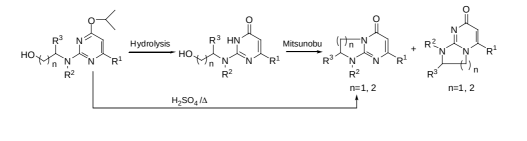
Regioselective electrophilic substitutions of fulvenes with ethyl glyoxylate and subsequent **Diels-Alder reactions**

Hsing-Chang Tseng, Arun Kumar Gupta, Bor-Cherng Hong* and Ju-Hsiou Liao



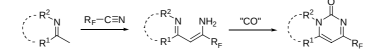
A simple approach for the regioselective synthesis of imidazo[1,2-a]pyrimidiones and pyrimido[1,2-a]pyrimidinones

David Font, Anthony Linden, Montserrat Heras* and José M. Villalgordo*



New fluorinated 1,3-vinylogous amidines as versatile intermediates: synthesis of fluorinated pyrimidin-2(1H)-ones

Santos Fustero,* Julio Piera, Juan F. Sanz-Cervera,* Raquel Román, Benjamin H. Brodsky, María Sánchez-Roselló, José Luis Aceña and Carmen Ramírez de Arellano

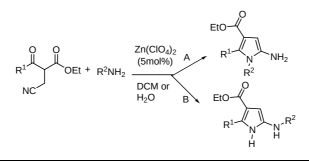


Zinc perchlorate catalyzed one-pot amination–annulation of α -cyanomethyl- β -ketoesters in water. Regioselective synthesis of 2-aminopyrrole-4-carboxylates

pp 1452-1458

pp 1444-1451

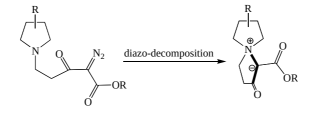
Ayhan S. Demir* and Mustafa Emrullahoglu



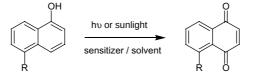
pp 1425-1432

pp 1433-1443

Stable chiral spirocyclic [5,5]-ammonium ylides using a metallo carbenoid approach Daniele Muroni, Antonio Saba* and Nicola Culeddu

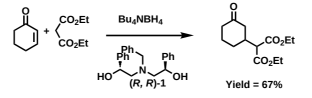


Photooxygenations of 1-naphthols: an environmentally friendly access to 1,4-naphthoquinones Oliver Suchard, Ronan Kane, Bernard J. Roe, Elmar Zimmermann, Christian Jung, Prashant A. Waske, Jochen Mattay^{*} and Michael Oelgemöller^{*}



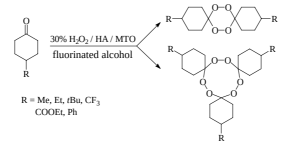
Investigation of the active species in a Michael addition promoted by chirally modified tetrahydroborate

Susan Abraham and G. Sundararajan*



Fluorinated alcohol directed formation of dispiro-1,2,4,5-tetraoxanes by hydrogen peroxide under acid conditions

Katja Žmitek, Stojan Stavber, Marko Zupan, Danièle Bonnet-Delpon and Jernej Iskra*



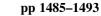
1323

pp 1479–1484

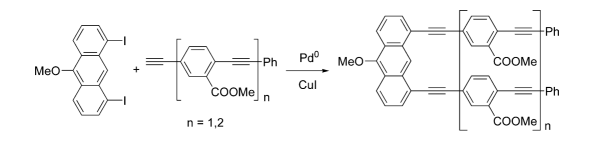
pp 1467-1473

pp 1474–1478

The synthesis of bis(oligophenyleneethynylenes): novel potential nonlinear optical materials David J. Armitt* and Geoffrey T. Crisp

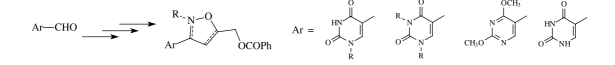


pp 1494-1501



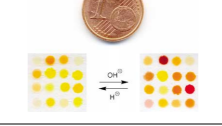
1,3-Dipolar cycloaddition approach to isoxazole, isoxazoline and isoxazolidine analogues of *C*-nucleosides related to pseudouridine

Evdoxia Coutouli-Argyropoulou,* Pygmalion Lianis, Marigoula Mitakou, Anestis Giannoulis and Joanna Nowak



Optical sensor arrays: one-pot, multiparallel synthesis and cellulose immobilization of pH and pp 1502–1507 metal ion sensitive azo-dyes

Tommaso Carofiglio,* Carlo Fregonese, Gerhard J. Mohr, Federico Rastrelli and Umberto Tonellato



Synthesis of alkoxymethyl derivatives of resorcinarene via the Mannich reaction catalysed with pp 1503 iminodiacetic acid

Mariusz Urbaniak and Waldemar Iwanek*

pp 1508-1511

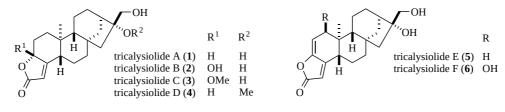
OR

-1507



Tricalysiolides A-F, new rearranged ent-kaurane diterpenes from Tricalysia dubia

Koichi Nishimura, Yukio Hitotsuyanagi, Noriko Sugeta, Kei-ichi Sakakura, Kazuya Fujita, Haruhiko Fukaya, Yutaka Aoyagi, Tomoyo Hasuda, Takeshi Kinoshita, Dong-Hui He, Hideaki Otsuka, Yoshio Takeda and Koichi Takeya*



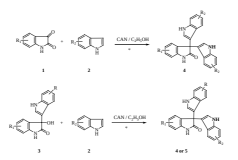
pp 1520-1526 Reactions of (1R,2S)-1,2-di(2-furyl)-1,2-di(3-guaiazulenyl)ethane and (1R,2S)-1,2-di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane with tetracyanoethylene (TCNE) in benzene: comparative studies on the products and their spectroscopic properties

Shin-ichi Takekuma,* Kenji Takahashi, Akio Sakaguchi, Masato Sasaki, Toshie Minematsu and Hideko Takekuma

Reactions of the title meso forms 1 and 2 with TCNE in benzene at 25 °C for 5 h (for 1) and 48 h (for 2) under oxygen give new compounds 3 and 4, respectively, in 74 and 21% isolated yields. The title studies, affording interesting molecular structures 3 and 4, are reported.

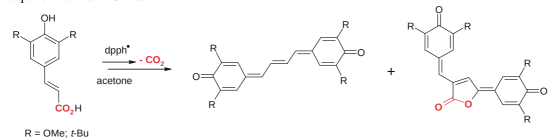
pp 1527-1535 Facile synthesis of 3,3-di(heteroaryl)indolin-2-one derivatives catalyzed by ceric ammonium nitrate (CAN) under ultrasound irradiation

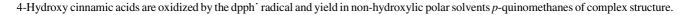
Shun-Yi Wang and Shun-Jun Ji*



Coupling and fast decarboxylation of aryloxyl radicals of 4-hydroxycinnamic acids with formation of stable *p*-quinomethanes

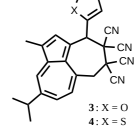
Carmelo Daquino and Mario C. Foti*





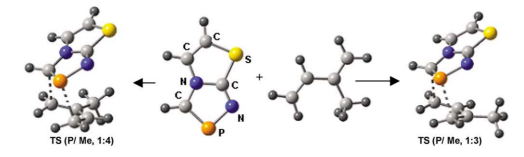
pp 1512-1519

pp 1536-1547



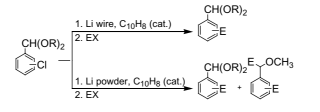
Origin of the stereo- and regioselectivities in the Diels–Alder reactions of azaphospholes: a DFT investigation

Raj K. Bansal,* Neelima Gupta and Surendra K. Kumawat

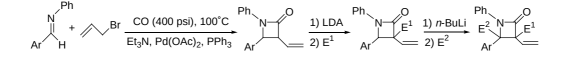


Regioselectivity in arene-catalyzed reductive lithiation of acetals of chlorobenzaldehydes Ugo Azzena,* Giovanna Dettori, Giuseppe Sforazzini, Miguel Yus* and Francisco Foubelo pp 1557-1563

pp 1564-1574

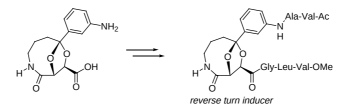


Stereoselective synthesis and functionalization of 4-heterosubstituted β-lactams Luigino Troisi,* Ludovico Ronzini, Catia Granito, Luisella De Vitis and Emanuela Pindinelli



3-Aza-8,10-dioxa-bicyclo[5.2.1]decane (9-*exo* BTKa) carboxylic acid as a new reverse turn inducer: pp 1575–1582 synthesis and conformational analysis of a model peptide

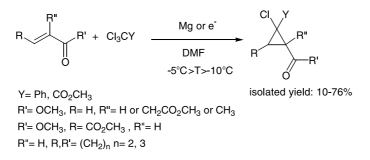
Dina Scarpi, Daniela Stranges, Andrea Trabocchi and Antonio Guarna*



pp 1548-1556

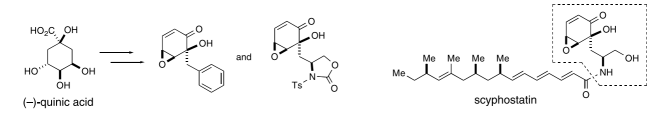
Formation of polysubstituted chlorocyclopropanes from electrophilic olefins and activated trichloromethyl compounds

Sylvain Oudeyer, Eric Léonel, Jean Paul Paugam,* Christine Sulpice-Gaillet and Jean-Yves Nédélec



Enantiocontrolled synthesis of the epoxycyclohexenone moieties of scyphostatin, a potent and specific pp 1590–1608 inhibitor of neutral sphingomyelinase

Tadashi Katoh,* Takashi Izuhara, Wakako Yokota, Munenori Inoue, Kazuhiro Watanabe, Ayaka Nobeyama and Takeyuki Suzuki



OTHER CONTENTS

Erratum	р 1609
Corrigendum	p 1610
Instructions to contributors	pp I–IV

Corresponding author () Supplementary data available via ScienceDirect

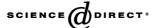
Indexed/Abstracted in: AGRICOLA, Beilstein, BIOSIS Previews, CAB Abstracts, Chemical Abstracts. Current Contents: Life Sciences, Current Contents: Physical, Chemical and Earth Sciences, Current Contents Search, Derwent Drug File, Ei Compendex, EMBASE/Excerpta Medica, Medline, PASCAL, Research Alert, Science Citation Index, SciSearch



pp 1583-1589



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Synthetic approaches to ingenol

Jin Kun Cha* and Oleg L. Epstein

Department of Chemistry, Wayne State University, Detroit, MI 48202, USA

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Contents

1.	Introduction	329
2.	Structural relationship between ingenanes and tiglianes 1.	330
3.	Inside–outside stereochemistry 1.	330
4.	Synthetic approaches to 2	331
	4.1. Funk's Ireland–Claisen rearrangement approach 12	331
	4.2. Rigby's [1,5]-hydrogen sigmatropic rearrangement approach 1.	332
	4.3. Cha's semi-pinacol rearrangement approach 1.	334
5.	Total syntheses of 2	335
	5.1. Winkler's synthesis of 2 1.	335
	5.2. Tanino–Kuwajima's synthesis of 2 12	336
	5.3. Kigoshi's formal synthesis of 2 1.	339
	5.4. Wood's synthesis of $(+)-2$ 1.	339
6.	Conclusion 1.	340
	Acknowledgements 13	341
	References and notes 1.	341
	Biographical sketch	343

1. Introduction

Some species of the *Euphorbiaceae* plant family were known to produce milky, often toxic latex, which was blamed for the poisoning of livestock. At the same time, several species were used in folk medicine for treatment of a variety of ailments. Studies to identify the active principles of these plants led to the isolation and characterization of 12,13-diesters of the tetracyclic diterpene, phorbol (1), from the *E. Croton tiglium* (Scheme 1).¹ From the *E. lathyris* and *E. ingens* species was subsequently isolated the 3-hexadecanoyl ester of ingenol (2).² Certain lipophilic long chain esters of 1 and 2, along with bryostatin, debromoaplysiatoxin, and teleocidin, are known to be highly potent tumor

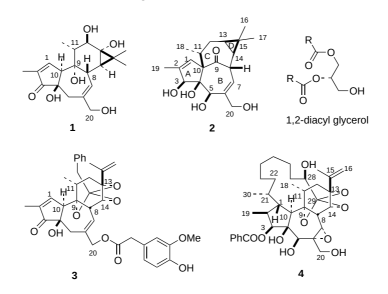
* Corresponding author. Fax: +1 313 577 8822;

promoting agents. Their mode of action is putatively associated with binding to and activation of protein kinase C (PKC) by mimicking the function of 1,2-diacyl glycerol, the endogenous PKC activator.³ However, a key pharmacophore common to a structurally diverse group of these tumor promoters has not yet been established. Two representative members of the structurally related daphnane family are resiniferatoxin (3) and gnidimacrin (4);⁴ there is a conspicuous similarity of the oxygenation pattern in the lower subunit between 1 and 3 and also 2 and 4. However, 3 and 4 are devoid of cocarcinogenic activity, but instead exhibit analgesic and antitumor activity, respectively. It is interesting to note that certain ester derivatives of 2 have recently been reported to possess anti-leukemic and anti-HIV activity.⁵ Biological activity of these natural products is thus significantly altered by subtle, yet little-understood, structural modifications. A unified synthetic strategy for tiglianes, ingenanes, and daphnanes would be highly desirable to shed light on the structure-activity relationships, which in turn could lead to the development of useful

Keywords: Ingenol; In, out-configuration; Photocycloaddition; Retro-aldol; Nicholas reaction; Semi-pinacol rearrangement; Ring-closing olefin metathesis; Ireland-Claisen rearrangement; High-order cycloaddition; [1,5-Hydrogen sigmatropic rearrangement.

e-mail: jcha@ chem.wayne.edu

^{0040–4020/\$ -} see front matter @ 2005 Elsevier Ltd. All rights reserved. doi:10.1016/j.tet.2005.10.035



Scheme 1.

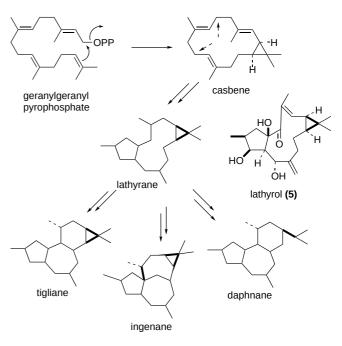
biochemical tools and new therapeutic agents, presumably by selective modulation of PKC isozymes.

The biological activity and the structural complexity of these diterpenes have attracted numerous studies directed toward total syntheses over two decades. A landmark synthesis of phorbol (1) was first reported by Wender and co-workers.⁶ The broad applicability of the key oxidopyrylium cycloaddition approach was demonstrated by subsequent synthesis of resiniferatoxin (3).⁷ Our laboratory achieved a formal synthesis of 1 by intersecting with Wender's advanced intermediate.⁸ More recently appeared the first total synthesis of ingenol (2) by Winkler and co-workers who devised an ingenious application of intramolecular dioxenone photocycloaddition.⁹ Soon thereafter followed three notable syntheses of 2 by the Tanino–Kuwajima, Wood, and Kigoshi groups.^{10–12} This review delineates the highlights of the total syntheses of the ingenanes, together with our own work.¹³

2. Structural relationship between ingenanes and tiglianes

A cursory look at the three diterpenoid families suggests that they are derived biosynthetically in plants from geranylgeranyl pyrophosphate, probably via macrocyclic precursors (Scheme 2). Casbene- and lathyrane-type macrocyclic diterpenes might serve as suitable biogenetic precursors.14 For example, a transannular aldol condensation of lathyrol (5), a prototypical lathyrane, could result in the C8-C9 bond formation to provide the tigliane skeleton. Although no details are known, a 1,2-alkyl shift (e.g., Wagner-Meerwein rearrangment) connects tiglianes to ingenanes. During the course of structural identification studies with 3,4;5,20-diisopropylideneingenol (6; structure not shown), treatment of 9(R)-alcohol 7 with MsCl yielded the tigliane skeleton 8 (Scheme 3).^{2b} On the other hand, migration of a different C–C bond (i.e., C4–C10) was observed with 9(S)-alcohol 9 to furnish 10 due to the well-defined (anti-periplanar) stereoelectronic requirements. Similarly, treatment of ingenol (2) itself with aqueous HClO₄ in methanol triggered a vinylogous

retro-pinacol rearrangement to give **11** in 49% yield (84% based on consumed starting material).¹⁵ Particularly note-worthy are the mild conditions and good yield for the pivotal rearrangement, which might well be of biogenetic significance.

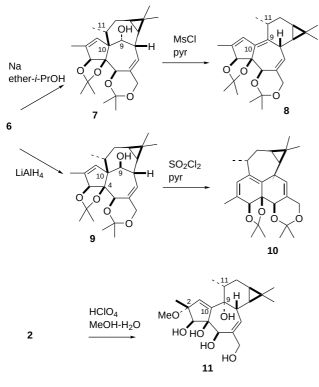


Scheme 2.

The facility of these skeletal rearrangements can be attributed to relief of sizeable strain associated with the trans intrabridgehead stereochemistry of 2.

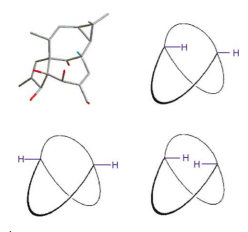
3. Inside-outside stereochemistry

The formidable challenges in synthetic studies of **2** arise primarily from the highly strained *in–out* stereochemistry, the most distinctive structural characteristic. The *in–out* nomenclature was first introduced for bridged bicyclic compounds by Simmons.¹⁶ As denoted in simple, graphic



Scheme 3.

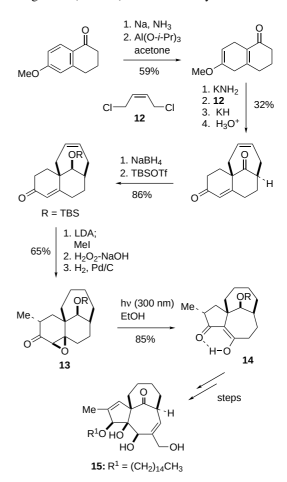
representations, the *in-out* nomenclature refers to the location of the bridgehead hydrogen atoms or other substituents (Scheme 4): typically, the *in-in* configuration is the least stable owing to the inevitable H–H repulsive interaction. The energy difference between *in-out* and *out-out* arrangements depends on ring sizes. In the case of ingenol, the natural *in-out* configuration was calculated to be more strained by 5.9 kcal/mol than the corresponding *out-out* isomer, that is, the C-8 epimer (isoingenol).^{17a}



Scheme 4.

The importance of the distinctive in-out stereochemical facet was clearly underscored by Paquette, as a suitably functionalized isoingenol analog 15, having the fully elaborated AB ring of 2, was completely lacking in the biological activity related to the esters of 2 (Scheme 5): the synthesis began with Birch reduction of 6-methoxy-

1-tetralone and subsequent double alkylation with **12**. Photoisomerization of α , β -epoxyketone **13** induced ring transposition to afford the isoingenol skeleton **14** with cis intrabridgehead (*out–out*) stereochemistry.¹⁸



Scheme 5.

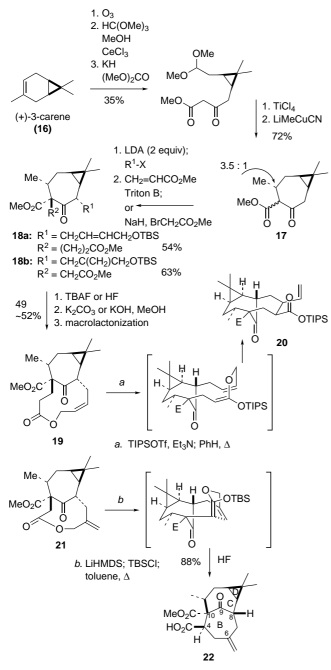
Therefore, a successful synthesis must address the rare *in–out* stereochemical issue, along with efficient installation of the densely positioned hydroxyl groups and stereoselective introduction of the methyl group at C11.

4. Synthetic approaches to 2

Inasmuch as the *in–out* stereochemistry has been shown to be indispensable to biological activity, the otherwise attractive approaches to the isoingenanes are not covered herein. Readers are instead referred to two excellent reviews on these previous studies.^{13,19}

4.1. Funk's Ireland–Claisen rearrangement approach

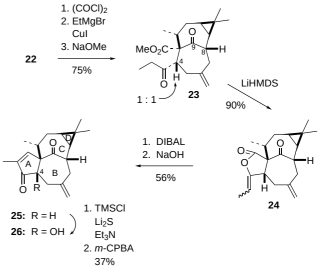
Funk found an incisive solution to the principal stereochemical issue in an Ireland–Claisen rearrangement of a considerably less strained macrobicyclic lactone, which proceeds with ring contraction to furnish a more strained *trans*-fused bicyclo [4.4.1] system.¹⁷ In his CD \rightarrow BCD \rightarrow ABCD sequence, the requisite *trans* configuration at C8 and C10 was established at an early stage by sequential diastereoselective alkylation reactions of ketoester **17** to provide **18a** and **18b** (Scheme 6). Starting with (+)-3carene (**16**), **17** was prepared by standard methods; conjugate addition of LiMeCuCN to the enone (structure not shown) took place with ~3.5:1 diastereoselectivity at C-11. Alkylation of the dianion of **17** occurred opposite to the cyclopropane ring and subsequent Michael reaction delivered **18a** as a single isomer. The dominant stereocontrol element in the last step is believed to be the methyl group at C-11. Following straightforward functional group elaboration, macrolactone **19** was subjected to the key rearrangement that took place via a boatlike transition state to deliver **20** possessing the BCD ring skeleton of **2**. The indicated stereochemical assignment was verified by singlecrystal X-ray analysis.^{17a}



Scheme 6.

The Ireland–Claisen rearrangement-induced ring contraction strategy was next extended to **21** containing suitably placed functionalities so as to facilitate the A ring construction. As one of the reacting termini in the [3,3]sigmatropic rearrangement is exocyclic to the macrocyle, a chairlike transition state was found to be operative, and single-crystal X-ray analysis (of the corresponding bromo lactone) indicated that the major rearrangement product **22** arose from the indicated transition state.^{17b}

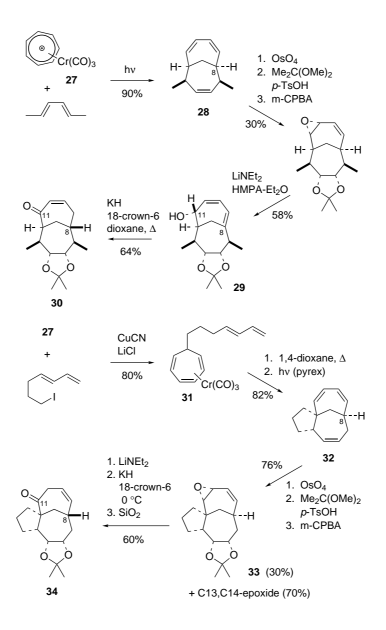
Toward completion of the ingenane tetracyclic ring system, inversion of configuration of the C4 carboxylic acid was necessary: the requisite epimerization was dealt with by base-catalyzed equilibration of the ketone intermediate to deliver 23 as a 1:1 mixture (Scheme 7). It should be noted that enolization of the C9 carbonyl is precluded by poor overlap between the inside C8 hydrogen atom and the carbonyl p-orbitals. The lithium enolate of ketone 23 underwent clean O-acylation to give enol lactone 24 as an inconsequential 2:1 mixture of the Z/E isomers. The desired aldol product 25 was then obtained by DIBAL reduction of 24 and subsequent treatment with NaOH in MeOH. Finally, the C4 hydroxyl group was introduced by the Rubottom oxidation of the trimethylsilyl enol ether of 25 to afford the fully assembled and enantiomerically pure ingenol derivative 26.17b



Scheme 7.

4.2. Rigby's [1,5]-hydrogen sigmatropic rearrangement approach

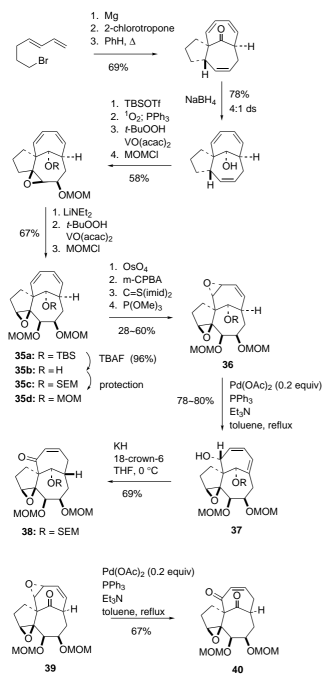
Rigby reported an ingenious solution for conversion of a readily accessible *cis*-intrabridgehead bicyclo[4.4.1]undecane compound to its highly strained trans (*in-out*) isomer: alkoxide-accelerated [1,5]-hydrogen shift was utilized to countermand the otherwise adverse thermodynamics.²⁰ Chromium(0)-mediated $[6\pi + 4\pi]$ cycloaddition between **27** and *E,E*,-2,4-hexadiene delivered **28** as a single (*endo*) diastereomer in excellent yield (Scheme 8).²¹ By taking advantage of the well-defined facial bias inherent in bicyclo[4.4.1] derivatives, the necessary bridgehead double bond was introduced by straightforward elaboration to give



Scheme 8.

29. The pivotal alkoxide-mediated [1,5]-hydrogen sigmatropic rearrangement²² of **29** yielded the *in–out* enone **30** to establish the correct C8 bridgehead stereochemistry as a consequence of a suprafacial [1,5] hydrogen shift.^{20a} This useful protocol was next extended to the intramolecular cycloadduct **32** to afford a functionalized ingenane tricycle **34**.^{20b} The structures of **30** and **34** (as its α,β -conjugated enone isomer) were confirmed by single crystal X-ray analysis. It is worth noting that a high level of convergency is possible by an intramolecular [6+4] cycloaddition reaction to construct the ABC ring system. In contrast, it is not feasible in intermolecular processes to directly introduce substituents at the incipient bond forming centers in the 6π component.

The general applicability of the key sigmatropic rearrangement, along with the compatibility with highly functionalized substrates (e.g., $35^{20d,e}$ including an epoxide functionality), was also demonstrated with an advanced intermediate **37** for the preparation of 38: chemoselective epoxide ring opening was achieved by palladium-promoted isomerization of an allylic epoxide, that is, $36 \rightarrow 37$, in the presence of a nonreactive C3,C4-epoxide (Scheme 9).^{20c} The mechanism for this interesting dienol formation was proposed to involve antielimination by the action of a base from a π -allyl-Pd intermediate. Interestingly, the reaction was found to be sensitive to steric effects, as the bulky TBS group at C9 (i.e., 36 where R = TBS) failed to react even in the presence of a stoichiometric amount of Pd(OAc)₂. The requisite transintrabridgehead compound 38 (where R = SEM) was readily obtained by applying the above-mentioned conditions to 37. It is noteworthy that the respective ketone 39, possessing a keto group at C9, underwent a different isomerization reaction, presumably via syn-elimination of a π -allyl-Pd intermediate, to give 40 in 67% yield: this remarkable divergence between 36 and 39 could be attributed to conformational changes attendant to the slight, yet significant, structural or functional group changes.^{20c}

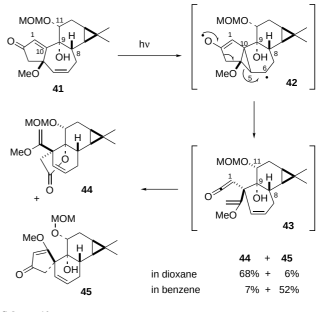


Scheme 9.

4.3. Cha's semi-pinacol rearrangement approach

Following up on our formal, enantioselective synthesis of (+)-phorbol (1),⁸ we were interested in the development of a unified approach to tiglianes, daphnanes, and ingenanes. As delineated in Scheme 3, a missing link between tiglianes and ingenanes could be found in an appropriate 1,2-alkyl shift, which might well be involved in their biogenesis and could also provide an efficient, integrated synthetic strategy. Other laboratories, not surprisingly, explored this simple, yet attractive, tactic. Recently, it came to our attention that several years ago the Wender group had examined the photochemical ring transposition of a C9, C10 epoxide as a logical extension of their first total synthesis of phorbol (1) toward the

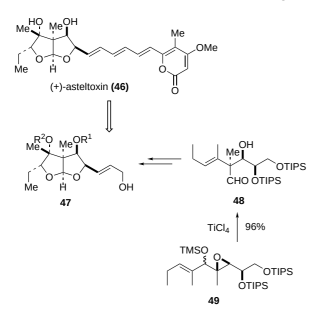
ingenanes: irraditation of a C9, C10 epoxide resulted in exclusive migration of the undesired C8–C9 bond despite the fact that the C9–C11 bond is aligned perfectly *anti*periplanar to the C10-oxygen bond of the epoxide.²³ More recently, Paquette and co-workers explored a possible photochemical entry to **2**. Instead of the photoinduced 1,2-shift in the anticipated vinylogous α -ketol rearrangement, however, they observed a deepseated rearrangement presumably due to the presence of the C5–C6 double bond (Scheme 10).²⁴ Photoexcitation of **41** likely generates the triplet state of the cyclopentenone, which next produces the cyclopropylcarbinyl biradical **42**. Subsequent formation of the ketene **43** accounts for the observed formation of **44** and **45**.





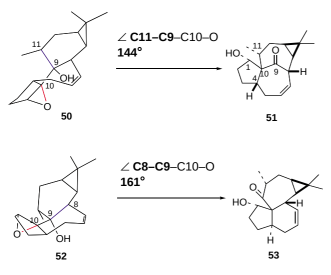
Nonetheless, a 1,2-alkyl shift was deemed by us to provide a unique solution to the challenging in-out stereochemistry of 2. Additionally, we reasoned that the irreversible semi-pinacol rearrangement of an epoxy alcohol would provide the necessary driving force for the 1,2-alkyl shift in the otherwise contra-thermodynamic direction. The choice of the Tsuchihashi–Suzuki rearrangement protocol^{25–27} was further reinforced by our successful synthesis of (+)-asteltoxin (46): a highly functionalized bis(tetrahydrofuran) 47 was readily prepared by the semi-pinacol rearrangement of an epoxy alcohol derivative 49 to enantioselectively provide the aldehyde 48 possessing the requisite quaternary center (Scheme 11).^{28,29} Other benefits in the $A + C'D \rightarrow$ $AB'C'D \rightarrow ABCD$ approach are convergence and projected ease in forming a rigid, yet strain-free, seven-membered B'-ring (to set the stage for the key semi-pinacol rearrangement).³⁰

Inspection of molecular models indicated that the epoxide **50** is conformationally rigid and that the desired migration of the C9–C11 bond would be most probable to deliver **51** in light of the stereoelectronic requirements: the C9–C11 bond could be aligned antiperiplanar to the C10-the epoxide



Scheme 11.

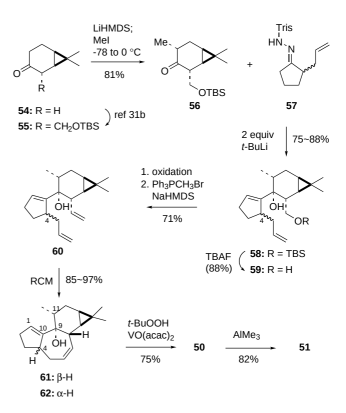
oxygen bond, whereas the C8–C9 bond is all but orthogonal (MM2 calculation results of the core skeleton are shown in Scheme 12). In the case of an isomeric epoxide **52** with the unnatural configuration at C4, it is noteworthy that migration of only the undesired C8–C9 bond could take place and that formation of **53** would be most likely.





Starting with (+)-3-carene (16), we first prepared the known, enantiomerically pure ketone 54,^{31a} which was then converted to 56 by adaptation of Shibasaki's method and subsequent methylation of 55 (Scheme 13).^{31b} The Shapiro reaction of racemic hydrazone 57 gave convenient access to the required cyclopentenyllithium and the adduct 58 was obtained in excellent yield. In a preliminary investigation, racemic 57 was employed for convenience. The vinyl group was then installed by standard methods to set the stage for ring closing olefin metathesis of 60,³² which proved to be remarkably efficient (refluxing CH₂Cl₂, 5–5.5 mM concentration) to

afford a separable 1:1 mixture of **61** and **62**. Hydroxyldirected epoxidation of allylic alcohol **61** and subsequent semi-pinacol rearrangement of the resulting epoxy alcohol **50** gave the tetracyclic core **51** bearing *in-out* intrabridgehead stereochemistry.



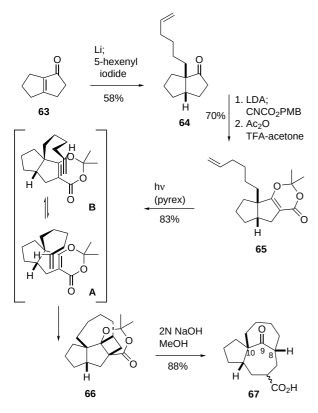
Scheme 13.

Our future plan is to complete a concise, convergent synthesis of ingenol (2) by pre-installation of all the necessary functionalities in fragment 57 prior to its coupling to 56. We also hope to develop a unified approach to the syntheses of the ingenane, tigliane, and daphnane diterpenes.

5. Total syntheses of 2

5.1. Winkler's synthesis of 2

An elegant synthetic methodology to establish the *in–out* intrabridgehead stereochemistry, concurrent with rapid increase of molecular complexity, was devised by Winkler and co-workers by utilizing a modified de Mayo photocycloaddition–retroaldol fragmentation;^{33,34} an intramolecular version was adapted to control the regio- and diastereoselectivity of the key photocycloaddition and provided the first preparation of a tricyclic ingenane system **67** having the correct *in–out* configuration (Scheme 14): preparation of the photocycloaddition resulted **65** began with reductive alkylation of **63**.^{13,35a,b} The intramolecular dioxenone [2+2] photocycloaddition resulted in exclusive formation of the *in–out* isomer **66** in 83% yield. This attractive approach is a striking example of the diastereo-control exerted by the preferred folding of the nascent (seven-membered) ring, in which conformer A would encounter the least nonbonded interactions (vis-à-vis B).



Scheme 14.

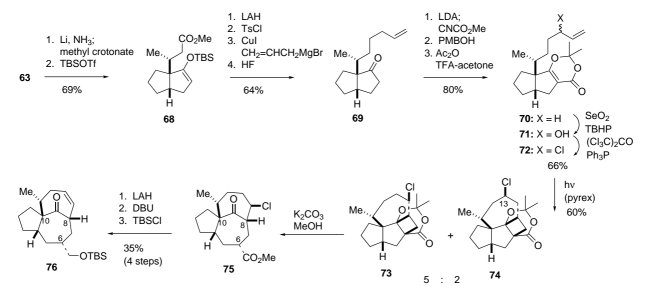
The synthetic utility of the intramolecular dioxenone photocycloaddition has been amply demonstrated by the Winkler group, including an imaginative synthesis of manzamine.³⁴

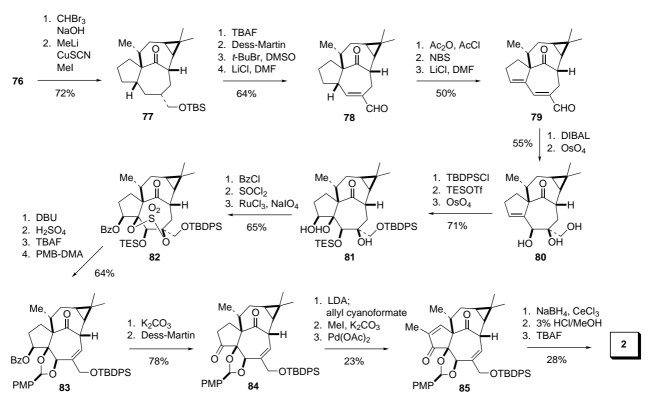
Extensive investigation for the introduction of all functionalities of the ABC rings of 2 finally culminated in its first total synthesis in 2002.⁹ The synthesis began with a highly diastereoselective (14:1) Michael addition of the enolate derived from enone 63 to establish the C11 methyl group at an early stage (Scheme 15). Dioxenone 70 was next prepared by following straightforward functional group elaboration of cyclopentanone 69. To facilitate the introduction of the cyclopropane D ring via the corresponding olefin, allylic chloride 72 was then secured as a 1:1 mixture of the C14 chloro epimers. Irradiation of 72 gave a 5:2 mixture (60% yield) of the C14 β chloro product **73** and the C13 β chloro isomer **74**. The selective formation of the former, presumably arising from the 72β isomer, closely mirrors clean conversion of 65 to 66. On the other hand, the bewildering formation of 74 has been rationalized by a series of transannular hydrogen atom abstractions initiated by the dioxenone triplet from the 72α epimer.³⁶ Retroaldol fragmentation of 73 with methanolic K_2CO_3 afforded 75, as a 7:1 ratio of C6 epimers, which was then converted to 76 by standard methods.

The tetracyclic core **77** was next obtained by dibromocarbene addition and reductive methylation. With **77** in hand, the remaining task for the completion of the synthesis entailed functionalization of the AB rings by relying on the C6 hydroxymethyl group as the sole linchpin. Diene aldehyde **79** was prepared via $\Delta^{5,6}$ unsaturated aldehyde **78** for the subsequent challenging introduction of the triol functionalities at C3, C4, and C5. Two consecutive dihydroxylation reactions occurred from the sterically less hindered β face to deliver **81**. The requisite elimination of the C6 tertiary alcohol was next accomplished via cyclic sulfate **82** by the action of DBU, and subsequent protection as a *p*-methoxybenzylidene acetal afforded **83**. Finally, the first total synthesis of **2** was completed through the intermediacy of ketone **84**.

5.2. Tanino-Kuwajima's synthesis of 2

The Tanino and Kuwajima group designed an efficient tandem cyclization–rearrangement approach to the *in–out* intrabridgehead stereochemistry by adaptation of the Nicholas reaction.^{10,37,38} An intramolecular, Lewis-acid mediated variant of *trans*-decalin **86**, initiated by a stabilized propargyl cation, provided excellent diastereocontrol at C11 as a consequence of the *E*-ethylidene

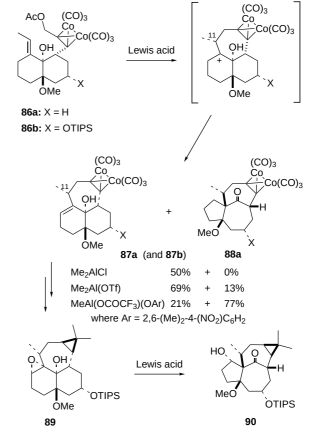




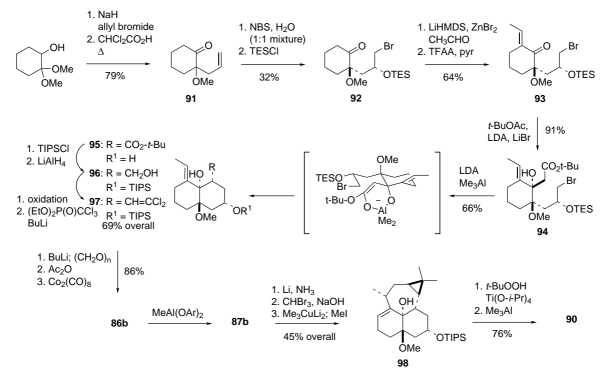
Scheme 16.

geometry (Scheme 17).³⁷ The final product distribution (i.e., elimination vs rearrangement leading to 87a and 88a, respectively) depended on the nature of aluminum-based Lewis acids, possibly due to their interaction with the tertiary hydroxyl group. Ultimately, the semi-pinacol rearrangement of the epoxy alcohol derived from 87b was successfully utilized for a total synthesis of 2. This rearrangement sequence $(89 \rightarrow 90)$ parallels the reaction pathway $9 \rightarrow 10$ (Scheme 3), but in the reverse direction. In the synthesis of 2 or analogs, therefore, the semi-pinacol rearrangement of epoxy alcohols could be profitably employed to reverse both transformations described in Scheme 3; the Tanino-Kuwajima synthesis of 2, along with the above-mentioned Cha's approach (Section 4.3), underscores the synthetic power of the semi-pinacol rearrangement of epoxy alcohols.²⁵⁻²⁸

The isolated methoxy group in the initial study (e.g., 87a and 88a) proved to be insufficient for introduction of the requisite functionalities in the AB rings. The total synthesis of 2 was made possible by way of 86b and 87b and began with a Claisen rearrangement of 2,2-dimethoxycyclohexanol, followed by bromoetherification, to afford 92 (Scheme 18). An aldol condensation of 92 with acetaldehyde and diastereoselective addition of an acetate enolate anion in the presence of LiBr (presumably to form a five-membered chelate) delivered 94. Trans-decalin 95 was then obtained by intramolecular alkylation by the action of trimethylaluminum and the observed stereochemistry is in accord with the indicated transition model. Subsequent chain elongation gave 97, which was next converted to a dicobalt-acetylene complex 86b. Under the influence of methylaluminum bis(2,6dimethyl-4-nitrophenoxide), 86b underwent exceptionally diastereoselective cyclization to afford 87b. The dicobalt

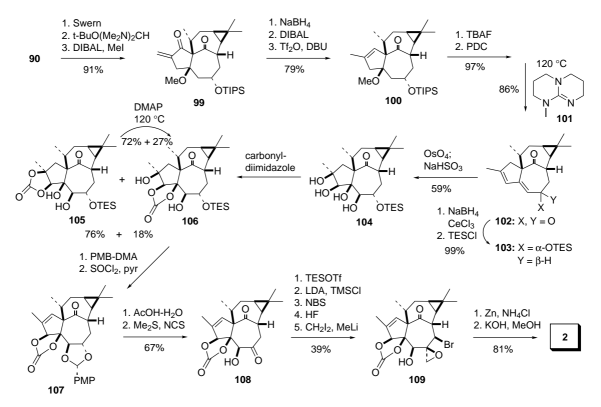






Scheme 18.

cluster was thus found useful for facile annulation of the seven-membered ring and also subsequent installation of the D ring. Hydroxy-directed epoxidation of 98 and the key semipinacol rearrangement of 89 by the action of Me₃Al yielded 90 possessing the ingenane skeleton with the correct stereochemistry. The remaining steps were directed at the taxing functionalization of the AB rings: oxidation of the secondary alcohol, use of Bredereck's reagent, and subsequent elaboration afforded cyclopentene **100** (Scheme 19). The fully conjugated dienone **102** was prepared to later introduce the triol functionalities; following the Luche reduction and



1338

silylation, dihydroxylation of **103** with an excess of osmium tetroxide gave **104** as a single isomer. Subsequent A ring functionalization was accomplished via the cyclic carbonate **106**; unfortunately, unfavorable regioselectivity in carbonate formation marred the protection sequence. As delineated in Winkler's first synthesis (Scheme 16), installation of the $\Delta^{6,7}$ double bond proved to be far from trivial. Ultimately, the allylic alcohol moiety in the B ring was introduced by reductive elimination of epoxide **109** to complete a total synthesis of **2**.

5.3. Kigoshi's formal synthesis of 2

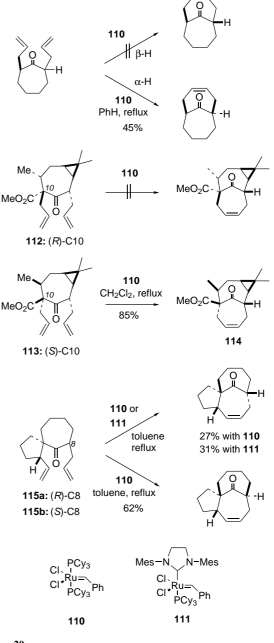
The Kigoshi group reported direct cyclization of a sevenmembered ring to construct the *in,out*-bicyclo[4.4.1]undecane skeleton of **2** by olefin metathesis.³⁹ Central to successful annulation of a highly strained seven-membered ring is the presence of the A ring, which would help constrain the otherwise flexible conformation of the pendant side chain and also bring closer the terminal olefins. Without the A ring, for example, ring-closing olefin metathesis (RCM) failed to afford the desired cyclization product; instead oligomerization was observed (Scheme 20). The prerequisite of the A ring for RCM was independently demonstrated by Wood and co-workers with **112** and **113**.⁴⁰

Starting with Funk's keto ester 17, the A ring in 118 was first constructed by intramolecular alkylation of 117 (Scheme 21). After considerable experimentation, use of a sterically hindered base was found to be essential for high diastereoselectivity of the spirocyclization step. The key substrates 119 and 120 were then prepared by straightforward allylation; RCM investigations showed that the second-generation catalyst 111 was more efficacious than 110 to provide 121 in 53% yield under optimized conditions [refluxing toluene, shorter reaction time (30 min), 1.5 mM concentration]. Under identical conditions, 120 afforded 122 in impressive (87%) yield, which is undoubtedly attributable to the well-known stability of the trisubstituted olefin toward the catalyst 111 to thwart competing ring opening reactions. Nonetheless, it is worth mentioning that a relatively high temperature is necessary for formation of 121 and 122 (compared to 61 and 62). Finally, allylic oxidation of 122 with SeO_2 gave 78; since the latter had been converted to 2 by Winkler, this work constitutes a formal synthesis of **2**.

Particularly noteworthy in Kigoshi's synthesis of **122** is the facility of RCM to effect closure to strained sevenmembered carbocycles. This example is another testimonial to the distinctive utility of RCM in organic synthesis.

5.4. Wood's synthesis of (+)-2

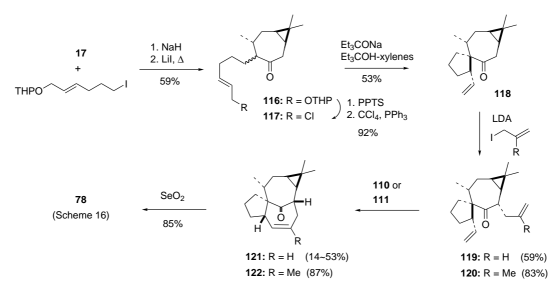
Independent of Kigoshi's work, Wood and co-workers reported a closely related RCM strategy to build the strained *in-out* ABCD ring system $(A + CD \rightarrow ABCD)$.⁴⁰ As pointed out in Scheme 20, successful cyclization is predicated on incorporation of the five-membered A ring, which was conveniently installed by a Diels–Alder reaction of cyclopentadiene (vide supra). However, tandem ringopening and ring-closing metathesis of **123** did not occur; instead only ring-opening metathesis was observed to



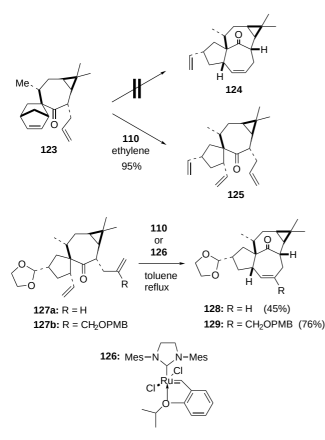
Scheme 20.

provide triene **125** in excellent yield under an atmosphere of ethylene (Scheme 22). A simple solution to circumvent competing reversion of **125** to **123** was to block the C2 olefin (e.g., in **125**) prior to the RCM step. As observed by Kigoshi,¹² the formation of a more robust trisubstituted olefin product (**129** vs **128**) benefited from improved yield (76 vs 45%) and lower catalyst loading (25 vs 80 mol%). Although the Hoveyda catalyst **126** was required for the formation of **129** in acceptable yields,^{11,32e} **129** has the built-in advantage of possessing the requisite C20 hydroxymethyl group.

Making use of Funk's ketoester **17**, the Lewis acidcatalyzed Diels–Alder reaction between **131** and cyclopentadiene gave a 20:8:1 mixture of three diastereomers; the major diastereomer (*endo* cycloadduct) **132** possessed the requisite stereochemistry (Scheme 23).^{11,40} Subsequent



Scheme 21.



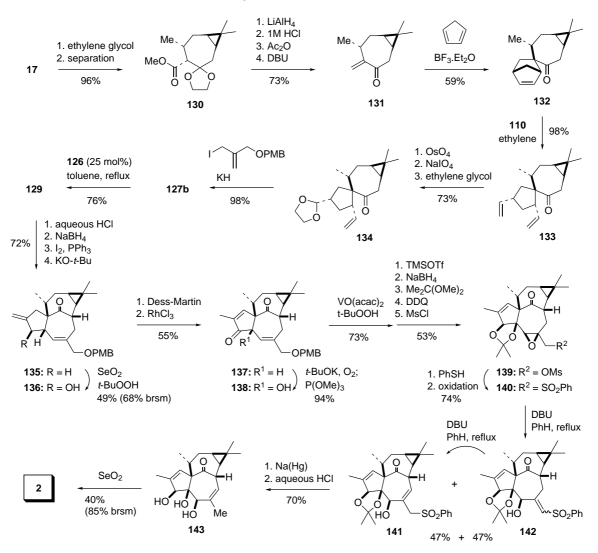


RCM of 132 with ethylene, selective functionalization of the desired olefin of 133, and allylation of 134 then delivered 127b. As mentioned above, RCM of 127b gave the key tetracycle 129 in good yield. Next, straightforward elaboration, including regioselective allylic oxidation of 135, afforded 137, and oxidation of its enolate cleanly yielded α -hydroxy ketone 138. Reduction of 137 or 138 proved to be futile, since it occurred from the less hindered, convex face to give the undesired *anti*-diol derivatives. The taxing reduction was, therefore, deferred to a later stage. Hydroxyl-directed epoxidation at the $\Delta^{5,6}$ double bond of **138** was then undertaken. Interestingly, protection of the C4 tertiary alcohol as the TMS ether and subsequent reduction of the C3 keto group stereoselectively afforded the desired β -alcohol. Just what subtle factors influence the preferred conformation and reactivity of the ingenane skeleton has not been determined.

The remaining task involved functionalization of the B ring. As was the case with both previous syntheses by Winkler and Tanino–Kuwajima, another obstacle had to be overcome to introduce the $\Delta^{6,7}$ double bond. For example, **138** proved to be recalcitrant toward oxidation by singlet oxygen. Ultimately, the unusually sluggish ring opening of the C5, C6 epoxide succeeded by use of DBU on the C20 sulfoxide; both β , γ and α , β -unsaturated sulfones **141** and **142** were obtained in a 1:1 ratio. The final conversion of the allylic sulfone **141** to the corresponding primary alcohol at C20 was achieved by a reductive removal–oxidation sequence in order to complete a total synthesis of **2**.

6. Conclusion

Recently emerged three ground-breaking total syntheses of ingenol (2) and a formal synthesis: these syntheses highlight attractive approaches to the highly strained inside, outside topography of the ingenane diterpenes, the principal synthetic challenge. Also significant are resourceful maneuvers that were deployed for elaboration of rather under-functionalized advanced intermediates to stereoselectively install the dazzling array of dense functionalities on the southern periphery. These total syntheses of 2, while stunning feats in natural product synthesis, were somewhat hamstrung by a linear sequence of multi-step transformations to hurdle the latter challenge: the unique intricacy, posed by the high degree of oxygenation and the surprisingly difficult introduction of the $\Delta^{6,7}$ double bond. should be addressed in future studies for more convergent, step-economical syntheses to be reduced to practice. New powerful methodology (e.g., ring-closing olefin metathesis) will undoubtedly aid in streamlining the total syntheses of



Scheme 23.

structurally complex target molecules such as **2**. Also likely is the development of unified strategies for synthesizing ingenanes, tiglianes, and daphnanes.

It is hoped that synthetic studies will help shed light on the structure–activity relationships of these biologically potent natural products, elucidate the molecular basis for their biological activity, and eventually lead to the development of useful biochemical tools and new therapeutic agents.

Acknowledgements

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Biographical sketch





Jin K. Cha received a BS degree in chemistry from Seoul National University in 1975. He received his DPhil under the direction of Professor Sir Jack E. Baldwin at Oxford University in 1981. Following two-year postdoctoral training under Professor Y. Kishi at Harvard University, he held faculty appointments at Vanderbilt University and the University of Alabama, Tuscaloosa. In Fall 2002 he moved to Wayne State University, where he is currently Professor of Chemistry.

Oleg L. Epstein graduated from Belarussian State University in Minsk, Belarus in 1993. He stayed on the same university and obtained his PhD degree in 2001 under the supervision of Professor Oleg G. Kulinkovich. In 2002 he joined Professor Cha's group as a postdoctoral fellow.



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Synthesis of stable azomethine ylides by the rearrangement of 1,3-dipolar cycloadducts of 3,4-dihydroisoquinoline-2-oxides with DMAD

Necdet Coşkun* and Selen Tunçman

Department of Chemistry, Uludağ University, 16059 Bursa, Turkey

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Abstract—1-Aryl-6,7-dimethoxy-1,2,3,4-tetrahydroisoquinolines were prepared according to a one-pot procedure involving the reaction of 2-(3,4-dimethoxyphenyl)-ethylamine with aromatic aldehydes in TFA at reflux. The tetrahydroisoquinolines were treated with $H_2O_2-WO_4^2$ in methanol at room temperature to give the corresponding 3,4-dihydroisoquinoline-2-oxides. Treatment of these cyclic nitrones with DMAD in toluene at room temperature gave the corresponding isoxazolo[3,2-*a*]isoquinolines. These compounds were heated in toluene at reflux to give the corresponding yildes (Method A). The effect of the substituents on the rate of the rearrangement of such compounds prompted us to discuss a new mechanism involving consecutive C–C bond heterolysis and 1,3-sigmatropic shift. A one-pot reaction involving the treatment of the nitrones with equimolar amounts of DMAD in refluxing toluene also gave the ylides (Method B). The structures of the prepared compounds were elucidated by spectral means and elemental analyses. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

The synthetic utility of the 1,3-dipolar cycloaddition reaction is evident from the number and the scope of targets that can be prepared by this chemistry. Nitrones are the most useful through their ability to generate nitrogenand oxygen based functionality from the cycloadducts.¹ The cycloadducts of di- and triarylimidazoline 3-oxides² with a variety of dipolarophiles³ give bicyclic compounds with potentially interesting biological activity.⁴ On the other hand, they are a source of new heterocyclic compounds via interesting ring-opening reactions.⁵

Previously, we reported the synthesis of stable adducts of Δ^3 -imidazoline 3-oxides with DMAD^{3d,e} and 3-phenylpropanoic acid alkyl esters.^{3f} Thermally and base-induced ringopening reactions of these adducts were demonstrated. As a continuation of our interest in the ring-opening reactions of 4-isoxazolines,^{3d,e} we prepared 1-aryl-3,4-dihydroisoquino-line-2-oxides from the oxidation of 1-aryl-1,2,3,4-tetrahydroisoquinolines under the conditions recently reported⁶

and their adducts with DMAD. It is known that nitrones react with alkynes to give generally unstable adducts or those, which are stable can be subjected to rearrangements under thermal conditions. Rearrangements of DMAD adducts of some heterocyclic *N*-oxides has been reviewed.^{1a} 4,5-Dihydroimidazole *N*-oxides undergo 1,3-dipolar cyclo-addition with alkyne dipolarophiles and the cycloadducts were shown to convert to the corresponding ene-1,1-diamines.⁷ The thermal reaction of some 4-isoxazoline derivatives leading to isoquinoline-fused pyrroles has been investigated and it was found that the pathway of the rearrangement to pyrroles is consistent with a route involving an acylaziridine.⁸

2. Results and discussion

We report herein the synthesis of 1-aryltetrahydroisoquinolines **2a–e** and their oxidation with $H_2O_2-WO_4^{2-}$ in methanol at room temperature to give cyclic nitrones **3a–e**. Isolated or in situ formed 8,9-dimethoxy-10b-aryl-6,10bdihydro-5*H*-isoxazolo[3,2-*a*]isoquinoline-1,2-dicarboxylic acid dimethyl esters **4a–e** were shown to undergo substituent–dependent rearrangement to novel stable 3,4dihydroisoquinolinium *N*-ylides **5a–e** (Scheme 1). The results are presented in Table 1. A new mechanism involving consecutive C–C bond heterolysis and 1,3-sigmatropic shift is discussed.

Keywords: Isoquinoline; 1-Aryl-1,2,3,4-tetrahydroisoquinoline; THI; Pictet–Spengler; Oxidation with H₂O₂–tungstate; 3,4-Dihydroisoquinoline-2-oxide; Rearrangement; Isoxazoloisoquinoline; Stable azomethine ylide; 4-Isoxazoline rearrangement mechanism; Alkyne; DMAD; Dipolar cycloaddition; Synthesis; Heterocycles.

^{*} Corresponding author. Tel.: +90 44292561308; fax: +90 4428022; e-mail: coskun@uludag.edu.tr

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1–5	Ar	Yield (%	6)	
		2	3	4
a	Ph	70 ^a	43 ^b	95°
b	$3,4 (MeO)_2C_6H_3$	85	50	97
с	$3-NO_2C_6H_4$	80	40	96
d	$4-ClC_6H_4$	70	69	98
e	3,4 (OCH ₂ O)C ₆ H ₃	60	45	97

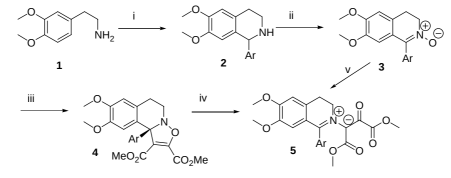
Table 1. Synthesis of compounds 2a-e, 3a-e and 4a-e

^a The reaction times were 3, 5.5, 2.5, 5, 5.5 h for **2a,b,c,d,e**, respectively.

^b The reaction times were 5.5, 17.5, 23, 21, 19 for **3a,b,c,d,e**, respectively.

^c The reaction times were 18, 5.5, 24, 18, 15 for **4a,b,c,d,e**, respectively.

corresponding nitrones 3a-e in toluene in the presence of DMAD (see Table 2, Method B). It was shown that isoxazolo[3,2-*a*]isoquinolines convert at different rates to the corresponding ylides 5a-e. The structure of stable 3,4-dihydroisoquinolinium *N*-ylides 5a-e was deduced from their elemental analyses and spectral data. The compounds are highly coloured and soluble in diluted acids with loss of their colours. The extraction of the acidic water solutions of ylides 5 with CHCl₃ again affords the free ylides 5. Our preliminary experiments show that they react, as expected, with dipolarophiles such as phenyl isocyanate.



Scheme 1. Reagents and conditions: (i) ArCHO; TFA; reflux; (ii) H₂O₂-Na₂WO₄; MeOH; rt; (iii) DMAD; toluene; rt; (iv) toluene; reflux; (v) DMAD; toluene; reflux.

2-(3,4-Dimethoxyphenyl)-ethylamine **1** was reacted with an equimolar amount of the corresponding aromatic aldehyde in refluxing TFA to give in good yields the corresponding 1-aryl-1,2,3,4-tetrahydroisoquinolines **2a–e**.

Compounds **2** were treated with $H_2O_2-WO_4^{2-}$ in methanol according to a method we have recently reported⁶ to give 3,4-dihydroisoquinoline-2-oxides **3a–e**. The products were purified by chromatographic methods and were recrystal-lized from ethanol–ether (1/3).

Nitrones **3a–e** were reacted with DMAD in toluene at room temperature to give quantitatively the corresponding isoxazolo[3,2-*a*]isoquinolines **4a–e**. The products were purified by recrystallization from ethanol in the cases of **4a,c,e** and preparative TLC in the cases of **4b,d**. The NMR as well as the infra red spectral data for compounds **4a–e** are in good agreement with those we have previously reported for similar adducts.^{3d–f} Isolated **4a–e** were refluxed in toluene for the times specified in Table 2 (Method A) to give heretofore unreported exclusively stable azomethine ylides **5a–e**. The methods available for generating azomethine ylides, were discussed in a resent review.⁹ The same products resulted from the direct heating of the

Table 2. Synthesis of N-ylides 5a-e

	Yield	Yield		
	Method A ^a	Method B ^b	A	В
5a	93	75	11	12
5b	100	74	1.5	1.5
5c	82	95	7	8
5b 5c 5d 5e	91	71	13	14
5e	87	96	4	4.2

^a Yields are based on the starting 4.

^b Yields based on the starting **3**.

On the other hand their reactions with amines as diethylamine lead to the formation of corresponding 3,4-dihydroisoquinoline. The ¹³C NMR spectroscopic assignments specifically for **5e** are shown in Figure 1.

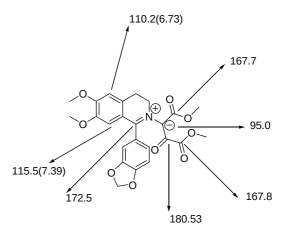
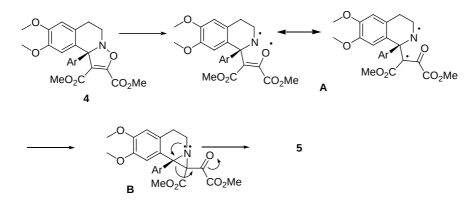


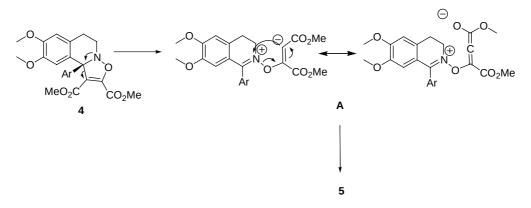
Figure 1. Some ¹H and ¹³C NMR assignments for compound 5e.¹⁰

Electron donating groups on the aromatic ring at C-10b of compounds **4** increase the rate of rearrangement to ylides **5** while electron-withdrawing groups (see Table 2 for the reaction times) decrease it.

Aziridines are generally assumed to be involved in the rearrangements of 4-isoxazolines.⁹ A similar approach could be assumed for the conversion of compounds 4 to 5 as depicted in Scheme 2. The homolysis of N–O bond in compounds 4 could give diradicals A, which could cyclize to the corresponding aziridines B. Thermal ring-opening of aziridine part of B could give ylides 5 (see Scheme 2).



Scheme 2. Probable aziridine involving mechanism for the rearrangement of isoxazoloisoquinolines 4.



Scheme 3. Probable C-C bond heterolysis involving mechanism for the rearrangement of isoxazoloisoquinolines 4.

However, the pronounced substituent effects discussed above do not support the acylaziridine intermediate in the rearrangement of 4-isoxazolines. It is expected that the substituents on the 10b-phenyl will affect neither homolysis nor heterolysis of the N–O bond in the isoxazoline part of compounds **4**. This prompted us to consider an alternative mechanism outlined in Scheme 3. Electron donating groups on the aromatic ring of **4** probably favour the C-3, C-4 bond heterolysis to give zwitter ions **A** stabilised by resonance, which in turn undergo 1,3-sigmatropic rearrangement to give ylides **5a–e**. The electron donating groups on the aromatic ring at C-10b could stabilise the forming azomethine ylides by their +R effects.

Thus, 1-aryltetrahydroisoquinolines prepared according to Pictet–Spengler procedure from 2-(3,4-dimethoxyphenyl)ethylamine and the corresponding aromatic aldehydes were oxidized to nitrones **3** the 1,3-dipolar cycloaddition products of which with DMAD were shown to afford previously unknown and stable 3,4-dihydroisoquinolinium N-ylides when heated in toluene. A plausible mechanism involving consecutive C–C bond heterolysis and 1,3-sigmatropic shift was discussed.

3. Experimental

Melting points were recorded on an Electrothermal Digital melting point apparatus. Infrared spectra were recorded on a Mattson 1000 FTIR. NMR spectra were recorded on a Mercury Plus 400 MHz spectrometer. UV/vis spectra of compounds **5a–e** were recorded on a Shimadzu UV-2100 spectrophotometer. TLC controls were performed using silica gel coated aluminium sheets. Chloroform, petroleum ether, methanol and acetone (45:40:10:5) solvent mixture was used as an eluent system. Visualisation was effected with UV light. The elemental analyses were performed on a EuroEA 3000 CHNS analyser.

3.1. Synthesis of 1-aryl-1,2,3,4-tetrahydroisoquinolines 2. General procedure

To a solution of 2-(3,4-dimethoxyphenyl)-ethylamine (5 mmol, 0.9062 g) in TFA (3 mL) the corresponding aldehyde (5 mmol) was added and the solution was refluxed for the time specified in Table 1. The reaction mixture was poured onto ice and basified with sodium hydroxide. The mixture was extracted with chloroform (3×10 mL) and the combined extracts were dried over anhydrous Na₂SO₄. The organic solvent was evaporated under vacuum and the residue was crystallized from ethanol.

3.1.1. 6,7-Dimethoxy-1-phenyl-1,2,3,4-tetrahydroisoquinoline 2a. $R_{\rm f}$ =0.31; yield 0.943 g, 70%; mp 110–111 °C; IR (KBr) $\nu_{\rm NH}$ 3328 cm⁻¹. (400 MHz, CDCl₃): δ 1.87 (1H, s), 2.64–2.71 (1H, m), 2.82–2.90 (1H, m), 2.94–2.99 (1H, m), 3.11–3.17 (1H, m), 3.55 (3H, s), 3.79 (3H, s), 4.97 (1H, s), 6.17 (1H, s), 6.56 (1H, s), 7.17–7.26 (5H, m). ¹³C NMR (100 MHz, CDCl₃): δ 29.7; 42.3; 56.3; 56.4; 61.9; 111.4; 111.9; 127.8; 128.1; 128.8; 129.3; 130.3; 145.3; 147.5; 148.1. Anal. Calcd for $C_{17}H_{19}NO_2$ (269.34) C, 75.81; H, 7.11; N, 5.20; Found C, 75.75; H, 7.20; N, 5.30.

3.1.2. 1-(3,4-Dimethoxyphenyl)-6,7-dimethoxy-1,2,3,4-tetrahydroisoquinoline 2b. $R_{\rm f}$ =0.23; yield 85%; mp 88–89 °C; IR (KBr) $\nu_{\rm NH}$ 3567 cm⁻¹. (400 MHz, CDCl₃): δ 1.83 (1H, s), 2.71–2.77 (1H, m), 2.92–2.99 (1H, m), 3.03–3.1 (1H, m), 3.22–3.28 (1H, m), 3.66 (3H, s), 3.84 (3H, s), 3.88 (3H, s), 3.89 (3H, s), 4.99 (1H, s), 6.28 (1H, s), 6.63 (1H, s), 6.78–6.83 (3H, m). ¹³C NMR (100 MHz, CDCl₃): δ 29.3; 42.2; 55.8; 55.9; 56.0; 61.5; 110.7; 110.9; 111.4; 111.8; 121.3; 127.6; 130.1; 137.3; 147.0; 147.6; 148.3; 149.0. Anal. Calcd for C₁₉H₂₃NO₄ (329.39) C, 69.28; H, 7.04; N, 4.25; Found C, 69.35; H, 6.88; N, 4.23.

3.1.3. 6,7-Dimethoxy-1-(3-nitrophenyl)-1,2,3,4-tetrahydroisoquinoline 2c. $R_{\rm f}$ =0.15; yield 1.257 g, 80%; mp 109–111 °C; IR (KBr) $\nu_{\rm NH}$ 3312 and 3256 cm⁻¹. (400 MHz, CDCl₃): δ 1.77 (1H, s), 2.73–2.79 (1H, m), 2.91–2.98 (1H, m), 3.04–3.10 (1H, m), 3.14–3.20 (1H, m), 3.64 (3H, s), 3.89 (3H, s), 5.16 (1H, s), 6.17 (1H, s), 6.66 (1H, s), 7.49 (1H, t, *J*=7.6 Hz), 7.61 (1H, d, *J*=7.6 Hz), 8.12–8.17 (2H, m). ¹³C NMR (100 MHz, CDCl₃): δ 29.1; 41.6; 55.8; 55.9; 60.7; 110.7; 111.8; 122.5; 123.8; 127.9; 128.2; 129.3; 135.2; 147.2; 147.3; 148.1; 148.4. Anal. Calcd for C₁₇H₁₈N₂O₄ (314.34) C, 64.96; H, 5.77; N, 8.91; Found C, 64.94; H, 5.75; N, 9.02.

3.1.4. 1-(4-Chlorophenyl)-6,7-dimethoxy-1,2,3,4-tetrahydroisoquinoline 2d. $R_f = 0.54$; yield 1.063 g, 70%; mp 103–105 °C; IR (KBr) $\nu_{\rm NH}$ 3242 cm⁻¹. (400 MHz, CDCl₃): δ 1.81 (1H, s), 2.71–2.77 (1H, m), 2.88–2.96 (1H, m), 3.01–3.07 (1H, m), 3.16–3.22 (1H, m), 3.64 (3H, s), 3.87 (3H, s), 5.02 (1H, s), 6.20 (1H, s), 6.63 (1H, s), 7.20 (2H, d, J=8.0 Hz), 7.29 (2H, d, J=8.0 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 29.2; 41.8; 55.8; 55.9; 60.8; 110.8; 111.5; 127.7; 128.5; 129.3; 130.3; 133.1; 143.4; 147.1; 147.8. Anal. Calcd for C₁₇H₁₈CINO₂ (303.78) C, 67.21; H, 5.97; N, 4.61; Found C, 67.10; H, 5.97; N, 4.75.

3.1.5. 1-Benzo[**1**,**3**]**dioxol-5-yl-6**,**7-dimethoxy-1**,**2**,**3**,**4-tetrahydroisoquinoline 2e.** $R_{\rm f}$ =0.37; yield 0.940 g, 60%; mp 133–134 °C; IR (KBr) $\nu_{\rm NH}$ 3252 cm⁻¹. (400 MHz, CDCl₃): δ 1.76 (1H, s), 2.70–2.75 (1H, m), 2.88–2.95 (1H, m), 3.0–3.06 (1H, m), 3.19–3.24 (1H, m), 3.67 (3H, s), 3.87 (3H, s), 4.97 (1H, s), 5.94 (2H, s), 6.28 (1H, s), 6.62 (1H, s), 6.71–6.77 (3H, m). ¹³C NMR (100 MHz, CDCl₃): δ 29.3; 41.9; 55.8; 55.9; 61.2; 101.0; 107.9; 109.2; 110.9; 111.42; 122.2; 127.7; 129.9; 139.1; 146.8; 147.1; 147.7; 147.7. Anal. Calcd for C₁₈H₁₉NO₄ (313.35) C, 68.99; H, 6.11; N, 4.47; Found C, 68.90; H, 5.99; N, 4.55.

3.2. Synthesis of 1-aryl-6,7-dimethoxy-1,2,3,4-tetrahydroisoquinoline-2-oxides 3a–e. General procedure

To a solution of tetrahydroisoquinoline **2** (0.5 mmol) in methanol (10 mL) H_2O_2 (35%, 2 mmol) was added in the presence of $Na_2WO_4 \cdot H_2O$ (0.025 mmol, 8.3 mg). The reaction mixture was stirred at room temperature for the specified time. The solvent was evaporated and water (15 mL) was added to the residue and extracted with chloroform (3×10 mL). The combined extracts were dried

and the solvent evaporated. The purification was performed by preparative TLC using silica gel as adsorbent and chloroform, petroleum ether, methanol and acetone (45:40:10:5) solvent mixture as an eluent.

3.2.1. 6,7-Dimethoxy-1-phenyl-1,2,3,4-tetrahydroisoquinoline-2-oxide 3a. $R_{\rm f}$ =0.51; yield 0.061 g, 43%; mp 156–157 °C; IR (KBr) $\nu_{\rm C=N}$ 1590 cm⁻¹; $\nu_{\rm N-O}$ 1286 cm⁻¹. (400 MHz, CDCl₃): δ 3.15 (2H, t, J=7.6 Hz), 3.62 (3H, s), 3.91 (3H, s), 4.26 (2H, t, J=7.6 Hz), 6.36 (1H, s), 6.76 (1H, s), 7.43–7.49 (3H, m), 7.56 (2H, d, J=7.02 Hz). (100 MHz, CDCl₃): δ 27.9; 56.3; 56.4; 59.8; 110.5; 110.6; 123.6; 125.7; 128.5; 129.6; 130.4; 131.4; 142.3; 147.9; 149.6. Anal. Calcd for C₁₇H₁₇NO₃ (283.32) C, 72.07; H, 6.05; N, 4.94; Found C, 72.05; H, 5.99; N, 4.97.

3.2.2. 1-(3,4-Dimethoxyphenyl)-6,7-dimethoxy-1,2,3,4-tetrahydroisoquinoline *N*-**oxide 3b.** $R_f = 0.28$; yield 0.086 g, 50%; mp 165–166 °C; IR (KBr) $\nu_{C=N}$ 1590 cm⁻¹; ν_{N-O} 1283 cm⁻¹. (400 MHz, CDCl₃): δ 3.14 (2H, t, J=7.2 Hz), 3.65 (3H, s), 3.87 (3H, s), 3.91 (3H, s), 3.93 (3H, s), 4.24 (2H, t, J=7.2 Hz), 6.45 (1H, s), 6.75 (1H, s), 6.95 (1H, d, J=8.4 Hz), 7.13 (1H, d, J=8.4 Hz) 7.20 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 27.9; 56.1; 56.2; 56.3; 56.4; 59.8; 110.6; 110.8; 111.8; 113.5; 123.5; 123.6; 123.7; 125.9; 142.3; 147.9; 148.7; 149.6; 149.9. Anal. Calcd for C₁₉H₂₁NO₅ (343.37) C, 66.46; H, 6.16; N, 4.08; Found C, 66.40; H, 6.34; N, 4.06.

3.2.3. 6,7-Dimethoxy-1-(3-nitrophenyl)-1,2,3,4-tetrahydroisoquinoline *N*-oxide **3c**. $R_{\rm f}$ =0.54; yield 0.066 g, 40%; mp 172–173 °C; IR (KBr) $\nu_{\rm C=N}$ 1584 cm⁻¹; $\nu_{\rm N-O}$ 1284 cm⁻¹. (400 MHz, CDCl₃): δ 3.20 (2H, t, *J*=7.6 Hz), 3.65 (3H, s), 3.95 (3H, s), 4.29 (2H, t, *J*=7.6 Hz), 6.31 (1H, s), 6.81 (1H, s), 7.69 (1H, t, *J*=8.0 Hz), 8.01 (1H, d, *J*= 8.0 Hz), 8.30 (1H, d, *J*=8.0 Hz), 8.51 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 27.6; 56.2; 56.3; 59.9; 109.8; 110.8; 122.2; 124.2; 125.6; 125.7; 129.3; 132.8; 136.6; 139.7; 148.0; 148.1; 149.9. Anal. Calcd for C₁₇H₁₆N₂O₅ (328.32) C, 62.19; H, 4.91; N, 8.53; Found C, 62.10; H, 4.95; N, 8.66.

3.2.4. 1-(4-Chlorophenyl)-6,7-dimethoxy-1,2,3,4-tetrahydroisoquinoline *N*-oxide **3d.** $R_{\rm f}$ =0.56; yield 0.110 g, 69%; mp 216–217 °C; IR (KBr) $\nu_{\rm C=N}$ 1595 cm⁻¹; $\nu_{\rm N-O}$ 1284 cm⁻¹. (400 MHz, CDCl₃): δ 3.15 (2H, t, *J*=7.6 Hz), 3.65 (3H, s), 3.92 (3H, s), 4.25 (2H, t, *J*=7.6 Hz), 6.35 (1H, s), 6.76 (1H, s), 7.45 (2H, d, *J*=8.4 Hz), 7.57 (2H, d, *J*= 8.4 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 27.9; 56.3; 56.4; 59.9; 110.3; 110.7; 123.1; 125.8; 128.8; 129.7; 132.0; 135.5; 141.3; 148.1; 149.7. Anal. Calcd for C₁₇H₁₆ClNO₃ (317.77) C, 64.26; H, 5.08; N, 4.41; Found C, 64.40; H, 5.03; N, 4.42.

3.2.5. 1-Benzo[**1**,3]dioxol-5-yl-6,7-dimethoxy-1,2,3,4tetrahydroisoquinoline *N*-oxide **3e**. R_f =0.56; yield 0.074 g, 45%; mp 169–170 °C; IR (KBr) $\nu_{C=N}$ 1590 cm⁻¹; ν_{N-O} 1288 cm⁻¹. (400 MHz, CDCl₃): δ 3.13 (2H, t, *J*= 8.0 Hz), 3.68 (3H, s), 3.91 (3H, s), 4.23 (2H, t, *J*=8 Hz), 6.02 (2H, s), 6.44 (1H, s), 6.74 (1H, s), 6.89 (1H, d, *J*=8.0 Hz), 7.16 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 27.9; 56.3; 56.4; 59.8; 101.6; 108.4; 110.5; 110.7; 111.0; 123.6; 123.7; 124.7; 125.8; 141.9; 147.7; 147.9; 148.6; 149.6. Anal. Calcd for C₁₈H₁₇NO₅ (327.33) C, 66.05; H, 5.23; N, 4.28; Found C, 66.00; H, 5.20; N, 4.08.

3.3. Synthesis of isoxazolo[3,2-*a*]isoquinolines 4a–e. General procedure

To a solution of nitrone **3** (0.15 mmol) in toluene (10 mL) DMAD (0.225 mmol, 0.032 g) was added and the reaction mixture stirred for the specified time. The solvent was evaporated under vacuum and the residue crystallized from ethanol in the cases of **4a,c,e**. Compounds **4b,d** were purified by preparative TLC.

3.3.1. 8,9-Dimethoxy-10b-phenyl-6,10b-dihydro-5*H***-isoxazolo[3,2-***a***]isoquinoline-1,2-dicarboxylic acid dimethyl ester 4a. R_f=0.22; yield 0.061 g, 95%; mp 124–125 °C; IR (KBr) \nu_{C=0} 1758; 1712 cm⁻¹; \nu_{C=C} 1626 cm⁻¹. (400 MHz, CDCl₃): \delta 2.64–2.70 (1H, m), 3.14–3.22 (1H, m), 3.26–3.33 (1H, m), 3.64–3.72 (1H, m), 3.66 (3H, s), 3.68 (3H, s), 3.87 (3H, s), 3.89 (3H, s), 6.63 (1H, s), 6.99 (1H, s), 7.27–7.37 (5H, m). ¹³C NMR (100 MHz, CDCl₃): \delta 23.8; 47.1; 52.2; 53.4: 56.0; 56.1; 77.3; 110.8; 112.3; 114.9; 126.6; 126.8; 128.2; 128.3; 129.1; 142.8; 147.7; 148.5; 153.5; 159.8; 163.6. Anal. Calcd for C₂₃H₂₃NO₇ (425.43) C, 64.93; H, 5.45; N, 3.29; Found C, 65.10; H, 5.55; N, 3.40.**

3.3.2. 10b-(**3,4**-Dimethoxyphenyl)-**8**,9-dimethoxy-**6**,10**b**-dihydro-5*H*-isoxazolo[**3**,2-*a*]isoquinoline-**1**,2dicarboxylic acid dimethyl ester **4b**. R_f =0.89; yield 0.071 g, 97%; oil; IR (KBr) $\nu_{C=0}$ 1758; 1712 cm⁻¹; $\nu_{C=C}$ 1626 cm⁻¹. (400 MHz, CDCl₃): δ 2.60–2.70 (1H, m), 3.15–3.22 (1H, m), 3.24–3.32 (1H, m), 3.64–3.72 (1H, m), 3.67 (3H, s), 3.71 (3H, s), 3.79 (3H, s), 3.85 (3H, s), 3.86 (3H, s), 3.89 (3H, s), 6.62 (1H, s), 6.76 (1H, d, *J*=8.4 Hz), 6.82 (1H, dd, *J*=8.4, 2.0 Hz), 6.96 (1H, d, *J*=2.0 Hz), 7.04 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 23.7; 46.9; 52.3; 53.3: 56.0; 56.1; 77.0; 110.4; 110.8; 112.3; 112.4; 115.7; 121.9; 126.6; 126.7; 135.0; 147.6; 148.4; 148.7; 149.0; 152.9; 159.7; 163.8 Anal. Calcd for C₂₅H₂₇NO₉ (485.48) C, 61.85; H, 5.61; N, 2.89; Found C, 61.80; H, 5.63; N, 2.89.

3.3.3. 8,9-Dimethoxy-10b-(3-nitrophenyl)-6,10b-dihydro-*5H*-isoxazolo[3,2-*a*]isoquinoline-1,2-dicarboxylic acid dimethyl ester 4c. R_f =0.79; yield 0.068 g, 96%; mp 123–124 °C; IR (KBr) $\nu_{C=0}$ 1758; 1719 cm⁻¹; $\nu_{C=C}$ 1644 cm⁻¹. (400 MHz, CDCl₃): δ 2.72–2.77 (1H, m), 3.15–3.24 (1H, m), 3.25–3.31 (1H, m), 3.64–3.72 (1H, m), 3.67 (3H, s), 3.70 (3H, s), 3.87 (3H, s), 3.89 (3H, s), 6.65 (1H, s), 6.89 (1H, s) 7.50 (1H, t, *J*=8.2 Hz), 7.8 (1H, d, *J*= 8.2 Hz), 8.16 (1H, d, *J*=8.2 Hz), 8.25 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 23.9; 47.3; 52.5; 53.5: 56.1; 77.2; 111.2; 111.6; 114.1; 123.3; 124.1; 125.2; 126.8; 129.3; 135.2; 145.5; 148.1; 148.3; 148.9; 153.9; 159.5; 163.4. Anal. Calcd for C₂₃H₂₂N₂O₉ (470.43) C, 58.72; H, 4.71; N, 5.95; Found C, 58.80; H, 4.90; N, 6.10.

3.3.4. 10b-(4-Chlorophenyl)-8,9-dimethoxy-6,10b-dihydro-5*H*-isoxazolo[3,2-*a*]isoquinoline-1,2-dicarboxylic acid dimethyl ester 4d. $R_{\rm f}$ =0.74; yield 0.068 g, 98%; oil; IR (KBr) $\nu_{\rm C=0}$ 1755; 1709 cm⁻¹; $\nu_{\rm C=C}$ 1638 cm⁻¹. (400 MHz, CDCl₃): δ 2.63–2.0 (1H, m), 3.12–3.20 (1H, m), 3.23–3.30 (1H, m), 3.64–3.72 (1H, m), 3.67 (3H, s), 3.69 (3H, s), 3.87 (3H, s), 3.89 (3H, s), 6.62 (1H, s), 6.95 (1H, s), 7.28–7.32 (4H, m). ¹³C NMR (100 MHz, CDCl₃): δ 23.8; 47.2; 52.3; 53.4; 56.0; 56.1; 76.8; 110.9; 112.1; 114.6; 126.2; 126.8; 128.5; 130.5; 134.2; 141.5; 147.9; 148.7; 153.6; 159.7; 163.5. Anal. Calcd for $C_{23}H_{22}CINO_7$ (459.88) C, 60.07; H, 4.82; N, 3.05; Found C, 60.10; H, 4.83; N, 3.10.

3.3.5. 10b-Benzo[**1**,**3**]**dioxol-5-yl-8**,**9**-**dimethoxy-6**,**10b-dihydro-5***H*-**isoxazolo**[**3**,**2**-*a*]**isoquinoline-1**,**2**-**dicarboxylic acid dimethyl ester 4e.** $R_{\rm f}$ =0.86; yield 0.068 g, 97%; mp 122–123 °C; IR (KBr) $\nu_{\rm C=0}$ 1749; 1716 cm⁻¹; $\nu_{\rm C=C}$ 1637 cm⁻¹. (400 MHz, CDCl₃): δ 2.61–2.67 (1H, m), 3.12–3.18 (1H, m), 3.25–3.32 (1H, m), 3.66–3.77 (1H, m), 3.67 (3H, s), 3.71 (3H, s), 3.84 (3H, s), 3.87 (3H, s), 5.93 (2H, s), 6.60 (1H, s), 6.75 (1H, d, *J*=8.4 Hz), 6.78 (1H, d, *J*=8.4 Hz), 6.88 (1H, s) 7.02 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 23.7; 46.9; 52.2; 53.3: 56.0; 56.1; 77.1; 101.5; 107.8; 109.7; 110.8; 112.2; 114.9; 122.9; 126.6; 126.7; 136.8; 147.5; 147.7; 147.8; 148.5; 153.4; 159.8; 163.6 Anal. Calcd for C₂₄H₂₃NO₉ (469.44) C, 61.40; H, 4.94; N, 2.98; Found C, 61.50; H, 5.10; N, 3.10.

3.4. Synthesis of azomethine ylides 5a–e. Method A; General procedure

A solution of compound 4 (0.1 mmol) in toluene (5 mL) was refluxed for the specified time (see Table 2). The solvent was evaporated under vacuum and the residue subjected to a silica gel coated TLC plate and eluted with chloroform, petroleum ether, methanol and acetone (45:40:10:5) solvent mixture. The isolated product was crystallized from ethanol ether mixture (1:5).

3.5. Synthesis of azomethine ylides 5a–e. Method B; General procedure

To a solution of nitrone 3 (0.2 mmol) dissolved in toluene (10 mL) DMAD was added and the mixture refluxed for the specified time. The solvent was evaporated and the mixture was subjected on a preparative TLC plate coated with silica gel. The isolated coloured compounds were crystallized from ethanol ether mixture (1:5).

3.5.1. Azomethine ylide 5a. R_f =0.5; yield; Method A, 0.040 g, 93%; Method B, 0.064 g, 75%; light red coloured crystals; mp 228–229 °C; IR (KBr) $\nu_{C=0}$ 1726; 1664 cm⁻¹. UV/vis λ_{max} CHCl₃ nm: 256.5, 313.5, 361.5, 455.1; (400 MHz, CDCl₃): δ 3.25 (2H, t, *J*=7.4 Hz), 3.52 (3H, s), 3.63 (3H, s), 3.69 (3H, s), 4.16 (3H, s), 4.01–4.21 (1H, m), 4.25–4.30 (1H, m), 6.57 (1H, s), 6.86 (1H, s), 7.44–7.53 (5H, m). (100 MHz, CDCl₃): δ 26.9; 50.8; 52.0; 54.3; 56.3; 56.8; 95.5; 110.5; 115.5; 121.0; 128.2; 128.5; 131.8; 131.9; 134.7; 148.3; 156.1; 164.3; 168.6; 171.1; 174.9. Anal. Calcd for C₂₃H₂₃NO₇ (425.43) C, 64.93; H, 5.45; N, 3.29; Found C, 64.98; H, 5.60; N, 3.40.

3.5.2. Azomethine ylide 5b. $R_f = 0.47$; yield; Method A, 0.049 g, 100%; Method B, 0.072 g, 74%; dark orange crystals; mp 128–129 °C; IR (KBr) $\nu_{C=0}$ 1725; 1665 cm⁻¹. UV/vis λ_{max} CHCl₃ nm: 255.5, 391.5; (400 MHz, CDCl₃): δ 3.13 (2H, t, J = 6.4 Hz), 3.37 (3H, s), 3.71 (3H, s), 3.87 (6H, s), 3.88 (3H, s), 3.98 (3H, s), 4.16 (2H, t, J = 6.4 Hz), 6.76 (1H, s), 6.86 (1H, d, J = 9.2 Hz), 6.91 (1H, d, J = 9.2 Hz), 6.98 (1H, s), 7.45 (1H, s). (100 MHz, CDCl₃): δ 26.6; 50.5; 51.8; 53.1; 56.0; 56.1; 56.2; 56.4; 96.1; 107.5; 109.9; 110.8; 115.3; 115.6; 121.9; 132.2; 139.1; 148.1; 148.6; 148.9; 154.8; 167.2; 167.6; 172.1; 180.3. Anal. Calcd for

C₂₅H₂₇NO₉ (485.48) C, 61.85; H, 5.61; N, 2.89; Found C, 61.90; H, 5.75; N, 3.00.

3.5.3. Azomethine ylide 5c. $R_{\rm f}$ =0.59; yield; Method A, 0.039 g, 82%; Method B, 0.090 g, 95%; dark red crystals; mp 213–214 °C; IR (KBr) $\nu_{\rm C=0}$ 1735; 1688 cm⁻¹. UV/vis $\lambda_{\rm max}$ CHCl₃ nm: 228.0, 233.0, 259.0, 315.5, 372.0, 469.0; (400 MHz, CDCl₃): δ 3.16–3.24 (1H, m), 3.31–3.40 (1H, m), 3.59 (3H, s), 3.63 (3H, s), 3.65 (3H, s), 4.03 (3H, s), 4.15–4.22 (1H, m), 4.24–4.32 (1H, m), 6.49 (1H, s), 6.89 (1H, s), 7.65 (1H, t, *J*=8.0 Hz), 7.88 (1H, d, *J*=8.0 Hz), 8.32 (1H, s), 8.37 (1H, d, *J*=8.0 Hz). (100 MHz, CDCl₃): δ 26.8; 51.1; 52.1; 54.3; 56.5; 56.9; 92.1; 110.9; 114.5; 119.9; 123.9; 126.2; 129.4; 133.4; 134.3; 135.1; 147.6; 148.7; 156.9; 164.3; 168.1; 170.3; 171.5. Anal. Calcd for C_{23H222}N₂O₉ (470.43) C, 58.72; H, 4.71; N, 5.95; Found C, 58.58; H, 4.80; N, 6.10.

3.5.4. Azomethine ylide 5d. $R_{\rm f}$ =0.67; yield; Method A, 0.042 g, 91%; Method B, 0.066 g, 71%; dark red crystals; mp 178–179 °C; IR (KBr) $\nu_{\rm C=0}$ 1727; 1665 cm⁻¹. UV/vis $\lambda_{\rm max}$ CHCl₃ nm: 258.0, 262.0, 366.5, 461.5; (400 MHz, CDCl₃): δ 3.22–3.29 (2H, m), 3.54 (3H, s), 3.65 (3H, s), 3.71 (3H, s), 4.01 (3H, s), 4.17–4.25 (2H, m), 6.53 (1H, s), 6.86 (1H, s), 7.42 (4H, br s). ¹³C NMR (100 MHz, CDCl₃): δ 26.8; 51.0; 52.1; 54.4; 56.4; 56.9; 96.3; 110.6; 115.1; 120.6; 128.6; 130.1; 130.3; 134.9; 138.1; 148.4; 156.4; 164.3; 168.4; 170.9; 173.7. Anal. Calcd for C₂₃H₂₂CINO₇ (459.88) C, 60.07; H, 4.82; N, 3.05; Found C, 60.15; H, 5.01; N, 3.30.

3.5.5. Azomethine ylide 5e. $R_{\rm f}$ =0.59; yield; Method A, 0.41 g, 87%; Method B, 0.090 g, 96%; dark red crystals; mp 123–124 °C; IR (KBr) $\nu_{\rm C=O}$ 1734; 1718 cm⁻¹. UV/vis $\lambda_{\rm max}$ CHCl₃ nm: 255.0, 297.5, 386.0; (400 MHz, CDCl₃): δ 3.08 (2H, t, *J*=7.0 Hz), 3.40 (3H, s), 3.70 (3H, s), 3.84 (3H, s), 3.96 (3H, s), 4.08 (2H, t, *J*=7.0 Hz), 5.98 (2H, s), 6.73 (1H, s), 6.78 (1H, d, *J*=8.4 Hz), 6.84 (1H, d, *J*=8.4 Hz), 6.86 (1H, s), 7.40 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 26.9; 50.9; 52.1; 53.3; 56.43; 56.6; 95.0; 102.1; 105.5; 108.4; 110.2; 115.5; 117.7; 122.2; 132.7; 140.7; 147.3; 148.1; 148.3; 155.0; 167.7; 167.8; 172.5; 180.5. Anal. Calcd for C₂₄H₂₃NO₉ (469.44) C, 61.40; H, 4.94; N, 2.98; Found C, 61.50; H, 5.10; N, 3.10.

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Tetrahedron

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Dipolar cycloadditions of imidazoline 3-oxides with N-arylmaleimides. Synthesis and diethylamine induced ring-opening of *exo* and *endo* hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3-diones

Necdet Coşkun,* Habibe Mert and Nevin Arıkan

Department of Chemistry, Uludağ University, 16059 Görükle Bursa, Turkey

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Abstract—1,4-Diarylimidazoline 3-oxides react with *N*-arylmaleimides in benzene to give predominantly the corresponding *endo* adducts. Chiral imidazoline 3-oxides react diastereospecifically (cis configuration of the tetrahydroimidazo ring) and diastereoselectively to give *cis—endo* adducts. The effects of substituents on the aromatic ring of the maleimide was investigated. The presence of electron-withdrawing or releasing groups have minor effect on the total yields but more pronounced is the effect on the ratio of *exo* and *endo* diastereomers. The adducts undergo an interesting and unprecedented ring-opening in the presence of secondary amines to give deoxygenated 3-imidazoline 3-oxides instead of the expected double cis elimination products. Tertiary amines did not induce any reaction. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Nitrones are well-known 1,3-dipoles in thermal cycloaddition reactions with multiple bond systems to provide various heterocyclic five membered ring systems.¹ The cycloadducts of di- and triarylimidazoline 3-oxides² with a variety of dipolarophiles³ give bicyclic compounds with potentially interesting biological activity.⁴ On the other hand, they are source of new heterocyclic compounds via interesting ring-opening reactions.⁵ In our previous work, the 1,3-dipolar cycloadditions of imidazoline 3-oxides was shown to proceed regio- and diastereoselectively and interesting reactions of these adducts under a variety of conditions especially the double cis elimination they undergo in the presence of dialkylamines was reported.^{3d–e,5} exo Adducts of N-methyl and N-phenylmaleimides with chiral 1-benzyl-4-phenyl-2-imidazoline 3-oxide were reported recently.⁶ As a continuation of our interest in the synthesis of imidazoisoxazolidines with potential anticancer activity and in the stereochemistry of dipolar cycloadditions of 1,4-diaryl and 1,2,4-triarylimidazoline 3-oxides with different dipolarophiles, we planned to react a series of *N*-arylmaleimides with imidazoline 3-oxides⁷ $\mathbf{1}$ and to

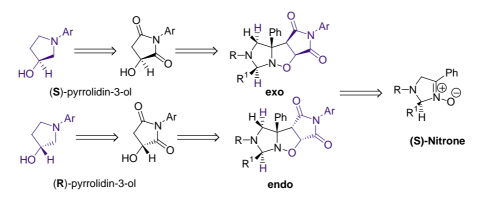
subject them to ring-opening in the presence of secondary and tertiary amines.⁸ The latter reaction would serve as an important entry into the synthesis of chiral 3-hydroxypyrrolidines, which have attracted attention after the discovery of the glycosidase inhibitor activity of the natural product nojirimycin.⁹ The retrosynthetic plan related to the synthesis of chiral pyrrolidin-3-ols is depicted in Scheme 1. (*S*)-Nitrone would give the *exo* and *endo* adducts; the ring-opening of the *exo* adduct would give (*S*)-pyrrolidin-2,5-diones while *endo* would give (*R*), and the reduction of both would give the corresponding chiral pyrrolidin-3-ols. The reverse will be true if we start from (*R*)-nitrone.

For the most widely studied nitrone, *C*-phenyl-*N*-methylnitrone, the frontier orbital energies indicate HOMO control for electron-deficient dipolarophiles.¹⁰ Our observations on the cycloadditions of compounds 1 with electron deficient dipolarophiles corroborate the conclusion that the process is HOMO controlled. In this investigation we were also interested in the effect of substituents on the *N*-aryl group of the maleimide on the reaction yield and the *exo–endo* selectivity of the cycloaddition reaction with cyclic nitrones 1. The problem of *endo–exo* selectivity in 1,3-dipolar cycloadditions is far from definitively assessed and the *endo–exo* selectivity of the cycloaddition of 3,4-dihydroisoquinoline 2-oxide with different types of dipolarophiles was reported.^{11a} The *exo–endo* selectivity of 1,3-dipolar cycloaddition of *C*,*N*-diphenylnitrone to

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^{*} Corresponding author. Tel.: +90 44292561308; fax: +90 4428022; e-mail: coskun@uludag.edu.tr

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Scheme 1. Retrosynthetic analysis for the asymmetric synthesis of pyrrolidin-3-ols.

tert-butyl vinyl ether in the presence of chiral Ti(IV) species was recently reported.¹¹

We report herein the synthesis and ring-opening reactions of a new class of compounds, namely hexahydro-7-oxa-2,5,6atriaza-cyclopenta[a]pentalene-1,3-diones. The reaction of nitrones **1a**-i with *N*-arylmaleimides **2** in benzene and toluene was shown to proceed selectively to give the *endo* adducts as major products. The *exo–endo* ratio increases when electron-donating groups are present on the N-2 aryl, and decreases when the groups are electron-withdrawing. The reaction of adducts **3** and **4** separately or their mixture with diethylamine led to a so far unobserved interesting ring-opening to give di- and triaryl-3-imidazolines **5** instead of the expected double cis elimination products. The mechanism of this reaction is also briefly discussed.

2. Results and discussions

To elucidate the solvent effect on the rate and product ratio of the dipolar cycloaddition, nitrone **1a** was refluxed in different solvents in the presence of 4 equiv of *N*-phenylmaleimide (Scheme 2 and Table 1). The reaction was observed to proceed much faster in solvents such as benzene, acetonitrile and toluene. The reaction proceeds with higher *endo* selectivity in toluene while the cycloaddition in dichloromethane, THF and acetonitrile was unselective. The reaction is too slow in DCM, 39% yield was achieved after 48 h reflux.

At first we decided to develop the model reaction starting with racemic nitrones 1. Compounds 1a–i were reacted with maleimides 2 in benzene to give adducts 3 and 4 in high total yields (Scheme 2 and Table 2).

The cycloaddition nearly completes within 10 h in the cases where C-2 of the nitrone is unsubstituted, while in the cases of C-2 aryl substituted nitrones 1c-e the reaction time was five times longer to achieve the same yields due to the steric hindrance of the aryl groups. The exo-endo ratio is approximately the same in the cycloaddition of nitrones 1a-d with N-phenylmaleimide. The ratio is close to 1:1 in the case of 1e (C-2 substituent is 3-nitrophenyl group) the steric hindrance of which probably does not support the formation of the transition state leading to the endo adduct. To understand the role of the substituents on the N-aryl group of 2 it is useful to compare the exo-endo ratio of cycloadditions with 1a,f-i (Table 2). It is seen that the electron-donating groups favor the formation of endo adducts, while electron-withdrawing groups do not. Beside the steric effects contributing to the exo-endo ratio, secondary orbital interactions between the aryl rings at N-2 and N-5 and may be between N-5 and the carbonyls at the pyrrolidine ring are probably also responsible for the stabilization of the transition state leading to *endo* adduct. The effect of substituents on the total yields of adducts 3 and **4** are of the same magnitude independent of their nature. This means electron-donating groups somewhat increase the LUMO energy of the electron deficient maleimide and thus decelerate the exo adduct formation. Computations of the HOMO and LUMO energies for maleimides 2 confirmed this. On the other hand, computation of the HOMO and LUMO energies for nitrone 1a and comparison with the corresponding HOMO and LUMO energies of maleimides 2 clearly revealed that the cycloaddition should be a HOMO controlled process. The same electron-donating substituent probably raises the energy of N-2 phenyls HOMO to give a better π interaction between the N-5 aryl. Conversely, electron-withdrawing groups decrease the LUMO energy of the electron deficient maleimide thus accelerating the exo

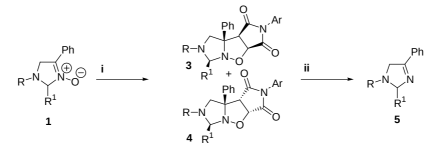


Table 1. Solvent effect on the 1,3-dipolar cycloaddition of 1a with
N-phenylmaleimide

Solvent	Reaction	Total yield	Yield (%)
	time (h)	(%)	3 a	4a
Benzene	10	100	35	65
Toluene	10	80	24	57
THF	10	56	23	33
DCM	48^{a}	39	19	20
Acetonitrile	10	86	45	41

^a The yield of the reaction for 10 h reaction time is 12% and the ratio of *exo* and *endo* isomers is 1:2.

adduct formation but lower the π interaction between the N-5 aryl.

Some characteristic assignments for adducts **3** and **4** based on extensive 1D and 2D NMR experiments are given in Table 3.

The *exo* stereochemistry of adducts **3a–b,f–i** was confirmed by NOESY1D experiments performed on compound **3a** (Fig. 1) as follows:

Irradiation of proton at C-7a enhanced the signal of 3aH (1%). Irradiation of the doublet of 6Ha enhanced the signals of 6Hb (12.72%) and *ortho* protons of *N*-tolyl group (6.66%). The irradiation of 6Hb enhances the signal of 6Ha (13.62%) and the *ortho* protons' signals of *N*-tolyl (3.92%) and 3b-phenyl (2.93%). Irradiation of 4Hb, enhanced the signals of 4Ha, and the *ortho* protons of both phenyls at N-5 and C-3b by 19.0, 6.81, and 9.13%, respectively. The irradiation of 3aH enhances the signals of 7aH (3.3%), 4Ha (3.09%) and *ortho* protons of 3b-phenyl by 0.5%. Irradiation of 4Ha enhances the signals of 4Hb (18.7%), 3aH (6.17%), 7a (1%), and *ortho* protons of *N*-tolyl group. 7aH was irradiated to give enhancement for 3aH (1.75%) and the *ortho* protons of *N*-phenyl group at 7.04 ppm (4.0%).

Table 2. Synthesis of hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[a]pentalene-1,3-diones 3a-i and 4a-i

Entry	R	\mathbb{R}^1	Ar	Total yield	Yield (%)	exo-endo ^a	
					3	4	
a	4-MeC ₆ H ₄	Н	Ph	100 ^b	35	65	1:1.86
b	4-MeOC ₆ H ₄	Н	Ph	100 ^b	38	62	1:1.63
с	$4 - MeC_6H_4$	4-MeOC ₆ H ₄	Ph	90°	33	57	1:1.73
d	4-MeOC ₆ H ₄	$4 - MeOC_6H_4$	Ph	92°	32	60	1:1.87
e	4-MeOC ₆ H ₄	3-NO ₂ C ₆ H ₄	Ph	$74^{\rm c}$	33	41	1:1.24
f	$4 - MeC_6H_4$	Н	4-MeOC ₆ H ₄	93 ^b	24	69	1:2.88
g	$4-MeC_6H_4$	Н	$4-NO_2C_6H_4$	85 ^b	35	50	1:1.43
ĥ	4-MeC ₆ H ₄	Н	$4-ClC_6H_4$	88^{b}	36	52	1:1.44
i	$4-MeC_6H_4$	Н	4-MeC ₆ H ₄	88^{b}	28	60	1:2.14

^a The ratio of the isolated adducts.

^b Reaction time 10 h.

^c Reaction time 51 h.

Table 3. Characteristic ¹H NMR spectroscopic data for *exo* and *endo* adducts 3 and 4

	exo							endo					
	ЗаН	4Ha	4Hb	6Ha	6Hb	7aH	_	3a	4Ha	4Hb	6Ha	6Hb	7aH
3a	3.98	3.80	4.17	4.79	4.53	5.17	4a	4.02	3.11	4.60	4.49	4.66	5.15
3b	3.98	3.79	4.12	4.76	4.53	5.17	4b	4.01	3.05	4.56	4.44	4.63	5.15
3c	3.92	3.96	4.61		5.81	5.17	4c	4.04	4.77	3.87		5.64	5.16
3d	3.92	3.96	4.61		5.76	5.19	4d	4.04	4.71	3.85		5.63	5.17
3e	3.92	3.96	4.70		5.85	5.21	4e	4.06	4.69	3.95		5.76	5.22
3f	3.96	3.79	4.16	4.78	4.52	5.15	4f	4.01	3.10	4.58	4.49	4.65	5.15
3g	4.02	3.81	4.19	4.79	4.54	5.21	4g	4.05	3.08	4.62	4.47	4.59	5.19
3h	3.97	3.79	4.16	4.78	4.52	5.16	4h	4.01	3.08	4.59	4.48	4.63	5.15
3i	3.95	3.78	4.15	4.78	4.52	5.14	4i	4.01	3.10	4.59	4.49	4.65	5.14

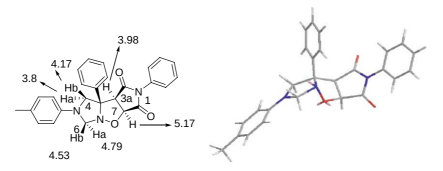


Figure 1. Some selected chemical shifts assignments for 3a and its energy minimised 3D model (total energy 99.4078 kcal/mol).

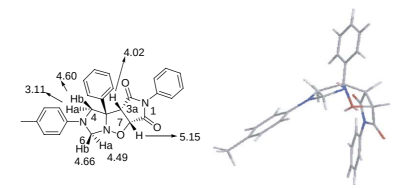


Figure 2. Some selected chemical shifts assignments for 4a and its energy minimised 3D model (total energy 99.0050 kcal/mol).

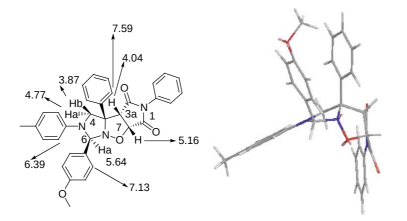


Figure 3. Some selected chemical shifts for *cis-endo* adduct 4c and its energy minimised 3D model (total energy 27.2092 kcal/mol).¹²

Finally, the *ortho* protons of 3b-phenyl were irradiated to give enhancements for the signals of 4Hb (2.11%) and 6Hb (0.5%). The energy minimised conformations of compounds **3a**, **4a** and **4c** (see Figs. 1–3) are supporting the observed correlations by NOESY1D experiments. On the other hand, the total energy of **3a** was by 0.4028 kcal/mol higher than that of **4a**.

The NOESY1D experiment results for *endo* adduct **4a** are as follows: irradiation of 6Ha enhanced the signals of 6Hb (17.2%), the *ortho* protons of *N*-tolyl group (3.07%) and 4Ha (0.66%). The irradiation of 4Ha enhanced the signals of 4Hb (19.18%), *ortho* protons of *N*-tolyl and 3b-phenyl by 1.69 and 0.9%, respectively. Irradiation of 3aH enhanced the signals of 7aH and 3b-phenyls *ortho* protons by 3.3 and 2.66%, respectively. Irradiation of 4Hb enhances the signal of 4Ha by 17.71% and the *ortho* protons of *N*-tolyl and 3b-phenyl by 7.30 and 2.59%. Irradiation of *ortho* protons of 3b-phenyl enhances the signals of 4Hb and 3aH by 1.5 and 1%, respectively.

To prove the cis orientation of the phenyls at C-3b and C-6 we have irradiated the corresponding protons at the imidazolidine and isoxazolidine rings of **4c** as follows: the proton at C-3a was irradiated to give enhancements for the C-3b-phenyls' *ortho* protons (3.61%) and for the 7aH (3.98%). 4Hb was irradiated to give enhancements for 4Ha (27.0%) and the *ortho* protons of the phenyls at C-3b, N-5 and C-6 by 2.84, 2.58 and 4.13%, respectively. Irradiation of

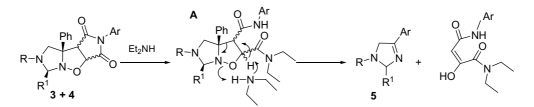
4Ha enhanced the signals of 4Hb (27.8%) and the signals of *N*- and C-3b-phenyls' *ortho* protons by 10.0 and 2.71%, respectively. The irradiation of C-6H enhanced the signals of C-6 phenyls and N-5 phenyls *ortho* protons. This unequivocally proves the *cis-endo* configuration of compounds 4c-e.

According to the developed procedure isolated, compounds **3** and **4** or their mixture were refluxed in diethylamine in order to prepare the racemic mixtures of pyrrolidin-3-ols as in Scheme 1, however, this treatment led to the formation of new 3-imidazolines **5a–e** (Scheme 2 and Table 4). The compounds were easily characterized by elemental analyses and spectral methods. The characteristic IR frequencies for C=N appears at ca.1630 cm⁻¹. The methylenes at C-2 and C-5 in the cases of **5a–b** appear as two proton triplets as a results of long range coupling between them. The long range

Table 4.	Synthesis	of	3-imidazolines	5а-е
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Starting material	Product	Yield (%)	Mp (°C)
3a+4a	5a	90	117–119
3b+4b	5b	100 ^a	129-130
4c	5c	92	176-178
3c	5c	98	176-178
3d+4d	5d	88	152-153
3e	5e	92	182-184
4e	5e	92	182-184

^a The reaction time was 23 h for all entries except for entry 2 where the reaction time is 39 h.



Scheme 3. Proposed mechanism for the conversion of adducts 3 and 4 into 5.

coupling is observed between the protons at C-2 and the AB system at C-5 in the cases of **5c-d**.

The probable mechanism for the ring-opening of compounds **3** and **4** in diethylamine is depicted in Scheme 3. The nucleophilic attack of diethylamine leads to intermediate **A** (isolated in the case of **4b**), which probably undergo diethylamine assisted synchronous ring-opening to give imidazolines **5** and the corresponding oxaloacetic acid amides. The isolated and characterized **A** (5-(4-methoxyphenyl)-3a-phenyl-hexahydroimidazo[1,5-*b*]isoxazole-2,3-dicarboxylic acid 2-diethylamide 3-phenylamide **4b**', was isolated from the reaction of **3b** and **4b** in diethylamine for 23 h) was refluxed in diethylamine for 23 h to give imidazoline **5b** in 92% yield.

To prove the structure of 5-(4-methoxyphenyl)-3a-phenylhexahydroimidazo[1,5-*b*]isoxazole-2,3-dicarboxylic acid 2-diethylamide 3-phenylamide **4b**' we have performed NOESY1D experiments as follows: irradiation of C-4Hb proton led to enhancement of the signals of C-4Ha (18.05%), *p*-anisyl (3.58%) and C-3b phenyls (4.27%) *ortho* protons. While the irradiation of C-4Ha enhances the signals of C-4Hb (16.34%), *N-p*-anisyl (7.20%) and C-3b phenyls *ortho* protons (1.3%). Irradiation of C-3H enhanced the signals of C-2H (4.09%), C-3b phenyls *ortho* protons (2%) and the amide proton at 8.89 ppm. The latter correlation was indicative for the determination of the right regioisomer. Thus, all these experiments allowed us to assign the configuration shown in Figure 4.

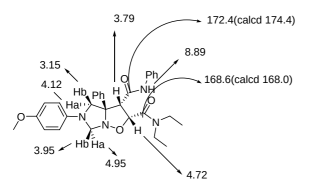


Figure 4. Some characteristic chemical shifts for intermediate bisamide 4b'.

exo Adduct **3a** was shown to give nucleophilic addition product faster than *endo* adduct **4a**. The reaction times for the disappearance of the corresponding adducts (TLC controls) were 1.5 and 6 h, respectively.

Compounds **3a**,**c** and **4a**,**c** were refluxed in triethylamine for 48 h but no conversion was observed, the starting materials were recovered unchanged.

3. Conclusions

In conclusion, we studied the reaction of imidazoline 3-oxides 1 with of N-arylmaleimides 2. The reactions of nitrones **1a–b,f–i** with *N*-arylmaleimides **2** in benzene give predominantly the corresponding *endo* adducts 4a-b,f-i. Chiral imidazoline 3-oxides **1c–e** react diastereospecifically with respect to the cis configuration in the tetrahydroimidazo ring and diastereoselectively to give cis-endo adducts 4c-e. The effect of substituents on the phenyl ring of the maleimide was investigated. The presence of electronwithdrawing or releasing groups have minor effects on the total yields but the effect on the ratio of exo and endo diastereomers is more pronounced. The exo-endo ratio increases when electron-donating groups are present on the N-2 aryl, and decreases when the groups are electronwithdrawing. Adducts 3 and 4 undergo an interesting ringopening in the presence of secondary amines to give the deoxygenated 3-imidazoline 3-oxides 5 instead of the expected double cis elimination products. This reaction will serve as a convenient method for the synthesis of otherwise inaccessible 3-imidazolines. Tertiary amines did not induce any reaction.

4. Experimental

4.1. General

Melting points were recorded on an Electrothermal Digital melting point apparatus. Infrared spectra were recorded on a Mattson 1000 FTIR. 1D and 2D NMR experiments were performed on a Varian Mercury Plus 400 MHz spectrometer. Visualisation was effected with UV light. Imidazoline 3-oxides **1a**–**e** were prepared according to the method we have recently reported.⁷ The elemental analyses were performed on a EuroEA 3000 CHNS analyser. The total energies of compounds **3a**, **4a**, **4c**, *cis–exo* **3c** and the FMO energy calculations for maleimides **2** and nitrone **1a** were performed using CS MOPAC Pro in ChemOffice 6.

4.1.1. 1,2-Bis-(4-methoxyphenyl)-4-phenyl-2,5-dihydro-*1H*-imidazole 3-oxide 1d. Yield, 2.0 g, 23%; white needles; mp 200–201.5 °C; IR (KBr) $\nu_{C=N}$ 1610 cm⁻¹; ¹H NMR δ 3.73 (3H, s), 3.80 (3H, s), 4.81 (1H, dd, *J*=14.0, 3.2 Hz), 5.14 (1H, dd, *J*=14.0, 5.6 Hz), 6.10 (1H, dd, *J*=5.6, 3.2 Hz), 6.57 (2H, d, *J*=8.8 Hz), 6.82 (2H, d, *J*=8.8 Hz), 6.94 (2H, d, J=8.4 Hz), 7.45–7.49 (3H, m), 7.56 (2H, d, J=8.4 Hz), 8.34 (2H, dd, J=7.6, 3.6 Hz). ¹³C NMR δ 53.4; 55.6; 55.9; 90.0; 113.9; 114.6; 115.3; 127.2; 128.9; 129.0; 129.6; 131.1; 134.5; 136.9; 138.8; 153.0; 161.2. Anal. Calcd for C₂₃H₂₂N₂O₃ (374.43) C, 73.78; H, 5.92; N, 7.48; found C, 73.75; H, 5.90; N, 7.45.

4.1.2. 1-(4-Methoxyphenyl)-2-(3-nitrophenyl)-4-phenyl-2,5-dihydro-1*H***-imidazole 3-oxide 1e.** Yield 2.38 g, 26%; yellow needles; mp 190–191 °C; $\nu_{C=N}$ 1610 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.80 (3H, s), 4.86 (1H, dd, J=14.0, 3.2 Hz), 5.26 (1H, dd, J=14.0, 5.6 Hz), 6.25 (1H, dd, J=5.6, 3.2 Hz), 6.55 (2H, d, J=9.2 Hz), 6.84 (2H, d, J=9.2 Hz), 7.48–7.52 (3H, m), 7.65 (1H, t, J=7.6 Hz), 8.06 (1H, d, J=8.0 Hz), 8.29–8.33 (3H, m), 8.51 (1H, t, J=2.0 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 53.9; 55.9; 89.1; 114.2; 115.5; 123.2; 123.4; 126.9; 127.2; 129.1; 130.1; 131.7; 134.9; 135.6; 138.3; 138.5; 148.0; 153.7. Anal. Calcd for C₂₂H₁₉N₃O₄ (389.40) C, 67.86; H, 4.92; N, 10.79; found C, 67.83; H, 4.90; N, 10.75.

The maleimides used were prepared according to a method known in the literature.^{13a} Maleimide **2g** was prepared according to a modified literature procedure:^{13b} to a mixture of 4-nitroaniline (5.1 mmol, 0.772 g) and maleic anhydride (6.04 mmol, 0.592 g) PPA (7 g) was added and the mixture stirred for 15 h at 80 °C on a water bath. The mixture was poured into cold water and the product precipitated was filtered and dried in a vacuum oven. Yield 0.598 g, 54%; yellow amorphous solid; mp 163–164 °C; IR (KBr) $\nu_{C=0}$ 1724 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 6.93 (2H, s), 7.68 (2H, d, J=9.6 Hz), 8.34 (2H, d, J=9.6 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 124.7; 125.7; 134.9; 137.3; 146.4; 168.8.

4.2. Synthesis of hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3-diones 3a–i and 4a–i. General procedure

To a solution of imidazoline 3-oxide 1 (0.12 mmol) in benzene (10 mL) maleimide (0.48 mmol) was added and the reaction mixture stirred for the specified time. The solvent was evaporated and the mixture was separated by column chromatography using silica gel as an adsorbent and petroleum ether ethyl acetate as a solvent mixture. The compounds were recrystallized from ether or ethanol.

4.2.1. *exo*-2,3**b**-Diphenyl-5-*p*-tolyl-hexahydro-7-oxa-**2,5,6a**-triaza-cyclopenta[*a*]pentalene-1,3-dione 3a. Yield 0.018 g, 35%; white needles; mp 177–178 °C; IR (KBr) $\nu_{C=0}$ 1713 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.24 (3H, s), 3.80 (1H, d, *J*=8.8 Hz), 3.98 (1H, d, *J*=7.6 Hz), 4.17 (1H, d, *J*=8.8 Hz), 4.53 (1H, d, *J*=11.2 Hz), 4.79 (1H, d, *J*=11.2 Hz), 5.17 (1H, d, *J*=7.6 Hz), 6.42 (2H, d, *J*=8.0 Hz), 7.00–7.05 (4H, m), 7.31–7.40 (6H, m), 7.57 (2H, d, *J*=7.2 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 56.7; 56.8; 71.0; 80.7; 112.3; 126.2; 126.6; 126.9; 129.0; 129.2; 129.3; 129.4; 130.1; 131.3; 136.0; 143.6; 171.2; 174.1. Anal. Calcd for C₂₆H₂₃N₃O₃ (425.48) C, 73.39; H, 5.45; N, 9.88; found C, 73.34; H, 5.40; N, 9.95. **4.2.2.** *exo*-5-(4-Methoxyphenyl)-2,3b-diphenyl-hexa-hydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3-dione 3b. Yield 0.020 g, 38%; white needles; mp 120–121 °C; IR (KBr) $\nu_{C=0}$ 1716 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.74 (3H, s), 3.79 (1H, d, J=8.4 Hz), 3.98 (1H, d, J=7.2 Hz), 4.12 (1H, d, J=8.8 Hz), 4.53 (1H, d, J=7.2 Hz), 4.76 (1H, d, J=11.2 Hz), 5.17 (1H, d, J=7.2 Hz), 6.46 (2H, d, J=8.8 Hz), 6.83 (2H, d, J=9.2 Hz), 6.99 (2H, d, J=7.6 Hz), 7.30–7.40 (6H, m), 7.56 (2H, d, J=7.2 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 56.1; 56.9; 57.2; 71.5; 77.3; 80.8; 113.3; 115.4; 126.2; 126.5; 129.0; 129.1; 129.2; 129.4; 131.3; 136.1; 140.5; 152.3; 171.2; 174.1. Anal. Calcd for C₂₆H₂₃N₃O₄ (441.48) C, 70.73; H, 5.25; N, 9.52; found C, 70.80; H, 5.40; N, 9.42.

4.2.3. *exo*-6-(4-Methoxyphenyl)-2,3b-diphenyl-5-*p*-tolylhexahydro-7-oxa-2,5,6a-triaza cyclopenta[*a*]pentalene-**1,3-dione 3c.** Yield 0.021 g, 33%; white needles; mp 167–168 °C; IR (KBr) $\nu_{C=0}$ 1712 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.23 (3H, s), 3.70 (3H, s), 3.92 (1H, d, *J*=7.6 Hz), 3.96 (1H, d, *J*=9.2 Hz), 4.61 (1H, d, *J*= 9.2 Hz), 5.17 (1H, d, *J*=7.6 Hz), 5.81 (1H, s), 6.43 (2H, d, *J*=8.8 Hz), 6.58 (2H, d, *J*=8.4 Hz), 6.89–7.18 (10H, m), 7.18–7.51 (4H, m). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 55.5; 57.5; 57.7; 77.0; 79.7; 85.5; 113.4; 113.5; 126.1; 127.0; 127.4; 128.2; 128.6; 128.9; 129.1; 129.3; 129.9; 130.2; 131.3; 135.3; 144.0; 159.5; 171.1; 174.1. Anal. Calcd for C₃₃H₂₉N₃O₄ (531.60) C, 74.56; H, 5.50; N, 7.90; found C, 74.60; H, 5.60; N, 7.78.

4.2.4. *exo*-5,6-Bis-(4-methoxyphenyl)-2,3b-diphenyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-**1,3-dione 3d.** Yield 0.021 g, 32%; white needles; mp 165–166 °C; IR (KBr) $\nu_{C=0}$ 1712 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.70 (3H, s), 3.72 (3H, s), 3.92–3.96 (2H, two overlapping d, J=7.6, 8.8 Hz), 4.61 (1H, d, J=8.8 Hz), 5.19 (1H, d, J=7.6 Hz), 5.76 (1H, s), 6.46 (2H, d, J=8.4 Hz), 6.58 (2H, d, J=8.0 Hz), 6.78 (2H, d, J=8.4 Hz), 6.95–6.99 (3H, m), 7.08–7.14 (3H, m), 7.21–7.25 (2H, m), 7.30–7.35 (4H, m). ¹³C NMR (100 MHz, CDCl₃): δ 55.5; 55.9; 57.6; 57.8; 77.0; 79.7; 85.4; 113.5; 114.4; 115.1; 126.1; 127.0; 127.4; 128.3; 128.7; 128.9; 129.1; 129.3; 129.6; 135.3; 144.7; 152.6; 159.5; 171.1; 174.1. Anal. Calcd for C₃₃H₂₉N₃O₅ (547.60) C, 72.38; H, 5.34; N, 7.67; found C, 72.32; H, 5.40; N, 7.60.

4.2.5. *exo-5-*(**4-Methoxyphenyl**)-**6-**(**3-nitrophenyl**)-**2,3bdiphenyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[***a***]-pentalene-1,3-dione 3e.** Yield 0.022 g, 33%; yellow needles; mp 173–174 °C; IR (KBr) $\nu_{C=0}$ 1715 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.73 (3H, s), 3.92 (1H, d, J=7.1 Hz), 3.96 (1H, d, J=8.8 Hz), 4.70 (1H, d, J=8.8 Hz), 5.21 (1H, d, J=7.1 Hz), 5.85 (1H, s), 6.44 (2H, d, J=8.8 Hz), 6.80 (2H, d, J=8.8 Hz), 7.01–7.09 (5H, m), 7.18–7.38 (6H, m), 7.52 (1H, d, J=7.6 Hz), 7.79 (1H, s), 7.96 (1H, d, J=7.6 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 55.9; 57.2; 57.4; 77.1; 80.0; 85.3; 114.5; 115.3; 122.9; 123.2; 125.9; 126.9; 128.5; 129.0; 129.2; 129.3; 129.4; 131.2; 134.1; 134.3; 140.0; 140.2; 148.2; 153.1; 170.8; 173.8. Anal. Calcd for C₃₂H₂₆N₄O₆ (562.57) C, 68.32; H, 4.66; N, 9.96; found C, 68.30; H, 4.60; N, 9.98. **4.2.6.** *exo*-2-(4-Methoxyphenyl)-3b-phenyl-5-*p*-tolylhexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-**1,3-dione** 3f. Yield 0.013 g, 24%; white needles; mp 187–188 °C; IR (KBr) $\nu_{C=0}$ 1716 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.23 (3H, s), 3.78 (3H, s), 3.79 (1H, d, *J*=8.6 Hz), 3.96 (1H, d, *J*=7.4 Hz), 4.16 (1H, d, *J*=8.6 Hz), 4.52 (1H, d, *J*=10.9 Hz), 4.78 (1H, d, *J*= 10.9 Hz), 5.15 (1H, d, *J*=7.4 Hz), 6.41 (2H, d, *J*=8.2 Hz), 6.87–6.94 (4H, m), 7.03 (2H, d, *J*=8.2 Hz), 7.31 (1H, t, *J*=7.2 Hz), 7.37 (2H, t, *J*=7.2 Hz), 7.55 (2H, t, *J*=7.2 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 55.7; 56.7; 56.8; 71.0; 77.2; 80.7; 112.3; 114.7; 123.8; 126.5; 126.8; 127.4; 129.2; 129.3; 130.1; 136.0; 143.6; 159.8; 171.5; 174.4. Anal. Calcd for C₂₇H₂₅N₃O₄ (455.51) C, 71.19; H, 5.53; N, 9.22; found C, 71.21; H, 5.50; N, 9.27.

4.2.7. *exo*-2-(4-Nitrophenyl)-3b-phenyl-5-*p*-tolyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3dione 3g. Yield 0.020 g, 35%; yellow needles; mp 175 °C; IR (KBr) $\nu_{C=0}$ 1728 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.24 (3H, s), 3.81 (1H, d, *J*=8.8 Hz), 4.02 (1H, d, *J*=7.2 Hz), 4.19 (1H, d, *J*=8.8 Hz), 4.02 (1H, d, *J*=10.8 Hz), 4.79 (1H, d, *J*=10.8 Hz), 5.21 (1H, d, *J*= 7.2 Hz), 6.40 (2H, d, *J*=8.4 Hz), 7.04 (2H, d, *J*=8.4 Hz), 7.24–7.26 (3H, m), 7.35–7.41 (2H, m), 7.54 (2H, d, *J*= 8.0 Hz), 8.23 (2H, d, *J*=9.2 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 56.8; 56.9; 71.0; 77.2; 80.8; 112.4; 124.6; 126.4; 126.6; 127.1; 129.3; 129.5; 130.2; 135.8; 136.6; 143.5; 147.3; 170.5; 173.4. Anal. Calcd for C₂₆H₂₂N₄O₅ (470.48) C, 66.37; H, 4.71; N, 11.91; found C, 66.40; H, 4.76; N, 11.90.

4.2.8. *exo*-2-(4-Chlorophenyl)-3b-phenyl-5-*p*-tolyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3dione 3h. Yield 0.020 g, 36%; white needles; mp 186–187 °C; IR (KBr) $\nu_{C=0}$ 1720 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.24 (3H, s), 3.79 (1H, d, *J*= 8.4 Hz), 3.97 (1H, d, *J*=7.6 Hz), 4.16 (1H, d, *J*=8.4 Hz), 4.52 (1H, d, *J*=11.2 Hz), 4.78 (1H, d, *J*=11.2 Hz), 5.16 (1H, d, *J*=7.6 Hz), 6.41 (2H, d, *J*=8.6 Hz), 6.95–6.98 (2H, m), 7.04 (2H, d, *J*=8.6 Hz), 7.31–7.40 (5H, m), 7.54 (2H, d, *J*=8.6 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 56.7; 56.8; 71.0; 77.2; 80.7; 112.3; 126.5; 127.0; 127.4; 129.2; 129.3; 129.6; 129.7; 130.1; 134.8; 135.9; 143.6; 171.0; 173.8. Anal. Calcd for C₂₆H₂₂ClN₃O₃ (459.92) C, 67.90; H, 4.82; N, 9.14; found C, 68.05; H, 4.96; N, 9.27.

4.2.9. *exo*-**3b**-**Phenyl-2,5-di**-*p*-**tolyl-hexahydro-7-oxa2,5,6a**-**triaza**-**cyclopenta**[*a*]**pentalene-1,3-dione 3i.** Yield 0.015 g, 28%; white needles; mp 194–195 °C; IR (KBr) $\nu_{C=0}$ 1716 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.23 (3H, s), 2.30 (3H, s), 3.78 (1H, d, *J*=8.19 Hz), 3.95 (1H, d, *J*=7.4 Hz), 4.15 (1H, d, *J*=8.19 Hz), 4.52 (1H, d, *J*=10.9 Hz), 4.78 (1H, d, *J*=10.9 Hz), 5.14 (1H, d, *J*=7.4 Hz), 6.41 (2H, d, *J*=8.0 Hz), 6.88 (2H, d, *J*=8.0 Hz), 7.03 (2H, d, *J*=8.0 Hz), 7.17 (2H, d, *J*=8.0 Hz), 7.30–7.37 (3H, m), 7.55 (2H, d, *J*=7.6 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 21.4; 55.7; 56.8; 71.0; 77.0; 80.7; 112.3; 126.0; 126.6; 126.8; 128.6; 129.1; 129.2; 130.0; 130.1; 136.0; 139.1; 143.6; 173.3; 174.2. Anal. Calcd for C₂₇H₂₅N₃O₃ (439.51) C, 73.78; H, 5.73; N, 9.56; found C, 73.75; H, 5.70; N, 9.50.

4.2.10. *endo*-**2,3b-Diphenyl-5**-*p*-tolyl-hexahydro-7-oxa-**2,5,6a-triaza-cyclopenta**[*a*]**pentalene-1,3-dione 4a.** Yield 0.033 g, 65%; white needles; mp 185–186 °C; IR (KBr) $\nu_{C=0}$ 1709 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.28 (3H, s), 3.11 (1H, d, J=10.1 Hz), 4.02 (1H, d, J=8.6 Hz), 4.49 (1H, d, J=9.4 Hz), 4.60 (1H, d, J=10.1 Hz), 4.66 (1H, d, J=8.6 Hz), 6.88 (2H, d, J=7.0 Hz), 7.08 (2H, d, J=8.6 Hz), 7.19–7.25 (3H, m), 7.35 (1H, t, J=7.4 Hz), 7.44 (2H, t, J=7.4 Hz), 7.66 (2H, d, J=7.4 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.7; 54.2; 59.8; 75.3; 80.6; 80.9; 115.0; 125.8; 126.5; 128.4; 128.9; 129.2; 129.3; 129.8; 130.2; 131.4; 141.5; 142.9; 173.0; 174.1. Anal. Calcd for C₂₆H₂₃N₃O₃ (425.48) C, 73.39; H, 5.45; N, 9.88; found C, 73.40; H, 5.33; N, 10.05.

4.2.11. *endo*-5-(4-Methoxyphenyl)-2,3b-diphenyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3dione **4b.** Yield 0.033 g, 62%; white needles; mp 182–183 °C; IR (KBr) $\nu_{C=0}$ 1712 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.05 (1H, d, J=10.4 Hz), 3.75 (3H, s), 4.01 (1H, d, J=8 Hz), 4.44 (1H, d, J=9.6 Hz), 4.56 (1H, d, J=10.0 Hz), 4.63 (1H, d, J=9.6 Hz), 5.15 (1H, d, J= 8.0 Hz), 6.57 (2H, d, J=8.5 Hz), 6.81 (2H, d, J=8.5 Hz), 6.90 (2H, d, J=7.4 Hz), 7.2–7.25 (3H, m), 7.34 (1H, t, J= 7.41 Hz), 7.43 (2H, t, J=7.4 Hz), 7.66 (2H, d, J=7.4 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 54.8; 55.9; 59.8; 75.9; 80.8; 81.0; 115.2; 116.3; 125.8; 126.6; 128.4; 128.9; 129.2; 129.3; 131.4; 139.3; 141.58; 154.1; 173.0; 174.3. Anal. Calcd for C₂₆H₂₃N₃O₄ (441.48) C, 70.73; H, 5.25; N, 9.52; found C, 70.85; H, 5.33; N, 9.48.

4.2.12. endo-6-(4-Methoxyphenyl)-2,3b-diphenyl-5-ptolyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[a]pentalene-1,3-dione 4c. Yield 0.036 g, 57%; white needles; mp 189–191 °C; IR (KBr) $\nu_{C=0}$ 1720 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.22 (3H, s), 3.74 (3H, s), 3.87 (1H, d, J = 10.0 Hz), 4.04 (1H, d, J = 8.4 Hz), 4.77 (1H, d, J =10.0 Hz), 5.16 (1H, d, J=8.4 Hz), 5.64 (1H, s), 6.38 (2H, d, J=8.6 Hz), 6.76 (2H, d, J=8.6 Hz), 6.80 (2H, d, J=7.4 Hz), 6.93 (2H, d, J=8.2 Hz), 7.12 (2H, d, J=8.6 Hz), 7.19-7.32 (4H, m), 7.37 (2H, t, J=7.4 Hz), 7.60 (2H, d, J=7.4 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 52.7; 55.4; 59.4; 79.9; 80.3; 84.6; 114.2; 114.3; 126.0; 126.3; 127.9; 128.3; 128.7; 128.8; 129.0; 129.2; 129.9; 131.2; 131.5; 141.3; 141.6; 159.6; 172.9; 174.0. Anal. Calcd for C₃₃H₂₉N₃O₄ (531.60) C, 74.56; H, 5.50; N, 7.90; found C, 74.45; H, 5.63; N, 7.85.

4.2.13. *endo*-5,6-Bis-(4-methoxyphenyl)-2,3b-diphenylhexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-**1,3-dione 4d.** Yield 0.039 g, 60%; white needles; mp 159–160 °C; IR (KBr) $\nu_{C=0}$ 1716 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.70 (3H, s), 3.73 (3H, s), 3.85 (1H, d, *J*=10.0 Hz), 4.04 (1H, d, *J*=8.4 Hz), 4.71 (1H, d, *J*=10.0 Hz), 5.17 (1H, d, *J*=8.4 Hz), 5.63 (1H, s), 6.41 (2H, d, *J*=9.0 Hz), 6.69 (2H, d, *J*=9.0 Hz), 6.74 (2H, d, *J*=8.6 Hz), 6.85–6.87 (2H, m), 7.09 (2H, d, *J*=8.6 Hz), 7.21–7.33 (4H, m), 7.38 (2H, t, *J*=7.8 Hz), 7.60 (2H, d, *J*=7.8 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 53.0; 55.4; 55.9; 59.4; 79.9; 80.4; 85.0; 114.1; 115.0; 115.6; 126.0; 126.2; 128.3; 128.8; 128.8; 129.1; 129.2; 131.2; 131.4; 137.7; 141.7; 152.9; 159.5; 172.9; 174.1. Anal. Calcd for $C_{33}H_{29}N_3O_5\ (547.60)\ C,\ 72.38;\ H,\ 5.34;\ N,\ 7.67;\ found\ C,\ 72.35;\ H,\ 5.53;\ N,\ 7.51.$

4.2.14. *endo*-5-(4-Methoxyphenyl)-6-(3-nitrophenyl)-**2,3b-diphenyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta-[a]pentalene-1,3-dione 4e.** Yield 0.028 g, 41%; yellow needles; mp 119–120 °C; IR (KBr) $\nu_{C=0}$ 1714; 1724 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.70 (3H, s), 3.95 (1H, d, J= 10.0 Hz), 4.06 (1H, d, J=8.0 Hz), 4.69 (1H, d, J=10.0 Hz), 5.22 (1H, d, J=8.0 Hz), 5.76 (1H, s), 6.41 (2H, d, J=8.8 Hz), 6.71 (2H, d, J=9.2 Hz), 6.90 (2H, d, J=8.8 Hz), 7.25–7.57 (10H, m), 8.05–8.08 (2H, m). ¹³C NMR (100 MHz, CDCl₃): δ 53.1; 55.9; 59.5; 80.0; 80.3; 84.5; 115.2; 115.7; 122.7; 123.5; 125.8; 126.1; 126.3; 129.0; 129.2; 129.3; 129.4; 131.1; 133.7; 134.4; 140.8; 141.5; 148.6; 153.4; 172.7; 173.6. Anal. Calcd for C₃₂H₂₆N₄O₆ (562.57) C, 68.32; H, 4.66; N, 9.96; found C, 68.43; H, 4.59; N, 9.90.

4.2.15. endo-2-(4-Methoxyphenyl)-3b-phenyl-5-p-tolylhexahydro-7-oxa-2,5,6a-triaza-cyclopenta[a]pentalene-1,3-dione 4f. Yield 0.038 g, 69%; white needles; mp 187–187.4 °C; IR (KBr) $\nu_{C=0}$ 1705 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.27 (3H, s), 3.10 (1H, d, J= 10.0 Hz), 3.75 (3H, s), 4.01 (1H, d, J = 8.4 Hz), 4.49 (1H, d, J=9.6 Hz), 4.58 (1H, d, J=10.0 Hz), 4.65 (1H, d, J=9.6 Hz), 5.15 (1H, d, J=8.4 Hz), 6.51 (2H, d, J=8.2 Hz), 6.71 (2H, d, J=8.9 Hz), 6.8 (2H, d, J=8.9 Hz), 7.05 (2H, d, J=8.2 Hz), 7.34 (1H, t, J=7.02 Hz), 7.43 (2H, t, J=7.4 Hz), 7.66 (2H, d, J=7.4 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.7; 54.2; 55.7; 59.7; 75.3; 80.5; 80.8; 114.5; 115.0; 123.9; 125.8; 127.8; 128.3; 129.3; 129.7; 130.2; 141.6; 143.0; 159.7; 173.2; 174.3. Anal. Calcd for C₂₇H₂₅N₃O₄ (455.51) C, 71.19; H, 5.53; N, 9.22; found C, 71.09; H, 5.70; N, 9.25.

4.2.16. *endo*-2-(4-Nitrophenyl)-3b-phenyl-5-*p*-tolyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3dione 4g. Yield 0.028 g, 50%; yellow needles; mp 176–177 °C; IR (KBr) $\nu_{C=0}$ 1716 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.29 (3H, s), 3.08 (1H, d, *J*= 10.0 Hz), 4.05 (1H, d, *J*=8.0 Hz), 4.47 (1H, d, *J*=9.6 Hz), 4.59 (1H, d, *J*=9.6 Hz), 4.62 (1H, d, *J*=10.0 Hz), 5.19 (1H, d, *J*=8.0 Hz), 6.50 (2H, d, *J*=8.2 Hz), 7.05 (2H, d, *J*= 8.2 Hz), 7.14 (2H, d, *J*=9.0 Hz), 7.36 (1H, t, *J*=7.4 Hz), 7.43 (2H, t, *J*=7.4 Hz), 7.65 (2H, d, *J*=7.4 Hz), 7.54 (2H, d, *J*=9.0 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.7; 54.2; 59.9; 75.4; 80.7; 81.4; 115.1; 124.4; 125.8; 127.1; 128.6; 129.4; 130.4; 130.5; 136.8; 141.0; 142.6; 147.2; 172.3; 173.5. Anal. Calcd for C₂₆H₂₂N₄O₅ (470.48) C, 66.37; H, 4.71; N, 11.91; found C, 66.30; H, 4.65; N, 11.85.

4.2.17. *endo*-2-(4-Chlorophenyl)-3b-phenyl-5-*p*-tolylhexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-**1,3-dione 4h.** Yield 0.029 g, 52%; white needles; mp 158–160 °C; IR (KBr) $\nu_{C=0}$ 1712 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.28 (3H, s), 3.08 (1H, d, J=10.0 Hz), 4.01 (1H, d, J=8.0 Hz), 4.48 (1H, d, J=9.6 Hz), 4.59 (1H, d, J=10.0 Hz), 4.63 (1H, d, J=9.6 Hz), 5.15 (1H, d, J=8.0 Hz), 6.50 (2H, d, J=8.4 Hz), 6.83 (2H, d, J=9.2 Hz), 7.05 (2H, d, J=8.4 Hz), 7.17 (2H, d, J=9.2 Hz), 7.31–7.38 (1H, m), 7.42–7.46 (2H, m), 7.64–7.67 (2H, m). ¹³C NMR (100 MHz, CDCl₃): δ 20.7; 54.2; 59.8; 75.3; 80.6; 81.0; 115.0; 125.8; 127.8; 128.4; 129.4; 129.4; 129.8; 130.0; 130.3; 134.7; 141.4; 142.8; 172.8; 174.0. Anal. Calcd for $C_{26}H_{22}ClN_3O_3$ (459.92) C, 67.90; H, 4.82; N, 9.14; found C, 68.00; H, 4.80; N, 9.22.

4.2.18. *endo*-3**b**-Phenyl-2,5-di-*p*-tolyl-hexahydro-7-oxa-2,5,6a-triaza-cyclopenta[*a*]pentalene-1,3-dione 4i. Yield 0.032 g, 60%; white needles; mp 178–179 °C; IR (KBr) $\nu_{C=0}$ 1706 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.28 (3H, s), 2.29 (3H, s), 3.10 (1H, d, *J*=10.0 Hz), 4.01 (1H, d, *J*=8.4 Hz), 4.49 (1H, d, *J*=9.2 Hz), 4.59 (1H, d, *J*=8.4 Hz), 6.52 (2H, d, *J*=8.0 Hz), 6.76 (2H, d, *J*=8.0 Hz), 7.06 (2H, d, *J*=8.0 Hz), 7.02 (2H, d, *J*=8.0 Hz), 7.06 (2H, d, *J*=8.0 Hz), 7.05 (1H, t, *J*=7.2 Hz), 7.43 (2H, t, *J*=7.2 Hz), 7.66 (2H, d, *J*=7.2 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 20.5; 21.1; 54.0; 59.5; 75.0; 80.4; 80.6; 114.8; 125.6; 126.1; 128.1; 128.5; 129.0; 129.5; 129.6; 130.0; 138.8; 141.4; 142.8; 172.9; 174.0. Anal. Calcd for C₂₇H₂₅N₃O₃ (439.51) C, 73.78; H, 5.73; N, 9.56; found C, 73.70; H, 5.70; N, 9.50.

4.2.19. 5-(4-Methoxyphenyl)-3a-phenyl-hexahydroimidazo[1,5-b]isoxazole-2,3-dicarboxylic acid 2-diethylamide 3-phenylamide 4b'. Yield 0.023 g, 18%; white needles; mp 166–167 °C; IR (KBr) $\nu_{\rm NH}$ 3445 cm⁻¹; $\nu_{\rm C=O}$ 1691 and 1620 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 0.92 (6H, t, J=7.6 Hz), 2.58 (4H, q, J=7.6 Hz), 3.17 (1H, d, J = 10.0 Hz), 3.70 (3H, s), 3.79 (1H, d, J = 7.2 Hz), 3.95 (1H, d, J=10.8 Hz), 4.12 (1H, d, J=10.0 Hz), 4.72 (1H, d, J=7.2 Hz), 4.95 (1H, d, J=10.8 Hz), 6.50 (2H, d, J=8.6 Hz), 6.70 (2H, d, J=8.6 Hz), 6.94 (1H, t, J=7.0 Hz), 7.09 (2H, t, J=7.0 Hz), 7.21-7.35 (6H, m), 7.59 (2H, d, J=7.8 Hz), 8.90 (1H, s). ¹³C NMR (100 MHz, CDCl₃): δ 11.3; 42.2; 55.8; 58.3; 66.7; 72.7; 79.4; 80.5; 115.0; 116.0; 120.1; 124.1; 125.9; 127.7; 128.7; 129.0; 138.2; 140.5; 144.7; 153.2; 168.6; 172.4. Anal. Calcd for C₃₀H₃₄N₄O₄ (514.62) C, 70.02; H, 6.66; N, 10.89; found C, 70.10; H, 6.60; N, 10.85.

4.3. Base catalysed ring-opening of compounds 3 and 4. Synthesis of 2,5-dihydro-1*H*-imidazole 5a–e. General procedure

Method A. Compound **3** or **4** (0.25 mmol) were refluxed in diethylamine (5 mL) for 23 h. The solvent was evaporated and the product was purified by preparative TLC using petroleum ether ethyl acetate as eluent (2:1). The products were recrystallized from ethanol or ether. Method B. The mixture of adducts **3** and **4** from the cycloaddition of nitrones **1** (0.25 mmol) with maleimides (1 mmol) **2** was dissolved in diethylamine (5 mL) and refluxed for 23 h. The solvent was evaporated and the isolation of the product **5** is as in Method A.

4.3.1. 4-Phenyl-1*-p***-tolyl-2,5-dihydro-1***H***-imidazole 5a.** Method B; yield 0.053 g, 90%; white needles; mp 117–119 °C; IR (KBr) $\nu_{C=N}$ 1631 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.28 (3H, s), 4.57 (2H, t, *J*=5.2 Hz), 5.43 (2H, t, *J*=5.2 Hz), 6.52 (2H, d, *J*=8.4 Hz), 7.11 (2H, d, *J*=8.4 Hz), 7.38–7.53 (3H, m), 7.82–7.89 (2H, m). ¹³C NMR (100 MHz, CDCl₃): δ 20.6; 55.3; 79.0; 112.2; 121.5; 127.6; 129.0; 130.3; 131.6; 136.0; 143.4; 169.3. Anal. Calcd for $C_{16}H_{16}N_2$ (236.31) C, 81.32; H, 6.82; N, 11.85; found C, 81.30; H, 6.85; N, 11.90.

4.3.2. 1-(4-Methoxyphenyl)-4-phenyl-2,5-dihydro-1*H***imidazole 5b.** Method B; yield 0.063 g, 100%; white needles; mp 129–130 °C; IR (KBr) $\nu_{C=N}$ 1631 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.78 (3H, s), 4.55 (2H, t, J=5.2 Hz), 5.41 (2H, t, J=5.2 Hz), 6.56 (2H, d, J=9.2 Hz), 6.91 (2H, d, J=8.4 Hz), 7.45–7.51 (3H, m), 7.87 (2H, d, J=8.0 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 55.7; 56.1; 79.4; 113.0; 115.5; 127.6; 129.0; 131.7; 132.5; 140.4; 152.0; 169.7. Anal. Calcd for C₁₆H₁₆N₂O (252.31) C, 76.16; H, 6.39; N, 11.10; found C, 76.20; H, 6.30; N, 11.05.

4.3.3. 2-(4-Methoxyphenyl)-4-phenyl-1-*p***-tolyl-2,5dihydro-1***H***-imidazole 5c. Method B; yield 0.080 g, 93%; white needles; mp 176–178 °C; IR (KBr) \nu_{C=N} 1623 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): \delta 2.23 (3H, s), 3.79 (3H, s), 4.72 (1H, dd,** *J***=15.2, 3.6 Hz), 4.99 (1H, dd,** *J***=15.2, 6.0 Hz), 6.45 (1H, dd,** *J***=6.0, 3.6 Hz), 6.51(2H, d,** *J***= 8.8 Hz), 6.89 (2H, d,** *J***=8.6 Hz), 7.02 (2H, d,** *J***=8.8 Hz), 7.39–7.48 (5H, m), 7.88 (2H, d,** *J***=7.8 Hz). ¹³C NMR (100 MHz, CDCl₃): \delta 20.5; 55.5; 57.5; 91.6; 113.0; 114.5; 126.6; 127.9; 128.3; 128.9; 130.0; 131.6; 132.4; 133.1; 143.3; 159.7; 166.8. Anal. Calcd for C₂₃H₂₂N₂O (342.43) C, 80.67; H, 6.48; N, 8.18; found C, 80.60; H, 6.40; N, 8.20.**

4.3.4. 1,2-Bis-(4-methoxyphenyl)-4-phenyl-2,5-dihydro-*1H*-imidazole 5d. Method B; yield 0.079 g, 88%; white needles; mp 152–153 °C; IR (KBr) $\nu_{C=N}$ 1623 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.73 (3H, s), 3.79 (3H, s), 4.69 (1H, dd, *J*=14.8, 4.0 Hz), 4.99 (1H, dd, *J*=14.4, 6.0 Hz), 6.40 (1H, dd, *J*=6.0, 4.0 Hz), 6.55 (2H, d, *J*=9.2 Hz), 6.81 (2H, d, *J*=9.2 Hz), 6.9 (2H, d, *J*=8.8 Hz), 7.4–7.5 (5H, m), 7.87–7.89 (2H, m). ¹³C NMR (100 MHz, CDCl₃): δ 55.5; 56.0; 57.9; 91.9; 113.8; 114.5; 115.2; 127.9; 128.3; 128.9; 131.5; 132.4; 133.2; 140.2; 152.0; 159.6; 166.9. Anal. Calcd for C₂₃H₂₂N₂O₂(358.43) C, 77.07; H, 6.19; N, 7.82; found C, 77.09; H, 6.10; N, 7.87.

4.3.5. 1-(4-Methoxyphenyl)-2-(3-nitrophenyl)-4-phenyl-2,5-dihydro-1*H***-imidazole 5e.** Method A (from **3e**) yield 0.086 g, 92%; Method A (from **4e**) yield 0.086 g, 92%; yellow needles; mp 182–184 °C; IR (KBr) $\nu_{C=N}$ 1623 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.74 (3H, s), 4.74 (1H, dd, *J*= 14.8, 4.0 Hz), 5.08 (1H, dd, *J*=14.8, 5.6 Hz), 6.49–6.54 (coincident 2H, d, *J*=9.0 Hz, 1H, dd, *J*=5.6, 4.0 Hz), 6.83 (2H, d, *J*=9.0 Hz), 7.44–7.57 (4H, m), 7.86–7.88 (3H, m), 8.16–8.19 (1H, m), 8.37 (1H, t, *J*=2.0 Hz). ¹³C NMR (100 MHz, CDCl₃): δ 55.6; 58.5; 91.5; 114.1; 115.4; 122.44; 123.5; 127.9; 129.1; 130.1; 131.9; 132.0; 133.5; 139.7; 143.5; 149.0; 152.7; 168.4. Anal. Calcd for C₂₂H₁₉N₃O₃ (373.40) C, 70.76; H, 5.13; N, 11.25; found C, 70.80; H, 5.10; N, 11.14.

4.4. The treatment of compounds 3 and 4 with triethylamine

Compound **3** or **4** (0.25 mmol) was dissolved in triethylamine (5 mL) and the mixture refluxed for 48 h. The solvent was evaporated and the starting material was recovered unchanged.

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Synthesis of chiral amino acids with metal ion chelating side chains from L-serine using Negishi cross-coupling reaction

Michael Kruppa, Giovanni Imperato and Burkhard König*

Institut für Organische Chemie, Universität Regensburg, Universitätsstrasse 31, 93040 Regensburg, Germany

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Abstract—The scope of the Negishi cross-coupling reaction of organozinc compounds derived from chiral amino acids was extended to electron rich iodoanilines and iodobenzylamines as coupling reagents. The protocol allows the direct modification of serine into phenylalanine derivatives bearing metal ion chelating ligands in their side chain, such as amino esters 6 and 7. The preparation of metal complex labeled peptides, the construction of synthetic receptors and hybrid materials are potential applications of the modified chiral amino acids.

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1. Introduction

The function of peptides and proteins is strongly correlated with their secondary and tertiary structure,¹ which is induced by side chain interactions and determined through the amino acid sequence. Non-proteinogenic amino acids²⁻⁴ and amino acid mimics are widely used to define,⁵ modulate⁶ or switch⁷ the structure of peptides, which may result in changes of their biological properties. One concept⁸ to control a peptide structure is the inter- or intramolecular formation of a metal complex, whereby parts of the peptide function as donor ligands. This requires modified amino acids bearing a strong metal ion chelating ligand for stable complex formation for incorporation into a peptide sequence.⁹ Different research groups used alkylation reactions of lysine or diaminopropionic acid to generate imino diacetic acid (IDA)¹⁰ or bis-(2-picolyl)amine (BPA)¹¹ chelates directly on protein surfaces.

As this modification of amino acid side chains is limited to a certain kind of chelates a more general solution to transform natural amino acids into metal ion chelating amino acids is desired.

Jackson already showed that palladium-catalyzed coupling of the organozinc compound derived from 1^{12} with aryl iodides is a feasible route to derivatives of phenylalanine (Scheme 1).¹³ We extend the scope of this reaction to

phenylalanine derivatives bearing p-NH₂ (**4**), p-NH-Boc (**5**), and p-Me (**3**) aryl substituents and report the efficient synthesis of the non-natural chiral amino acids **6** and **7** having protected IDA and cyclen metal ion chealting ligands in their side chain.

$$I \xrightarrow{H} PG \xrightarrow{1. Zn^{*}} Ar \xrightarrow{H} PG \xrightarrow{CO_2PG} 2. Arl, Pd_2(dba)_3, P(o-tol)_3} Ar \xrightarrow{H} PG \xrightarrow{CO_2PG} 1$$

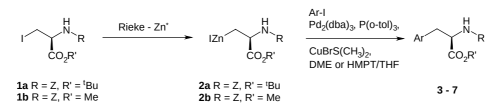
Scheme 1. Palladium-catalyzed coupling of aryl iodides with 1.

2. Results and discussion

So far, no Negishi coupling¹⁴ reaction has been reported for alkylorganozinc compounds with *p*-iodoaniline derivatives as reactants.¹⁵ Previous preparations of the unnatural amino acid H-*p*-NH₂-Phe-OH and protected derivatives of it used either the reduction of *p*-nitrophenyl alanine.¹⁶ or Staudinger reactions¹⁷ of *p*-azidophenyl alanine. The required iodo precursors $1a^{18,19}$ and $1b^{20}$ were synthesized from serine. The corresponding organozinc reagents 2a and 2b were obtained by direct insertion of freshly prepared Rieke zinc.²¹ The alternative method of zinc dust activation by Me₃SiCl in DMF was in our hands not as efficient as the use of Rieke zinc.²² The disappearance of the black zinc indicates the complete conversion by insertion reaction (Scheme 2).²³

Keywords: Amino acids; Metal chelates; Organozinc compounds; C–C bond formation; Negishi coupling.

^{*} Corresponding author. Tel: +49 941 943 4576; fax: +49 941 943 1717; e-mail: burkhard.koenig@chemie.uni-regensburg.de



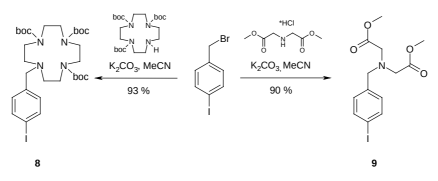
Scheme 2. Preparation of the organozinc reagents 2a and 2b and their coupling with aryl iodides.

Table 1. Palladium-catalyzed synthesis of the Phe derivatives 3-7

Entry	Iodoalanine	Ar-I	Solvent	Yield (%)	Product
1	2a		DME	66	
2	2a	NH ₂	HMPT/THF=1:1	73	
3	2b	H boc	HMPT/THF=1:1	75	× NH × NH × NH × S
4	2a	9	HMPT/THF=1:1	65	
5	2b	8	HMPT/THF=1:1	86	$z \sim N \rightarrow 0$ H $0 \sim 7$

The reaction conditions of the palladium-catalyzed coupling reaction were optimized using organozinc reagent 2a and *p*-iodotoluene and *p*-iodoaniline as aryl iodides. While *p*-iodotoluene gave 66% yield of the coupling

product (Table 1, entry 1) using DME as solvent with 0.1 equiv CuBr \cdot SMe₂,¹⁸ the more electron rich *p*-iodoaniline requires a solvent mixture of HMPT and THF (1:1) to give 73% of *Z*-*p*-NH₂-Phe-O^{*t*}Bu **4** (Table 1, entry 2). To



Scheme 3. Synthesis of aryl iodides 8 and 9.

apply the method to the synthesis of metal chelating chiral amino acids, aryl iodide substituted protected metal chelates were prepared starting from *p*-iodobenzyl bromide. Methyl imino diacetate and trifold Boc-protected cyclen²⁴ were alkylated in good yield giving aryl iodides **8** and **9** (Scheme 3).

Applying the same conditions as used for the coupling of *p*-iodoaniline, IDA derivative **6** was obtained in moderate 65% yield (Table 1, entry 4). Negishi coupling of aryl iodide **8** with iodoalanine **2a** gave cyclen-modified amino acid **7** in good yield of 86% (Table 1, entry 5).

3. Conclusion

We have extended the scope of the Negishi cross-coupling reaction of organozinc compounds derived from chiral amino acids to electron rich iodoanilines and iodobenzylamines as coupling reagents. This allows now the direct modification of serine into phenylalanine derivatives 4 and 5 and amino esters 6 and 7, bearing metal ion chelating ligands in their side chain. Such and similar modified chiral amino acids are useful for the preparation of peptides labeled with metal complexes as binding sites or probes, for the construction of synthetic receptors and hybrid materials having a peptide backbone and metal complex functionality.

4. Experimental

4.1. General procedure for the generation of organozinc reagents 2a and 2b

Rieke zinc was prepared from ZnCl₂ using a catalytic amount of naphthalene: to a Schlenk tube charged with finely divided lithium (1% Na) (3.0 mmol), naphthalene (0.3 mmol) and 2 mL of anhydrous THF (distilled from benzophenone-sodium ketyl under N2) or DME (in case of Z-*p*-Me-Phe-O^tBu **3**) under argon a solution of anhydrous zinc chloride [1.5 mmol in 2 mL of THF or DME (in case of Z-p-Me-Phe-O^tBu 3)] was transferred via syringe. The mixture was stirred vigorously until all lithium was consumed (0.5 h). To the Rieke zinc suspension under argon a solution of iodoalanine (1.28 mmol) in 3 mL of DME (in case of Z-p-Me-Phe-O^tBu 3) or THF was added via syringe. The exothermic reaction was completed in 5 min (the end of reaction is indicated when the black zinc disappeared). The mixture was stirred for an additional 0.5 h.

4.1.1. Tri-*tert*-butyl-10-(*p*-iodo-benzyl)-1,4,7,10-tetraaza cyclododecane-1,4,7-tricarbonate (8). *p*-Iodobenzylbromide (0.5 g, 1.7 mmol) was dissolved in 35 mL of MeCN. Trifold-Boc protected cyclen (1 g, 2.2 mmol) and K₂CO₃ (1.21 g, 8.8 mmol) were added, and after completion of the reaction the reaction mixture was filtered and concentrated. The crude product was purified by chromatography on silica gel (hexanes/dichloromethane = $5:1 \rightarrow$ AcOEt) giving **8** (1.09 g, 93%) as a colorless solid, mp = 75–77 °C. *R*_f (AcOEt) = 0.84; ¹H NMR (CDCl₃, 300 MHz): $\delta = 1.25-1.56$ (m, 27H), 2.39–2.78 (m, 4H), 3.11–3.73 (m, 14H), 7.00 (d, *J*=8.2 Hz, 2H), 7.61 (d, *J*=8.2 Hz, 2H); ¹³C NMR (CDCl₃, 75 MHz): δ =28.5, 28.7, 47.9, 50.0, 54.8, 55.5, 56.7, 79.5, 92.8, 132.2, 136.7, 137.2, 155.4, 155.7, 156.1; LR-MS (ESI/DCM/MeOH+10 mmol/L NH₄OAc): m/z=689 [(M+H)⁺].

4.1.2. Methyl [(4-iodo-benzyl)-methoxycarbonylmethylamino]-acetate (9). Dimethyl iminodiacetate hydrochloride (0.87 g, 4.4 mmol) and NaH (60% suspension) (0.18 g, 4.4 mmol) were dissolved in 35 mL of MeCN and stirred for 5 min. K₂CO₃ (2.42 g, 17.5 mmol) and *p*-iodobenzylbromide (1 g, 3.4 mmol) were added to this suspension. After completion of the reaction, the mixture was filtered and concentrated. The crude product was purified by chromatography on silica gel (hexanes/dichloromethane = $5:1 \rightarrow \text{AcOEt}$) yielding **9** (1.49 g, 90%) as a colorless oil. *R*_f (hexanes/dichloromethane = 5:1) = 0.44; ¹H NMR (CDCl₃, 300 MHz): δ = 3.52 (s, 4H), 3.68 (s, 6H), 3.84 (s, 2H), 7.13 (d, *J*=8.2 Hz, 2H), 7.62 (d, *J*=8.2 Hz, 2H); ¹³C NMR (CDCl₃, 75 MHz): δ =51.6, 54.1, 57.3, 92.9, 130.7, 137.2, 137.8, 171.5; LR-MS (CI/NH₃): *m*/*z*=378 [(M+H)⁺].

4.1.3. Z-p-Me-Phe-O'Bu 3. A Schlenk flask was charged with p-iodotoluene (0.34 g, 1.5 mmol), $Pd_2(dba)_3$ (40 mg, 2.5 mol%), P(o-tol)₃ (47 mg, 10 mol%), CuBr · DMS (32 mg, 0.1 equiv) and 4 mL of DME. At -10 °C a solution of the organozinc derivate of Z-I-Ala-O'Bu (1.2 mmol) in 4 mL of DME was added. The reaction mixture was slowly allowed to warm to room temperature over night. The reaction mixture was concentrated under argon and the solvent was removed in vacuum. AcOEt was added and the organic phase was washed twice with water and brine, the combined organic layers were dried over MgSO4 and concentrated. The crude resulting oil was purified by chromatography on silica gel (hexanes/AcOEt=4:1) affording 3 (0.31 g, 66%) as a colorless solid, mp = 51–52 °C. $R_{\rm f}$ (hexanes/AcOEt=4:1)=0.28; $[\alpha]_D^{20}$ +17.0 in CHCl₃; ¹H NMR (CDCl₃, 300 MHz): δ =1.41 (s, 9H), 2.31 (s, 3H), 3.04 (d, J = 5.5 Hz, 2H), 4.51 (dt, J = 5.9, 8.1 Hz, 1H), 5.10(s, 2H), 5.21 (d, J=8.2 Hz, 1H), 7.00–7.11 (m, 4H), 7.30–7.39 (m, 5H); ¹³C NMR (CDCl₃, 75 MHz): $\delta = 21.1$, 28.0, 37.9, 52.2, 66.8, 82.3, 128.1, 128.5, 129.1, 129.4, 132.9, 136.4, 136.5, 143.4, 155.6, 170.7; LR-MS (ESI/DCM/MeOH+10 mmol/L NH₄OAc): m/z = 756 $[(2 M + NH_4)^+], 387 [(M + NH_4)^+], 370 [(M + H)^+].$ Anal. Calcd for C₂₂H₂₇NO₄ (369.46): C 71.52, H 7.37, N 3.79; found C 71.60, H 7.14, N 3.60.

4.1.4. Z-p-NH₂-Phe-O'Bu 4. A Schlenk flask was charged with p-iodoaniline (0.34 g, 1.5 mmol), $Pd_2(dba)_3$ (40 mg, 2.5 mol%), P(o-tol)₃ (47 mg, 10 mol%), CuBr · DMS (32 mg, 0.1 equiv), and 4 mL of DME. At -10 °C a solution of the organozinc derivate of Z-I-Ala-O'Bu (1.2 mmol) in 4 mL of DME was added. The reaction mixture was slowly allowed to warm to room temperature over night. The reaction mixture was concentrated under argon and the solvent was removed under vacuum. AcOEt was added and the organic phase was washed twice with water and brine. The combined organic layers were dried over MgSO₄ and concentrated. The crude resulting oil was purified by chromatography on silica gel (3:2 hexanes/ AcOEt = 3:2) affording the pure product 4 (0.35 g, 73%) as a yellowish glass. $R_{\rm f}$ (hexanes/AcOEt=3:2)=0.2; $[\alpha]_{\rm D}^{20}$ +40.0 in CHCl₃; ¹H NMR (CDCl₃, 300 MHz): $\delta = 1.37$

(s, 9H), 2.92 (d, J=5.8 Hz, 2H), 3.56 (br s, 2H), 4.41 (dt, J=5.9, 8.1 Hz, 1H), 5.05 (s, 2H), 5.15 (d, J=8.0 Hz, 1H), 6.54 (d, J=8.2 Hz, 2H), 6.87 (d, J=8.2 Hz, 2H), 7.25–7.35 (m, 5H); ¹³C NMR (CDCl₃, 75 MHz): $\delta=28.0$, 37.5, 55.3, 66.8, 82.1, 115.2, 125.8, 128.1, 128.5, 128.6, 130.4, 136.5, 146.3, 155.7, 170.8; LR-MS (ESI/DCM/MeOH+10 mmol/L NH₄OAc): m/z=388 [(M+NH₄)⁺], 371 [(M+H)⁺]. Anal. Calcd for C₂₁H₂₆N₂O₄ (370.45): C 68.09, H 7.07, N 7.56; found C 68.23, H 7.20, N 7.38.

4.1.5. Z-p-Boc-NH-L-Phe-OMe 5. A Schlenk flask was charged with (4-iodo-phenyl)-carbamic acid tert-butyl ester (0.38 g, 1.2 mmol), Pd₂(dba)₃ (40 mg, 2.5 mol%), P(o-tol)₃ (47 mg, 10 mol%), CuBr · DMS (32 mg, 0.1 equiv), 2 mL of HMPT, and 2 mL of THF. At -10 °C a solution of the organozinc derivate of Z-I-L-Ala-OMe (1.1 mmol) in 2 mL of THF was added. The reaction mixture was slowly allowed to warm to room temperature over night. The reaction mixture was concentrated under argon and the solvent was removed under vacuum. AcOEt was added and the organic phase was washed twice with water and brine. The collected organic layers were dried over $MgSO_4$ and concentrated. The crude resulting oil was purified via chromatography on silica gel (hexanes/AcOEt = 2:1) affording the pure product 5 (0.39 g, 75%) as a solid, mp = 65–67 °C. $R_{\rm f}$ (hexanes/AcOEt=2:1)=0.2; $[\alpha]_{\rm D}^{20}$ +10.0 in CHCl₃; ¹H NMR (CDCl₃, 300 MHz): $\delta = 1.51$ (s, 9H), 2.97-3.13 (m, 2H), 3.71 (s, 3H), 4.57-4.68 (m, 1H), 5.09 (s, 2H), 5.21 (d, J=8.2 Hz, 1H), 7.00 (d, J=8.5 Hz, 2H), 7.27 (d, J = 8.2 Hz, 2H), 7.30-7.44 (m, 5H), 7.30-7.39 (m, 5H);¹³C NMR (CDCl₃, 75 MHz): $\delta = 28.3$, 37.5, 52.4, 54.8, 67.0, 80.6, 118.6, 128.1, 128.2, 128.6, 129.9, 130.1, 136.2, 137.4, 152.7, 155.6, 172.0; LR-MS (ESI/DCM/MeOH+ 10 mmol/L NH₄OAc): $m/z = 446 [(M + NH_4)^+], 429 [(M + NH_4)^+]$ H)⁺], 373 [(M+H-C₄H₈)⁺], 329 [(M+H-Boc)⁺]. Anal. Calcd for C₂₃H₂₈N₂O₆ (428.49): C 64.47, H 6.59, N 6.54; found C 64.30, H 6.27, N 6.19.

4.1.6. Z-p-Me(IDA-OMe)-Phe-O'Bu 6. A Schlenk flask was charged with 9 (0.30 g, 1.2 mmol), $Pd_2(dba)_3$ (40 mg, 2.5 mol%), P(o-tol)₃ (47 mg, 10 mol%), CuBr · DMS (32 mg, 0.1 equiv), 2 mL of HMPT, and 2 mL of THF. At -10 °C a solution of the organozinc derivate of Z-I-Ala-O'Bu (1.2 mmol) in 4 mL of THF was added. The reaction mixture was slowly allowed to warm to room temperature over night. The reaction mixture was concentrated under argon and the solvent was removed under vacuum. AcOEt was added and the organic phase was washed twice with water and brine. The collected organic layers were dried over MgSO₄ and concentrated. The crude resulting oil was purified via chromatography on silica gel (hexanes/ AcOEt = 7:3) affording the pure product 6 (0.41 g, 65%) as a glass. $R_{\rm f}$ (hexanes/AcOEt=7:3)=0.18; $[\alpha]_{\rm D}^{20}$ + 14.6 in CHCl₃; ¹H NMR (CDCl₃, 600 MHz): $\delta = 1.42$ (s, 9H), 3.09 (d, J = 5.7 Hz, 2H), 3.57 (s, 4H), 3.72 (s, 6H), 3.91 (s, 2H),4.51-4.57 (m, 1H), 5.12 (s, 2H), 5.25 (d, J=7.7 Hz, 1H), 7.13 (d, J=7.5 Hz, 2H), 7.31 (d, J=7.6 Hz, 2H), 7.32–7.40 (m, 5H), 7.30–7.39 (m, 5H); ¹³C NMR (CDCl₃, 150 MHz): $\delta = 27.9, 38.1, 51.5, 53.9, 55.2, 60.4, 66.8, 82.3, 128.1,$ 128.2, 128.5, 129.1, 129.6, 135.3, 136.0, 136.4, 155.6, 170.5, 171.5; LR-MS (ESI/DCM/MeOH+10 mmol/L NH₄OAc): $m/z = 529 [(M+H)^+], 473 [(M+H-C_4H_8)^+].$

Anal. Calcd for $C_{28}H_{36}N_2O_8$ (528.61): C 63.62, H 6.86, N 5.30; found C 63.79, H 6.91, N 5.20.

4.1.7. Z-p-Me(3-Boc-Cyc)-L-Phe-OMe 7. A Schlenk flask was charged with 8 (0.19 g, 0.28 mmol), Pd₂(dba)₃ (10 mg, 2.5 mol%), P(o-tol)₃ (12 mg, 10 mol%), CuBr · DMS (8 mg, 0.1 equiv), 2 mL of HMPT, and 2 mL of THF. At -10 °C a solution of the organozinc derivate of Z-I-L-Ala-OMe (0.35 mmol) in 2 mL of THF was added. The reaction mixture was slowly allowed to warm to room temperature over night. The reaction mixture was concentrated under argon and the solvent was removed under vacuum. AcOEt was added and the organic phase was washed twice with water and brine. The collected organic layers were dried over MgSO₄ and concentrated. The crude resulting oil was purified via chromatography on silica gel (hexanes/ AcOEt = 3:2) affording the pure product 7 (0.19 g, 86%)as a solid, mp = 56–57 °C. R_f (hexanes/AcOEt = 3:2) = 0.16; $[\alpha]_D^{20}$ +40.4 in CHCl₃; ¹H NMR (CDCl₃, 300 MHz): $\delta = 1.42$ (s, 9H), 1.47 (s, 18H), 2.53–2.79 (m, 4H), 3.03-3.11 (m, 2H), 3.11-3.50 (m, 8H), 3.57 (br s,4H), 3.69 (s, 3H), 4.57-4.70 (m, 1H), 5.08 (s, 2H), 5.21 (d, J = 7.7 Hz, 1H), 7.02 (d, J = 8.0 Hz, 2H), 7.16 (d, J = 8.0 Hz, 2H), 7.30–7.44 (m, 5H), 7.30–7.38 (m, 5H); ¹³C NMR (CDCl₃, 75 MHz): δ = 28.5, 28.7, 37.9, 47.3, 48.9, 49.5, 50.0, 52.3, 54.8, 56.9, 67.0, 79.3, 79.4, 128.1, 128.2, 128.6, 129.1, 130.3, 130.6, 134.7, 136.7, 155.6, 155.8, 156.1, 171.9; HR-MS (EI/70 eV): m/z = calcd for $C_{42}H_{63}N_5O_{10}$ 797.4575; found 797.4559 \pm 0.62 ppm.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11.034. HPLC analysis of compound **4** prepared from chiral and racemic **2a**.

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Tetrahedron

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A straightforward synthesis and structure of unprecedented iminium salts of dihydropyrido[3,2-*e*][1,3]thiazines

Margarita Suárez,^{a,*} Hector Novoa,^{b,*} Yamila Verdecia,^a Estael Ochoa,^a Amaury Alvarez,^{a,c} Rolando Pérez,^c Roberto Martínez-Alvarez,^d Dolores Molero,^e Carlos Seoane,^d Norbert M. Blaton,^b Oswald M. Peeters^b and Nazario Martín^{d,*}

^aLaboratorio de Síntesis Orgánica, Facultad de Química, Universidad de La Habana, 10400-Ciudad Habana, Cuba

^bLaboratorium voor Analytische Chemie en Medicinale Fysicochemie, Faculteit Farmaceutische Wetenschappen,

K.U. Leuven, Van Evenstraat 4, B-3000 Leuven, Belgium

^cInstituto Cubano de Investigaciones de los Derivados de la Caña de Azúcar, 10400-Ciudad Habana, Cuba ^dDepartamento de Química Orgánica, Facultad de Ciencias Químicas, Universidad Complutense, E-28040 Madrid, Spain ^cCAI de Resonancia Magnética Nuclear, Facultad de Ciencias Químicas, Universidad Complutense, E-28040 Madrid, Spain

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Abstract—Unprecedented 2-iminium chloride salts of 5,8-dihydro-2H-pyrido[3,2-e][1,3]thiazines derivatives (8) were easily synthesized in one step from the corresponding o-chloroformyl-1,4-dihydropyridine (2) and thiourea. The structural study has been carried out by X-ray crystallography and theoretical calculations at the B3LYP/6-31G* levels and reveal that the new salts exhibit appropriate structural features to behave as calcium channel modulators.

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1. Introduction

1,3-Thiazines are an important type of heterocycles showing a wide variety of pharmacological properties. Thus, 1,3thiazine derivatives have recently been reported as cholecystokinin antagonists¹ or antimycobacterial agents.² Fused thiazines such as those incorporating a pyridine nucleus have shown anticancer,³ antitumor,⁴ and antioxidant activities.⁵

On the other hand, 1,4-dihydropyridine derivatives (1,4-DHPs) form a class of heterocyclic compounds with interesting pharmacological and biological properties.⁶ The systematic structural modification of the 1,4-DHP ring yields different compounds used in the treatment of hypertension and angina pectoris.⁷ The most prominent of these compounds is Nifedipine, which was the first generation calcium channel blocker marketed by Bayer.⁸

Since then, a wide variety of novel compounds belonging to the second and third generations of new biological active substances from 1,4-DHP class have been developed in order to obtain larger bioavailability or greater tissue selectivity.⁹ Up to now, 1,4-DHPs are still the most potent group of calcium channel modulators and, therefore, the design and study of this class of compounds remains highly desirable.¹⁰

Very recently, new 4-aryl-1,4-dihydropyridines have been investigated as *P*-glycoprotein inhibitors¹¹ and *N*-alkoxy-carbonylmethyl derivatives of 1,4-dihydropyridine-3,5-dicarboxylate were reported to act as a new carrier system for delivering drugs to the brain.¹² The above studies clearly show that the 1,4-DHP nucleus appears to be a unique structure interacting with a wide variety of channels and receptors, thus being considered as a class of privileged pharmacophoric structures.⁹

Pyridothiazine derivatives have been synthesized by tedious multi-step procedures,¹³ and were patented as pharmacological agents.¹⁴ Recently, the antiepileptic properties of pyrido-1,3-thiazin-4-one derivative have been reported.¹⁵

In previous works we have reported the synthesis of 6-chloro-5-formyl-1,4-dihydropyridines (2), which were

Keywords: 2-Imino pyrido[3,2-*e*][1,3]thiazines; X-ray analysis; 1,4-Dihydropyridines; Iminium salts.

^{*} Corresponding authors. Tel.: +53 7 878 1398; fax: +53 7 873 5737 (M.S.); tel.: +32 16 323427; fax: +32 16 323469 (H.N.); tel.: +34 91 394 4332; fax: +34 91 394 4103 (N.M.); e-mail addresses: msuarez@fq.uh.cu; hector.novoa@pharm.kuleuven.be; nazmar@quim.ucm.es

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obtained from the corresponding 2(1H)pyridone **1**.¹⁶ 1,4-Dihydropyridine **2** proved to be an excelent candidate for further transformations into other heterocyclic-fused 1,4dihydropyridines such as pyrazolo[3,4-*b*]pyridines **3**¹⁶ and 4,7-dihydrothieno[2,3-*b*]pyridines **4**.¹⁷ Recently, we carried out the synthesis of novel fulleropyrrolidines **5** bearing biologically active 1,4-DHPs covalently connected to the fullerene core, from the corresponding 1,4-DHP **2**¹⁸ (see Chart 1).

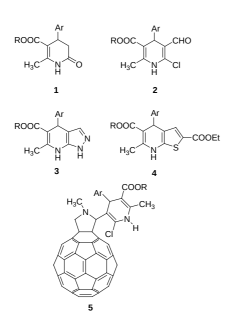
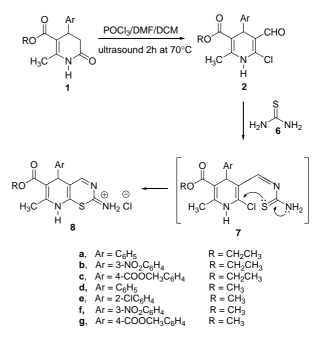


Chart 1.

2. Results and discussion

In this paper, we report on the synthesis of novel 2-iminodihyropyrido[3,2-e][1,3]thiazines (8), a class of unprecedented compounds, which are readily available from 6-chloro-5-formyl 1,4-dihydropyridines (2) by reaction with thiourea.

Compounds 2 were obtained as previously reported,¹⁶ from the corresponding methyl 2-oxo-1,2,3,4-tetrahydropyridine-5-carboxylate (1) via Vilsmeier-Haack chloroformylation. Even though this procedure allows the preparation of the desired 6-chloro-5-formyl 1,4-DHP in a straightforward fashion, we have made some modifications in order to simplify the experimental work. The new chloroformylation methodology involved the interaction of the halomethylenium salt, formed in situ from POCl₃ and DMF, with alkyl 4-aryl-6-methyl-2-oxo-1,3,3,4-tetrahydropyridine-5-carboxylate 1 in DCM at room temperature. Further irradiation of the reaction mixture in an ultrasonic bath at 70 °C for 2 h and subsequent hydrolysis afforded the desired 6-chloro-5formyl 1,4-DHP derivative 2 in good yields (Scheme 1). After washing twice with small portions of cold ether, the crude solid was obtained with excellent purity, as determined by TLC and RP-HPLC (>98%, based on the integration peak area at 226 nm).



Scheme 1. 2-Iminopyrido[3,2-e][1,3]thiazines synthesized 8a-g.

The ultrasonic-based methodology possesses important improvements over our previously reported conventional Vilsmeier-Haack chloroformylation. The reaction times were notably reduced, the final product was obtained with an excellent purity, and hence, it could be used in further synthetic steps without the need of wasteful purification. The structures of compounds **2** were confirmed by spectroscopic methods and are in agreement with previously reported data.¹⁶

The synthesis of the 2-iminopyrido[3,2-*e*][1,3]thiazines **8a–g** was readely accomplished by refluxing the appropriate alkyl 4-aryl-6-chloro-5-formyl-2-methyl-1,4-dihydropyridine-3-carboxylate **2a–g** with an equivalent amount of thiourea in dry ethanol under an inert atmosphere (Scheme 1). After a few minutes, the iminium chloride that precipitated from the reaction medium, was filtered off and washed with hot ethanol and dried.

Formation of the 1,3-thiazine ring can be accounted for by nucleophilic attack of the amino group in the thiourea at the formyl group in **2**, followed by a dehydration to yield the intermediate **7**. Subsequent attack of the sulfur atom with loss of a HCl molecule affords compounds **8**, via a 6-*endo*-*trig* cyclization,¹⁹ as stable crystaline solids in moderate to good yields (Scheme 1).

Compounds **8a–g** show a satisfactory analytical and spectroscopic data. The FTIR spectra of compounds **8** present the amino and carbonyl signals at 3360 and 3200 cm^{-1} (NH) and $1690-1700 \text{ cm}^{-1}$ (C=O). In addition, the C=N signal appears at 1645 cm^{-1} . The ¹H NMR (300 MHz) spectra of 2-iminiumpyrido[3,2-*e*][1,3]thiazine chloride derivatives **8a–g** show a very similar pattern. Assignment of the signals was ascertained by 1D and 2D NMR experiments. The NH and $-C=NH_2^+$ protons appear as broad singlets at ~12.6, 10.6 and ~10.3 ppm. The vinyl proton on C4, H5 and the methyl

group on C7 were observed as a singlet at δ 8.4–8.8, ~5 and ~2.4 ppm, respectively. The signals between 7.0–8.5 ppm correspond to the protons of the aromatic ring on C5, showing the characteristic multiplicity depending on the position of the substituent.

The ${}^{13}C$ NMR spectra of these compounds (8a-g) exhibit signals in the carbonyl, aromatic and aliphatic region. In order to unequivocally assign the signals corresponding to the heterocyclic ring, we used 1D and 2D techniques: DEPT(135), HMQC and HMBC. The signals corresponding to the carbon atoms of the bicyclic ring are relatively insensitive to the nature of the substituent on the phenyl ring. Around 165-166 ppm appears the signal corresponding to the carbonyl group, and the signal corresponding to C4 and C2 at 166–167 and ~155 ppm, respectively. C4a (106–107 ppm) and C6 (107–108 ppm) appear at lower δ values than those expected for typical olefinic carbon atoms, whereas C7 and C8a appear at higher δ values, 143–144 and \sim 167 ppm, respectively. These findings have been accounted for by the strong push-pull effect of the groups linked to the olefinic double bond, similarly to that observed previously in other related molecules.²⁰ C5 appears at 39–40 ppm and the rest of signals are in agreement with the nature of each particular aromatic or aliphatic carbon atoms (see Section 4).

The mass spectra under EI conditions of the 2-iminiumpyrido[3,2-*e*][1,3]thiazine chlorides (8) show the $[M-HCl]^+$ ions with low intensity. The loss of the aryl substituent on C5 gives the pyrylium ion as the base peak of the spectra in m/z=250 for 8a-c and m/z=236 for 8d-g. This spectrometric behaviour under EI conditions is similar to those reported for analogous structures.²¹

In order to gain a better understanding of the novel compounds, and to confirm their molecular structure, we have carried out X-ray diffraction studies. Single crystals of **8a** were obtained from slow evaporation in methanol solution. From a search in the latest version of the Cambridge Crystallographic Database (CSD, version 5.26),²² no crystal structure was found having the 2-imino-dihydropyrido[3,2-*e*][1,3]thiazine moiety, being the crystal structure of **8a** the first of this type of compounds. The molecules of **8a** crystallized as a hydrochloride salt. In the solid state, the DHP ring has a screw boat conformation

with puckering parameters (N8-C7-C6-C5-C4a-C8a)²³ Q = 0.185(6) Å, $\theta = 105.9(19)^{\circ}$ and $\varphi = 8(2)^{\circ}$ and a weighted average bond distance of 1.420(3) Å. The ring conformation represents 31% of puckering in ideal cyclohexane chair (17% chair with N8 pointing up, 24% twist boat with axis through C8A and C4A pointing up, and 59% boat with bowsprit at N8 pointing down). The bisection of the aromatic ring at C5 with respect to the DHP, described by torsion angle C6–C5–C1'–C2', displays a value of $71.2(8)^\circ$, showing that the plane of the phenyl ring bisects the pyridine ring. The torsion angle C1'-C5-C4a-C8a of $106.7(6)^{\circ}$ shows that the phenyl ring is in axial position. The ester group at C6 was found to be nearly coplanar with the nearest endocyclic double bond in the DHP ring and having the carbonyl group at C10 in a trans (ap) disposition with respect it.

The thiazine ring has puckering parameters (S1-C2-N3-C4-C4a-C8a):²³ Q=0.117(5) Å, $\theta=70(3)^{\circ}$ and $\varphi=$ $31(3)^{\circ}$ and can be described as screw boat with axis through N3 and C4 pointing up. This ring conformation represents 18% puckering of an ideal cyclohexane chair (22% chair with S1 pointing up, 76% twist boat with axis through N3 and C4 pointing up, and 2% boat with bowsprit at C2 pointing down). The weighted average bond distance in this ring is 1.526(3) Å. The distances involving the S atom in the heterocyclic ring are: $d(S1-C2(sp^2)) = 1.757(6)$ Å and $d(S1-C2(sp^2)) = 1.726(6)$ Å. The bond angles around C2 shows the sp² character of this atom [S1–C2–N3: 123.3(5)°, S1-C2-N9: 116.8(5)°, N3-C2-N9: 119.8(6)°. Figure 1 shows the solid state conformation of 8a, together with the atomic numbering scheme. Figure 2 shows the projection of the molecules of 8a in the unit cell, denoting the hydrogen bonds that involves the Cl atom. The molecules are packed by means of hydrogen bonds. The Cl atom is involved in three hydrogen bonds of the type $N \cdots$ Cl, one intramolecular (Fig. 1) and the others with two neighboring host molecules, resembling a propeller shape arrangement with the Cl atom in the centre and forming an infinite one-dimensional chain along the [001] direction. Intramolecular hydrogen bonds: N8-H8...Cl with N8... Cl = 3.145(5) Å and $N8-H8\cdots Cl = 171^{\circ}$. Intermolecular hydrogen bonds: N9–H9A····Cl (x+1, y, z) with N9···· Cl = 3.235(6) Å and N9-H9A···Cl = 171°, and N9-H9B··· Cl(x+1/2, -y-1/2, -z) with N9····Cl=3.126(5) Å and Å and N9–H9B····Cl = 170° .

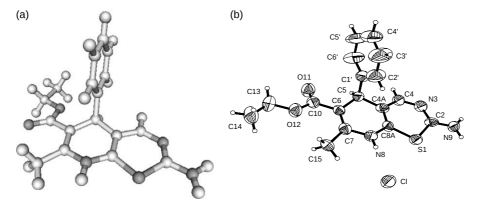


Figure 1. (a) Most stable conformation for compound 8a calculated by B3LYP/6-31G*. (b) ORTEP diagram of the crystal structure of 8a showing its atomic numbering scheme. Displacement ellipsoids are drawn at 50% probability level for non-H atoms.

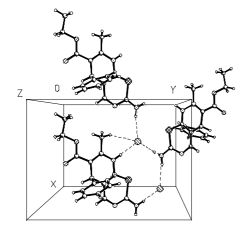


Figure 2. Projection of the molecules of **8a** in the unit cell along the [001] direction and denoting the hydrogen bonds that involves the Cl atom.

Also, the main geometrical characteristics of the synthesized compounds were obtained from $B3LYP/6-31G^{*24}$ ab initio calculations using the Gaussian 98^{25} program. The optimized geometry for compound **8a** is shown in Figure 1. Table 1 shows geometrical data calculated for compound **8a** (B3LYP/6-31G*) as well as those determined by X-ray

Table 1. Most relevant bond distances (Å) valence angles (°) and torsion angles (°) for the most stable conformation of compound **8a**, obtained by B3LYP/6-31G*, as well as by X-ray diffraction

	B3LYP/6-31G*	X-ray
Bond distrances (Å)		
S1-C2	1.783	1.757(6)
C2-N3	1.311	1.349(8)
N3-C4	1.340	1.332(8)
C4–C4a	1.393	1.388(8)
C4a–C8a	1.394	1.379(8)
C8a–S1	1.759	1.726(6)
C2-N9	1.337	1.292(8)
C4a–C5	1.532	1.530(8)
C8a–N8	1.344	1.349(7)
C5-C6	1.524	1.506(8)
C6–C7	1.353	1.351(9)
C7–N8	1.419	1.410(8)
C5-C1′	1.533	1.515(9)
Bond angles (°)		
C8a-S1-C2	100.3	101.6(3)
S1-C2-N3	125.3	123.3(5)
C2-N3-C4	122.2	122.1(6)
C4a-C5-C6	111.5	111.1(5)
C7–N8–C8a	123.7	122.2(5)
C4a–C5–C1′	110.3	111.4(5)
Torsion angles (°)		
N8-C8a-C4a-C5	7.3	7.4(9)
C8a-C4a-C5-C6	-18.0	-18.3(8)
C4a-C5-C6-C7	17.1	16.3(8)
C5-C6-C7-N8	-4.9	-3.1(9)
C6-C7-N8-C8a	-8.4	-10.4(9)
C7-N8-C8a-C4a	7.1	8.0(9)
$\Sigma \rho $	62.8	63.5(9)
C8a-S1-C2-N3	-1.8	-11.5(6)
S1-C2-N3-C4	2.7	9.3(8)
C2-N3-C4-C4a	-0.4	1.0(10)
N3-C4-C4a-C8a	-2.7	-6.0(10)
C4-C4a-C8a-S1	3.2	1.5(8)
C4a-C8a-S1-C2	-1.1	6.1(6)
C4a-C5-C1'-C2'	-52.5	-53.5(8)
C6-C5-C1'-C2'	72.6	71.2(8)
C5-C6-C10-O11	8.8	19.5(9)

diffraction. The data for the remaining compounds are collected in the Supplementary material.

In all cases, the distortion from planarity of the atoms comprising the 1,4-DHP ring can be clearly seen from the torsion angles calculated about the ring bonds. The greatest displacement from zero occurs from N1 and C4, indicating that the largest degree of puckering occurs at these positions, the distortion being greatest at the C4 position, as found in the solid state structure. The magnitude and sign of this torsion angles indicate that both C5 and N8 (Table 1) lie above of the plane formed by C8a, C4a, C6 and C7, which imparts a boat-like conformation to the DHP ring. Calculations predicted that the substituent on C5 is in a pseudoaxial position (torsion C8a-C4a-C5-C1' between 96 and 102°) and bisecting the plane containing the 1,4-DHP ring (torsion angles C4a–C5–C1 $^{\prime}$ –C2 $^{\prime}$ with values between 100 and 111°). It is important to note that B3LYP/6-31G* calculations predict the 1,3-thiazine ring planar as can be seen from the values of the corresponding dihedral angles. Also the condensation of the 1,4-DHP ring to the 1,3thiazine ring makes the former more planar in comparison to the monocyclic 1,4-DHP.²⁶ The sum, $\Sigma |\rho|$, of the absolute values of the internal torsion angles is a measure of the planarity of the 1,4-DHP ring. Different from the solid state structure, the geometry at the calculated minimum energy in the gas phase shows the exocyclic ester at C6 in a cis conformation with respect to the C6=C7 double bond (Fig. 1).

3. Conclusions

In summary, we have carried out the synthesis and characterization of unprecedented pyrido[3,2-*e*][1,3]thiazin-2-iminium chloride and determined their structure by X-ray analysis. It has been found that the 1,4-dihidropyridine shows a screw boat conformation with a pseudoaxial orientation of the aryl ring in C5 position. The geometrical features of the studied compounds are quite similar to the structurally related 1,4-dihydropyridines and, therefore, they exhibit appropriate structural features to act as potential calcium channel modulators.

4. Experimental

4.1. General

Reagents and solvents were purchased from Fluka or Aldrich. Alkyl 4-aryl-6-methyl-2-oxo-1,2,3,4-tetrahydropyridine-5-carboxylate were obtained as previously reported¹⁶ from commercial reagents. The progress of the reaction and the purity of compounds were monitored by TLC analytical silica gel plates (Merck $60F_{250}$). Column chromatographies were carried out with silica gel 60 (70–230 mesh ASTM). Melting points were determined in capillary tubes in an Electrothermal 9100 apparatus and are uncorrected. A Decon (U.K.) ultrasonic bath equipped with 40 KHz frequency transducer was used for ultrasonically irradiated reactions. Analytical HPLC runs were performed in an Amersham Bioscience Akta 10 equipment; gradient: 5–40% of ACN (0.05% TFA) in 15 min; acquisition/ processing data was accomplished with the UNICORN 4.11 program. Spectra were obtained as follow: FTIR were recorded on a FTIR 8300 spectrometer; ¹H NMR spectra were recorded at 300 MHz, and ¹³C NMR at 75.5 MHz, on a Bruker Avance-300 instrument, the one-bond heteronuclear correlation (HMQC) and the long range ¹H–¹³C correlation (HMBC) spectra were obtained using the inv4gs and the inv4gslpIrnd programs, respectively, with the Bruker software. Mass spectrawere obtained with a Hewlett Packard 5989A spectrometer. Microanalysis was performed in a Perkin-Elmer 2400 CHN by the Servicio de Microanálisis de la Universidad Complutense de Madrid. DFT calculations were performed using B3LYP/6-31G* basis set.²⁴ All calculations were carried out using the Gaussian 98 program.²⁵

4.2. Synthesis of ethyl 4-aryl-6-chloro-5-formyl-2methyl-1,4-dihydropyridine-3-carboxylates (2) under ultrasound irradiation

To a stirred mixture of ethyl 4-aryl-6-methyl-2-oxo-1,2,3,4tetrahydropyridine-5-carboxylate **1** (7 mmol) and 1.4 mL (18.2 mmol) of DMF in 15 mL of dry DCM under a nitrogen atmosphere, 1.1 mL (12.2 mmol) of POCl₃ was slowly added. The mixture was sonicated at 70 °C during 2 h. An aqueous sodium acetate solution was added (12 g in 21 mL of water) and stirred for 0.5 h. The reaction mixture was partitioned between water and DCM, and the aqueous phase was extracted with ethyl acetate. The organic phases were collected and dried under MgSO₄. The organic solvent was removed in vacuo and the solid residue was washed twice with small portions of cold ether to afford **2**.

4.3. General procedure for 6-alkoxycarbonyl-7-methyl-5-phenyl-5,8-dihydro-2*H*-pyrido[3,2-*e*][1,3]thiazin-2iminium chloride (8a–g)

A mixture of the appropriate alkyl 4-aryl-6-chloro-5formyl-2-methyl-1,4-dihydropyridine-3-carboxylate 2a-g(2 mmol) and thiourea 6 (2 mmol) in dry ethanol (10 ml) was refluxed for 30 min. The precipitated solid was filtered, washed with hot ethanol (3×5 mL) and dried.

4.3.1. 6-Ethoxycarbonyl-7-methyl-5-phenyl-5,8-dihydro-2H-pyrido[3,2-e][1,3]thiazin-2-iminium chloride (8a). Prepared from ethyl 6-chloro-4-phenyl-5-formyl-2-methyl-1,4-dihydropyridine-3-carboxylate (2a). Yield, 71%; mp 286–287 °C; IR (KBr) $\nu_{\rm max}$ 3360 and 3210 ($\nu \rm NH), 1692$ $(\nu C=0)$, 1643 $(\nu C=N)$ cm⁻¹; ¹H NMR (DMSO- d_6) δ : 12.72 (1H, s, NH), 10.58 (1H, s, NH), 10.30 (1H, s, NH), 8.39 (1H, s, H4), 7.33–7.19 (5H, m, Ph), 5.03 (1H, s, H5), 4.05-3.97 (2H, m, OCH₂), 2.41 (3H, s, CH₃), 1.07 (3H, t, CH₃); ¹³C NMR (DMSO-*d*₆) δ 167.2 (C8a), 166.3 (C4), 165.5 (COO), 155.4 (C2), 146.2 (C1[']), 143.3 (C7), 128.9 (C2['], C6[']), 127.2 (C4[']), 126.9 (C3['], C5[']), 108.4 (C6), 107.3 (C4a), 60.1 (OCH₂), 40.1 (C5), 17.8 (CH₃), 13.9 (CH₃); MSEI *m*/*z* (%): 327 ([M-HCl]⁺, 21), 312 (7), 298 (31), 250 (100), 223 (40), 195 (23). Anal. Calcd for C₁₇H₁₈-ClN₃O₂S (363.86): C: 56.12; H: 4.99; N: 11.55; S: 8.81. Found: C: 56.01; H: 4.91; N: 11.58; S: 8.97.

4.4. Crystal structure determination of 8a

Crystals suitable for X-ray diffraction were grown by slow evaporation from absolute methanol solution. The crystallographic and experimental data for these compounds are summarised in Table S1. Measurements were carried out using a Siemens P4 four-circle diffractometer with graphite monochromated and Cu K α_1 radiation. The intensity data were collected using $\omega - 2\theta$ scans, with ω scan width equal to the low range plus the high range plus the separation between the $K\alpha_1$ and $K\alpha_2$ positions; 4522 reflections measured. Empirical absorption correction, via ψ scan was applied.²⁷ Three standard reflections were monitored every 100 reflections (intensity decay: none). The structure was solved by direct methods and Fourier synthesis. Non-H atoms were refined anisotropically by full-matrix leastsquares techniques. H atoms were calculated geometrically and included in the refinement, but were restrained to ride on their parent atoms. The isotropic displacement parameters of the H atoms were fixed to 1.3 times U_{eq} of their parent atoms. Data collection: XSCANS.²⁸ Cell refinement: XSCANS.²⁸ Data reduction: XSCANS.²⁸ Program used to solve structure: SIR92.²⁹ Program used to refine structure: SHELXL97.³⁰ Molecular graphics: DIAMOND.³¹ Software used to prepare material for publication: PLATON.³² Detailed crystallographic data for have been deposited at the Cambridge Crystallographic Data Centre (CCDC 265438) and are available on request.

4.4.1. 6-Ethoxycarbonyl-7-methyl-5-(3'-nitrophenyl)-5,8-dihydro-2*H*-pyrido[3,2-*e*][1,3]thiazin-2-iminium chloride (8b). Prepared from ethyl 6-chloro-4-(3-nitrophenyl)-5-formyl-2-methyl-1,4-dihydropyridine-3-carboxylate (2b). Yield 53%; mp 273–275 °C; IR (KBr) ν_{max} 3392 and 3260 (vNH), 1690 (vC=O), 1645 (vC=N), 1530 and 1355 (ν NO₂) cm⁻¹; ¹H NMR (DMSO- d_6) δ 12. 74 (1H, s, NH), 10.68 (1H, s, NH), 10.37 (1H, s, NH), 8.47 (1H, s, H4), 8.16 (1H, t, H2', J=1.8 Hz), 8.09 (1H, ddd, H4', J=8.0, 1.8, 1.0 Hz), 7.60 (1H, dt, H6', J=8.0, 1.0 Hz), 7.62 (1H, t, H5['], J=8.0 Hz), 5.27 (1H, s, H5), 4.06–3.97 (2H, m, OCH₂), 2.43 (3H, s, CH₃), 1.08 (3H, t, CH₃); ¹³C NMR (DMSO-d₆) δ 167.9 (C8a), 167.1 (C4), 165.5 (COO), 156.0 (C2), 148.3 (C3'), 148.2 (C1'), 144.8 (C7), 134.5 (C2'),131.0 (C4'), 122.6 (C6'), 122.2 (C5'), 107.2 (C6), 106.7 (C4a), 60.6 (OCH₂), 40.0 (C5), 18.3 (CH₃), 14.2 (CH₃); MSEI *m*/*z* (%): 372 ([M-HCl]⁺, 12), 355 (34), 250 (100), 223 (40) 195 (25). Anal. Calcd for C₁₇H₁₇ClN₄O₄S (408.86): C: 49.94; H: 4.19; N: 13.70; S: 7.84. Found: C: 49.92; H: 4.18; N: 13.82; S: 7.88.

4.4.2. 6-Ethoxycarbonyl-5-(4-metoxycarbonylphenyl)-7-methyl-5,8-dihydro-2*H***-pyrido**[**3,2**-*e*][**1,3**]**thiazin-2-iminium chloride (8c).** Prepared from ethyl 6-chloro-4-(4'-methoxycarbonylphenyl)-5-formyl-2-methyl-1,4-dihydro-pyridine-3-carboxylate (**2c**). Yield 50%; mp 287–288 °C; IR (KBr) ν_{max} 3384 and 3250 (ν NH), 1728 and 1692 (ν C=O), 1647 (ν C=N) cm⁻¹; ¹H NMR (DMSO-*d*₆) δ : 12.72 (1H, s, NH), 10.63 (1H, s, NH), 10.34 (1H, s, NH), 8.41 (1H, s, H4), 7.89 (2H, d, *J*=7.5 Hz, H3', H5'), 7.42 (2H, d, *J*=7.5 Hz, H2', H4'), 5.15 (1H, s, H5), 4.05–3.95 (2H, m, OCH₂), 3.81 (3H, s, OCH₃), 2.42 (3H, s, CH₃), 1.06 (3H, t, CH₃); ¹³C NMR (DMSO-*d*₆) δ : 167.4 (C8a), 166.5 (C4), 165.9 (COO), 165.3 (COO), 155.6 (C2), 151.1 (C1'), 144.0 (C7), 129.9

(C3', C5'), 128.5 (C4'), 127.5 (C2', C6'), 107.6 (C6), 106.6 (C4a), 60.2 (OCH₂), 52.2 (OCH₃), 40.2 (C5), 17.9 (CH₃), 13.9 (CH₃); MSEI m/z (%): 385 ([M-HCI]⁺⁺, 16), 356 (29), 312 (25), 250 (100), 223 (37), 195 (25). Anal. Calcd for C₁₉H₂₀ClN₃O₄S (421.90): C: 54.09; H: 4.78; N: 9.96; S: 7.60. Found: C: 53.87; H:4.75; N: 10.07; S: 7.59.

4.4.3. 6-Methoxycarbonyl-7-methyl-5-phenyl-5,8-dihydro-2H-pyrido[3,2-e][1,3]thiazin-2-iminium chloride (8d). Prepared from methyl 6-chloro-4-phenyl-5-formyl-2methyl-1,4-dihydropyridine-3-carboxylate (2d). Yield, 73%; mp 241–243 °C; IR (KBr) v_{max} 3360 and 3220 (νNH) , 1690 $(\nu \text{C=O})$, 1645 $(\nu \text{C=N}) \text{ cm}^{-1}$; ¹H NMR (DMSO-d₆) δ: 12.61 (1H, s, NH), 10.57 (1H, s, NH), 10.32 (1H, s, NH), 8.45 (1H, s, H4), 7.35-7.22 (5H, m, Ph), 5.06 (1H, s, H5), 3.58 (3H, s, OCH₃); 2.43 (3H, s, CH₃); ¹³C NMR (DMSO-*d*₆) δ 167.3 (C8a), 166.4 (C4), 166.0 (COO), 155.3 (C2), 145.9 (C1[']), 143.5 (C7), 129.0 (C2['], C6[']), 127.3 (C4'), 126.7 (C3', C5'), 108.0 (C6), 107.4 (C4a), 51.5 (OCH₃), 39.7 (C5), 17.8 (CH₃); MSEI m/z (%): 313 ([M-HCl]⁺, 21); 298 (17), 236 (100), 209 (62), 177 (8). Anal. Calcd for $C_{16}H_{16}ClN_3O_2S$ (349.84): C: 54.93; H: 4.61; N: 12.01; S: 9.17. Found: C: 54.87; H: 4.65; N: 12.03; S: 9.19.

4.4.4. 5-(2-Chlorophenyl)-6-methoxycarbonyl-7-methyl-5,8-dihydro-2*H*-pyrido[3,2-*e*][1,3]thiazin-2-iminium chloride (8e). Prepared from methyl 6-chloro-4-(2-chlorophenyl)-5-formyl-2-methyl-1,4-dihydropyridine-3-carboxylate (2e). Yield 69%; mp 304–306 °C; IR (KBr) ν_{max} 3370 and 3225 (vNH), 1692 (vC=O), 1648 (vC=N) cm⁻¹; ¹H NMR (DMSO-*d*₆) δ: 12.68 (1H, s, NH), 10.58 (1H, s, NH), 10.35 (1H, s, NH), 8.15 (1H, s, H4), 7.44 (1H, m, H3'), 7.33-7.24 (3H, m, H5', H4', H6'), 5.50 (1H, s, H5), 3.51 (3H, s, OCH₃); 2.40 (3H, s, CH₃); ¹³C NMR (DMSO- d_6) δ 167.1 (C8a), 165.7 (COO), 165.3 (C4), 155.9 (C2), 144.3 (C7), 142.9 (C1'), 130.80 (C2'), 130.7 (C6'), 129.9 (C3'), 129.1 (C4'), 128.2 (C5'), 107.3 (C6), 106.6 (C4a), 51.4 (OCH₂), 37.3 (C5), 17.9 (CH₃); MSEI *m*/*z* (%): 347/349 ([M-HCl]⁺, 14/6); 332/334 (25/9), 288/290 (21/8), 236 (100), 209 (61), 177 (9). Anal. Calcd for C₁₆H₁₅Cl₂N₃O₂S (384.28): C: 50.01; H: 3.93; N: 10.93; S: 8.34. Found: C: 49.83; H: 3.92; N: 11.00; S: 8.32.

4.4.5. 6-Methoxycarbonyl-7-methyl-5-(3'-nitrophenyl)-5,8-dihydro-2*H*-pyrido[3,2-*e*][1,3]thiazin-2-iminium chloride (8f). Prepared from methyl 6-chloro-4-(3-nitrophenyl)-5-formyl-2-methyl-1,4-dihydropyridine-3-carboxylate (2f). Yield 58%; mp 250–252 °C; IR (KBr) ν_{max} 3390 and 3255 (ν NH), 1690 (ν C=O), 1645 (ν C=N), 1535 and 1350 (ν NO₂) cm⁻¹; ¹H NMR (DMSO- d_6) δ 12.70 (1H, s, NH), 10.64 (1H, s, NH), 10.38 (1H, s, NH), 8.52 (1H, s, H4), 8.13 (1H, t, H2', J = 1.8 Hz), 8.09 (1H, ddd, H4', J =7.9, 1.8, 1.0 Hz), 7.73 (1H, dt, H6', J=7.9, 1.0 Hz), 7.62 (1H, t, H5['], J=7.9 Hz), 5.28 (1H, s, H5), 3.57 (3H, m, OCH₃), 2.45 (3H, s, CH₃); ¹³C NMR (DMSO- d_6) δ 167.6 (C8a), 166.7 (C4), 165.8 (COO), 155.6 (C2), 148.1 (C3[']), 147.6 (C1'), 144.7 (C7), 133.9 (C2'), 130.6 (C4'), 122.3 (C6[']), 121.6 (C5[']), 106.9 (C6), 106.4 (C4a), 51.6 (OCH₃), 39.3 (C5), 18.0 (CH₃); MSEI *m*/*z* (%): 358 ([M-HCl]) 13); 341 (22), 236 (100), 209 (61), 177 (9). Anal. Calcd for C₁₆H₁₅ClN₄O₄S (394.83): C: 48.67; H: 3.83; N: 14.19; S: 8.12. Found: C: 48.56; H: 3.89; N: 14.22; S: 8.15.

4.4.6. 6-Methoxycarbonyl-5-(4-metoxycarbonylphenyl)-7-methyl-5,8-dihydro-2H-pyrido[3,2-e][1,3]thiazin-2iminium chloride (8g). Prepared from methyl 6-chloro-4-(4'-methoxycarbonylphenyl)-5-formyl-2-methyl-1,4-dihydropyridine-3-carboxylate (2g). Yield 52%; mp 242-244 °C; IR (KBr) ν_{max} 33890 and 3255 (ν NH), 1725 and 1690 (ν C=O), 1647 (ν C=N) cm⁻¹; ¹H NMR (DMSO- d_6) δ: 12.76 (1H, s, NH), 10.64 (1H, s, NH), 10.35 (1H, s, NH), 8.45 (1H, s, H4), 7.89 (2H, d, H3', H5', J=8.4 Hz), 7.42 (2H, d, H2['], H4['], J=8.4 Hz), 5.16 (1H, s, H5), 3.81 (3H, s, OCH₃), 3.56 (3H, s, OCH₃, overlaping with de H₂O signal), 2.42 (3H, s, CH₃); ¹³C NMR (DMSO-*d*₆) δ: 167.5 (C8a), 166.5 (C4), 165.9 (COO), 165.8 (COO), 155.5 (C2), 150.8 (C1'), 144.2 (C7), 129.9 (C3', C5'), 128.5 (C4'), 127.3 (C2', C6'), 107.2 (C6), 106.6 (C4a), 52.1 (OCH₃), 51.5 (OCH₃), 39.1 (C5), 17.9 (CH₃); MSEI *m*/*z* (%): 371 ([M-HCl]) 16); 356 (25), 236 (100), 209 (62), 177 (8). Anal. Calcd for $C_{18}H_{17}CIN_3O_4S$ (406.06): C: 53.14; H: 4.21; N: 10.33; S: 7.88. Found: C: 53.01; H: 4.17; N: 10.42; S: 7.89.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11. 054. The main geometrical characteristics of compounds **8b–g** obtained from B3LYP/6-31G*²⁴ ab initio are collected in the supplementary material.

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Photochemistry of a naphthalene–thymine dyad in the presence of acetone

Noureddine Belmadoui, María José Climent and Miguel A. Miranda*

Departamento de Química / Instituto de Tecnología Química UPV-CSIC, Universidad Politécnica de Valencia, Avda de los Naranjos s/n, 46022 Valencia, Spain

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Abstract—Irradiation of dyad 1 in aqueous acetone leads to the introduction of an acetonyl substituent at the naphthalene 5-position, to give photoproduct 2. The proposed reaction mechanism involves electron transfer from the naphthalene excited singlet state to the ketone. Neither thymine dimers, nor acetone photoadducts involving the thymine ring were detected. These photoproducts would arise from the thymine triplet excited state, which in dyad 1 must be efficiently depopulated via a fast intramolecular energy transfer to the naphthalene chromophore, due to the lower energy of its excited triplet state.

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1. Introduction

Exposure of living organisms to solar radiation may induce lethal mutagenic and carcinogenic effects as a result of photochemical modifications of DNA. The major lesions have been extensively characterized over the last decades and include pyrimidine cyclobutane type products, pyrimidine (6–4) pyrimidone adducts and Dewar valence isomers.¹

At wavelengths longer than 290 nm, where DNA is not absorbing, the bulk of photobiological effects is mediated by photosensitizers.² For example, upon irradiation of thymine³ or thymidine⁴ in aqueous acetone, the photosensitized formation of up to six diastereomeric cyclobutane dimers (together with two acetone adducts) has been observed.

On the other hand, the nonsteroidal anti-inflammatory 2-arylpropionic acids are known to photosensitize DNA damage.^{5–8} Specifically, naproxen (NAP, 6-methoxy- α -methyl-2-naphthaleneacetic acid) photoinduces single strand breaks;^{9,10} however, NAP is not able to mediate pyrimidine photodimerization in DNA.¹¹ Hence, modified nucleosides such as **1** (*S*-naproxen, 5'-ester with thymidine) containing key substructures present in drugs and nucleic acids can be relevant models to study the excited state interactions and the primary photophysical/photochemical processes underlying drug-mediated photoreactions of DNA and their photobiological consequences.¹²

The purpose of the present work was to investigate the photolysis of 1 in the presence of acetone. If this compound would follow the same behavior as thymine and thymidine, the corresponding dimerization products would be obtained, which would be potentially useful models for the study of cycloreversion via intramolecular electron transfer (related to DNA repair). However, the presence of the naphthalene chromophore could have a strong influence on the results, leading to diverging photochemistry and photophysics. As the excited states of NAP (singlet and triplet) are lower in energy than the corresponding states of the thymine unit, population of the former via intramolecular energy transfer could be envisaged. As a consequence, in the case of 1 the NAP chromophore could be dominating the photophysical and photochemical properties of this compound. Actually, it will be shown that naphthalene is indeed involved in the major photoreaction of 1, which occurs via electron transfer from the excited naphthalene-derived singlet state to ground state acetone. As the final result, an acetonyl substituent becomes attached to the naphthalene ring.

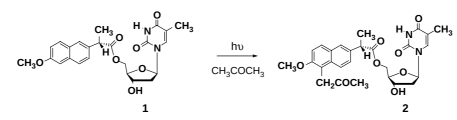
2. Results and discussion

2.1. Product studies

Steady-state photolysis of oxygen free solutions of 1 in acetone–water gave rise to its 5-acetonyl derivative (2) (23%) as the only product (Scheme 1). Its structure was assigned in a first stage on the basis of the ¹H and ¹³C NMR spectra.

Keywords: DNA; Nucleosides; Naproxen; Photoproducts; Acetone.

^{*} Corresponding author. Tel.: +34963877807; fax: +34963877809; e-mail: mmiranda@qim.upv.es



Scheme 1.

By contrast with the reported acetone photoaddition product of thymidine,⁴ where the acetonyl group becomes attached to position 5 of thymine ring with concomitant saturation of its carbon-carbon double bond, the pyrimidine moiety remained intact in the case of 2. This is supported by the presence of the typical signals at $\delta = 1.98$ and 7.15 ppm, safely assigned to the thymine methyl and olefinic proton, respectively. The fact that all the protons and carbons of the sugar were found in the NMR spectra of 2, following a pattern very similar to that observed in the starting material 1, pointed to the naphthalene ring as the possible reactive site. Actually, only five aromatic protons (instead of 6) were clearly distinguishable in the low field region $(7.8 > \delta >$ 7.2 ppm). Their multiplicities, chemical shifts and coupling constants were consistent with substitution at position 5. Moreover, a significant NOE enhancement was observed for the signal attributed to methylene protons of the acetonyl substituent (at $\delta = 4.23$ and 4.09 ppm) upon irradiation of the NAP methoxy group.

In order to confirm the structure assignment, irradiation of the closely related 6-methoxy- α -methyl-2-naphthaleneacetic acid methyl ester (3) was carried out under the same reaction conditions. As a result, the corresponding 5-acetonyl derivative was formed in low yield (<10%). In order to obtain the photoproduct in higher amounts, an analogous study was undertaken in the presence of chloroacetone, using acetonitrile as solvent, where compound 3 reacted much faster. Analysis of the photomixture revealed the presence of five photoproducts (4–8) that were isolated by semipreparative HPLC.

A comparison of the aromatic region in the ¹H NMR spectra of compounds 1-8 (see Fig. 1 and Supporting information) allowed us to confirm that the photoreaction occurred mainly at the naphthalene 5-position (to give the acetonyl or the chloro derivative), although detectable amounts of 4-monosubstituted, as well as 4,5 and 5,8 disubstituted products were also formed.

2.2. Reaction mechanism

A plausible mechanism to explain the obtained products is depicted in Scheme 2.

The key step would be photo-induced electron transfer from the naphthalene-like chromophore to (chloro)acetone. The different products would derive from the resulting radical cation, upon attack by the enol of acetone, chloride anion or the acetonyl radical (in the case of chloroacetone) within the solvent cage. All the other steps are rather obvious and do not deserve more detailed comments.

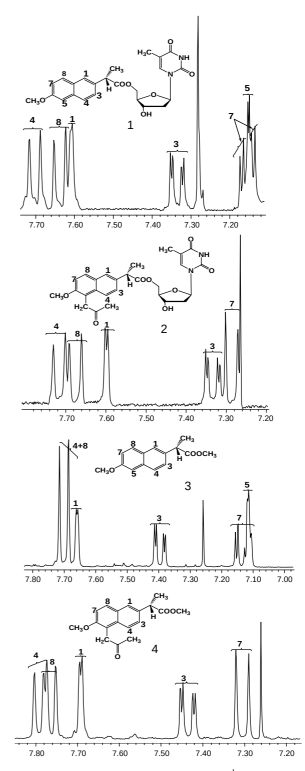
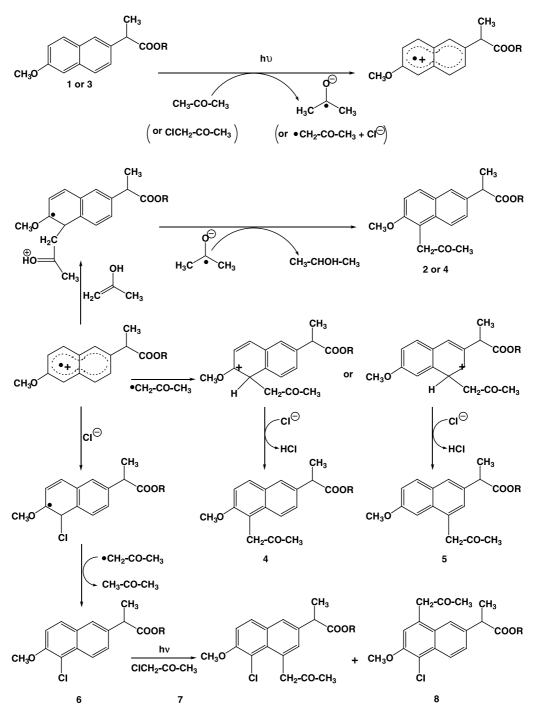


Figure 1. Comparison of the aromatic region in the ¹H NMR spectra of compounds 1–4.



Scheme 2.

Experimental evidence in favour of the proposed mechanism was obtained by means of steady-state and time-resolved fluorescence measurements. Upon excitation at 320 nm in acetonitrile, the typical emission of compound **1**, centered at 350 nm, was quenched upon addition of increasing amounts of aqueous acetone. The spectra are shown in Figure 2a.

The rate constant for this process was obtained from the Stern–Volmer plot (I_0/I vs acetone concentration) and found to be 2.7×10^{10} M⁻¹ s⁻¹. This quenching was dynamic in nature, as a similar rate constant (2.8×10^{10} M⁻¹ s⁻¹) was determined using lifetime measurements (τ_0/τ). Figure 2b

clearly shows the coincident trends of the experimental points obtained following both approaches.

According to the Weller equation $(Eq. 1)^{13}$ electron transfer from the NAP singlet excited state $(E^*=85-86 \text{ kcal/mol})^{14}$ to ground state acetone would be nearly thermoneutral.

$$\Delta G_{\rm ET} \text{ (kcal/mol)} = 23.06[E_{\rm D+\cdot/D} - E_{\rm A/A^{--}} + 2.6/\varepsilon - 0.13] -E^*(S_1 \text{ or } T_1)$$
(1)

Using the known redox potentials of NAP $(E_{D+,D}=1.4 \text{ V vs} \text{ SCE})^{15}$ and acetone $(E_{A/A^-}=-2.4 \text{ V vs SCE})^{16}$ the free energy change (ΔG) associated with this process in

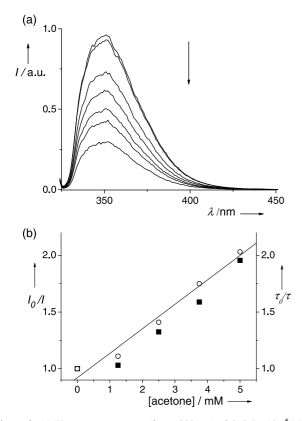


Figure 2. (a) Fluorescence spectra $(\lambda_{exc} = 320 \text{ nm})$ of $\mathbf{1}, 2.5 \times 10^{-5} \text{ M}$ in acetonitrile in the presence of increasing amounts of aqueous acetone $(0-1.25 \times 10^{-2} \text{ M})$. (b) Stern–Volmer plot for quenching of the fluorescence of $\mathbf{1}$ by aqueous acetone, based on emission intensities (\blacksquare) and lifetimes (\bigcirc).

acetonitrile (dielectric constant, $\varepsilon = 37$) is between +0.2 and +1.2 kcal/mol. Actually, no significant fluorescence quenching was observed when increasing amounts of acetone were added to an acetonitrile solution of **1** (see Fig. 3).

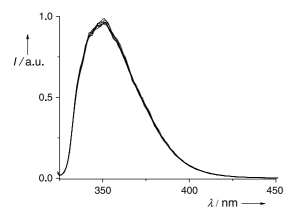


Figure 3. Fluorescence spectra (λ_{exc} =320 nm) of 1, 2.5×10⁻⁵ M in acetonitrile upon addition of dry acetone at different concentrations (0–7.50×10⁻² M).

Obviously, addition of water ($\varepsilon = 80$) to the reaction mixture should enhance charge separation, leading to negative free energy changes (ΔG between -0.7 and +0.3 kcal/mol). Also the presence of water would favor proton transferassisted electron transfer, which is well documented in the literature. This justifies the differences found between Figures 2a and 3.

On the other hand, with chloroacetone $(E_{A/A^{-}} = -1.4 \text{ V vs} \text{SCE})^{16}$ as electron acceptor instead of acetone, the reaction would be clearly exergonic ($\Delta G = -21.3 \text{ kcal/mol}$) even in non-aqueous medium. As a matter of fact, the fluorescence of **1** was efficiently quenched by chloroacetone in acetonitrile, with a rate constant $k_q = 3.1 \times 10^{11} \text{ M}^{-1} \text{ s}^{-1}$ (see Fig. 4 and inset).

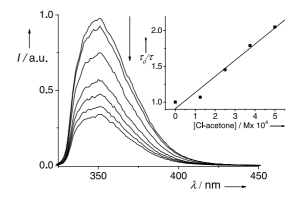


Figure 4. Fluorescence spectra of compound 1, 2.5×10^{-5} M in acetonitrile (λ_{exc} =266 nm), in the presence of increasing amounts of chloroacetone (0–7.50×10⁻⁴ M). Inset: Stern–Volmer plot for quenching of the fluorescence of 1 by chloroacetone, based on emission lifetimes.

At this stage, it is interesting to compare the photochemistry of **1** in aqueous acetone with that observed for thymidine under similar conditions. The reported photoproducts of thymidine were six cyclobutane dimers plus minor amounts of a C5–C6 hydrogenated derivative and two diastereomeric adducts with and acetonyl substituent attached to C5.⁴

The dimers would arise from triplet thymidine generated after triplet–triplet energy transfer from acetone. This would be an exergonic process in view of the relative excited state energies (acetone: $E_{\rm T}$ =78 kcal/mol,¹⁷ thymidine: $E_{\rm T}$ =74 kcal/mol¹⁸). The hydrogenated derivative and the acetone adduct would also be formed from the thymidine triplet via abstraction of the α -carbonyl hydrogen atom followed by well established radical reactions.¹⁹

In the case of 1, none of these reactions should occur, as triplet thymidine would be efficiently deactivated by NAP, due to the lower energy of the triplet state $(E_T=62 \text{ kcal/mol})$.¹⁴ Conversely, electron transfer from the NAP excited singlet state to acetone is feasible, as discussed above. This has been evidenced by the fluorescence quenching experiments shown in Figure 2a and b.

3. Conclusions

The photobehaviour of dyad 1 is dominated by the naphthalene-derived chromophore. Upon excitation, the typical naphthalene emission from its excited singlet state was observed at 350 nm. This is justified on the basis of the lower energy of the NAP excited state, as compared with

that of the thymine chromophore. In the presence of acetone, the NAP moiety was also found to be the photoreactive site; a 5-acetonyl derivative (2) was obtained as the only photoproduct. Neither thymidine dimers, nor acetone photoadducts involving the thymine ring were detected. These photoproducts would arise from the thymine triplet excited state, which in dyad 1 should be efficiently depopulated via a fast intramolecular energy transfer to the naphthalene chromophore.

4. Experimental

4.1. General

¹H and ¹³C NMR spectra were measured in CDCl₃ with a Varian Gemini 300 MHz instrument; chemical shifts are reported in δ (ppm) with TMS as internal standard.

Fluorescence measurements. Concentration was fixed by adjusting the absorbance of the solutions at a value lower than 0.2 at the excitation wavelength (320 nm). The steady-state fluorescence was obtained with a FS900 Edinburgh Analytical Instruments apparatus, equipped with a 450 W xenon lamp. The time-resolved fluorescence determinations were performed with a FL900 Edinburgh Analytical Instrument apparatus using a hydrogen lamp (1.0 ns pulse width) as the excitation source. The samples were placed into quartz cells of 1 cm pathlength, and deoxygenation was made by bubbling nitrogen.

4.2. Photolysis of S-naproxen, 5'-ester with thymidine (1)

A solution of 1^{12} (130 mg, 0.28 mmol) in acetone–water (30/70) was placed in a Pyrex test-tube sealed and purged for 30 min with argon. The solution was irradiated with a 125 W medium pressure Hg lamp for 41 h. The solvent was removed by rotary evaporation and the resulting mixture was purified by reverse phase HPLC on the RP-18 column, using water–acetonitrile (40/60) as the isocratic solvent, to give 2 (30 mg, 23%).

4.2.1. 5-Acetonyl-6-methoxy- α -methyl-2-naphthaleneacetic acid. 5'-ester with thymidine (2). ¹H NMR $(300 \text{ MHz}, \text{ CDCl}_3): \delta = 8.72 \text{ (br s, 1H, NH)}, 7.72$ (d, J=8.7 Hz, 1H, ArH-4), 7.67 (d, J=9.3 Hz, 1H, ArH-8), 7.60 (d, J=2.1 Hz, 1H, ArH-1), 7.33 (dd, J=8.7, 2.1 Hz, 1H, ArH-3), 7.31 (d, J=9.3 Hz, 1H, ArH-7), 7.15 (br s, 1H, H-6), 6.18 (dd, J=5.7, 7.8 Hz, 1H, H-1'), 4.49 (dd, J=12.1, 3.3 Hz, 1H, H-5'), 4.23 (d, J=16.2 Hz, 1H, H-5')CH₂CO), 4.14 (dd, J = 12.1, 3.0 Hz, 1H, H-5[']), 4.09 (d, J =16.2 Hz, 1H, CH₂CO), 4.04 (m, 1H, H-4'), 3.94 (s, 3H, OCH₃), 3.91 (m, 1H, H-3'), 3.83 (q, J=7.2 Hz, 1H, $CHCH_3$), 2.17 (s, 3H, COCH₃), 1.98 (d, J=1.2 Hz, 3H, CH=C-CH₃), 1.84 (ddd, J=13.6, 5.4, 2.1 Hz, 1H, H_β-2'), 1.60 (d, J=7.2 Hz, 3H, CH–CH₃), 1.13 (ddd, J=13.6, 7.8, 6.3 Hz, 1H, H_{α}-2'). ¹³C NMR (75 MHz, CDCl₃): δ =207.7 (CO), 174.1 (COO), 163.7 (C-4), 154.8 (C-Ar6), 150.2 (C), 135.2 (C), 135.1 (CH-6), 132.5 (C), 128.8 (CH-Ar8), 128.6 (C), 126.7 (CH-Ar1), 125.9 (CH-Ar3), 124.0 (CH-Ar4), 116.0 (C-Ar5), 113.6 (CH-Ar7), 110.7 (C-5), 84.9 (CH-1'), 84.4 (CH-4'), 71.6 (CH-3'), 64.2 (CH₂-5'), 56.3 (OCH₃), 45.6 (CH-CH₂), 40.5 (CH₂), 39.9 (CH₂), 29.3 (CH₃-CO),

18.2 (CH–*C*H₃), 12.8 (CH=C–*C*H₃). FAB m/z: calcd for C₂₇H₃₀N₂O₈ 510.2002, found 510.1994.

4.3. Irradiation of S-naproxen, methyl ester (3)

Chloroacetone (1.85 g, 20 mmol) was added to a solution of 6-methoxy- α -methyl-2-naphthaleneacetic acid methyl ester (1 mmol) in anhydrous acetonitrile. The resulting solution was placed into tubes surrounding a centrally positioned quartz cooling jacket containing a 125 W medium pressure Hg lamp. Prior to irradiation the solution was degassed for 30 min with a stream of argon, and then it was irradiated for 12 h, giving over 88% conversion of the starting material. The solvent was removed under reduced pressure and the residue was separated by HPLC (with an UV/vis absorption detector set at 280 nm), using normal phase silica column, hexane–ethyl acetate (70/30) as the isocratic eluent and a flow rate of 2.0 mL/min.

4.3.1. 5-Acetonyl-6-methoxy-α-methyl-2-naphthaleneacetic acid methyl ester. (Compound 4, yield 17%): ¹H NMR (300 MHz, CDCl₃): δ =7.78 (d, *J*=8.7 Hz, 1H, ArH-4), 7.77 (d, *J*=9.0 Hz, 1H, ArH-8), 7.69 (d, *J*= 1.8 Hz, 1H, ArH-1), 7.43 (dd, *J*=8.7, 1.8 Hz, 1H, ArH-3), 7.30 (d, *J*=9.0 Hz, 1H, ArH-7), 4.13 (s, 2H, CH₂CO), 3.96 (s, 3H, OCH₃), 3.88 (q, *J*=7.2 Hz, 1H, CHCH₃), 3.66 (s, 3H, COOCH₃), 2.10 (s, 3H, COCH₃), 1.57 (d, *J*=7.2 Hz, 3H, CH-CH₃). ¹³C NMR (75 MHz, CDCl₃): δ =207.5 (CO), 174.0 (COO), 154.6 (C-Ar6), 135.6 (C), 132.5 (C), 129.1 (CH-Ar8), 129.0 (C), 126.9 (CH-Ar1), 126.6 (CH-Ar3), 123.5 (CH-Ar4), 116.2 (C-Ar5), 113.3 (CH-Ar7), 56.4 (OCH₃), 52.0 (OCH₃), 45.2 (CH-CH₃), 40.9 (CH₂), 28.8 (CH₃-CO), 18.4 (CH-CH₃). EI-HRMS *m/z*: calcd for C₁₈H₂₀O₄ 300.1361, found 300.1359.

4.3.2. 4-Acetonyl-6-methoxy-α-methyl-2-naphthaleneacetic acid methyl ester. (Compound **5**, yield 15%): ¹H NMR (300 MHz, CDCl₃): δ = 7.73 (d, *J* = 8.7 Hz, 1H, ArH-8), 7.64 (d, *J* = 1.8 Hz, 1H, ArH-1), 7.35 (d, *J* = 1.8 Hz, 1H, ArH-3), 7.16 (dd, *J* = 8.7, 2.7 Hz, 1H, ArH-7), 7.12 (d, *J* = 2.7 Hz, 1H, ArH-5), 4.04 (s, 2H, CH₂CO), 3.90 (s, 3H, OCH₃), 3.84 (q, *J* = 7.2 Hz, 1H, CHCH₃), 3.67 (s, 3H, COOCH₃), 2.08 (s, 3H, COCH₃), 1.59 (d, *J* = 7.2 Hz, 3H, CH-*CH*₃). ¹³C NMR (75 MHz, CDCl₃): δ = 206.9 (CO), 175.0 (COO), 158.2 (C-Ar6), 135.3 (C), 132.5 (C), 130.3 (C), 130.1 (CH-Ar8), 129.3 (C), 128.6 (CH-Ar3), 125.9 (CH-Ar1), 118.8 (CH-Ar7), 102.4 (CH-Ar5), 55.3 (OCH₃), 52.1 (OCH₃), 49.9 (CH₂), 45.2 (*C*H–CH₃), 28.7 (*C*H₃–CO), 18.5 (CH–*C*H₃). EI-HRMS *m*/*z*: calcd for C₁₈H₂₀O₄ 300.1361, found 300.1349.

4.3.3. 5-Chloro-6-methoxy- α -methyl-2-naphthaleneacetic acid methyl ester. (Compound 6, yield 22%). This compound was previously reported in the literature.²⁰ ¹H NMR (300 MHz, CDCl₃): δ =8.19 (d, *J*=8.7 Hz, 1H, ArH-4), 7.76 (d, *J*=9 Hz, 1H, ArH-8), 7.71 (d, *J*=1.5 Hz, 1H, ArH-1), 7.54 (dd, *J*=8.7, 1.5 Hz, 1H, ArH-3), 7.32 (d, *J*=9 Hz, 1H, ArH-7), 4.05 (s, 3H, OCH₃), 3.90 (q, *J*=7.2 Hz, 1H, *CH*-CH₃), 3.69 (s, 3H, COOCH₃), 1.60 (d, *J*=7.2 Hz, 3H, CH-CH₃). ¹³C NMR (75 MHz, CDCl₃): δ =175.0 (COO), 153.0 (C-Ar6), 136.4 (C), 131.0 (C-Ar5),

129.5 (C), 127.8 (CH-Ar8), 127.4 (CH-Ar3), 126.1 (CH-Ar1), 124.0 (CH-Ar4), 113.9 (CH-Ar7), 56.9 (OCH₃), 52.1 (OCH₃), 45.1 (CH-CH₃), 29.7 (CH₃-CO), 18.5 (CH-CH₃).

4.3.4. 4-Acetonyl-5-chloro-6-methoxy-α-methyl-2naphthaleneacetic acid methyl ester. (Compound 7, yield 19%): ¹H NMR (300 MHz, CDCl₃): $\delta = 7.78$ (d, J=9.0 Hz, 1H, ArH-8), 7.64 (d, J=1.8 Hz, 1H, ArH-1), 7.27 (d, J=9.0. Hz, 1H, ArH-7), 7.18 (d, J=1.8 Hz, 1H, ArH-3), 4.55 (AB, 2H, CH₂CO), 3.98 (s, 3H, OCH₃), 3.80 (q, J=7.2 Hz, 1H, CHCH₃), 3.67 (s, 3H, COOCH₃), 2.29 (s, ¹³C NMR 3H, COCH₃), 1.56 (d, *J*=7.2 Hz, 3H, CH–CH₃). (75 MHz, CDCl₃): $\delta = 206.1$ (CO), 174.7 (COO), 153.7 (C-Ar6), 135.5 (C), 133.2 (CH-Ar3), 131.3 (C-Ar5), 130.7 (C), 129.9 (C), 129.5 (CH-Ar8), 128.7 (C-Ar4), 127.2 (CH-Ar1), 113.5 (CH-Ar7), 57.0 (OCH₃), 52.7 (CH₂), 52.1 (OCH₃), 44.7 (CH–CH₃), 29.7 (CH₃–CO), 18.2 (CH–CH₃). EI-HRMS m/z: calcd for C₁₈H₁₉ClO₄ 334.0971, found 334.0970.

8-Acetonyl-5-chloro-6-methoxy-α-methyl-2-4.3.5. naphthaleneacetic acid methyl ester. (Compound 8, yield 8%): ¹H NMR (300 MHz, CDCl₃): $\delta = 8.24$ (d, J=8.7 Hz, 1H, ArH-4), 7.67 (d, J=1.8 Hz, 1H, ArH-1), 7.55 (dd, J=8.7, 1.8 Hz, 1H, ArH-3), 7.20 (s, 1H, ArH-7), 4.12 (br s, 2H, CH₂CO), 4.02 (s, 3H, OCH₃), 3.88 (q, J=7.2 Hz, 1H, CHCH₃), 3.67 (s, 3H, COOCH₃), 2.18 (s, 3H, COCH₃), 1.57 (d, J=7.2 Hz, 3H, CH-CH₃). ¹³C NMR (75 MHz, CDCl₃): $\delta = 205.7$ (CO), 174.7 (COO), 151.9 (C-Ar6), 137.1 (C), 131.5 (C), 131.2 (C), 128.3 (C-Ar5), 127.3 (CH-Ar3), 124.9 (CH-Ar4), 122.5 (CH-Ar1), 116.5 (CH-Ar7), 57.1 (OCH₃), 52.1 (OCH₃), 48.7 (CH₂), 45.4 (CH-CH₃), 29.0 (CH₃-CO), 18.6 (CH-CH₃). EI-HRMS m/z: calcd for C₁₈H₁₉ClO₄ 334.0971, found 334.0975.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11.035.

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Tetrahedron

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Total synthesis of FR901464: second generation

Hajime Motoyoshi, Masato Horigome, Hidenori Watanabe and Takeshi Kitahara*

Department of Applied Biological Chemistry, Graduate School of Agricultural and Life Sciences, The University of Tokyo, 1-1-1 Yayoi, Bunkyo-ku, Tokyo 113-8657, Japan

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Abstract—FR901464, a potent cell cycle inhibitor, was synthesized in a convergent manner using natural chiral pool, L-threonine, ethyl (*S*)-lactate and 2-deoxy-D-glucose as starting materials. © 2005 Published by Elsevier Ltd.

1. Introduction

FR901464 (1), which has two highly functionalized tetrahydropyran rings linked by a diene chain (Fig. 1), was isolated from the culture broth of bacterium *Pseudomonas* sp. no. 2663.¹ This compound shows transcriptional regulating activity and induces characteristic G1 and G2/M phase arrest in the cell cycle. Related to these activities, it also exhibits a potent antitumor effect. The absolute configuration of 1, which was initially ambiguous, was elucidated by spectroscopic and chemical analysis.² Its unique structure as well as the significant biological activities prompted us to undertake the synthesis of this compound. We have previously communicated a synthesis of 1,³ and also the synthesis of biotinylated FR901464.⁴ In this paper, we wish to describe a new, improved synthesis of 1.

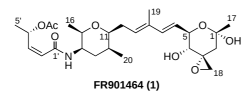


Figure 1.

Our basic synthetic strategy is illustrated in Scheme 1. From a synthetic perspective, we sought an efficient and convergent approach to our target, and decided upon disconnections at the diene and amide bonds generating segments **A**, **B** and **C** (or **C'**). Recently, Jacobsen et al. reported a total synthesis of **1** in which every chiral building

e-mail: kitahara-t@kitasato.or.jp

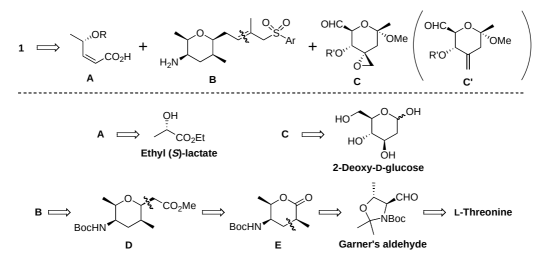
block was prepared using asymmetric catalysts.⁵ In contrast, we decided to synthesize all of the segments taking full advantage of materials from the chiral pool.³ Segment **A** would be prepared from commercially available ethyl (*S*)-lactate, and segment **C** would be synthesized from 2-deoxy-D-glucose. Segment **B**, in which all the substituents on the tetrahydropyran ring are cis-oriented, would be synthesized by stepwise carbon chain elongation via **D** and **E** starting from Garner's aldehyde⁶ derived from L-threonine.

In our first generation synthesis,³ we encountered difficulties at the assembly stage (Scheme 2). For example, (a) lower yield in Julia olefination (33%), (b) recyclingdemanding acetylation due to impossibility of either selective removal of the TBDPS group from 4 or selective acetylation of the corresponding diol and, most importantly, (c) poor regioselectivity in epoxidation of 5. Initially, we planned to construct the epoxide moiety on the right tetrahydropyran ring in the end of the synthesis because it was thought to be highly sensitive. However, concomitant epoxidation of the conjugated diene occurred in every reaction condition (mCPBA, DMDO, Sharpless' conditions,⁷ etc.) to give 7 and/or diepoxy compounds, and consequently lowered chemical yields of $6 (\sim 23\%)$. Similar results were also obtained by Jacobsen et al.,⁵ and we concluded that this side-reaction came from extraordinarily high reactivity of the diene and was essentially inevitable. In the second generation synthesis, therefore, we decided to install epoxide moiety in the segment C prior to coupling reaction. Other key modifications included (a) the use of a modified Julia olefination⁸ employing benzothiazolyl sulfone, and (b) replacing TBDPS group on the segment A with TBS group so as to allow removal of one of the silyl groups selectively and thus avoid a complicated acetylation-deacetylation sequence to obtain the monoacetate derivative.

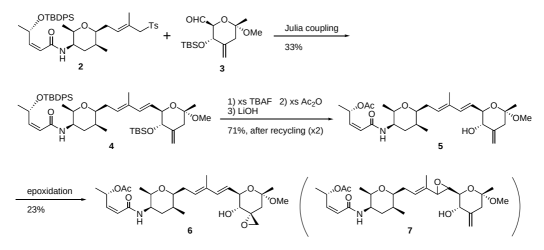
Keywords: FR901464; Asymmetric synthesis; Coupling reactions; Cell cycle inhibitor.

^{*} Corresponding author. Fax: +81 3 5841 8019;

^{0040–4020/\$ -} see front matter @ 2005 Published by Elsevier Ltd. doi:10.1016/j.tet.2005.11.031



Scheme 1. Synthetic plan for FR901464.



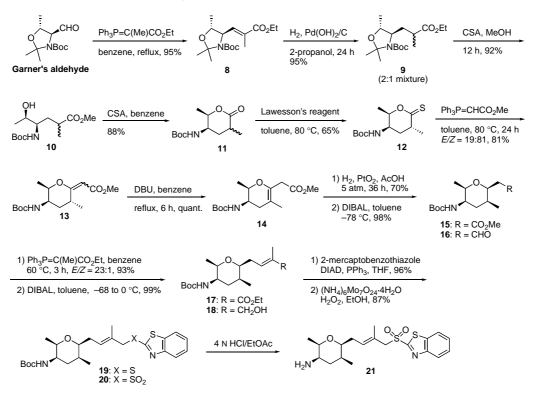
Scheme 2. Problems in our first generation synthesis.

The synthesis of segment **B** is shown in Scheme 3. Garner's aldehyde⁶ was subjected to a Wittig reaction followed by hydrogenation to give an inseparable 2:1 mixture of diastereoisomers. In this hydrogenation step, high stereoselectivity as described in the literature⁹ was not observed. This mixture was converted under general acidic conditions into lactone 11, which was then treated with Lawesson's reagent¹⁰ affording thionolactone **12**. Although complete epimerization of the stereocenter α to the thiocarbonyl was observed, this was of no consequence (vide infra). In order to install the other carbon chain, we attempted a Wittig reaction on thionolactone 12, and fortunately, the desired ester 13 was obtained in high yield. To the best of our knowledge, this is the first example of a successful Wittig reaction on a thionolactone.¹¹ Deconjugation of α , β unsaturated ester 13 was performed with DBU to furnish dihydropyran 14. The double bond of 14 was stereoselectively hydrogenated with platinum oxide to give the desired all-cis isomer 15 (70%) along with unassigned polar byproducts. No other stereoisomers were detected and the stereochemistry of 15 was confirmed by an NOE experiment (see Section 2). The ester moiety of 15 was reduced to an aldehyde with DIBAL and a three-carbon chain was added by the Wittig reaction. The resultant ethyl ester 17 was then

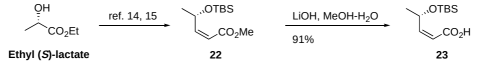
reduced to alcohol **18**. This allylic alcohol **18** was converted into benzothiazolyl sulfide **19** by Mitsunobu's method,¹² and then mildly oxidized with molybdenum catalyst¹³ to afford sulfone **20** in excellent yield. Acid treatment of **20** liberated the amino group to give **21**, which was used in the next reaction without purification.

The synthesis of segment **A** is displayed in Scheme 4. The known ester **22**,^{14,15} derived from ethyl (*S*)-lactate via the (*Z*)-selective Still–Gennari modification of the Horner–Emmons reaction,¹⁵ was hydrolyzed to furnish α , β -unsaturated carboxylic acid **23**.

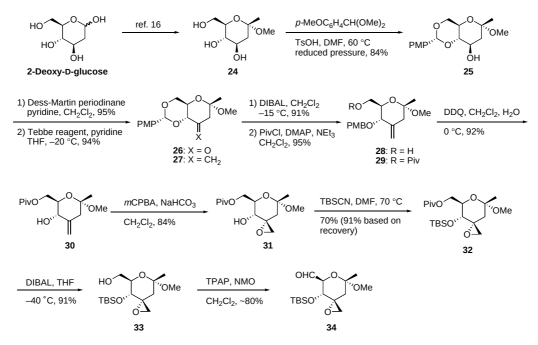
To synthesize segment C, 2-deoxy-D-glucose was first transformed into known triol 24^{16} in five steps (Scheme 5). Protection of the 1,3-diol as a *p*-methoxybenzylidene acetal furnished alcohol 25.¹⁷ Dess–Martin oxidation¹⁸ followed by Tebbe methylenation¹⁹ gave *exo*-olefin 27. Cleavage of the *p*-methoxybenzylidene acetal with DIBAL generated a primary alcohol 28, which in turn was protected as its pivalate ester. After removal of PMB group with DDQ,²⁰ hydroxyl group-directed epoxidation of 30 with *m*CPBA proceeded stereoselectively to give the desired epoxide 31. Silylation of 31 under usual conditions (TBSOTf or TBSCI



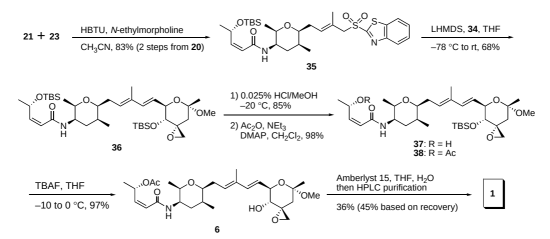
Scheme 3. Synthesis of segment B.



Scheme 4. Synthesis of segment A.



Scheme 5. Synthesis of segment C.



Scheme 6. Completion of the synthesis of FR901464.

in the presence of bases) gave only low yields, probably because the Lewis acidity of these reagents was incompatible with the labile epoxide of **31**. On the other hand, *t*-butyldimethylsilyl cyanide (TBSCN) was found to be a mild and effective silylating agent²¹ and afforded satisfactory yield (91% based on recovered **31**). Removal of pivaloyl group was successfully achieved with DIBAL in tetrahydrofuran as a solvent. When this reaction was executed in dichloromethane, reductive opening of the epoxide was observed even when stoichiometric amounts of DIBAL were used. We postulate that tetrahydrofuran solvated DIBAL and attenuated its reactivity to some degree. Finally, primary alcohol **33** was oxidized to aldehyde **34** by TPAP–NMO²² in good yield.

With these segments in hand, we set about the final coupling reactions (Scheme 6). Sulfone 35 was obtained from amine 21 and carboxylic acid 23 using HBTU.²³ Modified Julia olefination⁸ with sulfone 35 and epoxyaldehyde 34proceeded smoothly in a highly E-selective manner to furnish the desired diene **36** in 68% yield,²⁴ and with no observable side-reactions at the epoxide. Selective desilylation of 36 was achieved with a highly dilute solution (0.025%) of hydrochloric acid in methanol without any obvious damage to the epoxide moiety. The liberated hydroxyl group was acetylated to furnish 38 under standard conditions, and then the remaining TBS group was cleanly removed with TBAF. Interestingly, this methylated FR901464 (6) exhibited more powerful biological activity than that of natural 1, which was thought to come from its greater stability of methyl acetal in contrast to the labile hemiacetal of $1.^{3,4}$ The final hydrolysis of the methyl acetal moiety proved to be a most difficult problem because of extremely high sensitivity of **1** to acidic conditions.⁴ However, after substantial investigation, we found that Amberlyst 15 in wet THF gave the target molecule 1 as a single isomer, albeit in moderate yield. As a result of this revised synthetic route, we were able to realize an efficient and straightforward synthesis of FR901464.

In conclusion, we have accomplished a second generation total synthesis of FR901464 using materials derived from the natural chiral pool. Problems encountered in our first generation synthesis were solved by several revisions to the synthetic strategy. We have also synthesized biotinlabeled FR901464⁴ utilizing information gained from these synthetic studies. With increasing attention focused on this compound owing to its remarkable biological activity, our work may offer access to materials of value in these related fields.

2. Experimental

2.1. General

All of the solvents for non-aqueous reactions were dried before use. Benzene, toluene and THF were distilled from sodium benzophenone ketyl. Dichloromethane was distilled from phosphorus pentoxide. Acetonitrile and DMF were dried over molecular sieves (4 Å). Methanol was dried over molecular sieves (3 Å). Optical rotations were recorded with a JASCO DIP-1000 polarimeter. IR spectra were measured with a JASCO FT/IR-230 spectrophotometer. ¹H and ¹³C NMR were recorded on JEOL JNM AL300, Bruker AC-300, or JEOL JNM-A500. Mass spectra were recorded on JEOL JMS-700T. Column chromatography was performed using Merck silica gel 60 (0.060-0.200 mm). TLC was carried out on Merck glass plates precoated with silica gel 60 F₂₅₄ (0.25 mm). HPLC was performed using SHOWA DENKO shodex DS-4 equipped with Senshu Pak PEGASIL ODS ($4.6\phi \times 250$ mm), Senshu Pak PEGASIL ODS ($20\phi \times 250$ mm), Senshu Pak Silica-1251-N ($4.6\phi \times$ 250 mm) or Senshu Pak silica-5251-N ($20\phi \times 250$ mm), respectively.

2.1.1. *t*-Butyl (1'*E*,4*R*,5*R*)-4-(3'-ethoxy-2'-methyl-3'-oxoprop-1'-en-1'-yl)-2,2,5-trimethyl-1,3-oxazolidine-3-carboxylate (8). To a solution of Garner's aldehyde derived from L-threonine (50 mg, 0.206 mmol) in benzene (2 mL) was added (carbethoxyethylidene)triphenylphosphorane (82 mg, 0.226 mmol) and the mixture was stirred for 30 min at room temperature and then heated under reflux for 4 h. The solvent was removed under reduced pressure. The residue was purified by silica gel column chromatography (9:1 hexanes/EtOAc) to provide **8** as a colorless oil (64 mg, 95%); $n_D^{27} = 1.4602$; $[\alpha]_D^{25} + 17.7$ (*c* 0.45, CHCl₃); IR (film): ν_{max} 1703, 1658 cm⁻¹; ¹H NMR (300 MHz,

CDCl₃): δ (ppm) 1.27 (3H, d, J=6.0 Hz, 5-CH₃), 1.29 (3H, t, J=7.1 Hz, CH₃ of –CO₂Et), 1.35 (9H, s, 'Bu), 1.55, 1.61 (6H, two s, 2-CH₃), 1.89 (3H, s, 2'-CH₃), 3.86 (1H, dq, J=6.0, 8.3 Hz, 5-H), 4.0–4.15 (1H, m, 4-H), 4.20 (2H, q, J=7.1 Hz, CH₂ of –CO₂Et), 6.48 (1H, d, J=8.4 Hz, 1'-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 12.4, 14.0, 17.1, 25.0, 25.9, 27.9, 60.3, 61.8, 74.2, 79.4, 94.1, 128.6, 140.1, 151.4, 167.1. Anal. Calcd for C₁₇H₂₉NO₅: C, 62.36; H, 8.93; N, 4.28. Found: C, 62.39; H, 8.93; N, 4.48.

2.1.2. t-Butyl (2'RS,4R,5R)-4-(3'-ethoxy-2'-methyl-3'oxopropyl)-2,2,5-trimethyl-1,3-oxazolidine-3-carboxylate (9). To a solution of 8 (20.0 g, 61.1 mmol) in 2-propanol (100 mL) was added Pd(OH)₂/C (wet, Degussa type, 2.0 g) and the mixture was stirred for 24 h under H_2 atmosphere. The mixture was filtered through Celite[®] and concentrated under reduced pressure. The residue was purified by column chromatography on silica gel (10:1 hexanes/EtOAc) to yield 9 as a colorless oil (19.1 g, 95%). This material was obtained as an inseparable mixture of diastereomers (2:1); $n_{\rm D}^{26} = 1.4442$; $[\alpha]_{\rm D}^{25} - 23.3$ (*c* 0.87, CHCl₃); IR (film): $\nu_{\rm max}$ 1737, 1697 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.17 (1H, d, J = 6.9 Hz, 2'-CH₃ of minor isomer), 1.19 (2H, d, J=6.9 Hz, 2'-CH₃ of major isomer), 1.24 (3H, t, J=7.1 Hz, CH₃ of -CO₂Et), 1.28 (2H, d, J=6.3 Hz, 5-CH₃ of major isomer), 1.29 (1H, d, J = 6.3 Hz, 5-CH₃ of minor isomer), 1.46 (12H, br s, 2-CH₃ and ^tBu), 1.58 (3H, br s, 2-CH₃), 1.8–1.9, 2.1–2.25 (2H, two m, 1'-H), 2.35–2.55 (1H, br, 2'-H), 3.4-3.65 (1H, br, 4-H), 3.9-4.05 (1H, br, 5-H), 4.11 (2H, q, J=7.1 Hz, CH₂ of -CO₂Et). Anal. Calcd for C₁₇H₃₁NO₅: C, 61.98; H, 9.48; N, 4.25. Found: C, 61.73; H, 9.44; N, 4.44.

2.1.3. Methyl (2RS,4R,5R)-4-t-butoxycarbonylamino-5hydroxy-2-methylhexanoate (10). To a solution of 9 (1.04 g, 3.16 mmol) in methanol (20 mL) was added D-10camphorsulfonic acid (79 mg, 0.316 mmol) and the mixture was stirred for 12 h at room temperature. To the mixture was added triethylamine (88 µL, 0.631 mmol) and the solvent was removed under reduced pressure. The residue was purified by column chromatography on silica gel (2:1 hexanes/EtOAc) to yield **10** as a colorless oil (0.80 g, 92%). This material was obtained as an inseparable mixture of diastereomers (2:1); $n_{\rm D}^{25} = 1.4504$; $[\alpha]_{\rm D}^{23} + 10.8$ (c 0.34, CHCl₃); IR (film): ν_{max} 3382, 1715, 1690 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.1–1.3 (6H, m, 2-CH₃, 6-H), 1.42 (9H, s, ^tBu), 1.48–1.58, 1.83–2.04 (2H, two m, 3-H), 2.27 (1H, br s, -OH), 2.45-2.63 (1H, m, 2-H), 3.35-3.60 (1H, m, 4-H), 3.66 and 3.67 (3H, two s, -CO₂Me), 3.65-3.75 (2/3H, m, 5-H of major isomer), 4.05-4.15 (1/3H, m, 5-H of minor isomer), 4.6-4.8 (1H, m, -NH-). Anal. Calcd for C13H25NO5: C, 56.71; H, 9.15; N, 5.09. Found: C, 56.55; H, 9.13; N, 5.18.

2.1.4. δ -Lactone 11. To a solution of 10 (52.1 g, 0.189 mol) in benzene (1 L) was added D-10-camphorsulfonic acid (1.1 g, 0.0047 mol) and the mixture was stirred for 5 h at 45 °C. The solvent was removed under reduced pressure. The residue was purified by column chromatography on silica gel (2:1 hexanes/EtOAc) followed by recrystallization from hexanes–EtOAc to yield lactone 11 (40.5 g, 88%) as colorless crystals. This material was obtained as a mixture of diastereomers (2:1), and used in the next reaction without

separation. However, a small amount of this mixture was further purified and separated by preparative TLC (1:1 hexanes/EtOAc) for analytical purposes.

(2*R*,4*R*,5*R*)-4-*t*-Butoxycarbonylamino-2-methylhexan-5olide (11α, minor isomer, less polar). Mp 140–144 °C; $[α]_{D}^{25}$ +46.1 (*c* 0.45, CHCl₃); IR (KBr): ν_{max} 3394, 1712, 1514 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.32 (3H, d, *J*=6.9 Hz, 2-CH₃), 1.33 (3H, d, *J*=6.5 Hz, 6-H), 1.45 (9H, s, ^{*t*}Bu), 1.75 (1H, ddd, *J*=3.3, 12.0, 13.8 Hz, 3-H_a), 2.24 (1H, ddd, *J*=3.8, 6.9, 13.8 Hz, 3-H_b), 2.61 (1H, m, 2-H), 3.95 (1H, br, 4-H), 4.58 (1H, dq, *J*=2.2, 6.5 Hz, 5-H), 4.79 (1H, br d, *J*=8.7 Hz, -NH–); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 17.2, 17.6, 28.3, 30.9, 34.4, 47.4, 78.7, 80.2, 155.5, 173.5. Anal. Calcd for C₁₂H₂₁NO₄: C, 59.24; H, 8.70; N, 5.76. Found: C, 59.31; H, 8.70; N, 5.69.

(2*S*,4*R*,5*R*)-4-*t*-Butoxycarbonylamino-2-methylhexan-5olide (**11**β, major isomer, more polar). Mp 134–139 °C; $[\alpha]_D^{27}$ +75.2 (*c* 0.72, CHCl₃); IR (KBr): ν_{max} 3388, 1730, 1714, 1520 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.21 (3H, d, *J*=6.3 Hz, 2-CH₃), 1.34 (3H, d, *J*=6.4 Hz, 6-H), 1.3–1.4 (1H, m, 3-H_a), 1.43 (9H, s, ^{*T*}Bu), 2.5–2.7 (2H, m, H-2, 3-H_b), 4.11 (1H, br, 4-H), 4.50 (1H, dq, *J*=3.0, 6.4 Hz, 5-H), 4.75 (1H, br d, *J*=8.4 Hz, –NH–); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 15.5, 16.0, 28.3, 32.4, 35.5, 47.8, 75.5, 79.9, 155.5, 175.6. Anal. Calcd for C₁₂H₂₁NO₄: C, 59.24; H, 8.70; N, 5.76. Found: C, 59.19; H, 8.77; N, 6.01.

2.1.5. (3R,5R,6R)-5-t-Butoxycarbonylamino-3,6dimethyltetrahydropyran-2-thione (12). To a solution of 11 (38.1 g, 157 mmol) in toluene (1.2 L) was added Lawesson's reagent (63.3 g, 157 mmol) and the mixture was stirred for 7 h at 80 °C and for an additional 9 h at room temperature. The solvent was removed under reduced pressure. Two rounds of column chromatography on silica gel (3:1 hexanes/EtOAc first time; 2:80:20 MeOH/CHCl₃/ hexanes second time) gave thionolactone **12** as a yellow paste (26.5 g, 65%); $n_D^{24} = 1.5085$; $[\alpha]_D^{23} + 176$ (c 0.50, CHCl₃); IR (film): ν_{max} 3327, 2979, 1714, 1695, 1681, 1504, 1455, 1109, 1056, 1016 cm⁻¹; ¹H NMR (300 MHz, C₆D₆): δ (ppm) 0.45 (1H, ddd, J=4.7, 14.7, 16.2 Hz, 4-H_a), 1.10 (3H, d, *J*=6.1 Hz mm, 6-CH₃), 1.10 (3H, d, *J*=6.6 Hz, 3-CH₃), 1.38 (9H, s, ^tBu), 1.58–1.72 (2H, m, 3-H, 4-H_b), 3.61 (1H, dq, J=3.6, 6.1 Hz, 6-H), 3.79 (1H, m, 5-H), 4.07 (1H, d, J = 8.7 Hz, -NH-); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 15.8, 19.4, 28.2, 36.8, 41.3, 48.2, 78.7, 79.9, 155.5, 226.6; HRMS (FAB) *m/z* Calcd for C₁₂H₂₁NO₃S: 259.1242 (M⁺). Found: 259.1215.

2.1.6. Ester 13. To a solution of **12** (25.7 g, 99 mmol) in toluene (600 mL) was added methyl (triphenyl-phosphoranylidene)acetate (66.3 g, 198 mmol) and the mixture was stirred for 24 h at 80 °C. The solvent was removed under reduced pressure. Two rounds of column chromatography on silica gel (3:1 hexanes/EtOAc first time; 30:1-9:1 CHCl₃/EtOAc second time) gave **13**Z (19.4 g, 66%, colorless paste) and **13**E (4.56 g, 15%, colorless paste). Both of these compounds were used in the next reaction.

Methyl (*Z*)-(*3R*,5*R*,6*R*)-(5-*t*-butoxycarbonylamino-3,6dimethyltetrahydropyran-2-ylidene)acetate (**13***Z*, less polar). n_D^{27} =1.4809; $[\alpha]_D^{26}$ +78.5 (*c* 0.46, CHCl₃); IR (film): ν_{max} 3344, 2978, 1714, 1705, 1695, 1645, 1518, 1095 cm⁻¹; ¹H NMR (300 MHz, C₆D₆): δ (ppm) 0.51 (3H, d, *J*=6.3 Hz, 3-CH₃), 0.97 (1H, ddd, *J*=3.7, 12.1, 13.1 Hz, 4-H_a), 1.18 (3H, d, *J*=6.4 Hz, 6-CH₃), 1.45 (9H, s, ^{*t*}Bu), 1.52 (1H, ddd, *J*=3.5, 5.3, 13.1 Hz, 4-H_b), 1.62 (1H, m, 3-H), 3.48 (3H, s, -OMe), 3.53 (1H, dq, *J*=1.8, 6.4 Hz, 6-H), 3.68 (1H, m, 5-H), 4.67 (1H, d, *J*=8.7 Hz, -NH–), 5.14 (1H, d, *J*=1.2 Hz, =CHCO₂-); ¹³C NMR (75.5 MHz, C₆D₆): δ (ppm) 17.3, 17.4, 28.4, 28.9, 36.9, 48.5, 50.3, 78.0, 78.9, 99.0, 155.6, 165.1, 171.8; HRMS (FAB) *m/z* Calcd for C₁₅H₂₅NO₅: 299.1733 (M⁺). Found: 299.1752.

Methyl (E)-(3R,5R,6R)-(5-t-butoxycarbonylamino-3,6dimethyltetrahydropyran-2-ylidene)acetate (**13E**, more polar). $n_{\rm D}^{26}$ =1.4888; $[\alpha]_{\rm D}^{26}$ -19.4 (c 0.60, CHCl₃); IR (film): $\nu_{\rm max}$ 3340, 2978, 1714, 1705, 1697, 1633, 1520, 1167 cm⁻¹; ¹H NMR (300 MHz, C₆D₆): δ (ppm) 0.53 (3H, d, J=6.0 Hz, 3-CH₃), 0.61 (1H, ddd, J=5.1, 12.2, 13.4 Hz, 4-H_a), 1.19 (3H, d, J=6.2 Hz, 6-CH₃), 1.39 (9H, s, ¹Bu), 1.63 (1H, ddd, J=2.6, 6.8, 13.4 Hz, 4-H_b), 1.73 (1H, m, 3-H), 3.51 (3H, s, -OMe), 3.70 (1H, dq, J=3.5, 6.2 Hz, 6-H), 3.83 (1H, m, 5-H), 4.53 (1H, d, J=9.3 Hz, -NH-), 4.92 (1H, d, J=1.2 Hz, =CHCO₂-); ¹³C NMR (75.5 MHz, C₆D₆): δ (ppm) 15.7, 16.6, 28.4, 30.2, 36.0, 48.3, 50.2, 72.8, 78.9, 91.6, 155.6, 165.5, 172.4; HRMS (FAB) *m/z* Calcd for C₁₅H₂₅NO₅: 299.1733 (M⁺). Found: 299.1721.

2.1.7. Methyl (2R,3R)-(3-t-butoxycarbonylamino-2,5dimethyl-3,4-dihydro-2H-pyran-6-yl)acetate (14). To a solution of 13 (mixture of 13Z and 13E, 38.3 mg, 0.128 mmol) in benzene (2 mL) was added DBU (191 µL, 1.28 mmol) and the mixture was stirred for 6 h at 100 °C (bath temperature). The solvent was removed under reduced pressure. The residue was purified by column chromatography on silica gel (5:1 hexanes/EtOAc) to provide 14 as a colorless oil (38.3 mg, 100%); $n_{\rm D}^{25} = 1.4700$; $[\alpha]_{\rm D}^{24} + 29.2$ (c 0.22, CHCl₃); IR (film): ν_{max} 3410, 2979, 1745, 1714, 1695, 1505, 1366, 1162 cm⁻¹; ¹H NMR (300 MHz, C₆D₆): δ (ppm) 1.14 (3H, d, J = 6.3 Hz, 2-CH₃), 1.27 (3H, s, 5-CH₃), 1.45 (9H, s, ^tBu), 1.71 (1H, d, J = 17.1 Hz, 4-H_a), 2.03 (1H, dd, J=4.7, 17.1 Hz, 4-H_b), 2.88, 2.97 (2H, two d, J=16.7 Hz, $-CH_2CO_2$ -), 3.30 (3H, s, -OMe), 3.61 (1H, q, J=6.3 Hz, 2-H), 3.95 (1H, ddt, J=4.7, 9.6, 1.6 Hz, 3-H), 5.21 (1H, d, J=9.6 Hz, -NH-); ¹³C NMR (75.5 MHz, C_6D_6): δ (ppm) 17.4, 17.5, 28.4, 35.1, 36.2, 47.2, 51.4, 73.0, 78.6, 103.2, 142.1, 156.1, 170.4. Anal. Calcd for C₁₅H₂₅NO₅: C, 60.18; H, 8.42; N, 4.68. Found: C, 60.17; H, 8.38; N, 4.68.

2.1.8. Methyl (2*S*,3*S*,5*R*,6*R*)-(5-*t*-butoxycarbonylamino-3,6-dimethyltetrahydropyran-2-yl)acetate (15). To a solution of 14 (5.52 g, 18.4 mmol) in acetic acid (40 mL) was added PtO₂ (550 mg) and the mixture was stirred for 36 h under H₂ atmosphere (5 atm) using a Parr's hydrogenator. The mixture was filtered through Celite[®] and neutralized by addition of saturated NaHCO₃ solution and then extracted with EtOAc. The combined organic layer was washed with water and brine before drying with MgSO₄. The solvent was removed and the residue was purified by column chromatography on silica gel (5:1–1:1 hexanes/EtOAc) to give a mixture of compound 15 and intact 14. This mixture was again reacted and purified similarly to yield pure **15** as colorless crystals (3.89 g, 70%). Stereochemistry of this compound was confirmed by an NOE experiment; NOEs were observed between 2-H and 3-H, and between 2-H and 6-H; mp 69–71 °C; $[\alpha]_D^{27} - 20.6$ (c 0.56, CHCl₃); IR (KBr): v_{max} 3383, 2978, 1743, 1703, 1510, 1163 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.04 (3H, d, J=7.5 Hz, 3-CH₃), 1.13 (3H, d, J=6.4 Hz, 6-CH₃), 1.43 (9H, s, ^tBu), 1.81 (1H, m, 3-H), 1.95 (2H, t, J=3.5 Hz, 4-H), 2.35 (1H, dd, J=5.3, 15.2 Hz, $-CH_aCO_2-$), 2.56 (1H, dd, J=8.5, 15.2 Hz, -CH_bCO₂-), 3.57 (1H, m, 5-H), 3.65 (1H, dq, J=2.2, 6.4 Hz, 6-H), 3.69 (3H, s, -OMe), 3.99 (1H, ddd, J=2.9, 5.3, 8.5 Hz, 2-H), 4.72 (1H, d, J=9.2 Hz, -NH-); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 15.2, 17.6, 28.4, 29.1, 35.8, 38.4, 48.0, 51.7, 76.5, 77.3, 79.0, 155.8, 171.8. Anal. Calcd for C₁₅H₂₇NO₅: C, 59.78; H, 9.03; N, 4.65. Found: C, 59.91; H, 9.06; N, 4.64.

2.1.9. (2S,3S,5R,6R)-(5-t-Butoxycarbonylamino-3,6dimethyltetrahydropyran-2-yl)acetaldehyde (16). To a cooled $(-78 \,^{\circ}\text{C})$ solution of 15 (2.57 g, 8.53 mmol) in toluene (25 mL) was added diisobutylaluminum hydride (17.8 mL of 0.96 M in hexanes, 17.1 mmol) and the mixture was stirred for 30 min at this temperature. The reaction was quenched with MeOH (3.5 mL), poured into 1 N HCl solution (300 mL) and extracted with EtOAc. The combined organic layer was washed with saturated NaHCO₃ solution and brine, dried with MgSO4 and concentrated under reduced pressure. Purification on silica gel (3:1 hexanes/ EtOAc) gave aldehyde 16 as a colorless oil (2.29 g, 98%); $n_{\rm D}^{25} = 1.4665$; $[\alpha]_{\rm D}^{26} - 27.2$ (c 0.32, CHCl₃); IR (film): $\nu_{\rm max}$ 3460, 2976, 1722, 1714, 1496, 1171 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.03 (3H, d, J=7.2 Hz, 3-CH₃), 1.11 (3H, d, *J*=6.3 Hz, 6-CH₃), 1.42 (9H, s, ^{*t*}Bu), 1.78 (1H, m, 3-H), 1.95 (2H, m, 4-H), 2.34 (1H, ddd, J=2.2, 4.1,16.4 Hz, -CH_aCHO), 2.64 (1H, ddd, J=2.2, 9.3, 16.4 Hz, $-CH_bCHO$), 3.57 (1H, m, 5-H), 3.66 (1H, dq, J=2.3, 6.3 Hz, 6-H), 4.06 (1H, ddd, J = 3.1, 4.1, 9.3 Hz, 2-H), 4.69 (1H, d, J=9.1 Hz, -NH), 9.77 (1H, t, J=2.2 Hz, -CHO);¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 15.3, 17.5, 28.4, 29.3, 35.7, 46.9, 47.9, 75.8, 76.4, 79.1, 155.7, 201.1. Anal. Calcd for C₁₄H₂₅NO₄: C, 61.97; H, 9.29; N, 5.16. Found: C, 61.72; H, 9.30; N, 5.24.

2.1.10. Ethyl (2E, 2'S, 3'S, 5'R, 6'R)-4-(5'-t-butoxycarbonylamino-3',6'-dimethyltetrahydropyran-2'-yl)-2-methyl**but-2-enoate** (17). To a solution of aldehyde 16 (66.1 mg, 0.24 mmol) in benzene (1 mL) was added (carbethoxyethylidene)triphenylphosphorane (113 mg, 0.29 mmol) and the mixture was stirred for 3 h at 60 °C. The solvent was removed under reduced pressure. The residue was purified by silica gel column chromatography (4:1 hexanes/EtOAc) to furnish desired 17 (80.6 mg, 93%) and undesired Z-derivative (3.5 mg, 4%); $n_{\rm D}^{22} = 1.4755$; $[\alpha]_{\rm D}^{26} - 13.2$ (c 0.56, CHCl₃); IR (film): ν_{max} 3462, 3384, 2978, 1722, 1714, 1651, 1504, 1171 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.04 (3H, d, J=7.2 Hz, 3'-CH₃), 1.15 $(3H, d, J=6.3 \text{ Hz}, 6'-CH_3)$, 1.30 $(3H, t, J=7.0 \text{ Hz}, CH_3 \text{ of})$ -OEt), 1.44 (9H, s, ^tBu), 1.77 (1H, m, 3'-H), 1.85 (3H, d, J=1.4 Hz, 2-CH₃), 1.90–1.95 (2H, m, 4'-H), 2.25 (1H, m, 4-H_a), 2.40 (1H, m, 4-H_b), 3.5–3.7 (3H, m, 2'-H, 5'-H, 6'-H), 4.19 (2H, q, J=7.0 Hz, CH₂ of –OEt), 4.73 (1H, d, J=8.7 Hz, -NH-), 6.73 (1H, ddq, J=6.5, 7.8, 1.4 Hz,

3-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 12.6, 14.2, 15.0, 17.6, 28.4, 29.1, 32.4, 35.9, 48.2, 60.5, 76.4, 79.0, 79.8, 129.4, 137.8, 155.8, 167.9. Anal. Calcd for C₁₉H₃₃NO₅: C, 64.20; H, 9.36; N, 3.94. Found: C, 64.36; H, 9.38; N, 3.82.

2.1.11. (2E, 2'S, 3'S, 5'R, 6'R)-4-(5'-t-Butoxycarbonylamino-3',6'-dimethyltetrahydropyran-2'-yl)-2-methylbut-2-en-1-ol (18). A stirred solution of 17 (2.92 g, 8.21 mmol) in toluene (33 mL) was cooled to -68 °C, treated with diisobutylaluminum hydride (25.7 mL of 0.96 M in hexanes, 24.6 mmol) and warmed to 0 °C during 15 min. The reaction mixture was again cooled to -68 °C and quenched with MeOH (5 mL). The resultant mixture was poured into 1 N HCl solution (300 mL) and extracted with EtOAc. The combined organic layer was washed with water, saturated NaHCO₃ solution and brine, dried with MgSO₄ and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (1:1 hexanes/EtOAc) to give alcohol 18 as a colorless oil (1.1 hoxanes Level a) $[\alpha]_{D}^{24} = 1.4823; [\alpha]_{D}^{25} - 4.0 (c \ 0.75, CHCl_3); IR (film): <math>\nu_{max}$ 3458, 2976, 1714, 1504, 1169 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.00 (3H, d, J=7.2 Hz, 3'-CH₃), 1.12 (3H, d, J=6.4 Hz, 6'-CH₃), 1.41 (9H, s, ^tBu), 1.65 (3H, s, 2-CH₃), 1.73 (1H, m, 3^t-H), 1.85–1.90 (2H, m, 4'-H), 1.92 (1H, s, -OH), 2.11 (1H, m, 4-H_a), 2.26 $(1H, m, 4-H_b)$, 3.46 (1H, dt, J=2.7, 7.2 Hz, 2'-H), 3.52 (1H, dt, J=2.7, 7.2 Hz, 3'-H), 3.52 (1H, dt, J=2.7, 7.2 Hz), 3.52 (1H, dt, J=m, 5'-H), 3.58 (1H, dq, J=2.2, 6.4 Hz, 6'-H), 3.98 (2H, s, 1-H), 4.75 (1H, d, J=9.3 Hz, -NH-), 5.38 (1H, ddq, J=6.3, 7.6, 1.3 Hz, 3-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 13.9, 14.9, 17.7, 28.3, 28.8, 31.2, 35.9, 48.3, 68.5, 76.2, 79.0, 80.7, 121.2, 136.6, 155.8. Anal. Calcd for C₁₇H₃₁NO₄: C, 65.14; H, 9.97; N, 4.47. Found: C, 65.36; H, 10.12; N, 4.46.

 $(2'E, 2''S, 3''S, 5''R, 6''R) - 2 - \{[4' - (5'' - t - Butoxy - 1)]$ 2.1.12. carbonylamino-3",6"-dimethyltetrahydropyran-2"-yl)-2'-methylbut-2'-en-1'-yl]sulfanyl}-1,3-benzothiazole (19). To a cooled (0 °C) solution of 2-mercaptobenzothiazole (16 mg, 0.096 mmol) and triphenylphosphine (25 mg, 0.096 mmol) in THF (0.2 mL) was added a solution of the allylic alcohol **18** (20.0 mg, 0.064 mmol) in THF (0.2 mL) over 10 min. To the resulting solution was added diisopropyl azodicarboxylate (40% in toluene, 52 µL, 0.096 mmol) over 10 min. After stirring was continued at 0 °C for 3 h, the mixture was evaporated to dryness. The residue was dissolved in hexane and filtered through Celite® pad. Further purification of the resulting solution by silica gel column chromatography (10:1–3:1 hexanes/EtOAc) afforded sulfide **19** as a colorless viscous oil (28.4 mg, 96%); $n_{\rm D}^{25} = 1.5166$; $[\alpha]_{\rm D}^{30} - 7.1$ (*c* 0.51, CHCl₃); IR (film): $\nu_{\rm max}$ 2976, 1714, 1504 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.87 (3H, d, J=7.2 Hz, 3"-CH₃), 1.06 (3H, d, J=6.3 Hz, 6"-CH₃), 1.40 (9H, s, ^tBu), 1.45 (1H, m, 3"-H), 1.6-1.8 (2H, m, 4[#]-H), 1.75 (3H, s, 2'-CH₃), 2.05 (1H, m, $4'-H_a$), 2.21 (1H, m, $4'-H_b$), 3.32 (1H, dt, J=2.6, 7.4 Hz, 2"-H), 3.4–3.5 (2H, m, 5"-H, 6"-H), 3.90, 3.96 (2H, two d, J=13.1 Hz, 1'-H), 4.66 (1H, d, J=9.3 Hz, -NH-), 5.49 $(1H, t, J=6.6 \text{ Hz}, 3'-\text{H}), 7.24 (1H, br t, J=7.8 \text{ Hz}, H_{ar}),$ 7.36 (1H, br t, J=7.8 Hz, H_{ar}), 7.69 (1H, br d, J=7.8 Hz, H_{ar}), 7.82 (1H, br d, J=7.8 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 14.7, 15.4, 17.5, 28.3, 28.6, 31.7, 35.7, 43.0, 48.1, 76.1, 78.8, 80.3, 120.7, 121.4, 124.2, 125.9,

126.0, 131.0, 135.2, 153.0, 155.7, 166.3; HRMS (FAB) m/zCalcd for C₂₄H₃₅N₂O₃S₂: 463.2089 (M⁺ + H). Found: 463.2083.

2.1.13. $(2'E, 2''S, 3''S, 5''R, 6''R) - 2 - \{[4'-(5''-t-Butoxy-t)]$ carbonylamino-3",6"-dimethyltetrahydropyran-2"-yl)-2'-methylbut-2'-en-1'-yl]sulfonyl}-1,3-benzothiazole (20). To a solution of sulfide 19 (20.6 mg, 0.045 mmol) in EtOH (0.3 mL) were added $(NH_4)_6Mo_7O_{24} \cdot 4H_2O$ (11 mg, 0.0089 mmol) and H_2O_2 (34%, 45 μ L, 0.445 mmol) at 0 °C. The mixture was stirred at room temperature for 2 h, and then treated with 10% sodium thiosulfate solution. The aqueous layer was extracted with EtOAc, and the resulting organic layer was washed with saturated NaHCO₃ solution and brine before drying with MgSO₄. The solvent was removed and the residue was purified by column chromatography on silica gel (2:1 hexanes/EtOAc) to yield sulfone 20 as a colorless viscous oil (19.2 mg, 87%); $n_{\rm D}^{25} = 1.5170$; $[\alpha]_{\rm D}^{29} - 20.3$ (c 0.38, CHCl₃); IR (film): $\nu_{\rm max}$ 3460, 2976, 1714, 1496, 1331 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.76 (3H, d, J=7.2 Hz, 3"-CH₃), 0.96 (3H, d, J=6.3 Hz, 6"-CH₃), 1.16 (1H, m, 3"-H), 1.38 (9H, s, ^tBu), 1.50 (1H, m, 4"-H_a), 1.64 (1H, m, 4"-H_b), 1.84 (3H, s, 2'-CH₃), 1.98 (1H, m, 4'-H_a), 2.12 (1H, m, 4'-H_b), 3.02 (1H, dt, J=2.5, 7.1 Hz, 2"-H), 3.19 (1H, dq, J=1.9, 6.3 Hz, 6"-H), 3.39 (1H, m, 5"-H), 4.12, 4.19 (2H, two d, J=13.8 Hz, 1'-H), 4.56 (1H, d, J=9.3 Hz, -NH-), 5.27 $(1H, t, J = 6.8 \text{ Hz}, 3'-\text{H}), 7.5-7.65 (2H, m, H_{ar}), 7.97 (1H, br)$ d, J=8.2 Hz, H_{ar}), 8.19 (1H, br d, J=8.2 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 14.7, 17.0, 17.5, 28.4, 28.6, 32.0, 35.7, 48.0, 64.3, 76.2, 79.0, 79.9, 122.2, 123.8, 125.4, 127.7, 128.0, 133.4, 136.8, 152.7, 155.7, 165.6; HRMS (FAB) m/z Calcd for C₂₄H₃₅N₂O₅S₂: 495.1987 (M⁺ + H). Found: 495.1979.

2.1.14. $(2'E, 2''S, 3''S, 5''R, 6''R) - 2 - \{[4'-(5''-Amino-3'', 6''$ dimethyltetrahydropyran-2"-yl)-2'-methylbut-2'-en-1'-yl] sulfonyl}-1,3-benzothiazole (21). A solution of 20 (82 mg, 0.166 mmol) in 4 N HCl/EtOAc (2 mL) was stirred for 90 min at room temperature. The reaction mixture was neutralized with saturated NaHCO3 solution and extracted with CHCl₃ (four times). The organic layer was dried with Na_2SO_4 and evaporated to provide amine 21 (65 mg) as a colorless oil. This crude product was used in the next reaction without further purification; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.84 (3H, d, J = 7.5 Hz, 3["]-CH₃), 1.03 (3H, d, J=6.4 Hz, 6"-CH₃), 1.14 (1H, m, 3"-H), 1.5-1.65 (4H, m, H-4", -NH₂), 1.86 (3H, s, 2'-CH₃), 2.04 (1H, m, 4'-H_a), 2.17 (1H, m, 4'-H_b), 2.53 (1H, m, 5"-H), 3.07 (1H, dt, J=2.8, 7.3 Hz, 2"-H), 3.21 (1H, dq, J=2.0, 6.4 Hz, 6"-H), 4.15, 4.21 (2H, two d, J=13.2 Hz, 1'-H), 5.31 (1H, t, J = 6.6 Hz, 3' -H, 7.55–7.67 (2H, m, H_{ar}), 8.00 (1H, br d, J = 7.4 Hz, H_{ar}), 8.22 (1H, br d, J = 7.4 Hz, H_{ar}).

2.1.15. (2Z,4S)-4-(*t*-Butyldimethylsilyloxy)pent-2-enoic acid (23). To a solution of methyl ester 22 (500 mg, 2.05 mmol) in MeOH–H₂O (20:1, 10 mL) was added LiOH·H₂O (430 mg) and the mixture was stirred overnight at room temperature. The reaction mixture was neutralized with 3 N HCl and extracted with EtOAc. The organic layer was washed with brine, dried with MgSO₄ and evaporated. Purification of the residue on silica gel (4:1 hexanes/EtOAc) provided carboxylic acid 23 as a colorless oil (431 mg,

1385

91%); $n_{\rm D}^{25} = 1.4465$; $[\alpha]_{\rm D}^{28} + 57.7$ (*c* 0.86, CHCl₃); IR (film): $\nu_{\rm max}$ 2956, 1695, 1645, 1254 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.05, 0.07 (6H, two s, -SiMe), 0.89 (9H, s, ¹Bu), 1.27 (3H, d, J = 6.6 Hz, 5-H), 5.40 (1H, m, 4-H), 5.69 (1H, dd, J = 1.2, 11.8 Hz, 2-H), 6.35 (1H, dd, J = 7.7, 11.8 Hz, 3-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) -4.8, -4.8, 18.1, 23.4, 25.8, 65.6, 116.4, 157.2, 171.4. Anal. Calcd for C₁₁H₂₂O₃Si: C, 57.35; H, 9.63. Found: C, 57.23; H, 9.58.

2.1.16. (1S,2R,4S,6R,9R)-4-Methoxy-9-(4'-methoxyphenyl)-4-methyl-5,8,10-trioxabicyclo[4.4.0]decan-2-ol (25). To a solution of triol 24 (523 mg, 2.72 mmol) in DMF (10 mL) were added p-toluenesulfonic acid (5.2 mg, 0.027 mmol) and 4-methoxybenzaldehyde dimethyl acetal $(927 \ \mu L, 5.44 \ mmol)$ at room temperature. The mixture was stirred at 60 °C under slightly reduced pressure (water aspirator) for 40 min and then poured into water. The aqueous layer was extracted with EtOAc, and the resulting organic layer was washed with water, saturated NaHCO₃ solution and brine before drying with MgSO₄. The solvent was removed and the residue was purified by column chromatography on silica gel (3:1-1:1 hexanes/EtOAc) to vield alcohol 25 as a colorless amorphous solid (710 mg, 84%); $[\alpha]_D^{23} + 72.2$ (*c* 0.91, CHCl₃); IR (film): ν_{max} 3496, 1616, 1518, 1250 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.34 (3H, s, 4-CH₃), 1.61 (1H, dd, J=11.4, 13.1 Hz, $3-H_a$), 2.23 (1H, dd, J=5.1, 13.1 Hz, $3-H_b$), 2.71 (1H, br s, -OH), 3.20 (3H, s, -OMe of acetal), 3.38 (1H, t, J=9.5 Hz, 1-H), 3.63 (1H, dt, J=4.4, 9.5 Hz, 6-H), 3.72 (1H, t, J=9.5 Hz, 7-H_a), 3.79 (3H, s, -OMe of PMP), 4.12 (1H, m, 2-H), 4.23 (1H, dd, J=4.4, 9.5 Hz, 7-H_b), 5.50 (1H, s, 9-H), 6.89 (2H, d, J = 8.4 Hz, H_{ar}), 7.42 (2H, d, J = 8.4 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 22.7, 42.9, 47.8, 55.2, 55.2, 63.6, 66.2, 69.0, 83.7, 100.1, 101.8, 101.9, 113.6, 127.5, 129.8, 160.1; HRMS (FAB) m/z Calcd for C₁₆H₂₃O₆: 311.1495 (M⁺ + H). Found: 311.1501.

2.1.17. (1*R*,4*S*,6*R*,9*R*)-4-Methoxy-9-(4'-methoxyphenyl)-4-methyl-5,8,10-trioxabicyclo[4.4.0]decan-2-one (26). To a solution of alcohol 25 (1.63 g, 5.25 mmol) and pyridine (5.10 mL, 63.0 mmol) in dichloromethane (52 mL) was added Dess-Martin periodinane (6.68 g, 15.8 mmol) and stirred for 18 h at room temperature. The reaction mixture was diluted with ether, washed twice with a 1:1 mixture of saturated NaHCO3 solution and 10% sodium thiosulfate solution and then washed with brine. The combined organic layer was dried with MgSO₄ and concentrated. After column chromatography on silica gel (1:1 hexanes/ EtOAc), ketone **26** was isolated as colorless crystals (1.54 g, 95%); mp 113–115 °C; $[\alpha]_D^{23} + 119$ (*c* 0.93, CHCl₃); IR (KBr): ν_{max} 3450, 1734, 1248, 1099 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.45 (3H, s, 4-CH₃), 2.64 $(1H, d, J = 14.3 \text{ Hz}, 3-H_a), 2.70 (1H, d, J = 14.3 \text{ Hz}, 3-H_b),$ 3.21 (3H, s, -OMe of acetal), 3.78 (3H, s, -OMe of PMP), $3.87 (1H, t, J=9.6 Hz, 7-H_a), 3.96 (1H, dt, J=4.0, 9.6 Hz,$ 6-H), 4.24 (1H, d, J=9.6 Hz, 1-H), 4.33 (1H, dd, J=4.0, 9.6 Hz, 7-H_b), 5.51 (1H, s, 9-H), 6.86 (2H, d, J=8.6 Hz, H_{ar}), 7.42 (2H, d, J=8.6 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 22.5, 48.2, 51.7, 55.2, 55.2, 66.0, 69.4, 82.6, 101.9, 102.0, 103.0, 113.5, 127.7, 129.1, 160.2, 198.2; HRMS (FAB) m/z Calcd for C₁₆H₂₁O₆: 309.1338 (M⁺ + H). Found: 309.1339.

2.1.18. (1*S*,3*R*,6*R*,9*S*)-8-Methoxy-3-(4'-methoxyphenyl)-8-methyl-10-methylene-2,4,7-trioxabicyclo[4.4.0]decane (27). To a cooled $(-20 \,^{\circ}\text{C})$ solution of ketone 26 (102 mg, 0.329 mmol) and pyridine (2.7 µL, 0.033 mmol) in THF (2 mL) was added Tebbe reagent (0.99 mL of 0.5 M in toluene, 0.494 mmol) under argon and stirred for 20 min. The reaction was quenched with 2 N NaOH solution (0.49 mL, 0.98 mmol) at this temperature. After gas evolution ceased, Celite[®] was added to the resultant solution and stirring was continued for a few hours at room temperature. This reaction mixture was filtered through Celite[®], and then the solvent was removed under reduced pressure. Purification of the residue on silica gel (9:1 hexanes/EtOAc) provided **27** as colorless crystals (95 mg, 94%); mp 98–100 °C; $[\alpha]_{D}^{19} + 112$ (*c* 1.46, CHCl₃); IR (KBr): ν_{max} 1614, 1520, 1385, 1248, 1088, 1045 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.38 (3H, s, 8-CH₃), 2.46 (1H, d, J = 14.1 Hz, 9-H_a), 2.59 (1H, d, J = 14.1 Hz, 9-H_b), 3.23 (3H, s, –OMe of acetal), 3.66 (1H, dt, J=4.4, 9.8 Hz, 6-H), 3.78 (1H, t, J=9.8 Hz, 5-H_a), 3.81 (3H, s, -OMe of PMP), 3.99 (1H, d, J=9.8 Hz, 1-H), 4.23 (1H, dd, J = 4.4, 9.8 Hz, 5-H_b), 4.91 (1H, d, J = 1.5 Hz, methylene H_a), 5.14 (1H, d, J=1.5 Hz, methylene H_b), 5.60 (1H, s, 3-H), 6.90 (2H, d, J = 8.7 Hz, H_{ar}), 7.46 (2H, d, J = 8.7 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 22.7, 44.8, 48.2, 55.2, 55.2, 66.7, 69.5, 79.5, 99.8, 101.4, 101.5, 107.1, 113.5, 127.5, 130.2, 139.3, 160.0; HRMS (FAB) m/z Calcd for $C_{17}H_{23}O_5$: 307.1546 (M⁺ + H). Found: 307.1547.

2.1.19. (2R,3S,6S)-[6-Methoxy-3-(4'-methoxybenzyloxy)-6-methyl-4-methylenetetrahydropyran-2-yl]methanol (28). To a cooled $(-15 \,^{\circ}\text{C})$ solution of 27 (290 mg, 0.947 mmol) in dichloromethane (5 mL) was added diisobutylaluminum hydride (4.5 mL of 0.95 M in hexanes, 4.26 mmol) under argon and the mixture was stirred for 3 h at this temperature. The reaction was quenched with MeOH (0.35 mL, 8.52 mmol) and then treated with saturated aqueous potassium sodium tartrate solution. The resultant solution was extracted with EtOAc. The combined organic layer was washed with saturated NaHCO₃ solution and brine and then dried with MgSO₄. The solvent was removed under reduced pressure and the residue was purified by silica gel column chromatography (3:1 hexanes/EtOAc) to give 28 as a colorless oil (265 mg, 91%); $n_{\rm D}^{21} = 1.5166$; $[\alpha]_{\rm D}^{21} + 185$ (c 0.78, CHCl₃); IR (film): v_{max} 3477, 1612, 1514, 1250 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.35 (3H, s, 6-CH₃), 1.94 (1H, t, *J*=6.3 Hz, -OH), 2.34 (1H, d, $J = 13.5 \text{ Hz}, 5-H_a$), 2.55 (1H, d, $J = 13.5 \text{ Hz}, 5-H_b$), 3.18 (3H, s, -OMe of acetal), 3.50 (1H, dt, J=9.5, 3.7 Hz, 2-H), 3.67-3.86 (3H, m, 3-H and -CH2OH), 3.80 (3H, s, -OMe of PMB), 4.46, 4.69 (2H, two d, J=11.1 Hz, CH₂ of PMB), 4.94 (1H, s, methylene H_a), 5.19 (1H, s, methylene H_b), 6.89 (2H, d, J=8.7 Hz, H_{ar}), 7.29 (2H, d, J=8.7 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 22.7, 45.0, 48.2, 55.2, 62.7, 72.6, 74.1, 76.0, 99.2, 108.3, 113.8, 129.6, 130.0, 141.3, 159.3; HRMS (FAB) *m/z* Calcd for C₁₇H₂₅O₅: 309.1702 (M⁺ + H). Found: 309.1702.

2.1.20. (2*R*,3*S*,6*S*)-[6-Methoxy-3-(4'-methoxybenzyloxy)-6-methyl-4-methylenetetrahydropyran-2-yl]methyl pivalate (29). To a cooled (0 °C) solution of alcohol 28 (42 mg, 0.136 mmol), triethylamine (57 μL, 0.409 mmol) and DMAP (1.7 mg, 0.014 mmol) in dichloromethane (0.5 mL) was added PivCl $(22 \mu \text{L}, 0.177 \text{ mmol})$ and this mixture was stirred for 12 h at room temperature. The reaction mixture was poured into water and extracted with EtOAc. The organic layer was washed with saturated NaHCO₃ solution and brine, dried with MgSO₄ and then evaporated. Purification of the residue on silica gel (10:1 hexanes/EtOAc) furnished 29 as a colorless oil (51 mg, 95%); $n_{\rm D}^{20} = 1.4926$; $[\alpha]_{\rm D}^{25} + 150$ (c 0.99, CHCl₃); IR (film): $\nu_{\rm max}$ 2962, 1732, 1612, 1514 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.20 (9H, s, ^{*t*}Bu), 1.33 (3H, s, 6-CH₃), 2.34 (1H, d, $J = 13.7 \text{ Hz}, 5-\text{H}_{a}$), 2.55 (1H, d, $J = 13.7 \text{ Hz}, 5-\text{H}_{b}$), 3.20 (3H, s, -OMe of acetal), 3.68 (1H, ddd, J=1.9, 5.9, 9.7 Hz, 2-H), 3.76 (1H, d, J=9.7 Hz, 3-H), 3.80 (3H, s, -OMe of PMB), 4.14 (1H, dd, J=5.9, 11.6 Hz, H_a of PivOCH₂-), 4.39 (1H, dd, J=1.9, 11.6 Hz, H_b of PivOCH₂-), 4.40, 4.66 (2H, two d, J = 10.7 Hz, CH₂ of PMB), 4.95 (1H, d, J = 1.4 Hz, methylene H_a), 5.18 (1H, d, J=1.4 Hz, methylene H_b), 6.88 $(2H, d, J=8.6 \text{ Hz}, H_{ar}), 7.28 (2H, d, J=8.6 \text{ Hz}, H_{ar}); {}^{13}\text{C}$ NMR (75.5 MHz, CDCl₃): δ (ppm) 22.6, 27.2, 38.8, 44.8, 48.0, 55.1, 63.7, 72.4, 72.5, 76.2, 99.0, 108.8, 113.8, 129.5, 129.8, 140.9, 159.3, 178.1; HRMS (FAB) m/z Calcd for $C_{22}H_{33}O_6$: 393.2277 (M⁺+H). Found: 393.2277.

2.1.21. (2R,3S,6S)-(3-Hydroxy-6-methoxy-6-methyl-4methylenetetrahydropyran-2-yl)methyl pivalate (30). To a cooled (0 °C) solution of 29 (51 mg, 0.13 mmol) in dichloromethane (1 mL) were added NaHCO₃ (66 mg, 0.79 mmol), DDQ (90 mg, 0.40 mmol) and water (30 $\mu L),$ respectively, in 3 portions at intervals of 3 h. Stirring was kept for 8 h at this temperature. The reaction mixture was poured into 10% sodium thiosulfate solution and extracted with EtOAc. The organic layer was washed with a 1:1 mixture of saturated NaHCO3 solution and 10% sodium thiosulfate solution and with brine, and then dried with MgSO₄. After removal of the solvent, purification of the residue on silica gel (6:1-2:1 hexanes/EtOAc) provided alcohol **30** as a colorless oil (33 mg, 92%); $n_{\rm D}^{19} = 1.4645$; $[\alpha]_{\rm D}^{15}$ +108 (c 1.02, CHCl₃); IR (film): $\nu_{\rm max}$ 3474, 2959, 1731, 1481, 1287 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.24 (9H, s, ^tBu), 1.36 (3H, s, 6-CH₃), 2.35 (1H, d, $J = 14.0 \text{ Hz}, 5-\text{H}_{a}$), 2.50 (1H, d, J = 6.3 Hz, -OH), 2.55 (1H, d, J = 14.0 Hz, 5-H_b), 3.21 (3H, s, -OMe), 3.51 (1H, ddd, J=2.3, 4.8, 9.8 Hz, 2-H), 3.85 (1H, br, 3-H), 4.31 (1H, dd, J=2.3, 12.1 Hz, H_a of PivOCH₂-), 4.47 (1H, dd, J=4.8, 12.1 Hz, H_b of PivOCH₂-), 4.93 (1H, d, J=1.5 Hz, methylene H_a), 5.17 (1H, d, J=1.5 Hz, methylene H_b); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 22.6, 27.2, 38.9, 44.7, 48.2, 64.2, 68.7, 73.9, 99.0, 107.9, 143.1, 179.3; HRMS (FAB) m/z Calcd for C₁₄H₂₅O₅: 273.1702 (M⁺ + H). Found: 273.1700.

2.1.22. (3R,4R,5R,7S)-(4-Hydroxy-7-methoxy-7-methyl-**1,6-dioxaspiro[2.5]oct-5-yl)methyl pivalate** (31). To a cooled (-15 °C) solution of allylic alcohol **30** (152 mg, 0.56 mmol) and NaHCO₃ (188 mg, 2.23 mmol) in dichloromethane (3 mL) was added *m*CPBA (275 mg of >70% purity, 1.12 mmol). The reaction mixture was gradually warmed up to 0 °C during 3 h with stirring. The resultant solution was diluted with EtOAc, washed successively with a 1:1 mixture of 3 N NaOH solution and 10% sodium thiosulfate solution and with brine, and then dried with MgSO₄. The solvent was removed under reduced pressure and the residue was purified by silica gel column chromatography (1:1 hexanes/EtOAc) to furnish **31** as colorless crystals (135 mg, 84%); mp 77–78 °C; $[\alpha]_D^{19} + 115$ (*c* 0.88, CHCl₃); IR (KBr): ν_{max} 3423, 2987, 1729 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.23 (9H, s, ^{*t*}Bu), 1.36 (3H, s, 7-CH₃), 1.73 (1H, d, J = 14.7 Hz, 8-H_a), 1.88 (1H, d, J = 10.3 Hz, -OH), 2.27 (1H, d, J = 14.7 Hz, 8-H_b), 2.50 (1H, d, J = 4.5 Hz, 2-H_a), 2.99 (1H, d, J = 4.5 Hz, 2-H_b), 3.27 (3H, s, -OMe), 3.67 (1H, t, J = 10.3 Hz, 4-H), 3.79 (1H, ddd, J = 2.2, 6.2, 10.3 Hz, 5-H), 4.23 (1H, dd, J = 6.2, 11.8 Hz, H_a of PivOCH₂–); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 22.7, 27.2, 38.8, 41.8, 47.0, 48.1, 56.2, 64.0, 64.1, 70.8, 98.4, 178.4. Anal. Calcd for C₁₄H₂₄O₆: C, 58.32; H, 8.39. Found: C, 58.20; H, 8.37.

2.1.23. (3R,4R,5R,7S)-(4-t-Butyldimethylsilyloxy-7methoxy-7-methyl-1,6-dioxaspiro[2.5]oct-5-yl)methyl pivalate (32). To a solution of epoxyalcohol 31 (58 mg, 0.20 mmol) in DMF (0.1 mL) was added TBSCN (510 mg in all, 3.61 mmol) in 3 equal portions at intervals of 20 h. Stirring was continued for 2 days at 70 °C. The reaction mixture was diluted with EtOAc, washed with saturated NaHCO₃ solution and brine, and then dried with MgSO₄. The solvent was removed under reduced pressure and the residue was purified by silica gel column chromatography (6:1-1:2 hexanes/EtOAc) to give 32 as a colorless oil (56 mg, 70%) and unreacted starting material (13 mg). The yield was calculated to be 91% based on the recovered starting material; $n_{\rm D}^{20} = 1.4586$; $[\alpha]_{\rm D}^{17} + 123$ (c 1.07, CHCl₃); IR (film): ν_{max} 2958, 1732, 1479, 1284, 1257 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.07, 0.10 (6H, two s, –SiMe), 0.90 (9H, s, -SiBu^t), 1.22 (9H, s, -COBu^t), 1.33 (3H, s, 7-CH₃), 1.72 (1H, d, J = 14.1 Hz, 8-H_a), 2.16 (1H, d, J=14.1 Hz, 8-H_b), 2.43 (1H, d, J=5.1 Hz, 2-H_a), 2.87 (1H, d, J=5.1 Hz, 2-H_b), 3.28 (3H, s, -OMe), 3.78 (1H, d, J=9.3 Hz, 4-H), 3.97 (1H, dd, J=7.4, 11.1 Hz, H_a of PivOCH₂-), 4.05 (1H, m, 5-H), 4.48 (1H, dd, J=1.4, 11.1 Hz, H_b of PivOCH₂-); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) -4.5, -4.5, 18.1, 22.9, 25.7, 27.2, 38.8, 42.7, 47.3, 48.0, 57.1, 63.7, 66.5, 70.5, 98.7, 178.2; HRMS (FAB) m/z Calcd for $C_{20}H_{39}O_6Si$: 403.2516 (M⁺+H). Found: 403.2523.

2.1.24. (3R,4R,5R,7S)-(4-t-Butyldimethylsilyloxy-7methoxy-7-methyl-1,6-dioxaspiro[2.5]oct-5-yl)methanol (33). To a cooled $(-40 \,^{\circ}\text{C})$ solution of 32 (15.4 mg, 0.038 mmol) in THF (0.3 mL) was added diisobutylaluminum hydride (121 µL of 0.95 M in hexanes, 0.115 mmol) under argon and the mixture was stirred for 2 h at this temperature. The reaction was quenched with MeOH (15 µL, 0.383 mmol) and then treated with saturated aqueous potassium sodium tartrate solution. The resultant solution was extracted with EtOAc. The combined organic layer was washed with saturated NaHCO₃ solution and brine and then dried with MgSO₄. The solvent was removed under reduced pressure and the residue was purified by silica gel column chromatography (5:1 hexanes/acetone) to yield alcohol 33 as colorless crystals (11.1 mg, 91%); mp 51–52 °C; $[\alpha]_{D}^{22}$ +138 (c 0.93, CHCl₃); IR (KBr): ν_{max} 3490, 2932, 1473, 1382, 1255 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.05, 0.11 (6H, two s, -SiMe), 0.88 (9H, s, ^{*t*}Bu), 1.34 (3H, s, 7-CH₃), 1.70 (1H, d, J = 14.1 Hz, 8-H_a), 1.92 (1H, dd, J=5.4, 7.5 Hz, -OH), 2.16 (1H, d, J=14.1 Hz, 8-H_b), 2.40 (1H, d, J=5.0 Hz, 2-H_a), 2.86 (1H, d, J=5.0 Hz, 2-H_b), 3.24 (3H, s, -OMe), 3.69 (1H, m, H_a of -CH₂OH), 3.79 (1H, m, H_b of -CH₂OH), 3.85 (2H, br s, 4-H, 5-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) -4.7, -4.5, 18.1, 23.0, 25.7, 42.7, 47.0, 48.1, 57.1, 61.8, 65.5, 72.1, 99.0; HRMS (FAB) *m*/*z* Calcd for C₁₅H₃₁O₅Si: 319.1941 (M⁺ + H). Found: 319.1940.

2.1.25. (3R,4R,5S,7S)-4-t-Butyldimethylsilyloxy-7-methoxy-7-methyl-1,6-dioxaspiro[2.5]octane-5-carbaldehyde (34). To a stirred solution of alcohol 33 (93 mg, 0.29 mmol) in dichloromethane (2 mL) were added powdered MS-4 Å (450 mg), NMO (44 mg, 0.38 mmol), and finally TPAP (10 mg, 0.029 mmol) in this order. The reaction mixture was stirred for 3 h at room temperature and then directly purified by column chromatography on silica gel (2:1 hexanes/EtOAc) to provide aldehyde 34 as a colorless oil (74 mg) containing a small amount of inseparable impurity. The yield was calculated to be ~80%. This product was used in the next reaction without further purification; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.00, 0.03 (6H, two s, -SiMe), 0.87 (9H, s, ^tBu), 1.40 (3H, s, 7-CH₃), 1.75 (1H, d, J = 14.3 Hz, 8-H_a), 2.18 (1H, d, J = 14.3 Hz, 8-H_b), 2.46 $(1H, d, J=5.1 Hz, 2-H_a), 2.88 (1H, d, J=5.1 Hz, 2-H_b),$ 3.22 (3H, s, -OMe), 3.98 (1H, d, J = 9.6 Hz, 4-H), 4.40 (1H, d, J = 9.6 Hz, 4.40 (1H, d, Jdd, J=1.3, 9.6 Hz, 5-H), 9.74 (1H, d, J=1.3 Hz, -CHO).

Benzothiazol-2^{///}-ylsulfonyl)-3^{//}-methylbut-2^{//}-en-1^{//}-yl]-2',5'-dimethyltetrahydropyran-3'-yl}-4-t-butyldimethylsilyloxypent-2-enamide (35). To a stirred solution of crude amine 21 (65 mg) in acetonitrile (1 mL) were added N-ethylmorpholine (23 µL, 0.182 mmol) and a solution of carboxylic acid 23 (42 mg, 0.182 mmol) in acetonitrile (1 mL). This mixture was cooled to 0 °C and HBTU (69 mg, 0.182 mmol) was added to the resultant solution. The reaction mixture was stirred for 18 h at room temperature and then diluted with EtOAc. After successive washing with saturated NaHCO₃ solution, saturated NH₄Cl solution, saturated NaHCO₃ solution and brine, the organic layer was dried with MgSO₄ and evaporated. Purification of the residue on silica gel (2:1 hexanes/EtOAc containing 1% of NEt₃) provided amide **35** as a colorless viscous oil (84 mg, 83% in two steps from 20); $n_{\rm D}^{26} = 1.5160$; $[\alpha]_{\rm D}^{27} - 39.4$ (c 1.11, CHCl₃); IR (film): ν_{max} 3390, 2928, 1732, 1667, 1633, 1331 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 0.02, 0.04 (6H, two s, -SiMe), 0.76 (3H, d, J=7.5 Hz, 5'-CH₃), 0.86 (9H, s, ^tBu), 0.97 (3H, d, J=6.4 Hz, 2'-CH₃), 1.15–1.25 (1H, m, 5'-H), 1.23 (3H, d, J=6.6 Hz, 5-H), 1.55 $(1H, dt, J = 14.5, 4.7 Hz, 4'-H_a), 1.65 (1H, br d, J = 14.5 Hz,$ 4'-H_b), 1.86 (3H, s, 3"-CH₃), 2.02 (1H, m, 1"-H_a), 2.16 (1H, m, 1''-H_b), 3.08 (1H, dt, J = 2.7, 7.2 Hz, 6'-H), 3.26 (1H, dq, J=2.0, 6.4 Hz, 2'-H), 3.76 (1H, m, 3'-H), 4.14, 4.20 (2H, two d, J = 13.8 Hz, 4"-H), 5.29 (1H, t, J = 6.8 Hz, 2"-H), 5.48 (1H, dd, J=1.4, 11.5 Hz, 2-H), 5.5–5.6 (2H, m, 4-H, -NH-), 5.98 (1H, dd, J=7.8, 11.5 Hz, 3-H), 7.55-7.65 (2H, m, H_{ar}), 7.99 (1H, br d, J=7.3 Hz, H_{ar}), 8.21 (1H, br d, J=7.3 Hz, H_{ar}); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) -4.8, -4.8, 14.8, 17.0, 17.7, 18.1, 23.7, 25.8, 28.6, 31.9,35.6, 46.7, 64.3, 65.3, 75.8, 80.0, 119.0, 122.2, 124.0, 125.3, 127.7, 128.0, 133.2, 136.8, 151.0, 152.6, 165.0, 165.5; HRMS (FAB) m/z Calcd for C₃₀H₄₇N₂O₅S₂Si: 607.2696 $(M^+ + H)$. Found: 607.2681.

2.1.27. (2Z,2'R,2''E,3'R,3'''R,4S,4''E,4'''R,5'S,5'''R,6'S,7'''S)-N-{6'-[5"-(4"'-t-Butyldimethylsilyloxy-7"'-methoxy-7^{///}-methyl-1^{///},6^{///}-dioxaspiro[2.5]oct-5^{///}-yl)-3^{//}-methylpenta-2'', 4''-dien-1''-vl]-2', 5'-dimethyltetrahydropyran-3'-vl}-4-t-butyldimethylsilyloxypent-2-enamide (36). To a cooled $(-78 \degree C)$ solution of sulfone 35 (95 mg, 0.16 mmol) in THF (1.3 mL) was added LHMDS (330 µL of 1.0 M in hexanes, 0.33 mmol) dropwise and the mixture was stirred for 10 min. Then a solution of aldehyde 34 (40 mg, 0.13 mmol) in THF (0.4 mL) was added to the resultant solution. Stirring was kept for 1 h at -78 °C and for additional 2 h at room temperature. The reaction was poured into saturated NH₄Cl solution. The aqueous layer was extracted with EtOAc, and the combined organic layer was washed with saturated NaHCO3 solution and brine before drying with MgSO₄. The solvent was removed and the residue was purified by column chromatography on silica gel (7:1–2:1 hexanes/EtOAc) followed by preparative HPLC (2.8:1 hexanes/EtOAc) to furnish diene 36 as a colorless oil (61 mg, 68%); $[\alpha]_{D}^{21}$ + 35.1 (*c* 1.04, CHCl₃); IR (CHCl₃ solution): v_{max} 3441, 2929, 1666, 1501, 1255 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) -0.05, 0.00, 0.03, 0.04 (12H, four s, -SiMe), 0.82, 0.86 (18H, two s, ^tBu), 0.97 (3H, d, J=6.9 Hz, 5'-CH₃), 1.13 (3H, d, J=6.6 Hz, 2'-CH₃), 1.25 (3H, d, J=6.3 Hz, 5-H), 1.36 (3H, s, $7'''-CH_3$, 1.70 (1H, d, J=14.4 Hz, $8'''-H_a$), 1.7–1.82 (1H, m, 5'-H), 1.76 (3H, s, 3"-CH₃), 1.91 (2H, br s, 4'-H), 2.2–2.45 (2H, m, 1"-H), 2.22 (1H, d, J=14.4 Hz, 8^{III}-H_b), 2.40 (1H, d, J=5.4 Hz, $2'''-H_a$), 2.87 (1H, d, J=5.4 Hz, $2^{\prime\prime\prime}$ -H_b), 3.23 (3H, s, -OMe), 3.52 (1H, dt, J=2.4, 7.1 Hz, 6'-H), 3.64 (1H, m, 2'-H), 3.67 (1H, d, J=8.9 Hz, 4^{III}-H), 3.89 (1H, m, 3'-H), 4.28 (1H, t, J=8.9 Hz, 5'''-H), 5.43 (1H, t)t, J=6.9 Hz, 2''-H), 5.48–5.61 (3H, m, 2-H, 4-H, 5''-H), 5.67 (1H, d, J=9.0 Hz, -NH-), 5.99 (1H, dd, J=7.7, 11.6 Hz, 3-H), 6.31 (1H, d, J = 15.6 Hz, 4"-H); ¹³C NMR $(75.5 \text{ MHz}, \text{ CDCl}_3)$: δ (ppm) -4.8, -4.4, -4.1, 12.5,15.1, 17.8, 18.0, 18.2, 23.1, 23.7, 25.6, 25.8, 28.8, 31.9, 35.8, 42.9, 47.0, 47.1, 48.2, 57.2, 65.3, 69.4, 73.8, 75.9, 80.8, 98.8, 119.1, 124.3, 128.7, 134.6, 138.5, 150.9, 165.1; HRMS (FAB) *m*/*z* Calcd for C₃₈H₇₀NO₇Si₂: 708.4691 $(M^+ + H)$. Found: 708.4694.

2.1.28. (2Z,2'R,2''E,3'R,3'''R,4S,4''E,4'''R,5'S,5'''R,6'S,7'''S)-N-{6'-[5"-(4"'-t-Butyldimethylsilyloxy-7"'-methoxy-7^{'''}-methyl-1^{'''},6^{'''}-dioxaspiro[2.5]oct-5^{'''}-yl)-3^{''}-methylpenta-2",4"-dien-1"-yl]-2',5'-dimethyltetrahydropyran-3'-yl}-4-hydroxypent-2-enamide (37). To a cooled (-20 °C) solution of diene **36** (13.3 mg, 0.019 mmol) in MeOH (0.5 mL) was added 0.05% HCl/MeOH solution (0.5 mL) and the mixture was stirred for 10 h at this temperature. The reaction was neutralized with triethylamine (0.1 mL) and diluted with EtOAc. This solution was washed with saturated NaHCO3 solution and brine, dried with MgSO₄ and then evaporated. Purification of the residue by preparative TLC (1:3 hexanes/EtOAc) gave alcohol 37 as a colorless oil (9.5 mg, 85%); $[\alpha]_{D}^{23} + 83.7$ (c 1.11, CHCl₃); IR (CHCl₃ solution): ν_{max} 3438, 3006, 1655, 1626, 1509, 1256 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) -0.04, 0.01 (6H, two s, -SiMe), 0.83 (9H, s, ^tBu), 1.00 (3H, d, J=7.2 Hz, 5'-CH₃), 1.14 (3H, d, J=6.3 Hz, 2'-CH₃), 1.34 $(3H, d, J = 6.9 \text{ Hz}, 5-\text{H}), 1.37 (3H, s, 7'''-CH_3), 1.71 (1H, d, J)$ $J = 14.3 \text{ Hz}, 8'''-H_a$, 1.75–1.85 (1H, m, 5'-H), 1.77 (3H, s, 3''-CH₃), 1.94 (2H, t, J=3.3 Hz, 4'-H), 2.23 (1H, d,

 $J=14.3 \text{ Hz}, 8'''-\text{H}_{b}), 2.2-2.4 (2\text{H}, \text{m}, 1''-\text{H}), 2.41 (1\text{H}, \text{d}, J=5.1 \text{ Hz}, 2'''-\text{H}_{a}), 2.88 (1\text{H}, \text{d}, J=5.1 \text{ Hz}, 2'''-\text{H}_{b}), 3.24 (3\text{H}, \text{s}, -OMe), 3.54 (1\text{H}, \text{dt}, J=2.5, 7.3 \text{ Hz}, 6'-\text{H}), 3.66 (1\text{H}, \text{m}, 2'-\text{H}), 3.68 (1\text{H}, \text{d}, J=8.2 \text{ Hz}, 4'''-\text{H}), 3.94 (1\text{H}, \text{m}, 3'-\text{H}), 4.29 (1\text{H}, \text{t}, J=8.2 \text{ Hz}, 5'''-\text{H}), 4.78 (1\text{H}, \text{m}, 4-\text{H}), 5.44 (1\text{H}, \text{t}, J=7.1 \text{ Hz}, 2''-\text{H}), 5.54 (1\text{H}, \text{dd}, J=8.2, 15.6 \text{ Hz}, 5''-\text{H}), 5.60 (1\text{H}, \text{br} \text{ d}, J=3.9 \text{ Hz}, -O\text{H}), 5.71 (1\text{H}, \text{d}, J=12.0 \text{ Hz}, 2-\text{H}), 5.93 (1\text{H}, \text{d}, J=9.3 \text{ Hz}, -\text{NH}-), 6.18 (1\text{H}, \text{dd}, J=5.4, 12.0 \text{ Hz}, 3-\text{H}), 6.32 (1\text{H}, \text{d}, J=15.6 \text{ Hz}, 4''-\text{H}); ^{13}\text{C} \text{ NMR} (75.5 \text{ MHz}, \text{CDCl}_3): \delta (\text{ppm}) - 4.4, -4.1, 12.5, 15.1, 17.8, 18.0, 22.7, 23.1, 25.6, 28.7, 31.8, 35.7, 42.9, 47.1, 47.5, 48.2, 57.2, 64.5, 69.5, 73.8, 75.8, 80.8, 98.8, 122.5, 124.4, 128.6, 134.6, 138.4, 150.8, 166.1; \text{HRMS} (\text{FAB})$ *m/z*Calcd for C₃₂H₅₆NO₇Si: 594.3826 (M⁺ + H). Found: 594.3818.

2.1.29. (2Z,2'R,2''E,3'R,3'''R,4S,4''E,4'''R,5'S,5'''R,6'S,7'''S)-N-{6'-[5''-(4'''-t-Butyldimethylsilyloxy-7'''-methoxy-7^{"//}-methyl-1^{"//},6^{"//}-dioxaspiro[2.5]oct-5^{"//}-yl)-3^{"/}-methylpenta-2",4"-dien-1"-yl]-2',5'-dimethyltetrahydropyran-3'-yl}-4-acetoxypent-2-enamide (38). To a cooled (0 °C) solution of alcohol 37 (19.8 mg, 0.033 mmol), triethylamine (28 µL, 0.200 mmol) and DMAP (4.1 mg, 0.033 mmol) in dichloromethane (0.4 mL) was added acetic anhydride $(9.4 \,\mu\text{L}, 0.100 \,\text{mmol})$ and this mixture was stirred for 1.5 h at this temperature. The reaction mixture was poured into water and extracted with EtOAc. The organic layer was washed with saturated NaHCO₃ solution and brine, dried with MgSO₄ and then evaporated. Purification of the residue by preparative TLC (1:1 hexanes/EtOAc) yielded acetate 38 as a colorless oil (20.7 mg, 98%); $[\alpha]_D^{21} + 24.6$ (c 1.00, CHCl₃); IR (CHCl₃ solution): *v*_{max} 3441, 2930, 1732, 1670, 1646, 1504 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) -0.03, 0.01 (6H, two s, -SiMe), 0.84 (9H, s, ^tBu), 1.00 (3H, d, J=7.5 Hz, 5'-CH₃), 1.15 (3H, d, J=6.3 Hz, 2'-CH₃), 1.37 (3H, s, 7''-CH₃), 1.39 (3H, d, J=6.6 Hz, 5-H), 1.71 (1H, d, J = 14.3 Hz, $8'''-H_a$), 1.7–1.8 (1H, m, 5'-H), 1.77 (3H, s, 3"-CH₃), 1.95 (2H, m, 4'-H), 2.04 (3H, s, -OCOCH₃), 2.23 (1H, d, J=14.3 Hz, $8'''-H_b$), 2.2–2.4 (2H, m, 1"-H), 2.41 (1H, d, J=5.1 Hz, $2'''-H_a$), 2.88 (1H, d, J=5.1 Hz, $2'''-H_b$), 3.25 (3H, s, -OMe), 3.52 (1H, dt, J=2.1, 7.1 Hz, 6'-H), 3.66 (1H, m, 2'-H), 3.68 (1H, d, J=8.3 Hz, 4'''-H), 3.94 (1H, m, 3'-H), 4.30 (1H, br t, J=8.3 Hz, 5^{*III*}-H), 5.45 (1H, t, J=6.9 Hz, 2^{*II*}-H), 5.54 (1H, dd, J = 8.3, 15.6 Hz, 5"-H), 5.70 (1H, d, J = 11.6 Hz, 2-H), 5.89 (1H, dd, J = 8.0, 11.6 Hz, 3-H), 5.98 (1H, d, J = 9.0 Hz)-NH-), 6.26 (1H, m, 4-H), 6.32 (1H, d, J=15.6 Hz, 4''-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) -4.4, -4.1, 12.5, 15.0, 17.8, 18.1, 19.9, 21.2, 23.1, 25.7, 28.8, 31.9, 35.8, 42.9, 47.0, 47.1, 48.2, 57.2, 68.9, 69.5, 73.8, 75.9, 80.8, 98.8, 122.4, 124.3, 128.8, 134.5, 138.5, 143.7, 164.8, 170.3; HRMS (FAB) *m/z* Calcd for C₃₄H₅₈NO₈Si: 636.3932 $(M^+ + H)$. Found: 636.3936.

2.1.30. $(2Z,2'R,2''E,3'R,3'''R,4S,4''E,4'''R,5'S,5'''R,6'S, 7'''S)-N-{6'-[5''-(4'''-Hydroxy-7'''-methoxy-7'''-methyl-1''',6'''-dioxaspiro[2.5]oct-5'''-yl)-3''-methylpenta-2'',4''-dien-1''-yl]-2',5'-dimethyltetrahydropyran-3'-yl}-4-acetoxypent-2-enamide (6, FR901464 methyl acetal). To a cooled (-10 °C) solution of 38 (55 mg, 0.086 mmol) in THF (1 mL) was added TBAF (260 µL of 1.0 M in THF, 0.260 mmol) and this mixture was stirred for 3 h at 0 °C. The resultant solution was poured into water and extracted with EtOAc. The combined organic layer was washed with$

saturated NaHCO₃ solution and brine, dried with MgSO₄ and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (1:2-1:15 hexanes/EtOAc) to give alcohol 6 as a white powder (44 mg, 97%); mp 65–70 °C; $[\alpha]_{D}^{21}$ +26.6 (c 0.87, CHCl₃); IR (KBr): ν_{max} 3449, 2933, 1737, 1669, 1638, 1523 cm⁻¹; ¹H NMR (300 MHz, CDCl₃): δ (ppm) 1.01 (3H, d, J=7.2 Hz, 5'-CH₃), 1.15 (3H, d, J=6.4 Hz, 2'-CH₃), 1.38 $(3H, s, 7'''-CH_3)$, 1.38 (3H, d, J=6.3 Hz, 5-H), 1.73 (1H, d, J=6.3 Hz, 5-H), 1.73 (1J = 14.7 Hz, $8^{\prime\prime\prime}$ -H_a), 1.74–1.85 (2H, m, 5'-H, –OH), 1.79 $(3H, s, 3''-CH_3)$, 1.95 (2H, m, 4'-H), 2.04 (3H, s, -OCOCH₃), 2.23 (1H, m, 1''-H_a), 2.30 (1H, d, J=14.7 Hz, $8'''-H_b$), 2.39 (1H, m, 1''-H_a), 2.49 (1H, d, J=4.5 Hz, 2^{*III*}-H_a), 2.99 (1H, d, J=4.5 Hz, 2^{*III*}-H_b), 3.27 (3H, s, -OMe), 3.52 (1H, dt, J=2.4, 7.1 Hz, 6'-H), 3.60 (1H, t, J=9.9 Hz, 4'''-H), 3.66 (1H, dq, J=2.0, 6.4 Hz, 2'-H), 3.93 (1H, m, 3'-H), 4.05 (1H, dd, J=6.9, 9.9 Hz, 5'''-H), 5.50 (1H, t, J=7.1 Hz, 2''-H), 5.65–5.73 (2H, m, 2-H, 5''-H), 5.88 (1H, dd, J=7.8, 11.7 Hz, 3-H), 6.00 (1H, d, J=9.0 Hz, -NH-), 6.25 (1H, m, 4-H), 6.40 (1H, d, J = 15.6 Hz, 4["]-H); ¹³C NMR (75.5 MHz, CDCl₃): δ (ppm) 12.6, 15.0, 17.8, 20.0, 21.2, 23.0, 28.8, 31.9, 35.8, 42.0, 47.1, 48.4, 56.5, 67.5, 68.9, 73.2, 75.9, 80.8, 98.5, 122.5, 124.1, 128.9, 134.7, 138.1, 143.6, 164.8, 170.3; HRMS (FAB) m/z Calcd for C₂₈H₄₄NO₈: 522.3067 (M⁺ + H). Found: 522.3070.

2.1.31. (2Z,2'R,2''E,3'R,3'''R,4S,4''E,4'''R,5'S,5'''R,6'S,7^{"'}S)-N-{6'-[5["]-(4^{"'},7^{"'}-Dihydroxy-7^{"'}-methyl-1^{"'},6^{"'}dioxaspiro[2.5]oct-5^{"//}-yl)-3["]-methylpenta-2["],4["]-dien-1["]yl]-2',5'-dimethyltetrahydropyran-3'-yl}-4-acetoxypent-2-enamide (1, FR901464). To a cooled (0 °C) solution of 6 (28.4 mg, 0.054 mmol) in THF-water (4:1, 3.5 mL) was added Amberlyst 15 (40 mg). This mixture was gradually warmed up to room temperature and stirred for 20 h. The reaction was neutralized with triethylamine (0.2 mL), dried briefly with Na₂SO₄ and directly purified by silica gel column chromatography (1:3 hexanes/EtOAc-EtOAc only) to some degree. This crude product was further purified by preparative HPLC (2:3 water/MeOH) to give 1 as a white powder (9.9 mg, 36%) and unreacted starting material (6.0 mg). The yield was calculated to be 45% based on the recovered starting material. Finally, a small amount of this material was finely purified by recrystallization from hexane for analytical purposes; mp 65–70 °C; $[\alpha]_D^{24}$ –14.1 (c 0.50, CH₂Cl₂); IR (KBr): v_{max} 3448, 2925, 1736, 1666, 1634, 1523 cm⁻¹; ¹H NMR (500 MHz, CD₂Cl₂): δ (ppm) 1.01 (3H, d, J=7.5 Hz, 5'-CH₃), 1.11 (3H, d, J=6.3 Hz, 2'-CH₃), 1.34 (3H, d, *J*=7.0 Hz, 5-H), 1.43 (3H, s, 7^{*III*}-CH₃), 1.61 (1H, d, J=9.6 Hz, 4^{*m*}-OH), 1.64 (1H, d, J=14.5 Hz, 8^{///}-H_a), 1.72–1.82 (1H, m, 5'-H), 1.78 (3H, s, 3^{//}-CH₃), 1.92 (2H, m, 4'-H), 2.01 (3H, s, $-OCOCH_3$), 2.23 (1H, m, 1"-H_a), 2.34 (1H, d, J = 14.5 Hz, 8"'-H_b), 2.36 (1H, m, 1"-H_b), 2.55 $(1H, d, J=4.5 \text{ Hz}, 2'''-H_a), 3.06 (1H, d, J=4.5 \text{ Hz}, 2'''-H_b),$ 3.33 (1H, s, 7^{III}-OH), 3.53 (1H, m, 6'-H), 3.57 (1H, t, J=9.6 Hz, 4'''-H), 3.66 (1H, dq, J=2.3, 6.3 Hz, 2'-H), 3.90 (1H, m, 3'-H), 4.24 (1H, dd, J=7.0, 9.6 Hz, 5^{*III*}-H), 5.54 (1H, br t, J=7.0 Hz, 2''-H), 5.65 (1H, dd, J=7.0, 15.5 Hz, 5''-H), 5.71 (1H, dd, J = 1.8, 11.6 Hz, 2-H), 5.90 (1H, dd, J=8.0, 11.6 Hz, 3-H), 5.98 (1H, d, J=8.5 Hz, -NH-), 6.26 (1H, m, 4-H), 6.38 (1H, d, J=15.5 Hz, 4"-H); ¹³C NMR (125 MHz, CD₂Cl₂): δ (ppm) 12.7, 15.2, 17.9, 20.2, 21.4, 29.1, 29.5, 32.3, 36.2, 41.8, 47.3, 48.1, 58.1, 68.1, 68.9, 73.8, 76.2, 81.1, 96.7, 122.8, 124.6, 129.9, 134.8, 138.3, 143.8, 164.9, 170.6; HRMS (FAB) m/z Calcd for C₂₇H₄₂NO₈: 508.2910 (M⁺ + H). Found: 508.2908.

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Synthesis of α-diazo-β-hydroxyesters through a one-pot protocol by phase-transfer catalysis: application to enantioselective aldol-type reaction and diastereoselective synthesis of α-amino-β-hydroxyester derivatives

Kazuya Hasegawa, Shigeru Arai* and Atsushi Nishida*

Graduate School of Pharmaceutical Sciences, Chiba University, 1-33 Yayoi-cho, Inage-ku, Chiba 263-8522, Japan

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Abstract—The one-pot synthesis of α -diazo- β -hydroxyesters from sodium azide under phase-transfer-catalyzed conditions has been achieved. This protocol includes three different chemical transformations promoted by a single catalyst in each step to give products in good to excellent yields. The reaction was applied to a catalytic asymmetric aldol-type reaction using α -diazoesters with aldehydes in the presence of a chiral quaternary ammonium salt and gave products with up to 81% ee. The diastereoselective transformation of the products to chiral α -amino- β -hydroxyester derivatives is also described.

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1. Introduction

Phase-transfer-catalyzed reactions are some of the most environmentally-friendly processes in synthetic organic chemistry due to their simplicity, mild conditions and high cost performance.¹ Particularly, chiral quaternary ammonium salts have been recognized as powerful asymmetric phase-transfer catalysts (PTCs) since Dolling and O'Donnell independently reported their pioneering studies.² In this paper, we report the PTC-promoted one-pot synthesis of α -diazo- β -hydroxyesters from sodium azide, acetoacetate and aldehydes in the presence of a single catalyst without isolation of intermediates. The application to enantioselective carbon–carbon bond-forming reactions and the diastereoselective synthesis of α -amino and α -hydrazino- β -hydroxyesters is also described in detail.³

2. Resullts and discussion

2.1. One-pot synthesis of α -diazo- β -hydroxyesters

 α -Diazo- β -hydroxyesters are useful as a potential source of amino alcohols or acids and their facile preparation using

the aldol-type reaction of α -diazoesters with aldehydes has been investigated.^{4,5} Since the starting α -diazoesters and the precursor, tosyl azide are both readily available under PTC conditions,⁶ we expected that all three sequences could be promoted by a single phase-transfer catalyst without isolation of any explosive intermediates.

First, we investigated a three-step protocol using tosyl chloride, *t*-butyl acetoacetate **1a** and benzaldehyde **3a** in the presence of tetrahexylammonium bromide (THAB, 10 mol%) as a PTC. The results are summarized in Table 1. The azidation of tosyl chloride (first step) in CH₂Cl₂ proceeded quantitatively (rt, 1 h), however, diazotransfer (second step) with aqueous 11% NaOH at rt was slow (85 h). Subsequent aldol-type reaction (third step) with **3a** (5 equiv) gave the desired product **4a** in 70% yield (entry 1). Ethylester 1b was more reactive in diazo-transfer step and the reaction was completed within 9 h, and the aldoltype reaction gave 5a in 87% (entry 2). The solvent influenced the rate of diazo-transfer, for example, the reaction in diethyl ether enabled rapid conversion in diazotransfer and 5a was obtained in 82% yield through three steps (entry 3). Although a longer reaction time was required, benzylester 1c was also transformed to 2c in 25 h and subsequent C-C bond formation gave 6 in 51% overall yield (entry 4).

Next, we applied this three-step protocol to various aldehydes under optimized conditions (Table 2). The

Keywords: Aldol-type reaction; α -Diazo- β -hydroxyester; α -Amino- β -hydroxyester.

^{*} Corresponding authors. Tel.: +81 432902908; fax: +81 432902909; e-mail: arai@p.chiba-u.ac.jp

Table 1. One-pot synthesis of α-diazo-β-hydroxyesters under PTC conditions

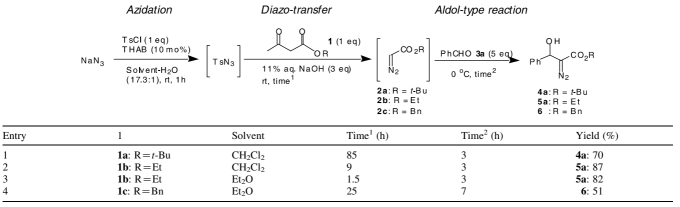


Table 2. One-pot reaction using various aldehydes

1 2

3

4

	1ststep:TsCI(1 eq),THAB (10 mol%) Et ₂ O - H ₂ O,rt,1 h 2ndstep: 1b (1 eq),11% aq.NaOH(3 eq) rt,1.5 h	OH → _ CO ₂Et
NaN ₃ -	➤ 3rd step : RCHO (3 , 5 eq),0 ^o C, time(h)	
		5

Entry	3: Aldehyde	Time (h)	Yield (%)
1	3b : $R = 4$ -MeO–C ₆ H ₄	2	5b : 73
2	3c : $R = 4 - CF_3 - C_6H_4$	12	5c : 76
3	3d : $R = 1$ -Napthyl	3	5d : 80
4	3e : $R = Ph(CH_2)_2$	3	5e : 86 ^a
5	3f: R = i-Bu	20	5f : 82
6	3g: R = i - Pr	3	5 g: 75
7	3h : $R = c$ -Hex	4	5h : 75
8	3i : $R = t$ -Bu	18	5i : 73

^a Three equivalent of **3e** was used.

aromatic aldehydes shown in entries 1-3 were smoothly transformed into the corresponding adducts **5b-d** in yields of 73-80%. In the case of aliphatic aldehydes including sterically hindered substrate such as 3i, the reactions also proceeded without any significant self-condensation with a range of 73–86% yield (entries 4–8).

2.2. Asymmetric aldol-type reaction using a chiral quaternary ammonium salt as a PTC

After succeeding with the one-pot synthesis of α -diazo- β hydroxyesters, we next investigated catalytic asymmetric synthesis. Only one example of a catalytic asymmetric

aldol-type reaction using α -diazoester has been reported.⁷ Initially, we surveyed this reaction using cinchoninium salts $(A-C^{9e})$ (Table 3). Although the enantioselectivities were poor, an N-anthracenyl group gave better results at 0 °C (entry 2). Moreover, a secondary hydroxy group was found to be essential in asymmetric induction, suggesting that hydrogen bonding between the catalyst and substrates is important (entry 2 vs 3). Since the cinchonidinium salt (PTC D) gave a slightly better result, the reaction conditions were further optimized using catalyst **D** (Table 4).

With a stronger base such as aqueous KOH, the reaction proceeded to give 4a at lower temperature (Table 4, entries

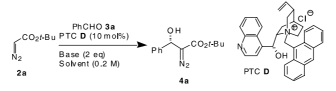
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Table 3. Catalytic asymmetric aldol-type reaction of 2a with 3a using various PTCs

	2a	CO2t-Bu PTC (10 mo%) 11% NaOH (2 Et ₂ O (0.2 M),) eq)	Ph ⁺ N ₂ ^{CO₂t-Bu N₂}	R ² O, R ¹ PTC A-C		
Entry	PTC	R^1	R^2	Х	Time (I	h) Yield (%)	ee (%) ^a
1	Α	Ph	Н	Br	2.5	51	6 (<i>S</i>)
2	В	9-Anthracenyl	Н	Cl	2	84	14 (R)
3	С	9-Anthracenyl	Allyl	Br	3	96	0
4	D		-		3	76	24 (S)

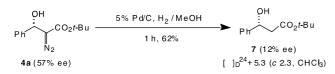
^a Determined by chiral HPLC analysis using DAICEL CHIRALCEL OD.

Table 4. Catalytic asymmetric aldol-type reaction of 2a with 3a using PTC D



Entry	Solvent	Base	3a (equiv)	Conditions	Yield (%)	ee (%)
1	Et ₂ O	25% KOH	5	-20° C, 8 h	70	25
2	Et ₂ O	50% KOH	5	-40° C, 2 h	74	39
3	Et ₂ O	50% KOH	5	-60° C, 16 h	65	20
4	Et_2O	50% KOH	1.5	−40°C, 14 h	96	48
5	Et ₂ O	50% RbOH	1.5	-40° C, 10 h	72	51
6	PhMe	50% KOH	1.5	−40°C, 38 h	83	45
7	PhMe	50% RbOH	1.5	−40°C, 10 h	96	56
8	PhMe	50% CsOH	1.5	-40° C, 20 h	76	56

1–3). Under biphase conditions of 50% KOH and Et₂O at -40 °C, 1.5 equiv of **3a** was enough for the reaction to proceed in 96% yield with moderate enantioselectivity (entry 4). The best result (96% yield, 56% ee)⁸ was obtained using 50% aqueous RbOH in toluene at -40 °C (entry 7). The absolute stereochemistry of **4a** was determined to be *S* by comparison with the reported optical rotation¹⁰ after diazo decomposition to β -hydroxyester **7** (Scheme 1 and Section 3).

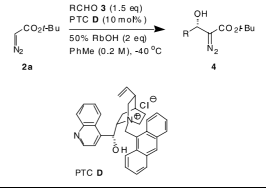


Scheme 1. Determination of the absolute stereochemistry of 4a.

Next, various aldehydes were used in this reaction under the optimized conditions. The electron density on the aromatic rings was found to strongly influence the enantioselectivity (Table 5). For example, **3b**, which has a 4-MeO group, gave a racemate, while **3c** which has an electron-withdrawing group 4-CF₃, gave **4c** in 81% yield with 73% ee (entries 1 and 2). 4-Alkylated substrates were also converted with lower ee (entries 4 and 5). 1- and 2-Naphthaldehydes were converted to **4d** and **4l** in respective yields of 86% (79% ee) and 94% (56% ee) (entries 3 and 6). In the case of aliphatic substrates, primary and secondary aldehydes such as **3e–h** gave 22–42% ee (entries 7–10), but pivalaldehyde **3i** was transformed to **4i** in 83% yield with 81% ee (entry 11).

The aldol-type reaction of α -diazoesters under basic media has been reported to include an equilibrium process,^{4f} so the time course of the chemical yield and ee of **4a** and **4i** during asymmetric reactions were investigated. As shown in Figure 1, in the case of aromatic aldehyde **3a** both the chemical yield and ee of **4a** gradually increased. The chemical yield reached equilibrium after 5 h and the ee remained at about 60% ee after 3 h. In the reaction of aliphatic aldehyde **3i**, the initial ee of **4i** was 70%, and this gradually increased to 80% ee after 5 h. The former result suggests the possibility of a retro-aldol reaction.¹¹ With regard to this reversible mechanism, enantioselection might occur in the differentiation of the carbonyl plane of

Table 5. Enantioselective synthesis of 4 with various aldehydes using PTC \boldsymbol{D}



Entry	Aldehyde 3	Time (h)	Yield (%)	ee (%)
1	3b : R=4-	120	4b : 56	0
	MeO-C ₆ H ₄			
2	3c : $R = 4 - CF_3 - C_6H_4$	140	4c: 81	73
3	3d : $R = 1$ -Napthyl	94	4d : 86	79
4	3j : $R = 4$ -Me $-C_6H_4$	18	4j : 66	39
5	3k : $R = 4$ -Bu–C ₆ H ₄	120	4k : 63	32
6	3l: R=2-Naphthyl	110	41 : 94	56
7	3e : $R = Ph(Ch_2)_2$	72	4e : 32	33
8	3f : $R = i$ -Bu	72	4f: 85	22
9	3g: R = i - Pr	20	4g: 53	42
10	$3\mathbf{\hat{h}}$: R = c-Hex	10	4h : 88	33
11	3i : $R = t$ -Bu	72	4i : 84	81

aldehydes in the C–C bond-forming step or the reversal retro-aldol step by kinetic resolution.

To test the latter possibility, (\pm) -4a was subjected to retroaldol conditions with 50% RbOH in toluene (10 h) in the presence of PTC **D**, and the formation of 2a and 3a was observed. However, the ee of the recovered 4a (72%) was very low (Scheme 2). This result suggests that the asymmetric induction of 4a occurs mainly not via kinetic resolution in the retro-aldol step but rather through the carbon–carbon bond-forming step. In the case of 4i, an alcoholic proton might be less acidic than benzyl alcoholic proton in 4a and the retro-aldol reaction seems to be disfavored. As outlined in Scheme 3, $k'_{\rm S}$ and $k'_{\rm R}$ are considered to be equal but the rate of C–C bond formation ($k_{\rm S}$) is greater than $k_{\rm R}$ in the reaction of 3a with 2a.

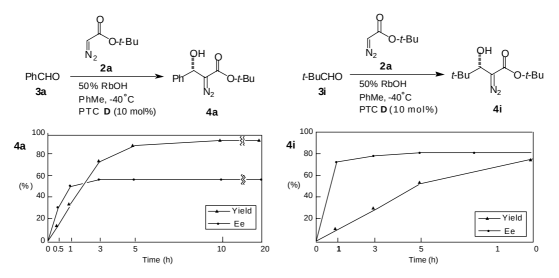
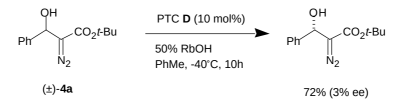
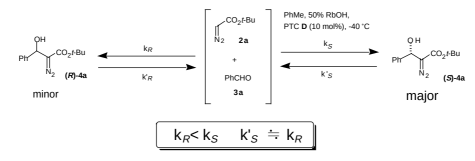


Figure 1. Time course of yield and ee in the reaction of 2a with 3a and 3i.



Scheme 2. Retro aldol-type reaction of (\pm) -4a with PTC D.



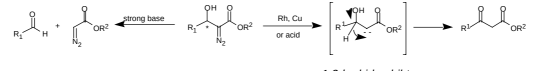
Scheme 3. Postulated reaction pathway for the catalytic asymmetric aldol-type reaction.

The presence of this retro-aldol process for racemic **4a** explains the increase in ee in the initial step of the reaction using **3a**. In the same way, a large excess of **3a** resulted in a lower ee (Table 4, entry 2 vs 4).

2.3. Transformation of optically active α-diazo-β-hydroxyesters to α-amino-β-hydroxyester derivatives

Many organic transformations using a diazo-functionality via diazo-decomposition have been reported¹² due to its

high reactivity with late transition metals. However, only limited examples of transformation using α -diazo- β hydroxyesters have been reported,¹³ since they are unstable under basic (retro-aldol reaction) and acidic (diazo decomposition) media. Furthermore, they gave simple 1,3dicarbonyl compounds via a 1,2-hydride shift by reacting with transition metals (Scheme 4). To establish a new synthetic transformation of α -diazo- β -hydroxyesters without any loss of chiral centers, we attempted the reduction of a diazo group to hydrazone or hydrazine as an amine



1,2-hydride-shiht

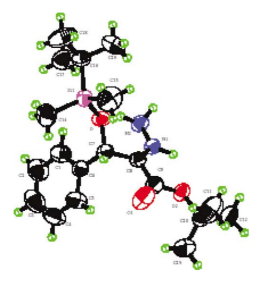


Figure 2. X-ray structure of (E)-10.

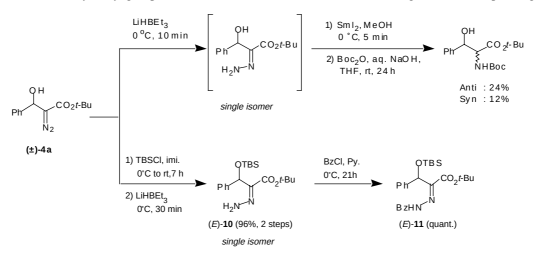
equivalent,^{14,15} which are known to be useful building blocks for the synthesis of polypeptides and biologically important molecules.^{16,17}

First, we attempted to convert (\pm) -4a to hydrazonoester without protection of its hydroxy group. Reduction with

LiHBEt₃ proceeded smoothly but the stability of the product was problematic and subsequent reduction with SmI₂ and protection with Boc₂O gave amino alcohols in low yields. Next, the initial protection of (\pm) -**4a** with TBSCl followed by reduction of the diazo group by LiHBEt₃ gave hydrazone **10** as a single isomer in 96% yield. Its configuration was confirmed by X-ray crystallographic analysis to be *E* (Fig. 2). Subsequent *N*-benzoylation (BzCl with pyridine in CH₂Cl₂ at 0 °C) gave (*E*)-**11** without significant isomerization¹⁸ (Scheme 5).

The further diastereoselective reduction of C=N bond was investigated (Table 6). The treatment of (*E*)-**11** with NaBH₄ at 0 °C in EtOH gave α -hydrazinoester *anti*-**12** in 92% yield, exclusively (entry 1). In the case of LiBH₄, a mixture of **12** and **13** was obtained with respective yields of 55 and 12%. However, no diastereoselectivities were observed in either product (entry 2). The reaction of Red-Al with **11** in toluene gave **13** as a separable diastereomixture in moderate yield, while no selectivity was observed (entry 3).

Further transformations of **12** and **13** are outlined in Scheme 6. N–N bond cleavage of *anti*-**12** with SmI₂ followed by protection with Boc₂O gave **14** (65% yield) as a mixture of *syn-* and *anti*-isomers (1:2.6) due to epimerization at the α -position.¹⁹ In the case of *syn-***13** and *anti-***13**, similar transformation gave the corresponding alcohols **15**

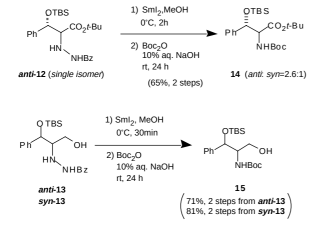


Scheme 5. Preparation of hydrazones.

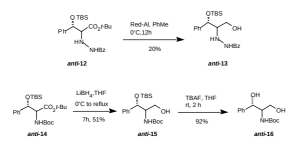
Table 6. Diastereoselective reduction of (E)-11

	OTBS Ph BzHN (<i>E</i>)-11	Reagents Conditions Solvent (0.2 M)	O TBS Ph HN NHBz 12	+ Ph OTBS HN NHBz 13	
Entry	Reagents (equiv)	Solvent	Conditions	$\frac{\text{Yield}}{12 (syn:anti)^{a}}$	d (%) 13 (syn:anti) ^a
1	NaBH ₄ (5)	EtOH	0°C, 2 h	92 (anti only)	0
2	$\text{LiBH}_4(5)$	THF	0° C to rt, 19 h	55 (1:1)	12 (1:1)
3	Red-Al (5)	PhMe	0°C, 2 h	0	60 (1:1)

^a Determined by ¹H MNR analysis.



Scheme 6. N-N bond cleavage of 12 and 13 by SmI₂.



Scheme 7. Determination of relative configuration.

Table 7. Diastereoselective reduction of hydrazones

	OTBS	1) Sml ₂ (6eq) Solvent (0.1M) 0 [.] C, 30min	01 	rBS CO₂t-Bu
	RHN ^N	2) Boc ₂ O,10%NaOH, rt, 24 h		NHBoc
	10 : R = H 11 : R = Bz			14
ntry	Substrat	e Solvent	Yield (%)	syn:anti
	10			

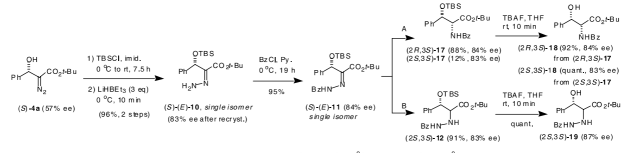
Entry	Substrate	Solvent	Yield (%)	syn:anti
1	10	MeOH	100	2:1
2	10	<i>i</i> -PrOH	100	3.3:1
3	11	MeOH	77	5.4:1
4	11	<i>i</i> -PrOH	100	6.7:1

in respective yields of 81 and 71%. The stereochemistry of *anti*-14 was determined by conversion to *anti*-16 (Scheme 7), the stereochemistry of which was confirmed by comparison to the literature via *anti*-15.²⁰ The relative stereochemistry of *anti*-12 was also confirmed by conversion to *anti*-13.

To enhance the utility of this synthetic protocol, we next investigated the diastereoselective one-pot transformation of hydrazones to amino groups (Table 7). For example, the reaction of (E)-10 with SmI_2^{19} in MeOH followed by protection with Boc₂O gave the desired product 14 in quantitative yield (syn/anti=2:1). Higher diastereoselectivity was observed when the reduction was carried out in isopropanol (entries 1 and 2). The reduction of *N*-benzoyl-hydrazone (*E*)-11 in isopropanol gave 14 in quantitative yield with better selectivity (syn/anti=6.7:1, entry 4).

After successfully developing an efficient transformation to aminoesters, we next investigated the synthesis of optically active aminoesters from (S)-4a (57% ee). Initial transformation gave siloxyhydrazone (E)-10, the ee of which was increased to 83% ee by recrystallization. Subsequent reduction of (S)-(E)-11 with SmI₂ followed by BzCl (condition A) gave (2R,3S)-17 and (2S,3S)-17 in respective yields of 88 and 12% without racemization. The reduction of (S)-(E)-11 with NaBH₄ (condition B) gave (2S,3S)-12 in 91% yield, exclusively, without any loss of optical purity. Treatment of these products with TBAF gave the corresponding hydroxyesters 18 and 19 (Scheme 8).

In summary, we have developed the one-pot synthesis of α -diazo- β -hydroxyesters with a single catalyst without any isolation of explosive intermediates. A PTC-catalyzed asymmetric aldol-type reaction using α -diazoester (up to 81% ee) with unique enantio enrichment and the transformations of α -diazo- β -hydroxyesters to α -amino- β -hydroxyesters in diastereoselective fashion were also established. This synthetic protocol provides a practical synthesis of optically active α -amino- β -hydroxyester derivatives, which have been recognized as useful building blocks for biologically important compounds or pharmaceuticals. Further studies of the application of this method are currently underway.



condition: A 1) Sml₂, *i*-PrOH, 0 ^oC, 30 min, 2) BzCl, Py. 0 ^oC, 10 min, separation

B NaBH₄, EtOH, 0 ^oC, 2h

3. Experimental

3.1. A general procedure for the one-pot synthesis of α -diazo- β -hydroxyester, synthesis of ethyl 2-diazo-3-hydroxy-3-phenylpropionate (5a) (Table 1, entry 3)

To a solution of TsCl (200 mg, 1.05 mmol) and THAB (45.5 mg, 0.1 mmol, 10 mol%) in diethylether (5.2 mL) was added NaN₃ (68.4 mg, 1.05 mmol) and water (0.3 mL) at rt. The mixture was stirred for 1 h and 1b (0.14 mL, 1.05 mmol) and 3 N NaOH (1.17 g, 3.15 mmol) were added with stirring for an additional 1.5 h. After α -diazoacetoacetate had diappeared, benzaldehyde **3a** (0.53 mL, 5.25 mmol) was added at 0 °C and the mixture was stirred for 3 h at 0 °C. The mixture was extracted with AcOEt (10 mL \times 3) and the combined organic layers were washed with brine, dried over Na₂SO₄ and concentrated in vacuo. Subsequent flash column chromatography (hexane/ AcOEt = 10:1) gave the desired product **5a** as a yellow oil (189.4 mg, 0.86 mmol, 82%) (reg.# 27262-59-5), ¹H NMR δ: (CDCl₃, 400 MHz) 1.28 (t, 3H, J = 6.8 Hz), 3.03 (br s, 1H), 4.29 (q, 2H, J=6.8 Hz), 5.92 (d, 1H, J=2.8 Hz), 7.31-7.49 (m, 5H).

3.1.1. *tert*-Butyl 2-diazo-3-hydroxy-3-phenylpropionate (4a). Yellow oil; IR (neat) ν : 3442, 2979, 2095, 1665 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.46 (s, 9H), 2.99 (br s, 1H), 5.87 (d, 1H, J=2.8 Hz), 7.30–7.46 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ : 165.7, 139.0, 128.5, 128.0, 125.6, 82.0, 68.5, 28.2; LRMS (FAB) *m/z*: 287 (M+K); HRMS (FAB) calcd for C₁₃H₁₆N₂O₃K 287.0798, found: 287.0807.

3.1.2. Benzyl 2-diazo-3-hydroxy-3-phenyl-propionate (6). Yellow oil; IR (neat) ν : 3425, 3032, 2101, 1682 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 2.90 (br s, 1H), 5.20 (s, 2H), 5.92 (d, 1H, J=3.2 Hz), 7.29–7.43 (m, 10H); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.1, 138.7, 135.6, 128.7, 128.5, 128.34, 128.32, 128.1, 125.6, 68.6, 66.6; LRMS (FAB) m/z: 321 (M+K); HRMS (FAB) calcd for C₁₆H₁₄N₂O₃K 321.0642, found: 321.0630.

3.1.3. Ethyl 2-diazo-3-hydroxy-3-(4-methoxyphenyl)propionate (5b) (reg.#39910-24-2). Yellow crystal; ¹H NMR (CDCl₃, 400 MHz) δ : 1.33 (t, 3H, *J*=7.2 Hz), 2.95 (br s, 1H), 3.84 (s, 3H), 4.32 (q, 2H, *J*=7.2 Hz), 5.90 (d, 1H, *J*=2.0 Hz), 6.94 (d, 2H, *J*=8.4 Hz), 7.38 (d, 2H, *J*=8.4 Hz).

3.1.4. Ethyl 2-diazo-3-hydroxy-3-(4-trifluoromethylphenyl)propionate (5c). Yellow crystal; mp 55–59 °C (hexane–CHCl₃); IR (thin film) ν : 3423, 3019, 2101, 1677 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.30 (t, 3H, J=7.2 Hz), 3.14 (br s, 1H), 4.28 (q, 2H, J=7.2 Hz), 5.97 (d, 1H, J=4.0 Hz), 7.57 (d, 2H, J=8.4 Hz), 7.65 (d, 2H, J= 8.4 Hz); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.2, 142.9, 130.3 (q, J=32.0 Hz), 126.0, 125.6 (q, J=3.3 Hz), 123.9 (q, J=270.7 Hz), 68.0, 61.4, 14.4; LRMS (FAB) *m/z*: 327 (M+K); HRMS (FAB) calcd for C₁₂H₁₁N₂O₃F₃K 327.0359, found: 327.0350. Anal. Calcd for C₁₂H₁₁F₃N₂O₃: C, 50.01; H, 3.85; N, 9.72. Found: C, 49.89; H, 3.80; N, 9.83. **3.1.5. Ethyl 2-diazo-3-hydroxy-(1-naphthyl)propionate** (5d). Yellow oil; IR (neat) ν : 3430, 3059, 2981, 2091, 1683 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.37 (t, 3H, J= 7.2 Hz), 3.09 (br s, 1H), 4.29–4.39 (m, 2H), 6.67 (d, 1H, J= 4.0 Hz), 7.55–7.66 (m, 3H), 7.87–8.02 (m, 4H); ¹³C NMR δ : (CDCl₃, 100 MHz) 166.3, 134.0, 133.5, 129.4, 128.86, 128.81, 126.3, 125.8, 125.2, 123.2, 122.4, 66.0, 61.2, 14.3; LRMS (FAB) m/z: 309 (M+K); HRMS (FAB) calcd for C₁₅H₁₄O₃N₂K 309.0642, found: 309.0672.

3.1.6. Ethyl 2-diazo-3-hydroxy-5-phenylpentanoate (5e). Yellow oil; IR (neat) ν : 3447, 3026, 2934, 2094, 1689 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.30 (t, 3H, J=7.2 Hz), 1.89–2.12 (m, 2H), 2.59 (br s, 1H), 2.70–2.88 (m, 2H), 4.26 (q, 2H, J=7.2 Hz), 4.66–4.70 (m, 1H), 7.19–7.31 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.4, 140.8, 128.3, 128.2, 125.9, 65.7, 60.9, 35.6, 31.7, 14.3; LRMS (FAB) *m/z*: 287 (M+K); HRMS (FAB) calcd for C₁₃H₁₆N₂O₃K 287.0798, found: 287.0779.

3.1.7. Ethyl 2-diazo-3-hydroxy-5-methylhexanoate (5f). Yellow oil; IR (neat) ν : 3433, 2959, 2094, 1685 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.95 (d, 6H, J=6.8 Hz), 1.28 (t, 3H, J=7.2 Hz), 1.36–1.43 (m, 1H), 1.59–1.68 (m, 1H), 1.72–1.82 (m, 1H), 2.42 (br s, 1H), 4.23 (q, 2H, J=7.2 Hz), 4.75–4.79 (m, 1H); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.6, 64.8, 60.9, 42.6, 24.5, 22.8, 22.0, 14.4; LRMS (FAB) m/z: 239 (M+K); HRMS (FAB) calcd for C₉H₁₆N₂O₃K 239.0798, found: 239.0780.

3.1.8. Ethyl 2-diazo-3-hydroxy-4-methylpentanoate (5g) (reg.# 38491-54-2). Yellow oil; ¹H NMR (CDCl₃, 400 MHz) δ : 0.95 (d, 3H, J=6.8 Hz), 1.07 (d, 3H, J=6.8 Hz), 1.29 (t, 3H, J=8.0 Hz), 1.85–1.94 (m, 1H), 2.48 (br s, 1H), 4.22–4.29 (m, 3H).

3.1.9. Ethyl 2-diazo-3-hydroxy-3-cyclohexylpropionate (5h) (reg.# 39910-21-9). Yellow oil; ¹H NMR (CDCl₃, 400 MHz) δ : 0.97–1.30 (m, 8H), 1.52–1.79 (m, 5H), 2.03 (d, 1H, J=12.4 Hz), 2.38 (br s, 1H), 4.24 (q, 2H, J=7.2 Hz), 4.30 (dd, 1H, J=5.2, 8.4 Hz).

3.1.10. Ethyl 2-diazo-3-hydroxy-4,4-dimethyl-valerate (5i) (reg.# 39910-22-0). Yellow oil; ¹H NMR (CDCl₃, 400 MHz) δ : 0.98 (s, 9H), 1.28 (t, 3H, J=6.8 Hz), 2.54 (br s, 1H), 4.19–4.27 (m, 3H).

3.2. Typical procedure for asymmetric synthesis of *tert*butyl 2-diazo-3-hydroxy-3-phenylpropionate (4a) using PTC D (Table 4, entry 7)

To a solution of benzaldehyde **3a** (56 µL, 0.53 mmol), *tert*butyl diazoacetate (50.0 mg, 0.35 mmol) and PTC **D** (18.2 mg, 0.035 mmol, 10 mol%) in toluene (1.8 mL) was added 50% RbOH (82.0 µL, 0.7 mmol) at -40 °C. The mixture was stirred for 10 h and partitioned between AcOEt and water. The aqueous layer was extracted with AcOEt (5 mL×3) and the organic layers were washed with brine, dried over Na₂SO₄ and concentrated in vacuo. Subsequent flash column chromatography (hexane/AcOEt=15:1) gave the desired product **4a** as a yellow oil (83.3 mg, 0.33 mmol, 96%). [α]_D²³ - 20.8 (*c* 1.06, CHCl₃, 56% ee); HPLC: DAICEL CHIRALCEL OD, 254 nm, flow rate 1.0 mL/min, hexane/*i*-PrOH=99:1, retention time: 20.7 min (major, *S*) and 23.1 min (minor, *R*).

3.2.1. *tert*-Butyl 2-diazo-3-hydroxy-3-(4-methoxyphenyl)propionate (4b). Yellow oil; IR (neat) ν : 3449, 2978, 2095, 1729, 1667 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.50 (s, 9H), 2.97 (br s, 1H), 3.81 (s, 3H), 5.82 (d, 1H, J= 2.0 Hz), 6.89–7.36 (m, 4H); ¹³C NMR (CDCl₃, 100 MHz) δ : 165.8, 159.4, 130.9, 127.0, 114.0, 82.0, 68.4, 55.2, 28.3; LRMS (FAB) *m/z*: 317 (M+K); HRMS (FAB) calcd for C₁₄H₁₈N₂O₄K 317.0904, found: 317.0875; HPLC: DAICEL CHIRALCEL OD, 254 nm, flow rate 0.5 mL/ min, hexane/*i*-PrOH=95:5, retention time: 20.8 and 24.2 min.

3.2.2. *tert*-Butyl 2-diazo-3-hydroxy-3-(4-trifluoromethylphenyl)propionate (4c). Yellow oil; IR (neat) ν : 3448, 2981, 2095, 1734, 1693 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.50 (s, 9H), 3.04 (br s, 1H), 5.91 (d, 1H, J=3.4 Hz), 7.55 (d, 2H, J=8.4 Hz), 7.65 (d, 2H, J=8.4 Hz); ¹³C NMR (CDCl₃, 100 MHz) δ : 165.5, 143.0, 130.3 (q, J=32.1 Hz), 126.2, 125.7 (q, J=4.1 Hz), 124.0 (q, J=271 Hz), 82.5, 68.3, 28.3; LRMS (FAB) *m/z*: 355 (M+K); HRMS (FAB) calcd for C₁₄H₁₅F₃N₂O₃K 355.0672, found: 355.0657; [α]_D²⁶ -21.7 (*c* 1.1, CHCl₃, 73% ee); HPLC: DAICEL CHIRALCEL OD, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=97:3, retention time: 14.2 min (major) and 16.1 min (minor).

3.2.3. *tert*-Butyl 2-diazo-3-hydroxy-3-(1-naphthyl)propionate (4d). Yellow solid (racemate); mp: 94 °C (hexane); IR (neat) ν : 3435, 2979, 2094, 1685 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.54 (s, 9H), 3.00 (br s, 1H), 6.59 (d, 1H J=2.4 Hz), 7.49–7.56 (m, 3H), 7.82–7.96 (m, 4H); ¹³C NMR (CDCl₃, 100 MHz) δ : 165.7, 134.1, 133.5, 129.5, 128.8, 128.7, 126.5, 125.7, 125.2, 123.2, 122.5, 82.1, 66.1, 28.5; LRMS (FAB) *m/z*: 337 (M+K); HRMS (FAB) calcd for C₁₇H₁₈N₂O₃K 337.0955, found: 337.0941. Anal. Calcd for C₁₇H₁₈N₂O₃: C, 68.44; H, 6.08; N, 9.39. Found: C, 68.30; H, 5.92; N, 9.51; optically active form (yellow oil), $[\alpha]_D^{25}$ –68.0 (*c* 0.7, CHCl₃, 79% ee); HPLC: DAICEL CHIRALCEL OD, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=97:3, retention time: 24.3 min (minor) and 37.4 min (major).

3.2.4. *tert*-Butyl 2-diazo-3-hydroxy-5-phenylpentanoate (4e). Yellow oil; IR (neat) ν : 3422, 3026, 2930, 2091, 1685 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.48 (s, 9H), 1.84–1.93 (m, 1H), 1.99–2.08 (m, 1H), 2.57 (br s, 1H), 2.67–2.75 (m, 1H), 2.79–2.86 (m, 1H), 4.61–4.65 (m, 1H), 7.19–7.31 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ : 165.9, 140.9, 128.4, 128.3, 125.9, 81.8, 65.7, 35.6, 31.8, 28.2; LRMS (FAB) *m*/*z*: 315 (M+K); HRMS (FAB) calcd for C₁₅H₂₀N₂O₃K 315.1111, found: 315.1100; $[\alpha]_D^{23}$ – 5.68 (*c* 0.3, CHCl₃, 33% ee); HPLC: DAICEL CHIRALCEL OD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=95:5, retention time: 16.7 min (major) and 19.0 min (minor).

3.2.5. *tert*-Butyl 2-diazo-3-hydroxy-5-methylhexanoate (**4f**). Yellow oil; IR (neat) ν : 3443, 2958, 2871, 2089, 1693 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.95 (d, 6H, J=6.4 Hz), 1.33–1.41 (m, 1H), 1.49 (s, 9H), 1.57–1.67 (m, 1H), 1.73–1.83 (m, 1H), 2.47 (br s, 1H), 4.70–4.75 (m, 1H); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.0, 81.7, 64.7, 42.6,

28.3, 24.5, 22.8, 21.9; LRMS (FAB) 267 (M+K); HRMS (FAB) calcd for $C_{11}H_{20}N_2O_3K$ 267.1111, found: 267.1097; $[\alpha]_D^{23} - 12.0$ (*c* 1.0, CHCl₃, 22% ee); HPLC: DAICEL CHIRALPAK AD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=99:1, retention time: 22.3 min (minor) and 23.9 min (major).

3.2.6. *tert*-Butyl 2-diazo-3-hydroxy-4-methylpentanoate (4g). Yellow oil; IR (neat) ν : 3448, 2966, 2090, 1669 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.94 (d, 3H, J=6.8 Hz), 1.06 (d, 3H, J=6.8 Hz), 1.49 (s, 9H), 1.82–1.94 (m, 1H), 2.58 (br s, 1H), 4.23 (dd, 1H, J=4.4, 8.0 Hz); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.1, 81.7, 72.3, 32.8, 28.3, 18.7, 18.6; LRMS (FAB) 253 (M+K); HRMS (FAB) calcd for C₁₀H₁₈N₂O₃K 253.0955, found: 253.0935; [α]_D²² – 10.3 (c0.6, CHCl₃, 42% ee); HPLC: DAICEL CHIRALPAK AD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH= 99:1, retention time: 32.2 min (minor) and 38.3 min (major).

3.2.7. *tert*-Butyl 2-diazo-3-hydroxy-3-cyclohexylpropionate (4h). Yellow oil; IR (neat) *v*: 3440, 2927, 2088, 1666 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.97–1.30 (m, 5H), 1.48 (s, 9H), 1.52–1.79 (m, 5H), 2.02 (d, 1H, *J*= 12.8 Hz), 2.38 (br s, 1H), 4.25 (dd, 1H, *J*=5.2, 8.4 Hz); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.1, 81.6, 71.2, 42.0, 29.1, 29.0, 28.3, 26.1, 25.8, 25.6; LRMS (FAB) *m/z*: 293 (M+K); HRMS (FAB) calcd for C₁₃H₂₂N₂O₃K 293.1268, found 293.1284; $[\alpha]_D^{19}$ – 3.4 (*c* 0.9, CHCl₃, 33% ee) HPLC: DAICEL CHIRALCEL OJ, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=99:1, retention time: 22.1 min (major) and 35.8 min (minor).

3.2.8. *tert*-Butyl 2-diazo-3-hydroxy-4,4-dimethylvalerate (4i). Yellow solid; mp 58–61 °C; IR (KBr) ν : 3469, 2960, 2103, 1647 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.97 (s, 9H), 1.47 (s, 9H), 2.75 (br s, 1H), 4.20 (s, 1H); ¹³C NMR (CDCl₃, 100 MHz) δ : 166.4, 81.6, 73.7, 38.3, 28.3, 25.6; LRMS (FAB) *m*/*z* 267 (M+K); HRMS (FAB) calcd for C₁₁H₂₀N₂O₃K 267.1111, found: 267.1104. Anal. Calcd for C₁₁H₂₀N₂O₃: C, 57.87; H, 8.83; N, 12.27. Found: C, 57.92; H, 8.83; N, 12.51; [α]_D²⁴ – 20.4 (*c* 1.1, CHCl₃, 81% ee); HPLC: DAICEL CHIRALPAK AD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=99:1, retention time: 25.7 min (minor) and 27.4 min (major).

3.2.9. *tert*-Butyl 2-diazo-3-hydroxy-3-(4-*tert* butylphenyl)propionate (4k). Yellow oil; IR (neat) ν : 3467, 2965, 2093, 1732, 1687 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.31 (s, 9H), 1.50 (s, 9H), 2.86 (br s, 1H), 5.83 (d, 1H, J= 3.6 Hz), 7.34 (d, 2H, J=8.0 Hz), 7.40 (d, 2H, J=8.0 Hz); ¹³C NMR (CDCl₃, 100 MHz) δ : 165.8, 151.1, 136.0, 125.5, 125.4, 81.9, 68.5, 34.5, 31.2, 28.3; LRMS (FAB) *m/z*: 343 (M+K); HRMS (FAB) calcd for C₁₇H₂₄N₂O₃K 343.1424, found: 343.1432; $[\alpha]_{D}^{26}$ -6.6 (*c* 0.9, CHCl₃, 32% ee); HPLC: DAICEL CHIRALCEL OD, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=97:3, retention time: 13.3 min (major) and 17.3 min (minor).

3.2.10. *tert*-Butyl 2-diazo-3-hydroxy-3-(2-naphthyl)propionate (41). Yellow oil; IR (neat) ν : 3432, 2978, 2095, 1667 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.51 (s, 9H), 3.21 (br s, 1H), 6.03 (d, 1H, J=2.4 Hz), 7.45–7.54 (m, 3H), 7.82–7.86 (m, 3H), 7.94 (s, 1H); ¹³C NMR (CDCl₃,

100 MHz) δ : 165.7, 136.2, 133.2, 133.1, 128.6, 128.1, 127.6, 126.3, 126.2, 124.7, 123.6, 82.2, 68.9, 28.3; LRMS (FAB) *m/z*: 337 (M+K); HRMS (FAB) calcd for C₁₇H₁₈N₂O₃K 337.0955, found: 337.0924; $[\alpha]_D^{25}$ – 28.2 (*c* 1.0, CHCl₃, 56% ee); HPLC: DAICEL CHIRALCEL OD, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=97:3, retention time: 26.7 min (major) and 28.4 min (minor).

3.3. Determination of the absolute configuration of 4a (Scheme 1)

Using a procedure similar to that of Wang et al.,⁷ **4a** was converted into the corresponding β -hydroxyester **7** by hydrogenation. To a solution of optically active **4a** (57% ee, 188.5 mg, 0.76 mmol) in MeOH (10.8 mL) was added 5% of palladium charcoal (54.0 mg) and the resulting suspension was stirred for 1 h under a hydrogen atmosphere. After being filtered through a Celite pad, the mixture was concentrated in vacuo. Purification of the crude mixture by flash column chromatography gave **7** as a colorless oil (104.3 mg, 0.47 mmol, 62%, 12% ee); $[\alpha]_{D}^{24} + 5.3$ (*c* 2.3, CHCl₃). Optical purity was determined by a chiral HPLC column using DAICEL CHIRALPAK AS, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=97:3, retention time: 15.9 min (major, *R*) and 17.3 min (minor, *S*).¹⁰

3.3.1. tert-Butyl 2-hydrazono-3-(tert-butyldimethylsilyloxy)-3-phenylpropionate (E)-10 (Schemes 5 and 8). To a solution of **4a** (1.80 g, 7.25 mmol) in DMF (24 mL) was added imidazole (2.16 g, 31.7 mmol) and TBSCl (1.62 g, 10.8 mmol) at 0 °C and the mixture was stirred for 1 h under the same conditions. After being stirred for 6 h at rt, the reaction mixture was diluted with H₂O (10 mL) and extracted with AcOEt ($20 \text{ mL} \times 3$). The combined organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Subsequent flash column chromatography (hexane/AcOEt=50:1) gave TBS ether as a yellow oil (2.52 g, 6.96 mmol, 96%). IR (neat) v: 2929, 2857, 2094, 1685 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.05 (s, 3H), 0.10 (s, 3H), 0.88 (s, 9H), 1.47 (s, 9H), 5.72 (s, 1H), 7.26–7.38 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ: 164.8, 141.6, 128.3, 127.5, 125.3, 81.4, 68.7, 28.3, 25.6, 18.1, -5.0, -5.3; LRMS (FAB) m/z: 401 (M+K); HRMS (FAB) calcd for $C_{19}H_{30}N_2O_3SiK$ 401.1663, found: 401.1659; $[\alpha]_D^{24}$ -15.1 (c 1.1, CHCl₃, 56% ee, for S isomer).

To a solution of the TBS ether (1.05 g, 2.9 mmol) in dry THF (41 mL) was added LiHBEt₃ (1 M solution of THF, 8.7 mL, 8.7 mmol) at 0 °C under an argon atmosphere, and the reaction mixture was stirred for 30 min. The reaction was quenched with cold water (10 mL) and the resulting organic layers were extracted with AcOEt (20 mL \times 3). The combined organic extracts were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt=5:1) gave (E)-10 as a white solid (1.06 g, 2.9 mmol, quant.); mp: 55-56 °C (nhexane, 83% ee); IR (neat) v: 3411, 3286, 2926, 1689, 1052 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.12 (s, 3H), 0.14 (s, 3H), 0.94 (s, 9H), 1.56 (s, 9H), 6.19 (s, 1H), 7.02 (br s, 2H), 7.23–7.40 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ: 164.0, 139.6, 136.6, 128.2, 127.2, 125.3, 81.2, 70.5, 28.1, 25.7, 18.2, -5.2, -5.3; LRMS (FAB) m/z: 365 (M+H); HRMS (FAB) calcd for $C_{19}H_{33}N_2O_3Si$ 365.2260, found: 365.2262. Anal. Calcd for $C_{19}H_{32}N_2O_3Si$: C, 62.60; H, 8.85; N, 7.68. Found: C, 62.63; H, 8.82; N, 7.77; $[\alpha]_{D}^{23} - 85.3$ (*c* 1.0, CHCl₃, 83% ee); HPLC DAICEL CHIRALPAK AD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=97:3, retention time: 16.3 min (major, *S*) and 18.5 min (minor, *R*).

3.3.2. tert-Butyl 2-(N-benzoylhydrazono)-3-(tert-butyldimethylsilyloxy)-3-phenylpropionate (11) (Scheme 5). To a solution of (E)-10 (298.5 mg, 0.82 mmol) in CH₂Cl₂ (8.2 mL) was added pyridine (0.40 mL, 4.9 mmol) and BzCl (0.19 mL, 1.6 mmol) at 0 °C and the reaction mixture was stirred for 21 h at 0 °C. After being diluted with water (10 mL), the mixture was extracted with CH_2Cl_2 (5 mL× 3). The resulting organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/ $Et_2O = 10:1$, hexane/AcOEt = 5:1) gave (E)-11 as a pale yellow oil (383.7 mg, 0.82 mmol, quant.). IR (neat) v: 3288, 2930, 1740, 1698, 1680, 1253, 1155, 836 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ: 0.14 (s, 3H), 0.18 (s, 3H), 0.95 (s, 9H), 1.58 (s, 9H), 6.31 (s, 1H), 7.23–7.54 (m, 9H), 7.83 (br s, 1H), 11.4 (br s, 1H); ¹³C NMR (CDCl₃, 100 MHz) δ: 162.9, 138.2, 132.1, 128.6, 128.4, 127.9, 127.0, 125.1, 82.7, 72.0, 27.7, 25.4, 17.9, -5.3, -5.4; LRMS (FAB) m/z: 469 (M+H); HRMS (FAB) calcd for C₂₆H₃₇N₂O₄Si 469.2523, found 469.2482: HPLC: DAICEL CHIRALCEL OD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=99:1, retention time: 18.8 min (major) and 23.7 min (minor); $[\alpha]_{\rm D}^{24}$ +4.60 (c 1.0, CHCl₃, 84% ee). All carbon signals were observed in DMSO at 120 °C, however, isomerization to (Z)-11 was observed.

Compound (*Z*)-**11** was obtained by acidic treatment of (*E*)-**11** as a colorless oil; IR (neat) ν : 3255, 2954, 2930, 2857, 1704, 1676, 1132 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 0.10 (s, 3H), 0.12 (s, 3H), 0.97 (s, 9H), 1.29 (s, 9H), 5.75 (s, 1H), 7.21–7.59 (m, 8H), 7.95 (d, 2H, J=7.2 Hz), 13.4 (br s, 1H); ¹³C NMR (CDCl₃, 100 MHz) δ : 163.7, 161.6, 142.5, 141.1, 132.4, 132.3, 128.6, 127.5, 126.7, 125.4, 84.0, 76.0, 27.5, 25.6, 18.0, -4.5, -5.2; LRMS (FAB) m/z: 469 (M+H); HRMS (FAB) calcd for C₂₆H₃₇N₂O₄Si 469.2523, found: 469.2494.

3.3.3. Reduction of (*E*)-**11**, *tert*-**butyl 2**-(*N*-**benzoyl-hydrazino**)-**3**-*tert*-**butyldimethylsilyloxy-3**-**phenylpropionate** (**12**) (**Table 6**, **entry 1**). To a solution of (*E*)-**11** (210.6 mg, 0.45 mmol) in EtOH (2.2 mL) was added NaBH₄ (83.6 mg, 2.2 mmol) at 0 °C and the solution was stirred for 2 h under the same conditions. The reaction was quenched with water (10 mL) and the mixture was concentrated in vacuo. The resulting mixture was extracted with AcOEt (5 mL×3) and the combined organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt=5:1) gave *anti*-**12** as a white amorphous solid (193.6 mg, 0.41 mmol, 92%). *syn*-**12** was synthesized under the conditions described in Table 6.

Compound *anti*-**12**. White amorphous solid; IR (KBr) ν : 3262, 2930, 2857, 1700, 1676, 1138 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.14 (s, 3H), 0.07 (s, 3H), 0.91 (s, 9H), 1.39 (s, 9H), 3.98 (t, J=4.4 Hz, 1H), 5.11 (d, J=

4.4 Hz, 1H), 5.36–5.39 (m, 1H), 7.26–7.68 (m, 11H); ¹³C NMR (CDCl₃, 100 MHz) δ : 169.4, 166.3, 140.3 132.9, 131.6, 128.5, 127.94, 127.90, 127.3, 126.7, 81.9, 74.9, 70.0, 27.9, 25.7, 18.1, -4.8, -5.1; LRMS (FAB) *m/z*: 471 (M+H); HRMS (FAB) calcd for C₂₆H₃₉N₂O₄Si 471.2679, found: 471.2671; HPLC DAICEL CHIRAPAK AS, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=95:5, retention time: 15.3 (minor: 2*R*,3*R*) and 23.9 (major: 2*S*,3*S*); [α]_D²⁴ + 10.1 (*c* 1.32, CHCl₃, 83% ee, (2*S*,3*S*)).

Compound *syn*-**12**. White amorphous solid; IR (neat) ν : 3306, 2923, 2853, 1717, 1669, 1097 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.23 (s, 3H), 0.07 (s, 3H), 0.90 (s, 9H), 1.20 (s, 9H), 3.92 (d, J=7.2 Hz, 1H), 4.88 (d, J=7.2 Hz, 1H), 5.66 (br s, 1H), 7.26–7.81 (m, 11H); ¹³C NMR (CDCl₃, 100 MHz) δ : 169.8, 166.5, 140.7, 132.8, 131.7, 128.6, 128.12, 128.10, 127.5, 126.8, 81.6, 75.6, 70.9, 27.7, 25.7, 18.1, -4.5, -5.0; LRMS (FAB) *m/z*: 471 (M+H); HRMS (FAB) calcd for C₂₆H₃₉N₂O₄Si 471.2679, found: 471.2639.

3.3.4. Reduction of (*E*)-11, synthesis of 2-(*N*-benzoyl-hydrazino)-1-*tert*-butyldimethylsilyloxy-1-phenylpropanol (13) (Table 6, entry 3). To a solution of (*E*)-11 (56.0 mg, 0.12 mmol) in toluene was added Red-Al (65% solution in toluene, 0.14 mL, 0.48 mmol) at 0 °C under an argon atmosphere and the solution was stirred for an additional 2 h. The reaction mixture was quenched with saturated Rochelle salt solution (2 mL) and MeOH (three portions). The resulting residue was extracted with CHCl₃ (5 mL×3) and the combined organic layers were dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt=3:1) gave a mixture of *syn*-13 and *anti*-13 isomers as a white amorphous solid (28.8 mg, 0.07 mmol, 60%). These isomers were separated by additional column chromatography (hexane:AcOEt).

Compound *anti*-**13**. White amorphous solid; IR (KBr) ν : 3256, 2927, 1635, 1251, 1065 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.21 (s, 3H), 0.06 (s, 3H), 0.86 (s, 9H), 3.08–3.13 (m, 1H), 3.71 (dd, 1H, *J*=6.4 Hz, 11.2 Hz), 3.89 (dd, 1H, *J*=2.8 Hz, 11.2 Hz), 4.66 (d, 1H, *J*=7.6 Hz), 7.29–7.41 (m, 8H), 7.47–7.52 (m, 3H); ¹³C NMR (CDCl₃, 100 MHz) δ : 167.7, 142.5, 132.2, 131.8, 128.5, 128.3, 127.8, 127.0, 126.7, 75.0, 68.6, 59.8, 25.6, 18.0, -4.6, -5.6; LRMS (FAB) *m/z*: 401 (M+H); HRMS (FAB) calcd for C₂₂H₃₃O₃N₂Si 401.2260, found: 401.2227.

Compound *syn*-**13**. White amorphous solid; IR (KBr) ν : 3301, 2929, 1635, 1061, 836, 701 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.18 (s, 3H), 0.04 (s, 3H), 0.87 (s, 9H), 3.07-3.11 (m, 1H), 3.44 (dd, 1H, *J*=6.8, 11.6 Hz), 3.60 (dd, 1H, *J*=2.8, 11.6 Hz), 3.83 (br s, 1H), 4.82 (d, 1H, *J*=6.0 Hz), 5.09 (br s, 1H), 7.27-7.70 (m, 11H); ¹³C NMR (CDCl₃, 100 MHz) δ : 167.7, 141.8, 132.5, 131.8, 128.5, 128.3, 127.8, 126.9, 126.7, 75.1, 68.5, 59.7, 25.7, 18.0, -4.5, -5.1; LRMS (FAB) *m/z*: 401 (M+H); HRMS (FAB) calcd for C₂₂H₃₃N₂O₃Si 401.2260, found: 401.2289.

3.3.5. *tert*-Butyl 2-(*tert*-buthoxycarbonylamino)-3-(*tert*-butyldimethylsilyloxy)-3-phenylpropionate (14) (Scheme 6). To a solution of *anti*-12 (89.3 mg, 0.19 mmol) in MeOH (1.9 mL) was added SmI₂ (0.1 M solution in THF, 4.2 mL,

0.42 mmol) at 0 °C under an argon atmosphere, and the solution was stirred for 2 h at 0 °C. The reaction mixture was quenched with water (1 mL) and concentrated in vacuo. To the crude residue was added THF (1.9 mL), Boc₂O (207 mg, 0.95 mmol) and 10% aqueous NaOH (380 mg, 0.95 mmol) and the resulting solution was stirred for 24 h at rt. The reaction mixture was then diluted with water (3 mL) and extracted with AcOEt (5 mL×3). The combined organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt = 100:1–20:1) gave *anti*-14 (less polar) as a colorless oil (40.9 mg, 0.09 mmol 47%) and *syn*-14 (more polar) as a colorless oil (15.6 mg, 0.03 mmol, 18%), respectively.

Compound *anti*-**14**. Colorless oil; IR (neat) ν : 3444, 2930, 2857, 1714 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.08 (s, 3H), 0.08 (s, 3H), 0.93 (s, 9H), 1.25 (s, 8/9H), 1.29 (br s, 1/9H) 1.46 (s, 9H), 4.25 (br s, 1/10H), 4.42 (dd, J=2.8, 7.6 Hz, 0.2/1H), 5.04 (br s, 0.8/1H), 5.15 (s, 8/9H), 5.42 (d, J=7.6 Hz, 1H); 7.20–7.41 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ : 168.1, 154.9, 141.0, 127.6, 127.0, 126.1, 81.7, 79.5, 75.5, 61.4, 28.3, 27.7, 25.7, 18.2, -4.8, -5.2; LRMS (FAB) *m/z*: 452 (M+H); HRMS (FAB) calcd for C₂₄H₄₂NO₅Si 452.2832, found: 452.2859.

Compound *syn*-**14**. Colorless oil; IR (neat) *v*: 3452, 2931, 2857, 1727, 1706 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.20 (s, 3H), 0.03 (s, 3H), 0.88 (s, 9H), 1.21 (br s, 2/9H), 1.35 (s, 7/9H) 1.45 (s, 7/9H), 1.49 (s, 2/9H), 4.07 (d, *J*= 9.6 Hz, 7/9H), 4.25 (dd, *J*=2.8, 9.6 Hz, 0.8/1H), 5.01 (d, *J*=9.6 Hz, 0.2/1H), 5.15 (d, *J*=2.8 Hz, 1H), 5.19 (d, *J*= 2.8 Hz, 0.8/1H), 7.22-7.30 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ : 169.8, 155.4, 141.0, 127.9, 127.6 126.5, 81.8, 79.3, 74.8, 61.0, 28.2 28.0, 25.7, 18.0, -4.4, -5.2; LRMS (FAB) *m/z*: 452 (M+H); HRMS (FAB) calcd for C₂₄H₄₂NO₅Si 452.2832, found: 452.2815.

2-tert-Buthoxycarbonylamino-1-tert-butyl-3.3.6. dimethylsilyloxy-1-phenylpropanol anti-15 (Scheme 6). To a solution of anti-13 (55.0 mg, 0.14 mmol) in MeOH (1.4 mL) was added SmI₂ (0.1 M solution in THF, 3.0 mL, 0.3 mmol) at 0 °C under an argon atmosphere and the reaction mixture was stirred for 30 min. After being quenched with water (1 mL), the resulting mixture was concentrated in vacuo. To the crude residue was added THF (1.4 mL), Boc₂O (90 mg, 0.41 mmol) and aqueous 10% NaOH (165 mg, 0.41 mmol), and the mixture was stirred for 24 h at rt. After the mixture was diluted with water (3.0 mL), it was extracted with AcOEt (5 mL \times 3). The combined organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt=5:1) gave anti-15 as a white solid (36.6 mg, 0.10 mmol, 71%). As described above, syn-15 was synthesized from syn-13 in 81% yield (two steps).

Compound *anti*-**15**. White solid; mp 136 °C (hexane); IR (neat) ν : 3341, 3232, 2926, 1671, 1056 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.09 (s, 3H), 0.08 (s, 3H), 0.93 (s, 9H), 1.47 (s, 9H), 2.97 (d, J=10.8 Hz, 1H), 3.39–3.46 (m, 1H), 3.60 (br s, 1H), 3.84 (d, J=10.8 Hz, 1H), 5.19 (s, 1H), 5.48 (d, 1H, J=8.0 Hz) 7.20–7.40 (m, 5H); ¹³C NMR

(CDCl₃, 100 MHz) δ : 155.6, 141.0, 128.2, 127.4, 125.8, 79.5, 77.6, 61.2, 56.5, 28.4, 25.8, 18.0, -4.9, -5.3; LRMS (FAB) *m*/*z*: 382 (M+H); HRMS (FAB) calcd for C₂₀H₃₆NO₄Si 382.2414, found: 382.2408. Anal. Calcd for C₂₀H₃₅NO₄Si: C, 62.95; H, 9.25; N, 3.67. Found: C, 62.74; H, 9.49; N, 3.59.

Compound *syn*-**15**. Colorless oil; IR (neat) *v*: 3447, 2929 2857, 1697 1496 1167 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.15 (s, 3H), 0.06 (s, 3H), 0.90 (s, 9H), 1.36 (s, 9H), 2.35 (br s, 1H), 3.60–3.77 (m, 3H), 4.90 (br s, 1H), 4.91 (d, 1H, *J*=3.6 Hz), 7.22–7.33 (m, 5H); ¹³C NMR (CDCl₃, 100 MHz) δ : 156.1, 141.5, 128.1, 127.5, 126.3, 79.5 73.7, 63.0, 58.5, 28.3, 25.8, 18.1, -4.6, -5.2; LRMS (FAB) *m/z*: 382 (M+H); HRMS (FAB) calcd for C₂₀H₃₆NO₄Si 382.2414, found: 382.2448.

3.4. Direct synthesis of 14 via a one-pot procedure (Table 7, entry 4)

To a solution of (E)-11 (75.0 mg, 0.16 mmol) in isopropanol (1.6 mL) was added SmI₂ (0.1 M solution in THF, 9.6 mL, 0.96 mmol) at 0 °C under an argon atmosphere and the mixture was stirred for 30 min. After being guenched with water (1.0 mL), the resulting mixture was concentrated in vacuo. To the crude residue was added THF (1.6 mL), Boc₂O (104 mg, 0.48 mmol) and 10% aqueous NaOH (192 mg, 0.48 mmol), and the mixture was stirred for 24 h at rt. After dilution with water (3.0 mL), the mixture was extracted with AcOEt (5 mL \times 3). The combined organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt = 100:1-20:1) gave anti-14 (less polar) as a colorless oil (9.3 mg, 0.02 mmol, 13%) and syn-14 (more polar) as a colorless oil (62.7 mg, 0.14 mmol, 87%), respectively.

3.4.1. Direct synthesis of 17 via a one-pot procedure (Scheme 8, condition A). To a solution of (S)-(E)-11 (113.0 mg, 0.24 mmol) in isopropanol (2.4 mL) was added SmI₂ (0.1 M solution in THF, 14.4 mL, 1.44 mmol) at 0 °C under an argon atmosphere and the mixture was stirred for 30 min. After being quenched with water (5.0 mL), the resulting mixture was extracted with AcOEt ($10 \text{ mL} \times 3$). The combined organic layers were washed with brine dry over MgSO₄ and concentrated in vacuo. To the crude residue was added CH₂Cl₂ (2.4 mL), BzCl (41 µL, 0.36 mmol) and pyridine (38 µL, 0.48 mmol), and the mixture was stirred for 10 min at 0 °C. After being diluted with water (1.0 mL), the mixture was extracted with AcOEt (5 mL \times 3). The combined organic layers were washed with brine, dried over MgSO₄ and concentrated in vacuo. Flash column chromatography (hexane/AcOEt = 100:1-15:1) gave (2S,3S)-17 (less polar) as a colorless oil (13.7 mg, 0.030 mmol, 12%) and (2R,3S)-17 (more polar) as a white solid (95.9 mg, 0.21 mmol, 88%), respectively.

3.4.2. *tert*-Butyl 2-benzoylamino-3-(*tert*-butyldimethylsilyloxy)-3-phenylpropionate (17) (Scheme 8). Compound (2*S*,3*S*)-17. Colorless oil; IR (neat) ν : 3434, 2929, 2857, 1725, 1661 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.10 (s, 3H), -0.01 (s, 3H), 0.93 (s, 9H), 1.30 (s, 9H), 4.88 (dd, 1H, J=2.4, 6.8 Hz), 5.33 (d, 1H, J=2.4 Hz), 7.13 (d, 1H, J=6.8 Hz), 7.24–7.85 (m, 10H); ¹³C NMR (CDCl₃, 100 MHz) δ : 168.2, 166.4, 141.1, 133.9, 131.7, 128.6, 127.7, 127.1 126.9, 126.1, 82.2, 75.3, 60.9, 27.8, 25.7, 18.2, -4.7, -5.1; LRMS (FAB) *m/z*: 456 (M+H); HRMS (FAB) calcd for C₂₆H₃₈NO₄Si 456.2570, found: 456.2615; HPLC: DAICEL CHIRALCEL OD-H, 245 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=96:4, retention time: 8.0 min (minor) and 14.8 min (major); $[\alpha]_{D}^{25}$ +40.6 (*c* 0.9, CHCl₃, 84% ee).

Compound (2*R*,3*S*)-**17**. White solid; mp: 68 °C; IR (neat) ν : 3448, 2954, 2929, 2856, 1728, 1668 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : -0.15 (s, 3H), 0.09 (s, 3H), 0.94 (s, 9H), 1.47 (s, 9H), 4.76 (dd, 1H, *J*=2.8, 8.8 Hz), 5.31 (d, 1H, *J*=2.8 Hz) 6.82 (d, 1H, *J*=8.8 Hz), 7.23–7.74 (m, 10H); ¹³C NMR (CDCl₃, 100 MHz) δ : 169.3, 167.0, 140.8, 134.4, 131.4, 128.5, 128.0, 127.8, 126.8, 126.2, 82.2, 74.4, 60.2, 28.0, 25.7, 18.0, -4.4, -5.2; LRMS (FAB) *m/z*: 456 (M + H); HRMS (FAB) calcd for C₂₆H₃₇NO₄Si; C, 68.53; H, 8.18; N, 3.07. Found: C, 68.42; H, 8.39; N, 2.99; HPLC: DAICEL CHIRALCEL OD-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=96:4, retention time: 9.9 min (major) and 13.2 min (minor); $[\alpha]_D^{25} + 52.3$ (*c* 1.0, CHCl₃, 83% ee).

3.4.3. tert-Butyl 2-(N-benzoylamino)-3-hydroxy-phenylpropionate (18) (Scheme 8). Compound (2R,3S)-18. To a solution of (2R,3S)-17 (71.0 mg, 0.16 mmol) in THF (0.75 mL) was added TBAF (1 M solution of THF, 0.23 mL, 0.23 mmol) at rt under an argon atmosphere and the reaction mixture was stirred for 10 min. After being quenched with water (1.0 mL), the reaction mixture was extracted with AcOEt (5.0 mL \times 3). The combined organic layers were washed with brine, dried over MgSO4 and concentrated in vacuo. Flash column chromatography gave (hexane/AcOEt = 2:1)(2R, 3S)-18 (49.2 mg, 0.14 mmol, 92%) as a white solid; mp: 134-138 °C (hexane-AcOEt); IR (neat) v: 3368, 2977, 1731, 1639, 1521, 1149 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.42 (s, 9H), 3.13 (br s, 1H), 4.96 (dd, 1H, J=3.6, 7.6 Hz), 5.26 (br s, 1H), 6.85 (d, 1H, J=7.6 Hz), 7.27–7.72 (m, 10H); ¹³C NMR (CDCl₃, 100 MHz) δ: 169.3, 167.7, 139.8, 133.8, 131.6, 128.4, 128.3, 128.0, 127.0, 126.1, 82.8, 74.5, 59.0, 27.8; LRMS (FAB) m/z: 342 (M+H); HRMS (FAB) calcd for C₂₀H₂₄NO₄ 342.1705, found: 342.1689. Anal. Calcd for C₂₀H₂₃NO₄: C, 70.36; H, 6.79; N, 4.10. Found: C, 70.34; H, 6.83; N, 4.10; HPLC: DAICEL CHIRALCEL OJ-H, 254 nm, flow rate 0.5 mL/min, hexane/i-PrOH=93:7, retention time: 19.7 min (minor) and 22.6 min (major); $[\alpha]_{\rm D}^{24}$ + 36.1 (*c* 1.04, CHCl₃, 98.1% ee).

Compound (2*S*,3*S*)-18. According to the procedure described above, (2*S*,3*S*)-18 was synthesized as a white solid from (2*S*,3*S*)-17. White solid; mp: 134–138 °C (hexane–AcOEt); IR (neat) ν : 3427, 3322, 2925, 1735, 1636, 1159 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.46 (s, 9H), 4.86 (br s, 1H), 5.15 (dd, 1H, *J*=2.4, 6.0 Hz), 5.39 (s, 1H), 6.95 (d, 1H, *J*=6.0 Hz), 7.24–7.63 (m, 10H); ¹³C NMR (CDCl₃, 100 MHz) δ : 168.7, 168.2, 139.3, 133.1, 132.1, 128.6, 128.1, 127.8, 127.1, 126.0, 83.6, 75.5, 60.2, 27.9; LRMS (FAB) *m/z*: 342 (M+H); HRMS (FAB) calcd for C₂₀H₂₄NO₄ 342.1705, found: 341.1676. Anal. Calcd for C₂₀H₂₃NO₄: C, 70.36; H, 6.79; N, 4.10. Found: C, 70.06; H,

6.82; N, 4.04; HPLC: DAICEL CHIRALCEL OJ-H, 254 nm, flow rate 0.5 mL/min, hexane/*i*-PrOH=95:5, retention time: 24.4 min (minor) and 28.5 min (major); $[\alpha]_D^{25}$ + 69.8 (*c* 0.13, CHCl₃, 96.5% ee).

3.4.4. (2*S*,3*S*) *tert*-Butyl 2-(*N*-benzoylhydrazino)-3hydroxy-phenylpropionate (19), (Scheme 8). According to the procedure described above, (2*S*,3*S*)-19 was synthesized as a white amorphous material from (2*S*,3*S*)-12. White amorphous; IR (neat) ν : 3296, 2977, 1721, 1641, 1152 cm⁻¹; ¹H NMR (CDCl₃, 400 MHz) δ : 1.32 (s, 9H), 3.94 (d, 1H, *J*=4.8 Hz), 5.11 (d, 1H, *J*=4.8 Hz), 5.25 (br s, 1H), 7.27–7.69 (m, 10H), 7.94 (br s, 1H); ¹³C NMR (CDCl₃, 100 MHz) δ : 169.5, 167.6, 139.6, 132.2, 132.0, 128.6, 128.1, 127.7, 126.9, 126.4, 82.6, 72.6, 69.3, 27.8; LRMS (FAB) *m/z*: 395 (M+K); HRMS (FAB) calcd for C₂₀H₂₄N₂O₄K 395.1373, found: 395.1383; HPLC: DAICEL CHIRALPAK AS-H, 254 nm, flow rate 1.0 mL/min, hexane/*i*-PrOH=80:20, retention time: 15.2 min (minor) and 28.2 min (major); [α]²³_D + 15.9 (*c* 0.65, CHCl₃, 87% ee).

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Lewis and protic acid mediated Nicholas reactions of 3-acetoxycyclohept-1-en-4-ynedicobalt hexacarbonyl: site selectivity of nucleophile incorporation

Joseph DiMartino and James R. Green*

Department of Chemistry and Biochemistry, University of Windsor, Windsor, Ont., Canada N9B 3P4

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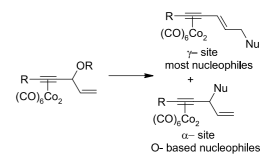
Abstract—Nicholas reactions on the cation derived from the cyclic allylic acetate alkynedicobalt complex **1** favour the γ -site kinetically for most nucleophiles, with increasing amounts of α -products in cases with greater nucleophilicity. Some regiocontrol in introduction of a specific nucleophilic fragment is possible by using different nucleophiles. Under conditions where reversibility is possible, the thermodynamically favoured site is exclusively γ -.

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1. Introduction

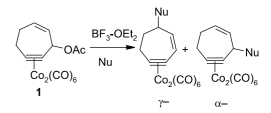
Propargyl cation dicobalt hexacarbonyl complexes are one of the most widely employed transition metal stabilized reactive intermediates in organic synthesis; their chemistry is often referred to as the Nicholas reaction.¹ These cations, which may stem from alkynedicobalt complexes with propargylic leaving groups and a protic or Lewis acid, or from enyne– $Co_2(CO)_6$ complexes and an electrophile,² normally substitute exclusively at the propargylic site, unless the cation is also allylic. In these allylic/propargylic situations, substitution has been found to occur predominantly at the site remote to the alkyne– $Co_2(CO)_6$ unit (γ site).³ Exceptions exist, however, particularly where intramolecular nucleophilic attack reactions are entropically driven towards the α -site;⁴ in some cases with nucleophiles, which are oxygen based, α -substitution is also observed (Scheme 1). 3a

While previous studies of Nicholas reactions of allylic substrates have been focussed on acyclic cations or cyclization reactions, the analogous question for cyclic cations has not been addressed to our knowledge. We have interest in this matter from several perspectives. Our group, and other groups, have been interested in the preparation and reactivity of cycloheptynedicobalt complexes.^{6,7,8} We have been able to incorporate nucleophiles γ - with



Scheme 1.

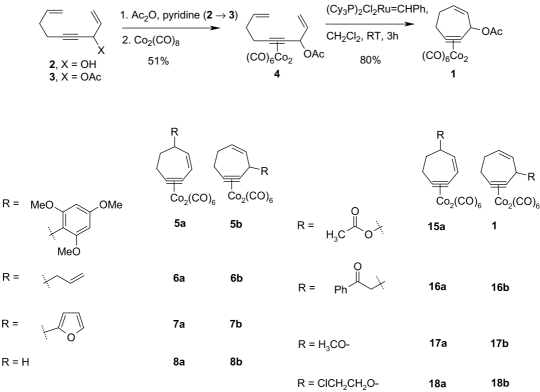
respect to the alkynedicobalt unit in tandem 4+3 cycloaddition/trapping reactions, but the list of participating nucleophiles in the process is quite restricted.^{6a} Substitution at the remote (γ -) position in the cycloheptenyne–Co₂(CO)₆ complexes (Scheme 2) would open up the ability to employ the now nucleophilic alkene function in annulation reactions with any highly electrophilic groups contained within the γ -substituent, ultimately giving fused 7,5- and 7,6- ring systems. In addition, we have an interest in clean





Keywords: Nicholas reaction; Cobalt–alkyne complexes; Cycloheptyne; Propargyl cations.

^{*} Corresponding author. Tel.: +519 253 3000x3545; fax: +519 973 7098; e-mail: jgreen@uwindsor.ca



10a

11a

12a

13a

X = OH

X = Cl

OAc

X = OAc

7b $R = H_{3}CO-$ 8b $R = CICH_{2}CH_{2}O-$ 17a 8b $R = CICH_{2}CH_{2}O-$ 18a 10b 11b R = CI- O19a 12b 13b R = O17a 20a

Figure 1. Nicholas reaction products of 1.

R =

R =

Scheme 3.

 α -substitution reactions on these complexes for facilitation of cycloaddition reactions employing the alkynedicobalt function.⁹ As a result, we have deemed it of importance to study the Nicholas substitution reactions of cycloheptyne–allyl acetate complex **1**, with a range of nucleophiles.

2. Results and discussion

Cycloheptyne–allyl acetate complex **1** was prepared in straightforward fashion from the known allyl propargyl alcohol **2** (Scheme 3).¹⁰ Standard acetylation of **2**, affording acetate **3**, followed by complexation with $Co_2(CO)_8$, gave **4** (51% yield, two steps). Ring closing metathesis, employing 10 mol% of $(Cy_3P)_2Cl_2Ru=CHPh$ (Grubbs' I catalyst), afforded **1** in 80% yield.¹¹

With the desired substrate in hand, we chose to investigate its reaction with 1,3,5-trimethoxybenzene in order to optimize the conditions of reaction. In CH₂Cl₂ solvent (0.05 M), and with excess BF₃–OEt₂ present (10 equiv), **1** underwent reaction with 1,3,5-trimethoxybenzene at temperatures as low as -30 °C to give mixtures of the γ -substitution (C-7 substitution) product **5a** and the α -substitution (C-3 substitution) product **5b** (Fig. 1). Variation of reaction temperature revealed that the γ -substitution product predominated in all cases, with optimal yields of condensation products realized at -10 °C (Table 1) with BF₃–OEt₂ as Lewis acid. Curiously, the amount of α -substitution decreased with increasing temperature, from 41% of the products -30 °C to 14% of the product composition at 23 °C. Changing the Lewis acid from BF₃–OEt₂ to SnCl₄ gave similar results at -10 °C, with a marginally inferior yield. Use of Bu₂BOTf as Lewis acid, however, caused extensive unproductive decomposition, even at -30 °C. As a result, the -10 °C, BF₃–OEt₂ combination was chosen as the standard set of conditions and applied in all other cases.

19b

20b

Table 1. Reaction of 1 with 1,3,5-trimethoxybenzene

Conditions	Yield 5a/5b (%)	γ-:α-Ratio
$BF_3-OEt_2, -30 \degree C$	70	59:41
BF_3-OEt_2 , -10 °C	86	70:30
BF ₃ –OEt ₂ , 0 °C	73	81:19
BF ₃ -OEt ₂ , 23 °C	52	86:14
$SnCl_4$, -10 °C	77	76:24
Bu₂BOTf, −30 °C	0	_

The change in isomer ratio towards increased amounts of the major, γ -substitution product at higher reaction temperatures suggested the possibility that the results with 1,3,5-trimethoxybenzene were not the consequence of purely kinetic reactivity of the propargyl allyl cation. Past work in our group has shown evidence of reversibility in Nicholas reactions involving this nucleophile,¹² and these results would be consistent with that feature here. In fact, subjecting purified α -substitution product **5b** to the 0 °C conditions of reaction (without added 1,3,5-trimethoxybenzene) afforded a **5a/5b** mixture (23:77, 67% recovery) along with some decomposition. By contrast, subjecting 5a to these conditions gave only recovered 5a. Consequently, allyltrimethylsilane was also investigated as a nucleophile with 1 under varying reaction temperatures (Table 2), as reversibility in this reaction is far less likely. Under analogous concentration and stoichiometry conditions, allyltrimethylsilane afforded γ -substitution product **6a** and α -substitution product **6b**. Once again the yield reached a maximum at -10 °C, but in these cases the α -: γ -product ratios remained relatively consistent (81:19-84:16) over the temperature range investigated.

Table 2. Reaction of 1 with allyltrimethylsilane

Conditions	Yield 6a/6b (%)	γ-:α-Ratio
BF_3 -OEt ₂ , -30 °C	68	82:18
BF_3 -OEt ₂ , -10 °C	83	84:16
BF ₃ -OEt ₂ , 0 °C	77	81:19
BF ₃ –OEt ₂ , 23 °C	56	83:17

Several other carbon and hydride based nucleophiles were investigated (Table 3). Allyltributylstannane gave 6a and 6b in good yield (74%), but with minimal γ -: α -selectivity (6a:6b=50:50). Conversely, furan gave condensation product 7a through its C-2 site, with almost none of α-condensation product 7b in evidence (62% yield, 7a:7b = >96:<4).¹³ The overall reduction products 8a and **8b** could be obtained in fair yield using triethylsilane (54%, **8a:8b**=63:37) or triisopropylsilane (62% yield, 8a:8b=84:16). The 2-hydroxymethyl-, 2-chloromethyl-, and 2-acetoxymethyl-substituted allylsilanes (9a, 9b, and 9c, respectively) (Fig. 2) afforded analogous products 10a/b, 11a/b, and 12a/b, respectively, with somewhat lower γ -: α -ratios (59:41–72:28) relative to allyltrimethylsilane itself. Homoenolate equivalent 1-trimethylsilylallyl acetate gave the enol acetate products 13a and 13b (as Z-/Eisomeric mixtures) with relatively high γ -selectivity (65%) yield, 13a:13b=89:11), along with small amounts of elimination product 14 (7%) and γ -acetoxy substitution

Table 3. Reaction of 1 with carbon and hydrogen nucleophiles^a

product 15a (7%). To our knowledge, this is the first example of a discrete homoenolate equivalent participating directly in a Nicholas reaction, although the cyclizationrearrangement processes of Tanino¹⁴ and Magnus' cycliza-tion-dyotropic rearrangements¹⁵ may be considered specialized cases of homoenolate equivalent reactivity. In addition, complexes with analogous functional group connectivity have been made by radical reactions on enyne complexes.¹⁶ Finally, two acetophenone enolate equivalents were introduced. The trimethylsilyl enol ether of acetophenone underwent reaction with 1 to give 16a and **16b** in good yield (74%), but the α -condensation product actually predominated slightly with this nucleophile (16a:16b=44:56). The enol acetate of acetophenone gave somewhat lower yields (61%, with 19% of 15a), with the γ -product once again as the major regioisomer (16a:16b=72:28).

Investigation of heteroatom based nucleophiles was also warranted due to the likelihood of reversibility in the substitution process (Table 4). Under standard conditions, acetic acid could be incorporated with great facility to give **15a** in good yield (79%) exclusively as the γ -substitution product. In this case, abandonment of the standard conditions in favour of neat acetic acid and H₂SO₄ gave superior results (97% yield) for 15a. Under the standard conditions, methanol, 2-chloroethanol, and 4-chloro-2buten-1-ol gave 17a (65%), 18a (59%), and 19a (68%), each exclusively as the γ -substitution products. The latter two cases also gave modest amounts of elimination product 14 and γ -acetoxy substitution product 15a. Again, use of a large excess of nucleophile and H₂SO₄ gave yield improvement for each of the commercially available alcohols (17a, 87%; 18a, 76%). Attempts to incorporate a nitrogen based nucleophile, acetamide, met with little success under the standard reaction conditions. While a small amount of γ -substitution product 20a could be obtained (12% yield), the major resulting product was γ -acetoxy substituted **15a** (83% yield); a small amount of elimination product 14 (5% yield) also could be isolated. Conversely, good yields of 20a (85%) could be realized by resorting to the addition of H_2SO_4 to a solution of 1 in CH₃CN. In no cases have we observed even traces of the heteroatom based α -condensation products 1, 17b–20b as a result of these protic- or Lewis acid mediated reactions.

Nucleophile	Product	Yield (%)	γ-:α-Ratio	15a (%)	14 (%)
1,3,5-Trimethoxybenzene	5a/5b	86	70:30		
Allyltrimethylsilane	6a/6b	83	84:16		
Allyltributylstannane	6a/6b	74	50:50		
Furan	7a/7b	62	>96:4		
Et ₃ SiH	8a/8b	54	72:28		
ⁱ Pr ₃ SiH	8a/8b	62	84:16	3.5	
9a	10a/10b	76	59:41		
9b	11a/11b	70	72:28		
9c	12a/12b	76	64:36		
1-Trimethylsilylallyl acetate	13a/13b	65	89 ^b :11 ^c	7	7
$H_2C = C(OSiMe_3)Ph$	16a/16b	74	44:56		
$H_2C = C(OAc)Ph$	16a/16b	61	72:28	19	

^a Reaction conditions: nucleophile, 1.5–2.0 equiv; solvent, CH_2Cl_2 (0.05 M); temperature, -10 °C; Lewis acid, BF_3 – OEt_2 (10 equiv); reaction time, 1 h. ^b Compound **13a** (*E*-:*Z*-)=38:62.

^c Compound **13b** (*E*-:*Z*-)=51:49.

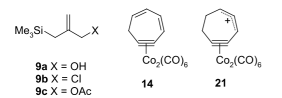


Figure 2.

Table 4. Reaction of 1 with heteroatom nucleophiles^a

Nucleophile	Product	Yield (%)	15a (%)	14 (%)
CH ₃ CO ₂ H	15a	79		
CH ₃ CO ₂ H	15a	97 ^b		
CH ₃ OH	17a	65		
CH ₃ OH	17a	87 ^b		
2-Chloroethanol	18a	59	15	15
2-Chloroethanol	18a	76 ^b		
4-Chloro-2-buten-1-ol	19a	68	13	4
CH ₃ C(O)NH ₂	20a	12	83	5
CH ₃ CN	20a	85 ^b		

^a Reaction conditions, unless otherwise stated: nucleophile, 1.5–2.0 equiv; solvent, CH₂Cl₂ (0.05 M); temperature, -10 °C; Lewis acid, BF₃–OEt₂ (10 equiv); reaction time, 1 h.

^b Using H₂SO₄ in place of BF₃–OEt₂ and excess nucleophile.

With the ready availability of γ -acetoxy substitution product **15a**, and the belief that the same cation could be generated from this compound as from **1**, we briefly explored its BF₃–OEt₂ induced Nicholas reactions. Under the otherwise standard conditions, allyltrimethylsilane reacted with **15a** to give **6a** and **6b** (81% yield) in the same ratio as from **1** (**6a**:**6b**=84:16), strongly suggesting an identical reactive intermediate from the two allyl acetate complexes. Compound **15a** also reacted with 1,3,5-trimethoxybenzene, affording **5a** and **5b** in 80% yield (**5a**:**5b**=76:24).

The distinction of γ - from α -adducts was readily apparent from the ¹H NMR spectra. Noteworthy in this respect were the resonances attributable to the vinyl proton adjacent to the alkyne–Co₂(CO)₆ unit in the γ -regioisomer, which appeared as a doublet $(J \approx 10 \text{ Hz})$ at 6.5–6.7 ppm, deshielded by ≥ 0.5 ppm relative to the other alkene protons. The most distinctive features of the analogous spectra of the α -isomers were the allylic and propargylic methine protons (or methylene in 8b), which resonated at 3.7-4.0 ppm (excepting **5b**). The ¹H NMR spectrum of **5b** was also noteworthy in that the resonances for two of the methoxy CH₃'s appeared as a broadened signal, which sharpened upon warming and decoalesced to two singlets at -20 °C. Variable temperature ¹H NMR studies established a coalescence T_c of 25 °C for these methyl group resonances, and a barrier at coalescence of $\Delta G_c = 15.2$ kcal/mol. This process was attributed to restricted rotation about the Ca- aryl C bond, which interchanged the two aryl ortho methoxy functions.

Our analysis of the reactivity patterns in this system is as follows. The allyl propargyldicobalt cation **21** generated from either **1** or **15a** reacts in a kinetic fashion with nucleophiles predominantly, but not exclusively, at the site γ - with respect to the alkynedicobalt unit (C-7). We find it particularly instructive that a comparison the γ -: α -selectivities with Mayr's published *N* (nucleophilicity) values¹⁷ reveals that greater nucleophilicity results in greater amounts of α - attack

(Table 5). While the exact correlation between N and γ -: α -ratios probably involves some coincidence and other factors likely contribute,¹⁸ a comparison between similar nucleophiles particularly supports this trend. For example, the less nucleophilic allyltrimethylsilane (N=1.79, γ -: α -= 84:16) has a much greater preference for the γ -site than allyltributylstannane (N=5.46, γ -: α -=50:50). In addition, the less nucleophilic acetophenone enol acetate¹⁹ reacts with greater γ -selectivity (γ -: α -=72:28) than the more nucleophilic trimethylsilyl enol ether ($N=6.22, \gamma$ -: α -=44:56). This is consistent with earlier work of Nicholas and Isobe on acyclic systems; low temperature reactions with alcohols and (to a small extent) enol acetates give α - attack kinetically, and these are the most reactive nucleophiles examined by these authors. The comparison of Et₃SiH and ¹Pr₃SiH suggests that increased γ -selectivity is encouraged by larger nucleophiles, likely as a consequence of the significant steric size of the alkyne–Co₂(CO)₆ unit.

Table 5. Nucleophile N values versus γ -: α -ratios

Nucleophile ^a	N value	γ-:α-Ratio
$H_2C = C(OSiMe_3)Ph$	6.22	44:56
Allyltributylstannane	5.46	50:50
Et ₃ SiH	3.64	72:28
Allyltrimethylsilane	1.79	84:16
Furan	1.36	>96:4

^a 1,3,5-Trimethoxybenzene (N=3.40) is excluded as it is likely not reacting at the kinetic limit.

Conversely, the product of thermodynamic reaction, as with the heteroatom based nucleophiles, is clearly exclusively γ -. This is supported by the results of reaction of **5b** and BF₃– OEt₂, and also by the fact that methyl ether **17a** underwent reaction with nucleophile **9a** (66%, 59:41 **10a**:10b) under the standard conditions. The conjugation between the alkene function and the complexed alkyne unit in the γ -products, and the assertion that the γ -products are more stable than the α -adducts, are also reflected by a shortened C-3/C-4 single bond length (1.450 Å) in **17a** and a 6.7 kcal/mol (28.0 kJ/mol) energy difference between **17a** and **17b** in DFT calculations (DFT B88-PW91, CAChe[®]).²⁰ The reaction of **1** with 1,3,5-trimethoxybenzene itself is neither at the kinetic nor thermodynamic limit.

In summary, the Nicholas reactions on the cation derived from the cyclic allylic acetate alkynedicobalt complex **1** kinetically favour the γ -site for most nucleophiles, with increasing amounts of α -products in cases with greater nucleophilicity. In the introduction of a specific nucleophilic fragment, some regiocontrol is possible through variation of the nucleophile. The thermodynamically favoured site is exclusively γ -. Work on employing some of the γ -adducts for access to 7,5- and 7,6- ring systems containing the alkynedicobalt unit, by way of cyclization reactions using the alkene function, is in progress and will be reported in due course.

3. Experimental

3.1. General methods

All reaction solvents were used after passage through a solvent purification system from Innovative Technologies.

Commercial BF₃–OEt₂ was distilled and stored under nitrogen. All reactions were conducted under a nitrogen atmosphere unless otherwise noted. Flash chromatography was performed as described by Still using silica gel 60 (230–400 mesh).²¹

All new compounds are >95% purity as determined by ¹H and ¹³C NMR spectroscopy. Reported regioisomeric ratios are on based on the ¹H NMR spectra of crude reaction products. NMR spectra were run at 500 or 300 MHz for ¹H and 125 or 75 MHz for ¹³C in CDCl₃; chemical shifts are given in ppm and coupling constants (*J*) are given in Hz. High resolution mass spectra were run at the McMaster Regional Centre for Mass Spectrometry and the Ohio State Chemistry Mass Spectrometry Facility.

3.1.1. Hexacarbonyl[µ-η⁴-(3-acetoxynona-1,8-dien-4yne)]dicobalt (4). To a mixture of alcohol 2 (0.3031 g, 2.23 mmol) and acetic anhydride (1 mL) at 0 °C was added pyridine (1 mL). The solution was stirred over a 6 h period and allowed to come to room temperature. The volatiles were removed under reduced pressure, and the resulting residue containing 3 was dissolved in Et_2O (15 mL). An excess amount of $Co_2(CO)_8$ was added and the solution stirred 12 h at room temperature. The removal of volatiles under reduced pressure followed by flash chromatography (100% petroleum ether—10:1 petroleum ether/Et₂O) gave acetate complex 4 (0.5239 g, 51% yield) as a red-brown oil. IR (neat, KBr, cm⁻¹): 3085, 2958, 2093, 2050, 2020, 1746; ¹H NMR δ : 6.48 (d, J=6.5 Hz, 1H), 5.92 (m, 2H), 5.42 (d, J=17.0 Hz, 1H), 5.28 (d, J=10.3 Hz, 1H), 5.16 (d, J=17.1 Hz, 1H), 5.09 (d, J = 10.3 Hz, 1H), 2.89 (m, 2H), 2.40 (m, 2H), 2.13 (s, 3H); 13 C NMR δ : 199.5, 169.8, 137.0, 135.3, 117.3, 115.9, 97.8, 94.5, 74.7, 35.5, 33.0, 20.6. MS EI m/e 408 (M⁺ - 2CO). HRMS m/e for C₁₇H₁₄Co₂O₈ calcd (M⁺-2CO) 407.9454, found 407.9455.

3.1.2. Hexacarbonyl[µ-η⁴-(3-acetoxycyclohept-1-en-4yne)]dicobalt (1). To a solution of 4 (0.0577 g, 0.124 mmol) in CH₂Cl₂ (5 mL) was added dichloro(phenylmethylene)bis(tricyclohexylphosphine)ruthenium (1st generation Grubbs' catalyst, 0.0102 g, 10.0 mol%) in CH_2Cl_2 (1 mL). The solution was stirred for 3 h, and subsequently concentrated under reduced pressure. Flash chromatography (20:1 petroleum ether:Et₂O) gave 1 (0.0436 g, 80%) as a red-brown oil. IR (neat, KBr, cm⁻¹) 3035, 2940, 2093, 2051, 2021, 1747; ¹H NMR δ: 6.70 (br s, 1H), 5.94 (m, 1H), 5.78 (dt, J=11.2, 2.2 Hz, 1H), 3.18 (dt, *J*=17.1, 4.3 Hz, 1H), 3.00 (ddd, *J*=3.7, 11.4, 17.1 Hz, 1H), 2.25–2.33 (m, 2H), 2.30 (s, 3H); ¹³C NMR δ: 199.3, 170.4, 134.3, 130.4, 98.0, 93.0, 73.9, 33.2, 27.2, 20.6. MS *m/e* 408 (M^+ – 1CO), 380 (M^+ – 2CO), 352 (M^+ – 3CO), 324 (M^+ – 4CO), 296 (M^+ – 5CO), 268 (M^+ – 6CO). HRMS m/e for C₁₅H₁₀Co₂O₈ calcd (M⁺ - 1CO) 407.9090, found 407.9103.

3.2. General procedure: reactions of the cycloheptenyne dicobalt complex with carbon- and heteroatom-based nucleophiles

To a solution of the nucleophile (1.5-2.0 equiv) and cycloheptenyne **1** in CH₂Cl₂ (0.05 M) at $-10 \degree \text{C}$ was added BF₃-OEt₂ (10 equiv) over 30 min as a solution in

 CH_2Cl_2 (1.0 M). The solution was stirred for 1 h and followed by addition of aqueous sodium bicarbonate. A typical workup was performed. The crude product was purified by flash chromatography.

3.2.1. Hexacarbonyl[μ - η^4 -(7-(2,4,6-trimethoxyphenyl)cyclohept-1-en-3-yne)]dicobalt (5a) and hexacarbonyl $[\mu-\eta^4-(3-(2,4,6-trimethoxyphenyl)cyclohept-1-en-$ 4-yne)]dicobalt (5b). A solution of cycloheptenyne 1 (0.0385 g, 0.0883 mmol) and 1,3,5-trimethoxybenzene (0.0297 g, 0.1766 mmol) in CH₂Cl₂ (2 mL) at $-10 \degree \text{C}$ was subjected to BF3-OEt2 (0.11 mL, 0.88 mmol) via the general procedure. The product was purified by flash chromatography (25:1 petroleum ether/Et₂O) gave 5a and **5b** (0.0412 g, 86%, **5a:5b**=70:30) as a red-brown oil. Careful repeated TLC afforded (in order of elution) 5b followed by **5a**. Compound **5a**. IR (neat, KBr, cm^{-1}): 2925, 2851, 2087, 2017, 1609, 1385; ¹H NMR δ : 6.46 (d, J =9.8 Hz, 1H), 6.14 (s, 2H), 5.97 (dd, J=2.7, 9.9 Hz, 1H), 4.03 (m, 1H), 3.79 (s, 9H), 3.35 (m, 1H), 3.16 (m, 1H), 2.19 (m, 1H), 1.82 (m, 1H); ${}^{13}C \delta$: 200.0, 159.0, 143.1, 123.7, 116.0, 99.3, 91.5, 89.7, 55.8, 55.5, 38.0, 35.9, 31.4, 24.3. MS EI *m/e*: 544 (M⁺), 516 (M⁺ – 1CO), 488 (M⁺ – 2CO), 460 (M⁺ – 3CO), 432 (M⁺ – 4CO), 404 (M⁺ – 5CO), 376 (M⁺ – 6CO). HRMS *m/e* for $C_{22}H_{18}Co_2O_9$ calcd (M⁺) 543.9615, found 543.9609. Compound 5b. IR (neat, KBr, cm⁻¹): 2926, 2085, 2043, 2014, 1733, 1609; ¹H NMR δ: 6.22 (m, 1H), 6.17 (s, 2H), 5.88 (m, 1H), 5.63 (s, 1H), 3.83 (s, 3H), 3.79 (br s, 6H), 3.24 (m, 1H), 3.03 (m, 1H), 2.41 (m, 2H); ¹³C NMR δ: 200.3, 160.4, 137.4, 128.4, 111.0, 101.0, 100.2, 91.2, 90.2, 55.5, 54.3, 38.5, 34.5, 27.3. MS EI $m/e: 544 (M^+), 516 (M^+ - 1CO), 488 (M^+ - 2CO), 460$ $(M^+ - 3CO), 432 (M^+ - 4CO), 404 (M^+ - 5CO),$ 376 (M⁺-6CO). HRMS *m/e* for $C_{22}H_{18}Co_2O_9$ calcd $(M^+ - CO)$ 515.9666, found 515.9666.

Reaction of **5b** *with* BF_3 – OEt_2 .

To a 0 °C solution of **5b** (0.0281 g, 0.0517 mmol) in CH₂Cl₂ (4 mL) was added BF₃–OEt₂ (65 μ L, 0.52 mmol). After stirring for 1 h at 0 °C, NH₄Cl_(aq) was added and the reaction was subjected to a conventional workup. Flash chromatography (20:1 petroleum ether/Et₂O) gave **5a** and **5b** (0.0189, 67% recovery, **5a:5b**=23:77).

3.2.2. Hexacarbonyl[μ - η^4 -(7-allylcyclohept-1-en-3-yne)] dicobalt (6a) and hexacarbonyl[μ - η^4 -(3-allylcyclohept-1en-4-yne)]dicobalt (6b). A solution if cycloheptenyne 1 (0.0817 g, 0.187 mmol) and allyltrimethylsilane (45 µL, 0.28 mmol) in CH₂Cl₂ (3.7 mL) at -10 °C was subjected to BF₃–OEt₂ (0.24 mL, 1.9 mmol) via the general procedure. Flash chromatography (25:1 petroleum ether/Et₂O) resulted in the co-elution of **6a** and **6b** (0.0650 g, 83%, **6a**:**6b** = 84:16) as a red-brown oil. IR (neat, KBr, cm⁻¹): 3015, 2926, 2854, 2089, 2046, 2017, 1641, 1582; ¹H NMR **6a** δ: 6.52 (d, J=9.9 Hz, 1H), 5.95 (dd, J=4.3, 9.9 Hz, 1H), 5.78 (m, 1H), 5.08 (m, 2H), 3.25 (m, 1H), 3.10 (m, 1H), 2.46 (m, 1H), 2.26 (m, 2H), 2.21 (m, 1H), 1.88 (m, 1H); resonances for **6b** could be observed at δ : 5.94 (m, 1H), 5.65 (m, 1H), 5.13 (m, 2H), 3.75 (m, 1H), 3.20 (m, 1H), 2.95 (m, 1H), 2.65 (m, 1H), 2.40 (m, 1H); 13 C NMR δ : 200.1, 139.7, 136.3, 126.4, 117.2, 98.1, 87.5, 41.0, 40.6, 33.4, 30.3; resonances for **6b** could be observed at 136.1, 131.5, 41.8, 34.3, 30.1, 27.1. MS EI *m/e*: 418 (M⁺), 390 (M⁺-1CO), 362 (M⁺-2CO), 334 (M⁺-3CO), 306 (M⁺-4CO), 278 (M⁺-5CO), 250 (M⁺-6CO). HRMS *m/e* for $C_{16}H_{12}Co_2O_6$ calcd (M⁺) 417.9298, found 417.9287.

3.2.3. Hexacarbonyl[µ-η⁴-(2-cyclohept-2-en-4-ynylfuran)]dicobalt (7a). A solution of cycloheptenyne 1 (0.0540 g, 0.124 mmol) and furan (0.136 g, 0.186 mmol) in CH₂Cl₂ (2.5 mL) at -10 °C was subjected to BF₃–OEt₂ (0.16 mL, 1.2 mmol) via the general procedure. The crude product was purified by flash chromatography (100% petroleum ether) to yield 7a (0.0341 g, 62%) as a redbrown oil. IR (neat, KBr, cm⁻¹): 2927, 2089, 2048, 2017, 1622, 1428; ¹H NMR δ : 7.35 (d, J = 1.8 Hz, 1H), 6.71 (d, J=9.9 Hz, 1H), 6.28 (dd, J=1.8, 3.1 Hz, 1H), 6.15 (dd, J=3.1, 9.9 Hz, 1H), 6.03 (d, J = 3.2 Hz, 1H), 3.89 (m, 1H), 3.17(m, 1H), 2.98 (m, 1H), 2.23 (m, 1H), 2.08 (m, 1H); ¹³C NMR δ: 199.9, 155.8, 141.7, 133.7, 127.8, 110.1, 106.3, 98.1, 86.8, 41.1, 32.2, 30.1. MS EI m/e: 444 (M⁺), 416 $(M^+ - 1CO)$, 388 $(M^+ - 2CO)$, 360 $(M^+ - 3CO)$, 332 $(M^+ - 4CO)$, 304 $(M^+ - 5CO)$, 276 $(M^+ - 6CO)$. HRMS m/e for C₁₇H₁₀Co₂O₇ calcd (M⁺) 443.9091, found 443.9082.

Hexacarbonyl[μ-η⁴-(cyclohept-1-en-3-yne)]-3.2.4. dicobalt (8a) and hexacarbonyl[µ-η⁴-(cyclohept-1-en-4yne)]dicobalt (8b). A solution of cycloheptenyne 1 (0.0500 g, 0.115 mmol) and triethylsilane (0.0200 g, 0.173 mmol) in CH₂Cl₂ (2.3 mL) at -10 °C was subjected to BF_3 -OEt₂ (0.15 mL, 1.1 mmol) via the general procedure. After flash chromatography (100% petroleum ether), an inseparable mixture of 8a and 8b (0.0235 g, 54%, **8a:8b**=72:28) was isolated. IR (neat, KBr, cm^{-1} 2928, 2089, 2046, 2016, 1581, 1385; ¹H NMR δ: 6.54 (d, J=9.7 Hz, 1H), 6.10 (m, 1H), 3.20 (t, J=5.6 Hz, 2H), 2.41 (m, 2H), 1.87 (m, 2H); peaks for **8b** could be observed at δ : 5.97 (m, 1H), 5.88 (m, 1H), 3.10 (m, 2H), 2.41 (m, 2H), 2.33 (m, 2H); ¹³C δ: 199.5, 135.1, 127.1, 97.9, 89.4, 35.7, 30.9, 24.9; resonances for **8b** could be observed at δ : 199.5, 132.4, 130.2, 98.1, 89.6, 34.5, 33.6, 27.2. MS EI *m/e*: 378 (M⁺), 350 (M⁺ - 1CO), 322 (M⁺ - 2CO), 294 (M⁺ - 3CO), 266 $(M^+ - 4CO)$, 238 $(M^+ - 5CO)$, 210 $(M^+ - 6CO)$. HRMS m/e for C₁₃H₈Co₂O₆ calcd (M⁺-CO) 349.9030, found 349.9008.

3.2.5. Hexacarbonyl[μ-η⁴-(2-cyclohept-2-en-4-ynylmethyl-prop-2-en-1-ol)]dicobalt (10a) and hexacarbonyl $[\mu-\eta^4-(2-cyclohept-2-ynyl-methyl-prop-2-en-1-ol)]dicobalt$ (10b). A solution of cycloheptenyne 1 (0.0776 g, 0.178 mmol) and 2-(trimethylsilylmethyl)-2-propen-1-ol (9a) (0.0384 g, 0.266 mmol) in CH_2Cl_2 (3.6 mL) at -10 °C was subjected to BF3-OEt2 (0.23 mL, 1.8 mmol) via the general procedure. Flash chromatography (3:1 petroleum ether/Et₂O) resulted in the isolation of 10a and **10b** (0.0607 g, 76%, **10a**: **10b** = 59:41) as a red-brown oil. Careful repeated TLC afforded (in order of elution) 10b followed by 10a. Compound 10a. IR (neat, KBr, cm^{-1}) 3354, 2923, 2086, 2047, 2021, 1608, 1435, 1384; ¹H NMR δ : 6.54 (d, J=9.9 Hz, 1H), 5.96 (dd, J=3.8, 9.9 Hz, 1H), 5.17 (s, 1H), 4.94 (s, 1H), 4.09 (s, 2H), 3.28 (m, 1H), 3.12 (m, 1H), 2.61 (m, 1H), 2.28 (m, 2H), 1.91 (m, 1H), 1.75 (m, 1H), 1.51 (br s, 1H); ¹³C NMR δ : 200.0, 146.1, 139.2, 126.3, 112.3, 98.0, 87.5, 65.6, 39.5, 38.7, 33.3, 30.3. MS EI

m/e: 448 (M⁺), 420 (M⁺ – 1CO), 392 (M⁺ – 2CO), 364 (M⁺ – 3CO), 336 (M⁺ – 4CO), 308 (M⁺ – 5CO), 280 (M⁺ – 6CO). HRMS *m/e* for C₁₇H₁₄Co₂O₇ calcd (M⁺ – 2CO) 391.9500, found 391.9513. Compound **10b**. IR (neat, KBr, cm⁻¹) 3385, 2925, 2088, 2046, 2016, 1608, 1506, 1093; ¹H NMR for the δ : 5.95 (m, 1H), 5.67 (m, 1H), 5.23 (s, 1H), 5.05 (s, 1H), 4.18 (s, 2H), 3.92 (m, 1H), 3.24 (m, 1H), 3.01 (m, 1H), 2.35 (m, 4H), 1.59 (br s, 1H); ¹³C NMR δ : 199.9, 146.1, 135.9, 131.4, 112.2, 100.9, 99.9, 65.9, 40.4, 39.3, 34.2, 26.9. MS EI *m/e*: 448 (M⁺), 420 (M⁺ – 1CO), 392 (M⁺ – 2CO), 364 (M⁺ – 3CO), 336 (M⁺ – 4CO), 308 (M⁺ – 5CO), 280 (M⁺ – 6CO). HRMS *m/e* for C₁₇H₁₄Co₂O₇ calcd (M⁺) 447.9403, found 447.9376.

3.2.6. Hexacarbonyl[μ - η^4 -(7-(2-chloromethylallyl)cyclohept-1-en-3-yne)]dicobalt (11a) and hexacarbonyl[μ - η ⁴-(3-(2-chloromethylallyl)cyclohept-1-en-4-yne)]dicobalt (11b). A solution of cycloheptenyne 1 (0.0477 g,0.109 mmol) and 2-chloromethyl-3-trimethylsilyl-1propene (9b) (0.030 mL, 0.17 mmol) in CH₂Cl₂ (2.5 mL) at -10 °C was subjected to BF₃–OEt₂ (0.14 mL, 1.1 mmol) via the general procedure. Flash chromatography (25:1 petroleum ether/Et₂O) resulted in the co-elution of 11a and 11b (0.0358 g, 70%, 11a:11b=72:28) as a red-brown oil. IR (neat, KBr, cm⁻¹): 2927, 2090, 2047, 2016, 2017, 1506, 1430; ¹H NMR δ : 6.55 (dd, J = 1.6, 9.9 Hz, 1H), 5.97 (dd, J = 4.1, 9.9 Hz, 1H), 5.27 (s, 1H), 5.02 (s, 1H), 4.05 (s, 2H), 3.28 (m, 1H), 3.18 (m, 1H), 2.68 (m, 1H), 2.37 (m, 2H), 1.89 (m, 1H), 1.87 (m, 1H); resonances for 11b could be observed at δ : 5.97 (m, 1H), 5.68 (dd, J = 3.3, 10.5 Hz, 1H), 5.31 (s, 1H), 5.14 (s, 1H), 4.13 (s, 2H), 3.26 (m, 2H), 3.14 (m, 1H), 2.45 (m, 1H), 2.33 (m, 2H), 1.71 (m, 1H); ¹³C NMR δ: 199.9, 142.5, 138.8, 126.7, 117.1, 96.3, 86.2, 47.8, 39.6, 38.5, 33.3, 30.3; resonances for **11b** could be observed at δ : 135.7, 133.0, 116.9, 96.3, 86.2, 48.0, 40.1, 39.1, 34.1, 27.2. MS EI m/e: 466 (M⁺), 438 (M⁺ - 1CO), 410 (M⁺ - 2CO), 382 (M⁺ - 3CO), 354 (M⁺ - 4CO), 326 (M⁺ - 5CO), 298 (M⁺ - 6CO). HRMS m/e for C₁₇H₁₃ $ClCo_2O_6$ calcd (M⁺) 465.9065, found 465.9038.

3.2.7. Hexacarbonyl[μ - η^4 -(acetic acid 2-cyclohept-2-en-4-ynylmethylallyl ester)]dicobalt (12a) and hexacarbonyl $[\mu-\eta^4-(acetic acid 2-cyclohept-2-en-6-ynylmethylallyl]$ ester)]dicobalt (12b). A solution of cycloheptenyne 1 (0.0706 g, 0.162 mmol) and 2-(acetoxymethyl)allyltrimethylsilane (9c) (0.0509 g, 0.274 mmol) in CH_2Cl_2 (3.5 mL) at -10 °C was subjected to BF₃-OEt₂ (0.205 mL, 1.62 mmol) via the general procedure. Flash chromatography (25:1 petroleum ether/Et₂O) resulted in the co-elution of **12a** and **12b** (0.0606 g, 76%, **12a**:**12b**=64:36) as a red-brown oil. Compound 12a. IR (neat, KBr, cm⁻ 1): 2927, 2089, 2048, 2018, 1747, 1053; ¹H NMR δ: 6.54 (dd, J = 1.9, 9.8 Hz, 1H), 5.94 (dd, J = 4.3, 9.8 Hz, 1H), 5.18 (s, 1H), 5.01 (s, 1H), 4.55 (1/2 ABq, J=13.5 Hz, 1H), 4.51 (1/2 ABq, J = 13.5 Hz, 1H), 3.28 (m, 1H), 3.13 (m, 1H), 2.61 (m, 1H), 2.27 (m, 2H), 2.22 (s, 3H), 2.09 (m, 1H), 2.06 (m, 1H); resonances for **12b** could be observed at ¹H NMR δ : 5.94 (m, 1H), 5.65 (br d, J = 10.5 Hz, 1H), 5.23 (s, 1H), 5.12 (s, 1H), 4.68 (1/2 ABq, J=13.2 Hz, 1H), 4.59 (1/2 ABq, J= 13.2 Hz, 1H), 3.87 (m, 1H), 3.22 (m, 1H), 2.98 (m, 1H), 2.71 (dd, J=4.1, 14.9 Hz, 1H), 2.33 (m, 2H), 2.28 (m, 1H), 2.11 (s, 3H); ¹³C NMR δ: 199.9, 170.7, 156.1, 141.2, 138.9, 126.5, 115.35, 97.9, 87.4, 66.6, 39.7, 38.6, 33.2, 30.1;

resonances for **12b** could be observed at δ : 170.7, 141.2, 135.40, 131.5, 115.4, 100.8, 99.8, 66.6, 40.1, 39.1, 34.1, 30.3, 27.0, 20.8. MS EI *m/e*: 434 (M⁺ - 2CO), 406 (M⁺ - 3CO), 378 (M⁺ - 4CO), 350 (M⁺ - 5CO), 322 (M⁺ - 6CO). HRMS *m/e* for C₁₉H₁₆Co₂O₈ calcd (M⁺ - 2CO) 433.9605, found 433.9636.

3.2.8. Hexacarbonyl[μ - η^4 -(7-(3-acetoxypropen-2-yl)cyclohept-1-en-3-yne)]dicobalt (13a) and hexacarbonyl $[\mu-\eta^4-(3-(3-acetoxypropen-2-yl)cyclohept-1-en-4-yne)]$ dicobalt (13b). A solution of cycloheptenyne 1 (0.0524 g, 0.120 mmol) and 1-trimethylsilylallyl acetate (0.0384 g, 0.223 mmol) in CH₂Cl₂ (2.4 mL) at -10 °C was subjected to BF₃-OEt₂ (0.15 mL, 1.2 mmol) via the general procedure. The crude product was purified by flash chromatography (25:1 petroleum ether/Et₂O) to yield of 13a and 13b (0.0369 g, 65%) as Z/E-isometric mixtures as a redbrown oil. IR (neat, KBr, cm⁻¹): 2926, 2089, 2047, 2016, 1760, 1673, 1217; **13a** ¹H NMR δ : 7.13 (d, J=6.8 Hz, 1H, Z-isomer) and 7.14 (d, J=12.3 Hz, 1H, E-isomer), 6.55 (d, J=9.9 Hz, 1H), 5.97 (dd, J=4.4, 10.0 Hz, 1H, Z-isomer) and 5.95 (dd, J=4.1, 9.9 Hz, 1H, E-isomer), 4.89 (apparent q, J = 6.8 Hz, 1H, Z-isomer) and 5.41 (dt, J = 12.3, 7.8 Hz, 1H, E-isomer), 3.28 (m, 1H), 3.12 (m, 1H), 2.40-2.50 (m, 1H), 2.34 (m, 1H), 2.19 (m, 1H), 2.15 (s, 3H, Z-isomer) and 2.13 (s, 3H, E-isomer), 1.86 (m, 1H), 1.73 (m, 1H); absorptions for 13b could be observed at 5.67 (m, 1H), 5.56 (dt, J=12.5, 7.5 Hz, 1H, E-isomer) and 5.08 (apparent q, J=7.0 Hz, 1H, Z-isomer), 3.22 (m, 1H), 3.00 (m, 1H); ¹³C NMR δ: 200.1, 168.4, 168.2, 139.3, 139.1, 137.2, 135.8, 126.9, 126.7, 112.3, 111.4, 98.3, 87.0, 41.3, 41.2, 34.1, 33.2, 30.9, 30.3, 30.1, 29.9, 20.9. MS EI m/e: 476 (M⁺), 448 $(M^+ - 1CO), 420 (M^+ - 2CO), 392 (M^+ - 3CO), 364$ $(M^+ - 4CO)$, 336 $(M^+ - 5CO)$, 308 $(M^+ - 6CO)$. HRMS m/e for C₁₈H₁₄Co₂O₈ calcd (M⁺ - 2CO) 419.9449, found 419.9455.

3.2.9. Hexacarbonyl[μ-η⁴-(2-cyclohep-2-en-4-ynyl-1phenylethanone)]dicobalt (16a) and hexacarbonyl- $[\mu-\eta^4-(2-cyclohept-2-en-6-ynyl-1-phenylethanone)]$ dicobalt (16b). A solution of cycloheptenyne 1 (0.0592 g, 0.135 mmol) and 1-phenyl-1-(trimethylsiloxy)ethane (0.0519 g, 0.270 mmol) in CH₂Cl₂ (6 mL) at $-10 \degree \text{C}$ was subjected to BF₃-OEt₂ (0.17 mL, 1.3 mmol) via the general procedure. The crude product was purified by flash chromatography (25:1 petroleum ether/Et₂O) to yield 16a+16b (0.0496 g, 74%, 44:56 ratio) as a red-brown oil. Repeated TLC (10:1 petroleum ether/Et₂O) allowed sequential isolation of α -16b and γ -16a. Compound 16a. IR (neat, KBr, cm⁻¹): 3018, 2927, 2089, 2047, 2017, 1683; ¹H NMR δ : 8.03 (d, J=7.8 Hz, 2H), 7.40–7.60 (m, 3H), 6.57 (dd, J=1.4, 9.8, Hz, 1H), 6.02 (dd, J=4.5, 9.8, Hz, 1H), 3.10–3.30 (m, 5H), 1.80–1.96 (m, 2H); ¹³C NMR 199.8, 198.3, 138.7, 136.9, 133.3, 128.7, 128.0, 126.7, 97.8, 87.2, 44.0, 36.7, 32.9, 30.3. MS EI *m/e*: 468 (M⁺ - 1CO), 440 (M^+ – 2CO), 412 (M^+ – 3CO), 384 (M^+ – 4CO), 356 (M^+-5CO) , 328 (M^+-6CO) . HRMS *m/e* for calcd (M^+-CO) 467.9454, found 467.9445. Compound **16b**. IR (neat, KBr, cm⁻¹): 3022, 2930, 2089, 2046, 2014, 1688; ¹H NMR δ : 7.96 (d, J=7.8 Hz, 2H), 7.40–7.60 (m, 3H), 5.94 (m, 1H), 5.65 (dd, J = 3.6, 9.8 Hz, 1H), 4.46 (m, 1H), 3.56 (dd, J=5.4, 17.3 Hz, 1H), 3.32 (dd, J=8.4, 17.3 Hz, 1H), 3.21 (m, 1H), 3.03 (m, 1H), 2.35–2.50 (m, 2H); ¹³C

NMR 199.9, 197.9, 136.7, 135.8, 133.3, 131.5, 128.7, 128.1, 100.3, 100.1, 45.7, 37.8, 34.0, 27.0. MS EI *m/e*: 496 (M⁺), 468 (M⁺ – 1CO), 440 (M⁺ – 2CO), 412 (M⁺ – 3CO), 384 (M⁺ – 4CO), 356 (M⁺ – 5CO), 328 (M⁺ – 6CO). HRMS *m/e* for $C_{21}H_{14}Co_2O_7$ calcd (M⁺) 495.9403, found 495.9401.

3.2.10. Hexacarbonyl[μ - η^4 -(7-acetoxycyclohept-1-en-3-yne)]dicobalt (15a). A solution of cycloheptenyne 1 (0.0540 g, 0.124 mmol) and glacial acetic acid (0.0149 g, 0.248 mmol) in CH₂Cl₂ (2.5 mL) at -10 °C was subjected to BF₃-OEt₂ (0.16 mL, 1.3 mmol) via the general procedure. The crude product was purified by flash chromatography (10:1 petroleum ether/Et₂O) to yield the 15a (0.0427 g, 79%) as a red-brown oil. IR (neat, KBr, cm⁻¹): 2923, 2850, 2092, 2051, 2021, 1740, 1238; ¹H NMR δ: 6.68 (d, J = 10.0 Hz, 1H), 6.06 (dd, J = 4.6, 10.0 Hz, 1H), 5.48 (m, 1H), 3.30 (m, 1H), 3.22 (m, 1H), 2.12 (m, 1H), 2.09 (s, 3H), 2.00 (m, 1H); ¹³C NMR δ: 199.4, 170.0, 133.2, 128.6, 96.6, 85.0, 72.4, 30.3, 30.1, 21.0. MS EI m/e: 436 (M⁺), 408 $(M^+ - 1CO), 380 (M^+ - 2CO), 352 (M^+ - 3CO), 324$ $(M^+ - 4CO)$, 296 $(M^+ - 5CO)$, 268 $(M^+ - 6CO)$. HRMS m/e for C₁₅H₁₀Co₂O₈ calcd (M⁺) 435.9040, found 435.9012.

 H_2SO_4 conditions. To a solution of cycloheptyne **1** (0.1681 g, 0.386 mmol) in acetic acid (5 mL) was added H_2SO_4 (5 drops). The solution was stirred 1 h, at which point NH₄Cl_(aq) was added and the mixture subjected to a conventional extractive workup. Flash chromatography as described above afforded **15a** (0.1631 g, 97%).

3.2.11. Hexacarbonyl[μ - η^4 -(7-methoxy-cyclohept-1-en-3-yne)]dicobalt (Co-Co) (17a). A solution of cycloheptenyne 1 (0.0623 g, 0.143 mmol) and methanol (7.0 μ L, 0.17 mmol) in CH₂Cl₂ (2.9 mL) at -10 °C was subjected to BF₃-OEt₂ (0.18 mL, 1.4 mmol) via the general procedure. The crude product was purified by flash chromatography (10:1 petroleum ether/Et₂O) to yield the 17a (0.0379 g, 65%) as a red-brown oil. IR (neat, KBr, cm⁻¹): 2923, 2090, 2048, 2017, 1615, 1430; ¹H NMR δ : 6.61 (d, J = 10.0 Hz, 1H), 6.17 (dd, J = 3.9, 10.0 Hz, 1H), 3.95 (m, 1H), 3.37 (s, 3H), 3.34 (m, 1H), 3.12 (m, 1H), 2.04 (m, 2H); 13 C NMR δ : 199.5, 136.6, 127.3, 97.2, 86.1, 79.8, 56.3, 30.8, 30.1. MS EI m/e: 408 (M⁺), 380 (M⁺ - 1CO), $352 (M^+ - 2CO), 324 (M^+ - 3CO), 296 (M^+ - 4CO), 268$ $(M^+ - 5CO)$, 240 $(M^+ - 6CO)$. HRMS *m/e* for C₁₄H₁₀Co₂O₇ calcd (M⁺) 407.9091, found 407.9080.

 H_2SO_4 conditions. To a solution of cycloheptyne **1** (0.0540, 0.124 mmol) in MeOH (2 mL) and CH₂Cl₂ (2 mL) at 0 °C was added H₂SO₄ (2 drops). The ice bath was removed and the reaction stirred for 1 h. NH₄Cl_(aq) was added and the reaction was subjected to a conventional workup. Flash chromatography as described above afforded **17a** (0.0442 g, 87%).

3.2.12. Hexacarbonyl[μ - η^4 -(7-(2-chloroethoxy)-cycloheptenyne]dicobalt (18a). A solution of cycloheptenyne **1** (0.0510 g, 0.117 mmol) and 2-chloroethanol (10.0 μ L, 0.150 mmol) in CH₂Cl₂ (2.3 mL) at -10 °C was subjected to BF₃–OEt₂ (0.15 mL, 1.2 mmol) via the general procedure. The crude product was purified by flash

1409

chromatography (20:1 petroleum ether/Et₂O) to yield the **18a** (0.0315 g, 59%) as a red-brown oil. IR (neat, KBr, cm⁻¹): 2927, 2856, 2091, 2050, 2021, 1612; ¹H NMR δ : 6.63 (d, J=9.9 Hz, 1H), 6.16 (dd, J=4.0, 10.0 Hz, 1H), 4.13 (m, 1H), 3.78 (m, 2H), 3.62 (t, J=5.9 Hz, 2H), 3.36 (m, 1H), 3.14 (m, 1H), 2.06 (m, 2H); ¹³C NMR δ : 199.6, 136.0, 127.8, 97.1, 85.8, 78.8, 68.9, 43.0, 30.6, 30.4. MS EI *m/e*: 456 (M⁺), 400 (M⁺ - 2CO), 372 (M⁺ - 3CO), 344 (M⁺ - 4CO), 316 (M⁺ - 5CO), 288 (M⁺ - 6CO). HRMS *m/e* for C₁₅H₁₁ClCo₂O₇ calcd (M⁺) 455.8857, found 455.8841.

 H_2SO_4 conditions. To a solution of cycloheptyne **1** (0.0858 g, 0.197 mmol) and 2-chloroethanol (1 mL) in CH₂Cl₂ (5 mL) at 0 °C was added H₂SO₄ (3 drops). The solution was stirred for 1 h, at which point NH₄Cl_(aq) was added and a standard workup performed. Flash chromatography as above afforded **18a** (0.0679 g, 76%).

3.2.13. Hexacarbonyl[μ - η^4 -(7-(4-chlorobut-2-enyloxy)cyclohept-1-en-3-yne)]dicobalt (19a). A solution of cycloheptenyne 1 (0.0589 g, 0.135 mmol) and 4-chloro-2buten-1-ol (0.022 g, 0.21 mmol) in CH₂Cl₂ (2.7 mL) at -10 °C was subjected to BF₃–OEt₂ (0.17 mL, 1.3 mmol) via the general procedure. The crude product was purified by flash chromatography (25:1 petroleum ether/Et₂O) to yield the 19a (0.0440 g, 68%) as a red-brown oil. IR (neat, KBr, cm⁻¹): 2925, 2091, 2051, 2021, 1457, 1054; ¹H NMR δ : 6.65 (d, J = 10.0 Hz, 1H), 6.15 (dd, J = 4.0, 10.0 Hz, 1H), 5.76 (m, 2H), 4.18 (d, J=5.7 Hz, 2H), 4.12 (d, J=7.4 Hz, 2H), 4.10 (m, 1H), 3.34 (m, 1H), 3.12 (m, 1H), 2.04 (m, 2H); ¹³C NMR δ: 199.7, 136.1, 131.0, 128.1, 127.9, 97.1, 85.9, 63.7, 48.6, 39.1, 30.6, 30.4. MS EI m/e: 482 (M⁺), 454 $(M^+ - 1CO), 426 (M^+ - 2CO), 398 (M^+ - 3CO), 370$ $(M^+ - 4CO)$, 342 $(M^+ - 5CO)$, 314 $(M^+ - 6CO)$. HRMS m/e for C₁₇H₁₃ClCo₂O₇ calcd (M⁺) 481.9014, found 481.9001.

3.2.14. Hexacarbonyl[µ-η⁴-(cyclohept-2-en-4-ynylacetamide)]dicobalt (20a). H_2SO_4 conditions. Concentrated sulfuric acid was added dropwise (3 drops) to a solution of cycloheptenyne 1 (0.0645 g, 0.148 mmol) in acetonitrile (5 mL). After 10 min aqueous sodium bicarbonate was added and a typical workup proceeded. The crude reaction product was purified by flash chromatography (1:2 petroleum ether/ethyl acetate) to yield 20a (0.0546 g, 85%) as a red-brown oil. IR (neat, KBr, cm⁻¹) 2927, 2091, 2048, 2021, 1651, 1548, 1431; ¹H NMR δ: 6.66 (dd, J=1.6, 9.9 Hz, 1H), 6.17 (dd, J=4.7, 9.9 Hz, 1H), 5.48 (br d, J=7.2 Hz, 1H), 4.75 (m, 1H) 3.15–3.25 (m, 2H), 2.05 (m, 1H), 1.99 (s, 3H), 1.96 (m, 1H); ¹³C NMR δ: 199.4, 168.9, 115, 128, 1, 97.1, 85.5, 50.6, 31.1, 23.2. MS EI m/e: 435 (M⁺), 407 (M⁺ - 1CO), 379 (M⁺ - 2CO), 351 (M⁺ - 3CO), 323 (M⁺ - 4CO), 295 (M⁺ - 5CO), 267 (M⁺ -6CO). HRMS *m/e* for $C_{15}H_{11}Co_2NO_7$ calcd $(M^+ - CO)$ 406.9250, found 406.9242.

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Cobalt- and rhodium-catalyzed cross-coupling reaction of allylic ethers and halides with organometallic reagents

Hiroto Yasui, Keiya Mizutani, Hideki Yorimitsu* and Koichiro Oshima*

Department of Material Chemistry, Graduate School of Engineering, Kyoto University, Kyoto-daigaku Katsura, Nishikyo-ku, Kyoto 615-8510, Japan

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Abstract—Reactions of 2-alkenyl methyl ether with phenyl, trimethylsilylmethyl, and allyl Grignard reagents in the presence of cobalt(II) complexes are discussed. The success of the reactions heavily depends on the combination of the substrate, ligand, and Grignard reagent. In the reaction of cinnamyl methyl ether, the formation of the linear coupling products predominates over that of the relevant branched products. In the cobalt-catalyzed allylation of allylic ethers, addition of a diphosphine ligand can change the regioselectivity, mainly providing the corresponding branched products. Rhodium complexes catalyze the reactions of allylic ethers and halides with allylmagnesium chloride and allylzinc bromide, respectively, in which the branched coupling product is the major product.

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1. Introduction

Palladium-, nickel-, and copper-catalyzed cross-coupling reactions of allylic substrates with organometallic reagents are recognized as one of the most useful reactions catalyzed by transition metals.¹ On the other hand, cobalt-catalyzed cross-coupling reactions of allylic substrates are quite rare.² We have been interested in cobalt-catalyzed cross-coupling reactions.³ Here we report the reactions of allylic ethers with phenyl, trimethylsilylmethyl, and allyl Grignard reagents in the presence of cobalt complexes.⁴ Rhodium-catalyzed coupling reactions are also disclosed herein.⁵

2. Results and discussions

2.1. Cobalt-catalyzed phenylation reaction of allylic ethers

The coupling reaction of cinnamyl methyl ether (1) with phenylmagnesium bromide was first performed (Table 1). A number of ligands were screened, and 1,5-bis(diphenyl-phosphino)pentane (DPPPEN) proved to be most effective for the phenylation reaction. 3,3-Diphenyl-1-propene was not detected at all. A small amount of β -methylstyrene was

the only byproduct in each experiment, along with untouched 1. The reaction of branched ether 3 with phenylmagnesium reagent under $CoCl_2(dpppen)$ catalysis provided linear 2 selectively in good yield (Eq. 1). The regioselectivity of the phenylations suggests that the reactions proceed via a π -allylcobalt intermediate. The phenylation reaction of 1 at 25 °C decreased the yield of 2. The choice of the solvent was essential to obtain 2 in satisfactory yield. A similar reaction in THF resulted in very low conversion of 1.

 $\label{eq:constraint} \begin{array}{c} \textbf{Table 1}. \ \textbf{Cobalt-catalyzed reaction of cinnamyl methyl ether (1) with } \\ \textbf{phenylmagnesium bromide} \end{array}$

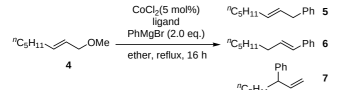
Ph. // OMe	CoCl ₂ (ligand) (5 mol%) PhMgBr (2.0 eq.)	Ph, 🥢 , Ph
1	ether, reflux, 16 h	2
Entry	Ligand	Yield (%)
1	None	29
2	PPh ₃ (10 mol%)	30
3	DPPM	24
4	DPPE	15
5	DPPP	27
6	DPPB	50
7	DPPPEN	72
8	DPPH	58

Ligands DPPM–DPPH represent $Ph_2P(CH_2)_nPPh_2$, n=1: DPPM; n=2: DPPE; n=3: DPPP; n=4: DPPB; n=5: DPPPEN; n=6: DPPH.

Keywords: Cross-coupling reaction; Cobalt; Grignard reagent; Rhodium; Allylzinc reagent.

^{*} Corresponding authors. Tel.: +81 75 383 2441; fax: +81 75 383 2438; e-mail addresses: yori@orgrxn.mbox.media.kyoto-u.ac.jp; oshima@orgrxn.mbox.media.kyoto-u.ac.jp

Table 2. Cobalt-catalyzed phenylation reaction of trans-2-octenyl methyl ether (4)

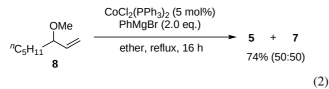


Entry	Ligand (amount)	Combined yield (%)	5/6/7
1	DPPPEN (5 mol%)	12	Not determined
2	None	47	58:10:32
3	DPPE (5 mol%)	32	10:53:37
4	PPh ₃ (10 mol%)	78	36:7:57
5	$P(2-MeC_6H_4)_3$ (10 mol%)	39	66:<1:33
6	$P(4-MeC_6H_4)_3$ (10 mol%)	49	42:6:52
7	$P[3,5-(CF_3)_2C_6H_3]_3$ (10 mol%)	Trace	Not determined
8	$P(4-MeOC_6H_4)_3$ (10 mol%)	41	31:16:53

It is worth noting that treatment of cinnamyl bromide under similar conditions furnished a mixture of dimeric compounds such as 1,6-diphenyl-1,5-hexadiene and 3,4diphenyl-1,5-hexadiene, in addition to a trace of 2. The formation of the dimeric products implies that single electron transfer from a cobalt complex would yield cinnamyl radical that is destined to dimerize.^{2a,c,d}

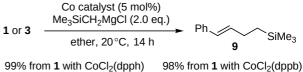
$$\begin{array}{c} OMe \\ Ph \\ 3 \end{array} \xrightarrow{\begin{array}{c} CoCl_2(dpppen) (5 mol\%) \\ PhMgBr (2.0 eq.) \\ ether, reflux, 16 h, 60\% \end{array}} \mathbf{2} \qquad (1)$$

The cobalt-catalyzed phenylation reaction of trans-2octenyl methyl ether (4) required triphenylphosphine as a ligand (Table 2, entry 4). A mixture of the corresponding coupling products 5, 6, and 7 was obtained. Under the reaction conditions, a part of 5 was transformed into 6. In contrast to the reaction of 1, the use of CoCl₂(dpppen) led to very poor conversion (entry 1). Without any phosphine ligand, coupling products were obtained in moderate combined yield (entry 2). Other monodentate phosphine ligands were inferior to triphenylphosphine (entries 5–8). Under $CoCl_2(PPh_3)_2$ catalysis, branched ether 8 was also converted into 5 and 7 (Eq. 2), in which no isomerization from 5 to 6 was observed.

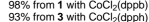


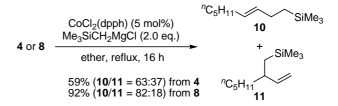
2.2. Cobalt-catalyzed trimethylsilylmethylation reaction of allylic ethers

Cross-coupling reaction with Me₃SiCH₂MgCl proceeded much more smoothly than that with PhMgBr (Scheme 1). Treatment of 1 with Me₃SiCH₂MgCl in the presence of CoCl₂(dpph) for 14 h at 20 °C afforded the corresponding



92% from 1 with CoCl₂



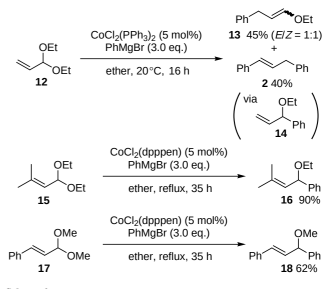


Scheme 1.

linear product 9 in 99% yield. Whereas the choice of the ligand was crucial to establish the phenylation, ligandless CoCl₂ and CoCl₂(dppb) also effected the allylation to afford 9 in 92 and 98% yields, respectively. Reactions of branched 3 with Me₃SiCH₂MgCl afforded 9 in excellent yield. On the other hand, alkyl-substituted allylic ethers 4 and 8 were converted into mixtures of regioisomers 10 and 11. The reaction required a higher temperature to complete the reaction within a satisfactory reaction time. Trimethylsilvlmethylation of branched ether 8 afforded a higher yield of 10 and 11 than that of 4.

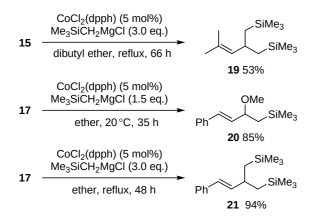
2.3. Cobalt-catalyzed reaction of α , β -unsaturated aldehyde dialkyl acetal

Treatment of acrolein diethyl acetal (12) with phenylmagnesium bromide in the presence of CoCl₂(PPh₃)₂ afforded a mixture of 2 and vinyl ether 13 (Scheme 2) Formation of doubly phenylated 2 would indicate a reaction path via the intermediate 14. Monophenylation of acetals 15 and 17 having substituents at the terminal olefinic positions was successful under CoCl₂(dpppen) catalysis. The dimethyl and phenyl groups of 16 and 18 would interfere with further phenylation.



Scheme 2.

In contrast to the reaction with phenylmagnesium bromide, bis(trimethylsilylmethylation) occurred in the reaction of **15** with 3 equimolar amounts of Me₃SiCH₂MgCl in refluxing dibutyl ether (Scheme 3). Intriguingly, in the reaction of **17**, we could completely control the distribution of the product by changing the amount of the Grignard reagent and reaction time. The reaction with 1.5 equimolar amounts of Me₃SiCH₂MgCl at ambient temperature for 35 h afforded monosubstituted product **20** exclusively in 85% yield. On the other hand, treatment of **17** with 3 equimolar amounts of the Grignard reagent in refluxing ether for 48 h furnished doubly substituted product **21** in 94% yield.



Scheme 3.

Table 3. Cobalt-catalyzed coupling reaction of 1 with allylmagnesium bromide

Dh

PhOMe	$ \begin{array}{c} \text{CoCl}_2 \text{ (5 mol}^9 \\ \text{CH}_2 = \text{CHCH}_2 \text{Mg} \\ \text{ether, refluence} \end{array} $	Br (2.0 eq.) x, 18 h	Pn 22 +
1		Ph	23
Entry	Ligand	Yield (%)	22/23
1	None	78	<1:99
2	NBu ₃	79	<1:99
3	TMEDA	75	<1:99
4	DPPE	57	51:49
5	DPPP	70	70:30
6	DPPB	54	19:81
7	DPPF	32	54:46

TMEDA and DPPF denote N,N,N',N'-tetramethylethylenediamine and 1,1'-bis(diphenylphosphino)ferrocene, respectively.

2.4. Cobalt-catalyzed cross-coupling reaction of cinnamyl methyl ether with allyl Grignard reagent

To extend the scope of the cobalt-catalyzed cross-coupling reactions, the allylation reaction of cinnamyl methyl ether was examined. The regioselectivity of the title reaction heavily depended on the ligand used (Table 3). Cobalt(II) chloride by itself catalyzed the cross-coupling to yield linear **23** exclusively (entry 1). Addition of amines as a ligand did not influence the regioselectivity (entries 2 and 3). Phosphine ligands allowed us to obtain significant amounts of branched **22**. Among them, DPPP exhibited the highest **22/23** selectivity, 70:30.

Judging from the results of Table 1, Scheme 1, and Table 3, trimethylsilylmethylmagnesium reagent proved to be the most reactive, and phenyl- and allylmagnesium reagents have similar reactivity. The low reactivity of allylmagnesium reagent may be due to the formation of π -allylcobalt that has less vacant coordination sites than phenyl- or trimethylsilylmethylcobalt has and that hence interacts weakly with the substrates at the initial oxidative addition stage.

The reactions of **1** and **3** with other Grignard reagents including vinylmagnesium bromide, methylmagnesium iodide, and alkynylmagnesium bromide failed to yield satisfactory amounts of the cross-coupling products.

2.5. Rhodium-catalyzed cross-coupling reaction of allylic ethers with allylmagnesium reagents

Although the catalytic activity of rhodium is lower than that of cobalt, rhodium complexes also catalyzed allylation of **1** (Scheme 4). Treatment of **1** with allylmagnesium chloride in the presence of [RhCl(nbd)]₂ (NBD=norbornadiene) in refluxing THF yielded the corresponding dienes in 47% combined yield. The branched form **22** was mainly obtained, and the selectivity is opposite to that of cobaltcatalyzed allylation. The use of [RhCl(cod)]₂ (COD=1,5cyclooctadiene) instead of [RhCl(nbd)]₂ slightly improved the efficiency and selectivity of the reaction. Other rhodium complexes such as Wilkinson's catalyst and rhodium(III) acetylacetonate as well as an iridium complex [IrCl(cod)]₂ exhibited no catalytic activity. Branched ether **3** yielded **22** and **23** in good yield in a similar ratio under the [RhCl(cod)]₂ catalysis.

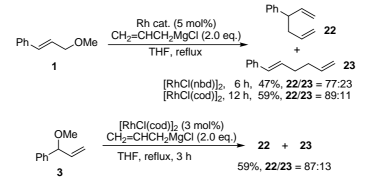
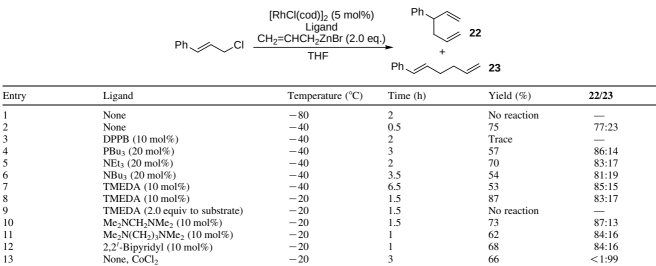


Table 4. Rhodium-catalyzed coupling reaction of cinnamyl chloride with allylzinc chloride



2.6. Rhodium-catalyzed cross-coupling reaction of cinnamyl chloride with allylzinc reagents

1

2

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Rhodium complexes also mediated the reaction of cinnamyl chloride with allylzinc bromide (Table 4). The reaction at -40 °C in the presence of [RhCl(cod)]₂ for 30 min furnished 22 and 23 in 75% yield in a ratio of 77:33 (entry 2). We screened many ligands to find that TMEDA is the best ligand with respect to the regioselectivity as well as the efficiency (entry 8). It is worth noting that a catalytic amount of diphosphine ligands such as DPPB (entry 3) and a stoichiometric amount of TMEDA (entry 9) completely inhibited the reaction. Interestingly, ligandless CoCl₂ effected the allylation to yield linear 23 exclusively (entry 13). An iridium complex $[IrCl(cod)]_2$ exhibited no catalytic activity.

3. Conclusion

The cobalt-catalyzed cross-coupling reaction with phenyl Grignard reagent proved to be a function of a substrate as well as of solvent and ligand. To attain high yields in the phenylation reaction, intensive tunings of variants are needed. In contrast, introduction of trimethylsilylmethyl group was facile and clean under cobalt catalysis. The reactions of cinnamyl methyl ether with both phenyl and trimethylsilylmethyl Grignard reagents yielded the corresponding linear products, irrespective of reaction conditions. The cross-coupling reactions of allylic ethers with allyl Grignard reagent with the aid of ligandless cobalt(II) chloride afforded the corresponding linear dienes. Interestingly, addition of DPPP could reverse the regioselectivity, leading to predominant formation of the branched dienes. Rhodium complexes catalyzed the reactions of allylic ethers and halides with allylmagnesium chloride and allylzinc bromide, respectively. Under rhodium catalysis, the branched coupling product was primarily formed. In both cobalt- and rhodium-catalyzed systems, π -allylmetal intermediates would be the key intermediates. The regioselectivity would depend on the ways how the carboncarbon bonds are formed, that is, via the outer-sphere mechanism or the inner-sphere mechanism. The exact mechanism is not clear at this stage.

4. Experimental

4.1. Instrumental

 $^1\mathrm{H}$ NMR (500 MHz) and $^{13}\mathrm{C}$ NMR (125.7 MHz) spectra were taken on Varian UNITY INOVA 500 spectrometers unless otherwise noted. ¹H and ¹³C NMR spectra were obtained in CDCl₃ with tetramethylsilane as an internal standard. Chemical shifts (δ) are in parts per million relative to tetramethylsilane at 0.00 ppm for 1 H and relative to CDCl₃ at 77.2 ppm for 13 C unless otherwise noted. IR spectra were determined on a SHIMADZU FTIR-8200PC spectrometer. Mass spectra were determined on a JEOL Mstation 700 spectrometer. TLC analyses were performed on commercial glass plates bearing 0.25-mm layer of Merck Silica gel 60F₂₅₄. Silica gel (Wakogel 200 mesh) was used for column chromatography. The analyses were carried out at the Elemental Analysis Center of Kyoto University.

4.2. Material

Unless otherwise noted, materials obtained from commercial suppliers were used without further purification. THF and ether were purchased from Kanto Chemical Co., stored under nitrogen, and used as they are. The starting materials 1, 3, 4, and 8 are prepared by the conventional Williamson ether synthesis.

4.3. General procedure for the cross-coupling reactions with Grignard reagents

The reaction of **1a** with trimethylsilylmethyl Grignard reagent is representative. Anhydrous CoCl₂ (7 mg, 0.05 mmol) was placed in a 50-mL two-necked flask and heated with a hair dryer in vacuo for 3 min. DPPH (27 mg, 0.06 mmol) and ether (1 mL) were sequentially added under argon. After the mixture was stirred for 30 min to obtain blue suspension, cinnamyl methyl ether (1a, 0.15 g,

1.0 mmol) and Me₃SiCH₂MgCl (1.0 M in ether, 2.0 mL, 2.0 mmol) were successively added to the reaction mixture at 0 °C. After being stirred for 14 h at 20 °C, the reaction mixture was poured into saturated NH₄Cl solution. The products were extracted with ethyl acetate (20 mL×3) and the combined organic layer was dried over sodium sulfate and concentrated. Silica gel column purification of the crude product provided **9** (0.20 g, 0.99 mmol) in 99% yield as colorless oil.

4.4. Rhodium-catalyzed cross-coupling reactions of cinnamyl chloride with allylzinc bromide

Zinc powder (2.94 g, 45 mmol) was placed in a 50-mL reaction flask under argon. THF (3.4 mL) was added. Chlorotrimethylsilane (0.1 mL, 0.8 mmol) and dibromoethane (0.1 mL, 2 mmol) were sequentially added at ambient temperature to activate zinc. After the mixture was stirred for 5 min, allyl bromide (2.6 mL, 30 mmol) in THF (24 mL) was added dropwise to the suspension with vigorous stirring over 15 min at 0 °C. The mixture was stirred for an additional 1 h at 25 °C. The gray supernatant liquid obtained was transferred to another flask filled with argon. The concentration of allylzinc bromide was determined by quantitative allylation reaction of an excess of benzaldehyde with allylzinc bromide prepared. The concentration was 0.87 M. [RhCl(cod)]₂ (25 mg, 0.05 mmol) was placed in another 50-mL two-necked flask under argon. THF (5 mL) and TMEDA (15 $\mu L,$ 0.10 mmol) were successively added. The resulting solution was stirred for 5 min. Cinnamyl chloride (153 mg, 1.0 mmol, dissolved in 5 mL of THF) was added. The solution was cooled at -20 °C, and allylzinc bromide (0.87 M in THF, 2.3 mL, 2.0 mmol) was added. After being stirred for 1.5 h at -20 °C, the reaction mixture was poured into 1 M hydrochloric acid. The product was extracted with ethyl acetate $(2 \times 20 \text{ mL})$. The combined organic phase was dried over sodium sulfate. Evaporation followed by silica gel column purification afforded a mixture of 22 and 23 (137 mg, 0.87 mmol, 87% combined yield) in a ratio of 83:17.

4.5. Characterization data

The spectral data of the products $5,^{6} 6,^{6} 7,^{7} 13,^{8} 18,^{9} 22,^{10}$ and 23^{10} are found in the literature.

4.5.1. (*E*)-4-Trimethylsilyl-1-phenyl-1-butene (9). IR (neat) 3061, 2953, 2903, 1497, 1248, 962, 862, 837, 692 cm⁻¹; ¹H NMR (CDCl₃) δ 7.27–7.35 (m, 4H), 7.17–7.20 (m, 1H), 6.37 (d, *J*=16.0 Hz, 1H), 6.27 (dt, *J*=16.0, 6.5 Hz, 1H), 2.23 (ddt, *J*=10.0, 1.0, 6.5 Hz, 2H), 0.68–0.71 (m, 2H), -0.10 to 0.16 (m, 9H); ¹³C NMR (CDCl₃) δ 137.98, 133.83, 128.45, 128.26, 126.66, 125.87, 27.39, 16.27, -1.59. Found: C, 76.27; H, 9.73%. Calcd for C₁₃H₂₀Si: C, 76.40; H, 9.86%.

4.5.2. (*E*)-1-(Trimethylsilyl)-3-nonene/3-(trimethylsilylmethyl)-1-octene (10/11=82:18). IR (neat) 2955, 2926, 1460, 1248, 968, 862, 835, 756, 691 cm⁻¹; ¹H NMR (CDCl₃) δ 5.52–5.59 (m, 0.18×1H), 5.35–5.46 (m, 0.82× 2H), 4.91 (ddd, *J*=17.0, 2.0, 0.5 Hz, 0.18×1H), 4.87 (ddd, *J*=10.0, 2.0, 0.5 Hz, 0.18×1H), 2.05–2.13 (m, 0.18×1H), 1.95–2.02 (m, 0.82×4 H), 1.23–1.37 (m, 0.82×6 H+ 0.18×8H), 0.88 (t, J=7.0 Hz, 3H), 0.55–0.59 (m, 2H), -0.01 (s, 9H); ¹³C NMR (CDCl₃). For major isomer, δ 113.03, 128.87, 32.48, 31.44, 29.35, 26.85, 22.57, 16.58, 14.08, -1.58. For minor isomer, δ 145.53, 112.56, 40.38, 38.55, 31.93, 23.26, 22.69, 14.12, -0.58. One of the sp³-hybridized carbons of **11** could not been observed, probably due to overlapping. Found: C, 72.34; H, 12.94%. Calcd for C₁₂H₂₆Si: C, 72.69; H, 13.21%.

4.5.3. 1-Ethoxy-3-methyl-1-phenyl-2-butene (16). IR (neat) 2974, 2930, 1425, 1086, 756, 698 cm⁻¹; ¹H NMR (CDCl₃) δ 7.31–7.35 (m, 4H), 7.26–7.23 (m, 1H), 5.35 (d, J=9.0 Hz, 1H), 5.01 (d, J=9.0 Hz, 1H), 3.45–3.51 (m, 1H), 3.35–3.42 (m, 1H), 1.79 (s, 3H), 1.74 (s, 3H), 1.22 (t, J=6.8 Hz, 3H); ¹³C NMR (CDCl₃) δ 142.81, 134.99, 128.40, 127.19, 126.59, 126.43, 78.13, 63.39, 25.91, 18.40, 15.36. Found: C, 81.84; H, 9.54%. Calcd for C₁₃H₁₈O: C, 82.06; H, 9.54%.

4.5.4. 2-Methyl-5-trimethylsilyl-4-(trimethylsilyl-methyl)-2-pentene (19). IR (neat) 2953, 2909, 1248, 837, 692 cm⁻¹; ¹H NMR (CDCl₃) δ 4.86 (d, J=10.0 Hz, 1H), 2.51–2.58 (m, 1H), 1.62 (s, 3H), 1.59 (s, 3H), 0.67 (dd, J=14.7, 5.3 Hz, 2H), 0.57 (dd, J=14.7, 8.5 Hz, 2H), -0.17 to 0.07 (m, 18H); ¹³C NMR (CDCl₃) δ 134.73, 126.19, 30.41, 28.54, 25.63, 18.19, -0.72. Found: C, 64.59; H, 12.24%. Calcd for C₁₃H₃₀Si₂: C, 64.38; H, 12.47%.

4.5.5. *(E)***-3-Methoxy-4-trimethylsilyl-1-phenyl-1-butene** (**20**). ¹H NMR (300 MHz, CDCl₃) δ 7.30–7.41 (m, 4H), 7.22–7.25 (m, 1H), 6.48 (d, *J*=15.9 Hz, 1H), 6.01 (dd, *J*=15.9, 8.4 Hz, 1H), 3.81 (q, *J*=7.8 Hz, 1H), 3.27 (s, 3H), 1.14 (dd, *J*=14.3, 6.8 Hz, 1H), 0.94 (dd, *J*=14.3, 7.7 Hz, 1H), 0.03 (s, 9H); ¹³C NMR (75 MHz, CDCl₃) δ 132.04, 131.15, 128.49, 127.51, 126.34, 106.68, 80.60, 55.72, 25.05, -0.62. Found: C, 71.79; H, 9.45%. Calcd for C₁₄H₂₂OSi: C, 71.73; H, 9.46%.

4.5.6. (*E*)-1-Phenyl-4-trimethylsilyl-3-(trimethylsilylmethyl)-1-butene (21). ¹H NMR (300 MHz, CDCl₃) δ 7.28–7.33 (m, 4H), 7.15–7.21 (m, 1H), 6.27 (d, *J*=15.6 Hz, 1H), 5.98 (dd, *J*=15.6, 9.0 Hz, 1H), 2.47–2.59 (m, 1H), 0.70–0.84 (m, 4H), -0.02 (s, 18H); ¹³C NMR (75 MHz, CDCl₃) δ 139.22, 128.37, 126.61, 126.54, 125.82, 106.68, 36.42, 28.13, -0.41. Found: C, 70.21; H, 10.28%. Calcd for C₁₇H₃₀Si₂: C, 70.26; H, 10.41%.

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Application of tri- and tetrasubstituted alkene dipeptide mimetics to conformational studies of cyclic RGD peptides

Shinya Oishi,^{a,b} Kazuhide Miyamoto,^a Ayumu Niida,^a Mikio Yamamoto,^c Keiichi Ajito,^c Hirokazu Tamamura,^a Akira Otaka,^a Yoshihiro Kuroda,^a Akira Asai^b and Nobutaka Fujii^{a,*}

^aGraduate School of Pharmaceutical Sciences, Kyoto University, Sakyo-ku, Kyoto 606-8501, Japan

^bGraduate School of Pharmaceutical Sciences, University of Shizuoka, Shizuoka, Shizuoka 422-8526, Japan

^cPharmaceutical Research Center, Meiji Seika Kaisha, Ltd., Kohoku-ku, Yokohama 222-0002, Japan

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Abstract—The first application of a combination of novel $\psi[(E)$ -CX=CX]-type alkene dipeptide isosteres to conformation studies of cyclic bioactive peptides was carried out (X=H or Me). For exploration of bioactive conformations of Kessler's cyclic RGD peptides, cyclo(-Arg-Gly-Asp-D-Phe-Val-) 1 and cyclo(-Arg-Gly-Asp-D-Phe-N-MeVal-) 2, D-Phe- ψ [(E)-CX=CX]-L-Val-type dipeptide isosteres were utilized having di-, tri- and tetrasubstituted alkenes containing the γ -methylated isosteres that have been reported to be potential type II' β -turn promoters. All of the (E)-alkene pseudopeptides 3-6 exhibited higher antagonistic potency against $\alpha_{\nu}\beta_{3}$ integrin than 1, although potencies were slightly lower than 2. Detailed structural analysis using ¹H NMR spectroscopy revealed that representative type II' β/γ backbone arrangements proposed for 1, were not observed in peptides 3-6. Rather on the basis of ¹H NMR data, the conformations of peptides 3-6 were estimated to be more analogous to those of the N-methylated peptide 2.

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1. Introduction

Use of both natural and artificial modifications of bioactive peptides and proteins provides opportunities to better understand the basis for bioactivities of the parent structures and to find novel functionality that may be applied for new purposes.¹ Application of unnatural amino acids and peptidomimetics constitutes one of the most powerful methodologies in such chemical approaches to understanding ligand-protein interactions.² Among large numbers of mimetics, (E)-alkene dipeptide isosteres that are designed as nonpolar alkene replacements of planar amide moieties within dipeptides, have been widely applied to bioand chemoactive peptides by us and others (Fig. 1).³ Gellman et al. reported that $Gly-\psi[(E)-CMe=CMe]-Gly$ type isostere **D** is a potential β -hairpin promoter.⁴ In addition, Wipf et al. have characterized D-Ala- $\psi[(E)$ -CMe=CH]-L-Ala- and L-Ala- $\psi[(E)$ -CCF₃=CH]-D-Alatype isosteres such as C as promoting β -turn formation in the solid state due to $A^{1,2}$ and $A^{1,3}$ -strain as opposed to L-Ala- $\psi[(E)$ -CH=CH]-D-Ala-type motifs exemplified by A that have a disubstituted alkene.⁵ These γ -methylated and

 γ -trifluoromethylated isosteres, which possess a carbon atom corresponding to a peptide bond carbonyl oxygen, are thought to be reasonable amide mimetics. Recent development of organocopper-mediated stereoselective synthesis of multi-substituted (E)-alkene isosteres⁶ allowed us to utilize a combination of these isosteres for practical structureactivity relationship (SAR) studies on bioactive peptides.^{5,6}

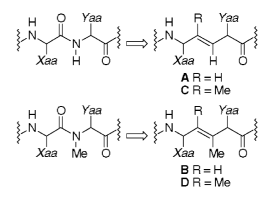


Figure 1. (E)-Alkene dipeptide isosteres having di-, tri- and tetrasubstituted alkenes; Xaa, Yaa=amino acid side chains.

As an exemplary application, we chose cyclic RGD peptides, cyclo(-Arg-Gly-Asp-D-Phe-Val-) 1⁷ and cyclo(-Arg-Gly-Asp-D-Phe-N-MeVal-) 2,⁸ which have

Keywords: (E)-Alkene dipeptide isostere; Cyclic RGD peptide; Integrin; Structure-activity relationship study.

^{*} Corresponding author. Tel.: +81 75 753 4551; fax: +81 75 753 4570; e-mail: nfujii@pharm.kyoto-u.ac.jp

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been shown to be highly potent and selective $\alpha_{v}\beta_{3}$ integrin antagonists (Fig. 2). It is well-known that $\alpha_{v}\beta_{3}$ integrin receptor and its ligands participate in many biological processes including tumor-induced angiogenesis and adhesion of osteoclasts to bone matrix and so on.⁹ Peptide 1 was originally reported to adopt two distinctive secondary structures in DMSO solution; a type II' β -turn with D-Phe at the i+1 position and a γ -turn with Gly at the i+1position.^{10,11} These structures allow two principal pharmacophores consisting of an Arg guanidino group and an Asp carboxylic acid to be located in close proximity. Among cyclic peptides, 2 is the most potent $\alpha_v \beta_3$ antagonist reported so far. It has been found to exhibit considerable conformational flexibility in water, including interconversion of two inverse γ turns (γ_i turns) and a γ turn, that do not afford identical topology of two closely located pharmacophores as observed in $1.^8$ On the other hand, the binding structure of 2 with $\alpha_{v}\beta_{3}$ integrin, which was recently disclosed by a crystal structure analysis of the ligandreceptor complex, is somewhat different from that proposed by Kessler et al.¹² The ligand binding seems to induce a structural change of the ligand binding domains of $\alpha_v \beta_3$ intergin, as well as a conformation change of ligand itself. As a result, peptide 2 exhibits distorted backbone conformations in the binding state to some extent, as compared with its calculated free-state conformations. Meanwhile, addition of an N-methyl group to the Val residue apparently improves $\alpha_{v}\beta_{3}$ antagonistic activity and $\alpha_{v}\beta_{3}/\alpha_{IIb}\beta_{3}$ selectivity, while the effect of *N*-methylation on the conformation of peptides as a whole as well as on the topology of the pharmacophores, especially in the neighbourhood of the D-Phe-Val/MeVal peptide bond, have not been discussed in detail. As such, it is difficult to rationalize structural and biological effects of certain characteristic functional groups in spite of such extensive research.

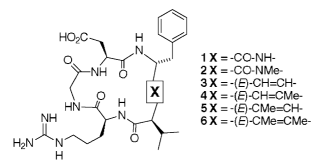


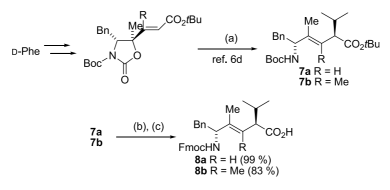
Figure 2. Structures of cyclic RGD peptides and peptidomimetics.

We recently reported the diastereoselective synthesis of γ -unmethylated D-Phe- $\psi[(E)$ -CH=CX]-L-Val- and D-Phe- $\psi[(Z)$ -CH=CMe]-L-Val-type alkene dipeptide isosteres (X=H or Me) with application to D-Phe-L-Val/N-MeVal moieties in peptides 1 and 2.^{6b} Both peptides 3 and 4 contain $\psi[(E)$ -CH=CH]- and $\psi[(E)$ -CH=CMe]-type isosteres, respectively, and exhibit potent antagonistic activity against $\alpha_{v}\beta_{3}$ integrin. In contrast, (Z)-congeners show extremely low $\alpha_v \beta_3$ and $\alpha_{IIb} \beta_3$ antagonist potency. This indicated that cis-conformation within the D-Phe-L-Val/N-MeVal peptide bond distorted the peptide bioactive conformations. On the other hand, slight differences between the potencies of 3 and 4, which are independent of the presence of a β -methyl group in 4 that corresponds to an N-methyl group of 2, support a conformational role for the N-methyl group of 2 beyond a simple steric one. To facilitate a deeper understanding of structure-activity relationships of cyclic RGD peptides, it was thought that utilization of highly functional β -turn promoters such as γ -methylated (E)-alkene isosteres, could be of value. Moreover, a D-Phe- ψ [(E)-CMe=CMe]-L-Val-type analogue could also be regarded as a D-Phe-L-N-MeVal dipeptide equivalent having reduced polarity, wherein the β - and γ -methyl groups could replicate allylic strain across peptide bonds between the D-Phe carbonyl oxygen and the N-MeVal side chain, as well as between the *N*-methyl group and the D-Phe side chain.¹³ With this in mind, the synthesis and bio-evaluation of isostere-containing cyclic peptides 5 and 6 was undertaken, along with 1 H NMR conformational analysis and comparison with the previous peptides 1-4. Reported herein are results of our application of γ -methylated alkene dipeptide isosteres to proposed type II' β -turn motifs in bioactive peptides. We also examined the structure-activity effects of N-methylation of Val in cyclic RGD peptides using (E)-alkene isosteres having differential substitution motifs.

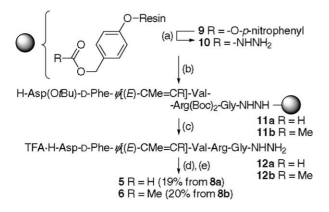
2. Results and discussion

2.1. Synthesis

Preparation of cyclic RGD peptides **5** and **6** that contain D-Phe- $\psi[(E)$ -CMe=CH]-L-Val- and D-Phe- $\psi[(E)$ -CMe=CMe]-L-Val-type alkene dipeptide isosteres, respectively, was performed according to the synthetic scheme utilized for the synthesis of peptide **4** (Schemes 1 and 2).^{6b} In this process, a combination of Fmoc-based solid phase peptide synthesis (SPPS) and cyclization of linear peptide



Scheme 1. (a) Organocopper reagents; (b) TFA; (c) Fmoc-OSu, Et₃N.



Scheme 2. (a) $NH_2NH_2 \cdot H_2O$; (b) Fmoc-based SPPS; (c) TFA; (d) HCl, isoamyl nitrite; (e) iPr_2NEt .

hydrazides **12a**,**b** without side-chain protecting groups was employed by an adapted azide method in order to avoid olefinic isomerization, which would otherwise be possible during final deprotection by strong acid treatment in Bocbased synthesis. For side-chain protection, tert-butyl ester for Asp and (Boc)₂ for Arg were employed, both of which are amenable to mild acidic deprotection. TFA-treatment of the *N*-Boc-protected isosteres **7a**,**b**, which were obtained by regio- and stereoselective alkylation of β -(1,3-oxazolidin-2-one)-5-yl- α , β -enoates by organocopper reagents,¹⁴ followed by Fmoc-reprotection, provided building blocks **8a,b** that were suitable for SPPS. Following the preparation of hydrazide linker 10 by treatment of *p*-nitrophenyl carbonate resin 9 with hydrazine hydrate in DMF, peptide chain elongation by Fmoc-based SPPS gave the expected protected peptide resins 11a,b. Side chain deprotection and TFA-mediated cleavage from resins 11a,b provided peptide hydrazides 12a,b, which were subjected to successive azide formation and cyclization in highly diluted DMF solution.¹⁵ The crude peptides were readily purified by reverse-phase HPLC to yield the expected cyclic peptides 5 and 6 in 19 and 20% yield, respectively, which were fully characterized by ¹H NMR and mass spectra.

2.2. Structure–activity relationships of cyclic RGD peptides and peptidomimetics

Integrin antagonistic activities of the resulting peptides **5** and **6** against $\alpha_v\beta_3$ and $\alpha_{IIb}\beta_3$ integrins were comparatively evaluated along with Kessler's RGD peptides **1** and **2**, and peptides **3** and **4** having γ -unmethylated D-Phe- $\psi[(E)$ -

CH=CX]-L-Val-type isosteres (X=H or Me). ELISA assays were performed using immobilized $\alpha_v \beta_3$ or $\alpha_{IIb} \beta_3$ integrin, according to the modified method of Kouns et al.¹ The results are shown in Table 1 as inhibition by peptides **1–6** of vitronectin or fibring to the respective integrins (n=8). Each of the isostere-containing peptides **3–6** showed strong $\alpha_{v}\beta_{3}$ integrin antagonistic activity within the range from $IC_{50} = 6.8$ nM for 1 to $IC_{50} = 1.4$ nM for 2. It appeared that the amide or olefinic moiety in the D-Phe-Val/ *N*-MeVal dipeptide portion of peptides **1–6** was not directly involved in recognition and binding to $\alpha_{v}\beta_{3}$ integrin. These data also support that trans-amide conformation within the D-Phe-Val/N-MeVal dipeptide was predominant in the bioactive conformations. This is consistent with a crystal structure analysis of an $\alpha_{\rm v}\beta_3$ integrin–ligand complex¹² and our previous research using a combination of (E)- and (Z)-alkene dipeptide isosteres.^{6b} Structure–activity relationship studies on cyclic RGD peptides investigating effects due to the *N*-methyl group of **2** using novel alkene dipeptide isosteres seemed to be highly appropriate.

It is noteworthy that only minimal differences were observed between the activities of peptides **3** and **5** having β -unmethylated isosteres and the respective β -methylated isostere-containing congeners **4** and **6**. In contrast, peptide **2**, having *N*-methyl valine, exhibited approximately five times higher potency than peptide **1**, similar to a previous report.⁸ If *N*-methylation is potency-enhancing in **2**, then either peptide **4** or **6**, which possesses β -methyl group isosteric to the *N*-methyl group of **2**, could also show potencies superior to **3** or **5**, respectively. This unexpected result demonstrated that a conformational transformation from **1** to **2** and the resulting improvement of $\alpha_v\beta_3$ integrin antagonism depend on factors other than simple steric properties of the *N*-methyl group.

It was also found that peptides **5** and **6**, containing an isostere γ -methyl group, had slightly higher potency against $\alpha_{\nu}\beta_{3}$ integrin than the γ -unmethylated congeners **3** and **4**, respectively. Interestingly, in a crystal structure of peptide **2** complexed to $\alpha_{\nu}\beta_{3}$ integrin, the carbonyl oxygen of D-Phe, to which the isostere γ -methyl group corresponds, is not directly associated with any polar interactions with integrin, such as hydrogen bonding.¹² In light of this, the improved potencies of **5** and **6** may potentially be derived from steric interactions, including allylic strain induced by the γ -methyl group. Similarly, it could be surmised that D-Phe carbonyl oxygens of **1** and **2** could partially contribute to

Table 1. Integrin antagonistic activities of cyclic RGD peptides and peptidomimetics

Peptide	Х	$\alpha_v \beta_3$	$\alpha_v \beta_3$		$\alpha_{IIb}\beta_3$	
		$IC_{50} (nM)^b$	Q ^c	$IC_{50} (nM)^{b}$	Q ^c	
RGDS ^d	_	98 ± 29	14	270 ± 41	0.35	2.7
1	-CO-NH-	6.8 ± 2.7	1	770 ± 120	1	110
2	-CO-NMe-	1.4 ± 0.31	0.20	280 ± 42	0.36	200
3	-CH=CH-	3.6 ± 1.3	0.53	140 ± 18	0.19	40
4	-CH=CMe-	3.3 ± 0.93	0.48	100 ± 42	0.13	30
5	-CMe=CH-	2.4 ± 0.33	0.35	81 ± 18	0.11	34
6	-CMe=CMe-	1.8 ± 0.51	0.27	48 ± 11	0.06	26

^a SI values were calculated as SI = IC₅₀($\alpha_{IIb}\beta_3$)/IC₅₀($\alpha_v\beta_3$).

^b The data for peptides **1–6** were obtained in comparative experiments using the same conditions.

^c Q values were calculated as $Q = IC_{50}(\text{peptide})/IC_{50}(1)$.

^d A linear peptide RGDS (H-Arg-Gly-Asp-Ser-OH) was used as a standard peptide.

appropriate dispositions of close functional groups, resulting in enhanced potencies.

In contrast, isostere-containing peptides 3-6 were less selective $\alpha_{v}\beta_{3}$ integrin antagonists than 1 or 2, due to their relatively high potency against $\alpha_{IIb}\beta_3$ integrin. These increased potencies against $\alpha_{IIb}\beta_3$ integrin resulted from substituting the amide bonds of 1 and 2 with alkene isosteres, indicated that distinct functional groups derived from the olefinic moieties may be compatible with structural features of $\alpha_{IIb}\beta_3$ integrin. Other independent factors of RGD motifs displayed by the ligands may contribute to selectivity in interaction with the two integrins. However, we failed to ascertain what characteristics could be associated with selective recognition by the respective integrins. Locardi et al. revealed that the conformations of $\alpha_{\text{IIb}}\beta_3$ antagonists are different in the presence and absence of the receptor.¹⁷ It is conceivable that the cyclic peptides may vary their shape by distinctive interactions in the binding state, even if the isostere moieties in 3-6 do not affect peptide conformations in the absence of the receptor.

2.3. Conformational aspects of cyclic peptidomimetics derived from ¹H NMR spectroscopy

Conformations of cyclic peptides have been intensively investigated using NMR spectroscopy and molecular dynamics calculations.¹⁸ In structure–activity relationship studies on cyclic RGD peptides under 'conformational control', Kessler et al. reported that replacement at either the D-Phe or Val positions did not induce changes in backbone conformations.^{10a} ¹H NMR parameters such as chemical shifts, temperature dependence of amide protons and ³*J*coupling constants support homogeneous families of cyclic peptide conformations. Based on similar concepts using alkene isosteres, we attempted to understand effects of the *N*-methyl groups or isostere β -methyl groups on conformations and their relationship to $\alpha_v \beta_3$ integrin antagonistic activity.¹⁹

In chemical shift data of peptides 1–6 in DMSO solution, downfield shifts of Arg H^N , Asp H^N , one Gly H^{α} (high field) and D-Phe H^{α} of peptides 2, 4 and 6 that possess N-MeVal *N*-methyl groups or corresponding β -methyl groups, were comparable to those of 1, 3 and 5, respectively, (see the Supporting information). On the other hand, Gly H^N, Arg H^{α} , the other Gly H^{α} (low field) and Asp H^{α} of 2, 4 and 6 were located at higher fields than those of 1, 3 and 5, respectively. For D-Phe H^N, no significant differences were found between 1 and 2, while similar upfield shift correlations were observed among the isostere-containing peptides 3-6. As such, the addition of a methyl group to the α -amino group of Val or to the isostere β -position, induced nearly equal chemical shift changes, although this may not necessarily indicate similar changes in peptide backbone conformation. These observations are in contrast to the fact that among peptides 2, 4 and 6, an increase in $\alpha_v \beta_3$ integrin antagonistic activity was observed only in 2.

In a sharp contrast to effects on the chemical shifts of amide and α -protons, the vicinal coupling constants between amide protons and α -protons of each residue of peptides **1–6** displayed no common tendency due to *N*-methylation or β-methylation. If anything, the values of each residue were similar among all the peptides **1–6**. This revealed that methylation did not result in drastic ϕ angle changes.

Temperature coefficients often indicate solvent accessibility of amide protons.^{18a} Kessler et al. previously reported that the temperature dependence of Arg H^N in cyclo(-Arg-Gly-Asp-D-Xaa⁴-Val-) and cyclo(-Arg-Gly-Asp-D-Phe-Yaa⁵-) is typically small, except in cases where cyclic amino acids such as proline are utilized for D-Xaa⁴ and Yaa⁵.^{10a} This data supports solvent shielding of Arg H^N and indicates the presence of a hydrogen bond corresponding to a type $II' \beta$ -turn substructure. On the other hand, only a small coefficient for Gly H^{N} is observed in peptide 2, although it has been reported that this has no relation to hydrogen bonding.⁸ If anything, peptide 2 appeared to exhibit conformational flexibility around the Gly residues. We examined this parameter comparatively in peptides 1-6, based on chemical shifts of amide protons in the range of 300-340 K (Table 2). Interestingly, among peptides **1–6**, a small coefficient for Arg H^N was observed in peptide **1** only, while the Gly H^N coefficients were small in the remaining peptides 2–6. Coefficients of other residues in 2–6 were over 2.0 ppb/K, although these varied somewhat for residue among the peptides. Thus, temperature dependence tendencies of amide protons in isostere-containing peptides **3–6** appeared to be nearly identical with **2**, but different from **1**. These observations implied that the conformations of 3-6may resemble one another in DMSO, and that these peptides may adopt flexible structures similar to 2, rather than the representative type II' β/γ arrangements seen with **1**.

Table 2. Temperature dependence of amide proton chemical shifts, $-\Delta\delta/\Delta T$ (ppb/K) of cyclic peptides 1–6^a

Peptide	Arg	Gly	Asp	D-Phe	Val
1	1.8	5.5	5.1	3.1	3.0
2	5.5	1.0	4.7	5.1	
3	5.4	2.2	3.0	3.5	
4	4.8	0.9	5.5	3.3	
5	5.7	2.5	5.5	2.7	_
6	6.8	-1.4	7.4	2.5	

^a The data for peptides **1–6** were obtained in comparative experiments using the same conditions.

Taking into account combined biological and ¹H NMR data, it is evident that the lack of a Val amide hydrogen incurred by *N*-methylation in **2**, may have contributed to conformational changes that increased $\alpha_v\beta_3$ antagonistic activity. This was also observed with peptides **3–6** having alkene isosteres as well. In other words, the Val amide proton in **1** may contribute unfavorably to bioactive conformations likely through intramolecular interactions, although such an amide proton originating from the *i*+2 residue of a β -turn would be indispensable for the distinctive type II' β/γ arrangement of cyclic pentapeptides. In contrast, it can be supposed that the carbonyl oxygen of D-Phe in **1** and **2** may be unrelated to significant interactions, since it has little apparent effect on conformation and bioactivity as compared to the amide proton of Val.

2.4. Structural calculations on cyclic peptidomimetics

To promote a better understanding of conformational aspects derived from ¹H NMR parameters, structural

calculations of the cyclic peptides 1-6 were carried out by simulated annealing molecular dynamics/energy minimization using dihedral constraints derived from ¹H NMR vicinal coupling constants and NOE distance constraints.²⁰ These calculations afforded well-converged conformations. Interestingly, calculated low-energy backbone structures of 1-6 are highly similar to each other (see Supporting information). Backbone structures based on the five α -carbons showed nearly symmetrical pentagonal shapes. In all cases, the olefinic moieties and peptide bonds were found to be vertical to the cyclic peptide plane, although some exhibited slightly differential rotations. Of note, the proposed type II' β/γ arrangement of **1** was not observed in either 2-6 or in 1 itself. This apparently reflects the fact that similar averaged parameters were used for structural calculations, as reported results by Nikiforovich et al.¹¹ This may indicate that it could be difficult to rationalize SAR studies on cyclic RGD peptides solely using structural calculation, unless the receptor-binding structures of ligands could be discussed. In practice, the presence of β - and/or γ -methyl groups in the isostere moiety appear to have little effect on the global backbone structures of the cyclic peptidomimetics, in spite of the fact that peptides **3–6** exhibited somewhat different bioactivities, respectively.

Ten superimposed low-energy structures of peptide 6 having the D-Phe- $\psi[(E)$ -CMe=CMe]-L-Val-type isostere are depicted in Figure 3 as representative of the isosterecontaining peptides **3–6**. Peptide **6** was the most potent $\alpha_{v}\beta_{3}$ integrin antagonist among 3-6. The root mean square deviation (RMSD) value for all backbone heavy atoms of 6 was below 0.22 Å, and the total energy values of the refined structures were in the range of 102–108 kcal/mol. The olefinic plane of the isostere was perpendicular to the plane of the cyclic peptide. This is an ideal substructural component for a type II' β -turn. In practice, the averaged dihedral ψ angle of D-Phe (-103.5°) and ϕ angle of Val (-92.9°) , were highly consistent with theoretical β -turn values. However, the expected β -turn hydrogen bond between the amide hydrogen of Arg and the α -carbonyl oxygen of Asp could not be identified, since the peptide bonds of Asp-D-Phe and Val-Arg were also oriented perpendicular to the cyclic peptide plane. The torsional angles, D-Phe ϕ and Val ψ , were apparently different from

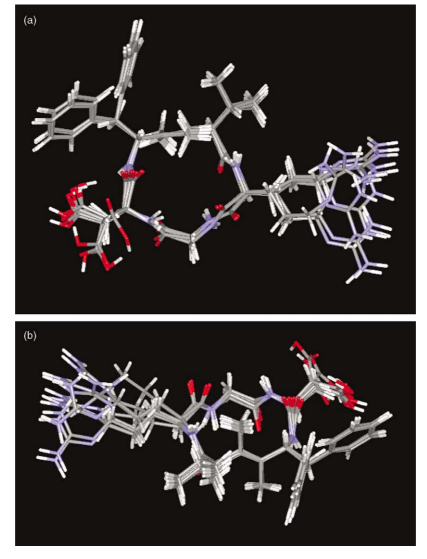


Figure 3. Overlay of ten low-energy structures of peptide 6. (a) Top view. (b) Side view.

those typically associated with a β -turn. This allows the side chains of all residues to exhibit pseudoequatorial conformations as derived from the conformational analysis of the peptide $2.^{8}$ Additionally, all isostere carbonyl oxygens and γ -methyl groups were commonly directed away from side chains of neighboring residues, most probably to avoid 1,3allylic strain across the peptide bonds. Similarly, isostere β-methyl groups were oriented upward so as to reduce steric interactions with D-Phe side chains. The averaged distance between the β -carbons of Arg and Asp of **6**, which provides topological orientation for two significant pharmacophores needed for bioactivity, was 9.0 Å. This distance was slightly longer than observed in 2, which had previously been determined in aqueous solution.8 These results indicated that the calculated conformation of 6 is more similar to that of the most potent peptide 2 having an N-methylvaline, rather than the proposed kinked conformation of 1, which is based on a type II' β/γ conformation.²¹

In $\alpha_{v}\beta_{3}$ integrin–ligand complexes, ligand 2 was reported to adopt a more distorted conformation as compared with structures in the absence of integrin.¹² In addition, it has been shown recently that cyclic RGD peptide ligands vary in conformations in the presence of integrins.^{17,23} Thus, it may be of significance to discuss the effects of the D-Phe-L-Val/N-MeVal moieties in 1-6 from the viewpoint of receptor-binding conformations, even if these moieties do not interact directly with the $\alpha_v\beta_3$ integrin in the crystal structure. Analyses of binding modes of 3-6 were not carried out. However, analogous conformations in the receptor-free state and the presence of common functional groups required for binding interactions, with the exception of the olefinic moiety, could enable an estimation of conformations of the most potent isostere-containing peptide 6 in the bound state. This presupposes that 6 can adhere to $\alpha_{\rm v}\beta_3$ integrin in a manner conformationally similar to 2.

3. Conclusion

In conclusion, SAR studies on cyclic RGD peptides were conducted using novel alkene dipeptide isosteres. Cyclic peptides 5 and 6, having D-Phe- ψ [(E)-CMe=CH]-L-Valand D-Phe- $\psi[(E)$ -CMe=CMe]-L-Val-type isosteres were designed and synthesized in order to investigate effects of the type II' β/γ arrangement found in **1** as well as the role of the N-methyl group of N-MeVal in 2 on conformation and biological activity. Evaluation of the biological activities of **1–6** against $\alpha_v \beta_3$ and $\alpha_{IIb} \beta_3$ integrin demonstrated that loss of the amide proton of Val in 1 by N-methylation led to a remarkable increase in $\alpha_v\beta_3$ antagonistic activity of 2, though this was not apparently due to steric factors arising from the methyl group. Structural analysis showed that γ -methylated isostere moieties would not be expected to serve as β -turn promoters, at least in these cyclic pentapeptides. Nevertheless, the calculated conformations of isostere-containing peptides 3-6 appeared to be analogous to those reported for the most potent peptide 2 rather than for 1. Taken together, these results indicate that influences of the N-methyl group on conformation and biological activity of 2 could be attributed mainly to loss of the amide hydrogen functionality in the D-Phe-N-MeVal

moiety, as opposed to steric factors such as allylic strain induced by the methyl group.

With advances in genome science, development of efficient methodologies for the rational design of therapeutically relevant agents from natural ligands is an area of increasing importance. As presented herein, alkene isosteres having differential methyl-substitutions could serve as practical tools to derive information concerning pharmacophores and bioactive conformations of bio- and chemoactive peptides and proteins.

4. Experimental

4.1. General synthetic

¹H NMR spectra were recorded using a Bruker AC 300 or a Bruker AM 600 spectrometer at 300 or 600 MHz. Chemical shifts of the compounds measured in CDCl₃ are reported in parts per million downfield from internal Me₄Si (s = singlet, d = doublet, dd = double doublet, t = triplet, m =multiplet). Those of the compounds measured in DMSO d_6 are calibrated to the solvent signal (2.50 ppm). Nominal (LRMS) and exact mass (HRMS) spectra were recorded on a JEOL JMS-01SG-2 or JMS-HX/HX 110A mass spectrometer. Optical rotations were measured with a Horiba highsensitive polarimeter SEPA-200 (Kyoto, Japan). For flash chromatographies, silica gel 60H (silica gel for thin-layer chromatography, Merck) and Wakogel C-200 (silica gel for column chromatography) were employed. For HPLC separations, a Cosmosil 5C18-ARII analytical $(4.6 \times$ 250 mm, flow rate 1 mL/min) column or a Cosmosil 5C18-ARII preparative $(20 \times 250 \text{ mm}, \text{ flow rate } 11 \text{ mL/}$ min) column was employed, and eluting products were detected by UV at 220 nm. A solvent system consisting of 0.1% TFA solution (v/v, solvent A) and 0.1% TFA in MeCN (v/v, solvent B) were used for HPLC elution.

4.1.1. (2R,5R,3E)-5-(9-Fluorenylmethoxycarbonyl)amino-2-isopropyl-4-methyl-6-phenylhex-3-enoic acid (Fmoc-D-Phe- ψ [(*E*)-CMe=CH]-L-Val-OH, 8a). After treatment of the ester 7a (108 mg, 0.258 mmol) with TFA (5 mL) for 1.5 h at room temperature, concentration under reduced pressure gave an oily residue. To a stirred solution of the above residue in MeCN-H₂O (2/1, 2.25 mL) were added Et₃N (0.072 mL, 0.517 mmol) and a solution of Fmoc-OSu (91 mg, 0.271 mmol) in MeCN (1.5 mL) at 0 °C. After being stirred for 3 h, the mixture was acidified with 0.1 N HCl and was extracted with EtOAc. The extract was washed with 0.1 N HCl and brine, and dried over MgSO₄. Concentration under reduced presssure followed by flash chromatography over silica gel with *n*-hexane–EtOAc (2/1)gave the title compound 8a (123 mg, 99% yield) as a colorless oil: $[\alpha]_{\rm D}^{20} - 16.6$ (c 0.542 in CHCl₃); ¹H NMR $(300 \text{ MHz}, \text{CDCl}_3, 55 \text{ °C}, \text{TMS}): \delta 0.70 \text{ (d}, J = 6.7 \text{ Hz}, 3\text{H}),$ 0.87 (d, J=6.6 Hz, 3H), 1.67 (s, 3H), 1.92 (m, 1H), 2.77-2.99 (m, 3H), 4.14 (t, J=6.6 Hz, 1H), 4.25-4.41 (m, 3H), 4.91 (m, 1H), 5.26 (d, J = 10.1 Hz, 1H), 7.08 (d, J = 6.9 Hz, 2H), 7.12–7.30 (m, 5H), 7.35 (t, J = 7.4 Hz, 2H), 7.49 (m, 2H), 7.72 (d, J=7.5 Hz, 2H). LRMS (FAB), m/z484 (MH⁺), 392, 260, 191, 179, 164, 154, 149, 143, 136,

91, 57, 43. HRMS (FAB), *m*/*z* calcd for C₃₁H₃₄NO₄ (MH⁺) 484.2488, found: 484.2477.

4.1.2. (2*R*,5*R*,3*E*)-5-(9-Fluorenylmethoxycarbonyl)amino-2-isopropyl-3,4-dimethyl-6-phenylhex-3-enoic acid (Fmoc-D-Phe- ψ [(*E*)-CMe=CMe]-L-Val-OH, 8b). By use of a procedure similar to that described for the preparation of the Fmoc-amino acid 8a from 7a, the ester 7b (138 mg, 0.319 mmol) was converted into the title compound 8b (131 mg, 83% yield) as a colorless oil: $[\alpha]_D^{24} - 70.9 (c 1.00 in CHCl_3); ^1H NMR (300 MHz, CDCl_3,$ $50 °C, TMS) <math>\delta$ 0.36 (m, 3H), 0.91 (d, *J*=6.4 Hz, 3H), 1.49 (m, 3H), 1.71 (d, *J*=1.4 Hz, 3H), 1.96 (m, 1H), 2.66 (m, 1H), 2.81 (m, 1H), 3.98 (m, 1H), 4.15 (t, *J*=6.4 Hz, 1H), 4.41 (m, 2H), 4.81 (br, 1H), 7.06 (br, 2H), 7.10–7.30 (m, 5H), 7.31–7.39 (m, 2H), 7.51 (m, 2H), 7.72 (m, 2H). LRMS (FAB), *m*/*z* 498 (MH⁺, base peak), 452, 406, 391, 274, 191, 179, 149, 136, 91, 69, 57, 43. HRMS (FAB), *m*/*z* calcd for C₃₂H₃₆NO₄ (MH⁺) 498.2644, found: 498.2641.

4.2. General procedure for assembly of the peptide chain

Protected peptide resins were manually constructed by Fmoc-based solid phase peptide synthesis. *t*Bu ester for Asp and (Boc)₂ for Arg were employed for side-chain protection. Fmoc-amino acids except for Fmoc-D-Phe- $\psi[(E)$ -CMe=CX]-Val-OH (X=H or Me) were coupled using 5 equiv of reagents [Fmoc-amino acid, *N*,*N*'-diisopropylcarbodiimide (DIPCDI), and HOBt·H₂O] to free amino group (or hydrazino group) in DMF for 1.5 h. Fmoc deprotection was performed by 20% piperidine in DMF (2×1 min, 1×20 min).

4.2.1. H-Asp(OtBu)-D-Phe-\psi[(E)-CMe=CH]-Val-Arg(Boc)₂-Gly-NHNHCO-Wang resin (11a). After treatment of *p*-nitrophenyl carbonate Wang resin **9** (0.93 mmol g⁻¹, 161 mg, 0.15 mmol) with NH₂NH₂·H₂O (0.046 mL, 0.75 mmol) in DMF (2 mL) at room temperature for 2 h, Gly and Arg(Boc)₂ residues were coupled by general coupling protocol. Fmoc-D-Phe- $\psi[(E)$ -CMe=CH]-Val-OH **8a** (48.3 mg, 0.100 mmol) was incorporated by double treatment with DIPCDI (0.018 mL, 0.120 mmol) and HOBt·H₂O (0.015 mg, 0.100 mmol) for 1.5 h each. After capping of the remaining free amino group with Ac₂O-pyridine, Asp(OtBu) residue was coupled by general coupling protocol to provide the title peptide resin **11a**.

4.2.2. H-Asp(OtBu)-D-Phe- $\psi[(E)$ -CMe=CMe]-Val-Arg(Boc)₂-Gly-NHNHCO-Wang resin (11b). By use of a procedure similar to that described for the preparation of the resin 11a, the title resin 11b was synthesized from *p*-nitrophenyl carbonate Wang resin 9 (0.15 mmol) and Fmoc-amino acid 8b (60 mg, 0.121 mmol).

4.2.3. Cyclo[-Arg-Gly-Asp-D-Phe- $\psi[(E)$ -CMe=CH]-Val-]·TFA (5). The protected peptide resin **11a** was treated with TFA for 1.5 h at room temperature. Removal of the resin followed by concentration under reduced pressure gave the colorless residue, which was purified by preparative HPLC (linear gradient of B in A, 15–20% over 45 min) to provide a peptide hydrazide **12a**. To a stirred solution of **12a** in DMF (12 mL) were added

a solution of 4 M HCl in DMF (0.075 mL, 0.300 mmol) and isoamyl nitrite (0.013 mL, 0.100 mmol) at -40 °C, and the mixture was stirred for 30 min at -20 °C. After dilution of the mixture with precooled DMF (68 mL), *i*Pr₂NEt (0.174 mL, 1.00 mmol) was added at $-40 \degree \text{C}$, and the mixture was stirred for 24 h at -20 °C. Concentration under reduced pressure and purification by preparative HPLC (linear gradient of B in A, 20-25% over 30 min) to give the cyclic pseudopeptide **5** (13.1 mg, 19% yield from **8a**) as freeze-dried powder: $[\alpha]_D^{20} - 59.4$ (*c* 0.656 in H₂O); $t_R =$ 33.4 min (linear gradient of B in A, 20-40% over 40 min); ¹H NMR (600 MHz, DMSO- d_6 , 25 °C) δ 0.46 (d, J = 6.6 Hz, 3H), 0.66 (d, J=6.6 Hz, 3H), 1.32–1.46 (m, 2H), 1.53 (m, 1H), 1.58 (s, 3H), 1.69 (m, 1H), 1.84 (m, 1H), 2.40 (dd, J = 16.2, 6.7 Hz, 1H), 2.56 (t, J = 9.1 Hz, 1H), 2.66 (dd, J)J = 16.2, 7.8 Hz, 1H), 2.74 (dd, J = 13.5, 9.7 Hz, 1H), 2.84 (dd, J = 13.5, 5.8 Hz, 1H), 3.07 (m, 2H), 3.26 (dd, J = 14.4)4.2 Hz, 1H), 3.98 (dd, J = 14.4, 6.8 Hz, 1H), 4.17 (m, 1H), 4.29 (m, 1H), 4.55 (m, 1H), 5.05 (d, J=9.4 Hz, 1H), 7.12-7.25 (m, 5H), 7.36 (d, J=8.3 Hz, 1H), 7.46 (m, 1H), 7.93–7.99 (m, 2H), 8.11 (d, J=8.3 Hz, 1H), 12.28 (br, 1H). LRMS (FAB), m/z 572 (MH⁺), 185, 154, 137, 93. HRMS (FAB), m/z calcd for C₂₈H₄₂N₇O₆ (MH⁺) 572.3197, found: 572.3208.

4.2.4. Cyclo[-Arg-Gly-Asp-D-Phe- ψ [(*E*)-CMe=CMe]-Val-]·TFA (6). By use of a procedure similar to that described for the preparation of the peptide 5 from the resin 11a, the resin 11b was converted into the title peptide 6 (16.9 mg, 20% yield): $[\alpha]_D^{22} - 62.6$ (c 0.846 in H₂O); $t_R =$ 36.3 min (linear gradient of B in A, 20-40% over 40 min); ¹H NMR (600 MHz, DMSO- d_6 , 25 °C) δ 0.26 (d, J = 6.5 Hz, 3H), 0.80 (d, J=6.4 Hz, 3H), 1.33–1.49 (m, 5H), 1.71 (s, 3H), 1.72–1.81 (m, 2H), 1.86 (m, 1H), 2.43 (dd, J=16.5, 6.8 Hz, 1H), 2.70–2.79 (m, 3H), 2.84 (dd, J=13.3, 5.1 Hz, 1H), 3.07 (m, 2H), 3.27-3.32 (m, 1H), 3.86 (m, 1H), 3.91 (dd, J = 14.4, 6.7 Hz, 1H), 4.52 (m, 1H), 4.92 (m, 1H),7.10-7.22 (m, 5H), 7.29 (m, 1H), 7.50 (br, 1H), 7.56 (d, J =6.9 Hz, 1H), 7.74 (d, J=7.4 Hz, 1H), 8.54 (d, J=8.2 Hz, 1H), 12.30 (br, 1H). LRMS (FAB), m/z 586 (MH⁺), 154 (base peak), 93, 91, 87, 70. HRMS (FAB), m/z calcd for $C_{29}H_{44}N_7O_6$ (MH⁺) 586.3353, found: 586.3368.

4.3. Integrin-binding assays

Compounds were evaluated for their inhibitory activities in $\alpha_{v}\beta_{3}$ and $\alpha_{IIb}\beta_{3}$ -ELISA (enzyme linked immunosorbent assay). $\alpha_{v}\beta_{3}$ was purified from human placenta, using RGDSPK-sepharose CL-4B affinity chromatography, followed by mono Q ion exchange chromatography, according to Pytela's protocol.²⁴ $\alpha_{IIb}\beta_3$ was purified from 24 human platelet by RGDSPK-sepharose CL-4B as well.²⁴ $\alpha_v \beta_3$ and $\alpha_{IIb} \beta_3$ binding assays were performed according to the modified method of Kouns et al.¹⁶ EIA plates were coated with $\alpha_v \beta_3$ or $\alpha_{IIb} \beta_3$, and blocked with bovine serum albumin. In each reaction, a test sample in the reaction mixture (20 mM Tris-HCl, 150 mM NaCl, 1 mM CaCl₂, 1 mM MgCl₂, pH 7.4, 0.100 mL) including vitronectin or fibrinogen, was added to the receptor-coated plate and incubated for 4 h at 25 °C. Thereafter the ligand binding was measured using anti-vitronectin rabbit antibody and peroxidase-conjugated anti-rabbit IgG antibody for $\alpha_{\rm v}\beta_3$, or peroxidase-conjugated anti-fibrinogen antibody for

 $\alpha_{IIb}\beta_3$, and 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) as the substrate of peroxidase. The IC₅₀ values were determined from measurement of absorbance at 415 nm.

4.4. NMR spectroscopy

The peptide sample was dissolved in DMSO- d_6 at concentration of 5 mM. ¹H NMR spectra of the peptides were recorded at 300 K using a Bruker AM 600 spectrometer at 600 MHz 1 H frequency. The chemical shifts were referenced to the residual DMSO (2.50 ppm). The assignments of the proton resonances were completely achieved by use of ${}^{1}H^{-1}H$ COSY spectra. ${}^{3}J(H^{N},H^{\alpha})$ coupling constants were measured from one-dimensional spectra. The mixing time for the NOESY experiments was set at 200, 300 and 400 ms. NOESY spectra were composed of 2048 real points in the F2 dimension and 512 real points, which were zero-filled to 1024 points in the F1 dimension, with 32 scans per t1 increment. The cross-peak intensities were evaluated by relative build-up rates of the cross-peaks. For the examination of the temperature dependence of the amide protons, the spectra of all peptides were also recorded at the every 10 K in the range of 300–340 K.

4.5. Calculation of structures

The structure calculations were performed on a Silicon Graphics Origin 2000 workstation with the NMR-refine program within the Insight II/Discover package using the consistent valence force field (CVFF). The prochiralities of two γ -methyl protons of Val were assigned based on the ${}^{3}J(\mathrm{H}^{\alpha},\mathrm{H}^{\beta})$ and the different NOE intensities in the NOESY spectra. On the other hand, the pseudoatoms were defined for the methylene protons of Arg, Asp and D-Phe, prochiralities of which were not identified by ¹H NMR data. The restraints, in which the Gly α -methylene participated, were defined for the separate protons without definition of the prochiralities. The dihedral ϕ angle constraints were calculated based on the Karplus equation: ${}^{3}J(\mathrm{H}^{\mathrm{N}},\mathrm{H}^{\alpha}) = 6.7 \cos^{2} (\theta - 60) - 1.3 \cos (\theta - 60) + 1.5.^{23}$ Lower and upper angle errors were set to 15°. The NOESY spectra with a mixing time of 200 ms were used for the estimation of the distance restraints between protons. The NOE intensities were classified into three categories (strong, medium and weak) based on the number of contour lines in the cross-peaks to define the upper-limit distance restraints (2.7, 3.5 and 5.0 Å, respectively). The upper-limit restraints were increased by 1.0 Å for the involved pseudoatoms. Lower bounds between nonbonded atoms were set to their van der Walls radii (1.8 Å). These restraints were included with force constants of $25-100 \text{ kcal mol}^{-1} \text{\AA}^{-2}$ for the distances and of $25-100 \text{ kcal mol}^{-1} \text{ rad}^{-2}$ for the dihedral angles. The 50 initial structures generated by the NMR refine program randomly were subjected to the simulated annealing calculations. Detailed protocols for the calculation are found in the Supporting information. The final minimization stage was achieved until the maximum derivative became less than 0.01 kcal mol⁻¹ Å⁻² by the steepest descents and conjugate gradients methods without any solvent matrix. The families of the preferred conformations were selected from the structures with energies not higher than 8 kcal mol^{-1} compared with the lowest energy.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11.033. ¹H NMR spectra for all new compounds; ¹H NMR data of **3–6**; protocols of structural calculations; calculated structures and averaged dihedral angles of **3–5**; and overlay of the representative structures of **3–6**.

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Regioselective electrophilic substitutions of fulvenes with ethyl glyoxylate and subsequent Diels–Alder reactions

Hsing-Chang Tseng, Arun Kumar Gupta, Bor-Cherng Hong* and Ju-Hsiou Liao

Department of Chemistry and Biochemistry, National Chung Cheng University, Chia-Yi 621, Taiwan, ROC

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Abstract—Highly regioselective electrophilic substitution of fulvenes with ethyl glyoxylate, catalyzed by EtAlCl₂ or Yb(OTf)₃ was achieved. Subsequent Diels–Alder reaction of the adduct with various dienophiles provides an efficient protocol toward highly functionalized indane and tricyclo[$5.2.1.0^{2.6}$]dec-8-ene systems.

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1. Introduction

For decades, Friedel-Crafts reactions of aromatic compounds have played important roles in organic synthesis.¹ Recent studies of electrophilic aromatic substitution with glyoxylate, catalyzed by Yb(OTf)3 or chiral bisoxazolinecopper (II) complexes, expand the applications in chemical synthesis.² Fulvenes, the benzene counterpart with nonbenzenoid aromaticity and high polarizability, usually have different reaction patterns with benzenes. Accordingly, very few examples of the Friedel-Crafts reactions of fulvenes with acyl chlorides have been reported.³ Yet, to the best of our knowledge, there are no reports of the direct Friedel-Crafts (or Alder-ene) reactions of fulvenes with aldehydes. In conjunction with our continuing efforts in fulvene chemistry,⁴ we reported herein a simple method for the direct Friedel-Craft reaction of fulvenes with glyoxylate and the subsequent regio- and stereoselective Diels-Alder cycloaddition. The sequence provides an efficient protocol toward the highly functional indane and tricyclo $[5.2.1.0^{2,6}]$ dec-8-ene system.

2. Results and discussion

2.1. Regioselective electrophilic substitutions of fulvenes with ethyl glyoxylate

Initially, a solution of 6,6-dimethylfulvene (1a) and ethyl glyoxylate in toluene was heated to reflux for 12 h, the

reaction affording trace amounts of the hydroxyester (2a) with complex mixtures and decomposition materials (Table 1, entry 1). Although reaction under microwave condition was accelerated (3 vs 12 h), decomposition with complex mixtures was still observed. Reaction under ultrasonic conditions did not proceed for 2 days, and the starting compounds were recovered (Table 1, entry 3). In recent studies, Lewis acid-catalyzed Friedel-Crafts reaction of aromatic compounds and ethyl glyoxylate have been shown to accelerate the reaction and improve yield.5 Accordingly, various Lewis acid catalysts were tested in the system (Table 1, entries 4-13). Among them, reaction with catalytic amounts of EtAlCl₂ in benzene gave the best result: 77% yield (Table 1, entry 12).⁶ A series of fulvenes (**1b–1g**) were reacted with ethyl glyoxylate to give the corresponding hydroxyesters (2b-2g) (Table 1, entries 14-21). In most cases, EtAlCl₂ was the best catalyst. However, Yb(OTf)₃ was more efficient than EtAlCl₂ in the reaction of dimethylamino fulvene (1b or 1c) and ethyl glyoxylate (Table 1, entries 14–17); the reaction afforded the two regioisomers 2b and 2b' in ca. 5:4 ratio. Interestingly, 2cwas obtained as one regioisomer. The regio-chemistry of these adducts was determined by NOE experiment, as depicted in the scheme, Table 2.

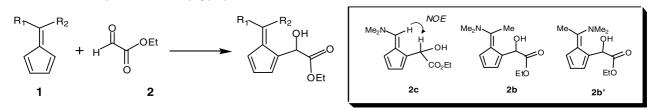
2.2. Diels-Alder reactions of adduct 2a

In order to expand the synthetic application of these hydroxyesterfulevenes, 2a was used as a representative example for the Diels–Alder reaction with dienophiles, such as maleic anhydride and maleic imide.⁷ Reaction of 2a with maleic anhydride in refluxing benzene yielded 85% of 3a and 3b in a 6:1 isomeric ratio (Table 2, entry 1). In contrast

Keywords: Diels–Alder; Friedel–Craft; Fulvene; Cycloaddition; Catalysis. * Corresponding author. Tel.: +886 52428174; fax: +886 52721040; e-mail: chebch@ccu.edu.tw

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Table 1. Reactions of alkylfulvenes with ethyl glyoxylate



Entry	Fulvene	Condition	Temperature (°C)	Time (h)	Yield (%) ^a
1	1a . $R_1 = R_2 = Me$	Toluene	110	12	5
2	1a . $R_1 = R_2 = Me$	Toluene, μwave ^b	100	3	5
3	1a . $R_1 = R_2 = Me$	Toluene, ultrasound ^c	25	48	NR
4	1a . $R_1 = R_2 = Me$	Cat. BF_3 – OEt_2 , toluene	-78	0.2	2
5	1a . $R_1 = R_2 = Me$	Cat. Yb(OTf) ₃ , THF	25	1	0^{d}
6	1a . $R_1 = R_2 = Me$	Cat. $Sc(OTf)_3$, toluene	25-60	0.5	18
7	1a . $R_1 = R_2 = Me$	Cat. Sc(OTf) ₃ , toluene, μ wave ^b	110	0.5	48
8	1a . $R_1 = R_2 = Me$	Cat. ZnCl ₂ , THF	25	0.5	0^{d}
9	1a . $R_1 = R_2 = Me$	Cat. AlCl ₃ , toluene	-30	0.5	0^{d}
10	1a . $R_1 = R_2 = Me$	Cat. Me ₂ AlCl, toluene	25	8	20
11	1a . $R_1 = R_2 = Me$	Cat. EtAlCl ₂ , toluene	0–25	3	40
12	1a . $R_1 = R_2 = Me$	Cat. EtAlCl ₂ , benzene	0–25	1.5	77
13	1a . $R_1 = R_2 = Me$	Cat. EtAlCl ₂ , CH ₂ Cl ₂	0–25	1.5	62
14	1b . $R_1 = NMe_2$; $R_2 = Me$	Cat. EtAlCl ₂ , benzene	0–25	3	22 ^e
15	1b . $R_1 = NMe_2$; $R_2 = Me$	Cat. Yb(OTf) ₃ , THF	25	3	73 ^e
16	1c . $R_1 = NMe_2; R_2 = H$	Cat. EtAlCl ₂ , benzene	0–25	3	5
17	1c . $R_1 = NMe_2; R_2 = H$	Cat. Yb(OTf) ₃ , THF	25	3	75
18	1d . $R_1 = R_2 = Et$	Cat. EtAlCl ₂ , benzene	0–25	1.5	72
19	1e . $R_1 = R_2 = n - Pr$	Cat. EtAlCl ₂ , benzene	0–25	1.5	75
20	1f . $R_1 = R_2 = -(CH_2)_{4-}$	Cat. EtAlCl ₂ , benzene	0–25	1.5	60
21	1g . $R_1 = R_2 = Ph$	Cat. EtAlCl ₂ , benzene	0–25	1.5	73

^a Isolated yield of **2**.

^a Isolated yield of 2. ^b Performed using a Synthewave S402 Prolabo microwave reactor (300 W; monomode system; 10-mL reactors) operated at 60% power. ^c Performed using an Elma Transsonic TP690-A operated at 35 kHz. ^d Decomposed into a complicated mixture.

^e Mixture of regioisomers (5:4).

Table 2.	Reaction	of fulvene	2a	with	dienophiles

Entry	Dienophile	Reaction conditions	Temperature (°C)	Time (h)	Products	Yield (%) ^a
1	°, °, °, °, °, °, °, °, °, °, °, °, °, °	Benzene	80	8	о H H H H CO ₂ Et H EO ₂ C O H H H H H H H H H H H H H	85 (6:1) ^b
2	0 0 0	Microwave, DMF ^c	130	0.5		77 ^d
3	CO ₂ Me	Benzene	80	8	MeO ₂ C HO Ga MeO ₂ C HO EtO ₂ C H HO EtO ₂ C H HO EtO ₂ C H HO EtO ₂ C H HO EtO ₂ C H HO EtO ₂ C H HO H HO H HO H HO H HO H HO H H HO H H HO H H HO H H H HO H	83 (1.6:1) ^b
4	CO ₂ Me	Microwave, toluene ^c	130	0.66		84 (2.1:1) ^b

Table 2 (continued)

Entry	Dienophile	Reaction conditions	Temperature (°C)	Time (h)	Products	Yield (%) ^a
5	O NPh O	Benzene	80	8	PhN H H H H H H H H	95 (1.7:1) ^b
6	O NPh O	Microwave, toluene ^c	130	0.66		93 (2.2:1) ^b
7	O NH O	Benzene	80	5	$HN \rightarrow HH \rightarrow CO_2Et \qquad HN \rightarrow HH \rightarrow HH \rightarrow HH \rightarrow HH \rightarrow HH \rightarrow HH \rightarrow HH$	87 (1.3:1) ^b
8	O NH O	Microwave, toluene ^c	130	0.66	60	92 (1.7:1) ^b
9	O NMe O	Benzene	80	5	$MeN \rightarrow H \rightarrow CO_2Et \qquad MeN \rightarrow H \rightarrow H \rightarrow CO_2Et \qquad H \rightarrow H \rightarrow CO_2C \rightarrow CO_2Et$	93 (1.9:1) ^b 1
10	O NMe O	Microwave, toluene ^c	130	1	34 9 0	85 (5:1) ^b

^a Isolated yield.

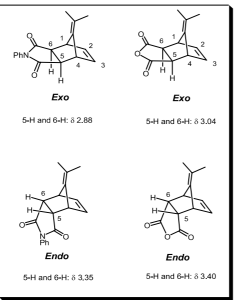
^b Ratio of isomers **a** and **b**.

^c Performed using a Synthewave S402 Prolabo microwave reactor (300 W; monomode system; 10-mL reactors) operated at 60% power.

^d Only one isomer observed.

to the endo-selectivity for the reaction of cyclopentadiene with maleic anhydride, reactions of 6,6-dimethylfulvene with N-phenylmaleic imide, maleic imide or maleic anhydride in toluene under reflux conditions have been reported to afford predominately the exo-adduct.⁸ Thus, we suspected that the two diastereoisomers are both exo adducts that differ in the stereochemistry of their CH(OH)CO₂Et units. Our hypothesis is consistent with the results provided by ¹H NMR spectroscopy. In general, the exo and endo 6,6dimethylfulvene adducts exhibit very different chemical shifts for their 5-H and 6-H protons (Scheme 1),⁹ but our pair of isomers display very similar chemical shifts for these protons. We confirmed the epimeric nature of 9a and 9b unambiguously through their oxidation with the Dess-Martin periodate; both diastereoisomers gave the same ketoester (4) as a single isomer.

Reaction of 2a with maleic anhydride under microwave conditions, however, afforded a different type of Diels-Alder adduct (5) (Table 2, entry 2). Maleic anhydride appears to add across the C-1 methyl and C-6 atom of fulvenes. The formation of 5 involves microwave-inducing isomerization of 2a to 2-isopropyenyl-cyclopenta-1,3diene, followed by trapping with maleic anhydride via [4+2] cycloaddition.¹⁰ The stereo- and regio-chemistry of 5 was determinate by NOE experiment, and the structure was concluded as depicted in entry 2 of Table 2. Reaction of 2a with other dienophiles gave the corresponding Diels-Alder adduct (Table 2, entries 3-10). It is noted that reactions under microwave conditions not only facilitate the reaction rate but also increase the diastereoselectivity, especially in entry 10 of Table 2. The structure and stereochemistry of compound 7b, the minor adduct, was





unambiguously assigned based on single crystal analysis (Fig. 1). 11

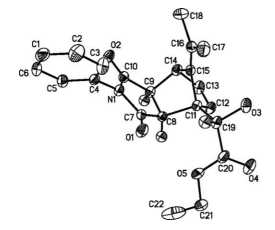
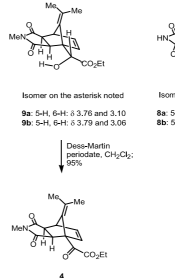


Figure 1. ORTEP plot for X-ray crystal structural of 7b.

The Diels–Alder reaction of fulvene (1a) with maleic anhydride usually requires a few days to reach completion. In comparison, the reaction of 2a with maleic anhydride is facile, presumably because of catalysis through intermolecular hydrogen bonding (Scheme 2). Such intermolecular hydrogen bonding not only facilitates the Diels–Alder reaction—acting as a Lewis acidic catalyst—but also plays a role in controlling the π -facial selectivity.¹² The observed periselectivity of the Diels–Alder reaction of 2a with dienophiles may arise from the fact that the reaction favors the transition state T1 over T2 because of its lesser degree of steric hindrance (Scheme 2), that is, the α -ester group in transition state T2 is eclipsed with the bridging carbon– carbon bond.

3. Conclusion

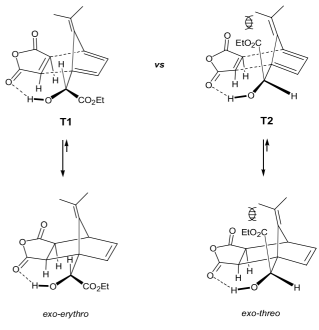
In summary, we have demonstrated that the Friedel-Crafts reactions of fulvenes with ethyl glyoxylate occur with



N O H H CO2Et

Isomer on the asterisk noted **8a**: 5-H, 6-H: δ 3.17 and 2.83 **8b**: 5-H, 6-H: δ 3.12 and 2.83

excellent regioselectivity when performed in the presence of catalytic amounts of either $EtAlCl_2$ or $Yb(OTf)_3$. In addition, the adducts obtained may be utilized in the construction of the frameworks of a number of natural products, such as those of the illudanes and FR182877. Further application of this methodology toward total synthesis of natural compounds is currently under investigation in our laboratory.



Scheme 2.

4. Experimental

4.1. General

All solvents were reagent grade. All chemicals were purchased from Aldrich Chemical Co. Reactions were normally carried out under argon atmosphere in flamedried glassware. Merck silica gel 60 (particle size 0.04–0.063 mm) was employed for flash chromatography. HPLC was equipped with the ultraviolet and refractive index detectors. The sample was analyzed and/or separated on a Spherisorb-Si column ($25 \text{ cm} \times 10 \text{ mm}$, particle size 8 μ , pore size 60 Å) or a μ -Porasil column (25 cm \times 1.0 cm) using a flow rate of 5 mL/min and ultraviolet and refractive index detectors (ethyl acetate and hexane eluents). The flow rate of the indicated elution solvent is maintained at 5 or 1 mL/min, and the retention time of a compound is recorded accordingly. Melting points are uncorrected. Most compounds were characterized by full spectroscopic (¹H, ¹³C, DEPT, HMQC, COSY, and NOESY) data. ¹H NMR, COSY and NOESY spectra were obtained in CDCl₃ unless otherwise noted at 400 MHz (Bruker DPX-400) or 500 MHz (Varian-Unity INOVA-500). ¹³C NMR spectra, HMBC, HMQC and DEPT experiments were obtained at 100 or 125 MHz.

4.1.1. Representative procedure for the synthesis of hydroxyesterfulvene, hydroxy-(5-isopropylidene-cyclopenta-1,3-dienyl)-acetic acid ethyl ester (2a). To a solution of dimethylfulvene 1a (27 mg, 0.25 mol) and ethyl glyoxylate (102 mg, 1.0 mmol) in dry benzene (1 mL) was added slowly a solution of ethylaluminum dichloride in C_6H_6 (0.05 mL, 1 M, 0.05 mmol) at 0 °C. The solution was stirred at 0 °C for 30 min and warm up slowly to 25 °C for 1.5 h. The reaction was quenched by the addition of H₂O (1.0 mL). The solution was extracted with EtOAc (25 mL), washed with brine, dried over Na₂SO₄ and concentrated in vacuo to give the crude product. The residue was purified by flash column chromatography with 25% EtOAc-hexane to give adduct **2a** (40 mg, 77% yield; $R_{\rm f}$ = 0.33 in 33% EtOAc-hexane) as a yellow oil. IR (neat): 3436, 2917, 1735, 1622, 1448, 1368, 1182, 1076, 829, 751 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 6.60 (dd, J = 5.5, 1.5 Hz, 1H), 6.36 (s, 1H), 6.29 (dd, J=5.5, 2.5 Hz, 1H), 5.28 (s, 1H), 4.26–4.32 (m 2H), 2.37 (s, 3H), 2.28 (s, 3H), 1.30 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.9 (C), 152.6 (C), 139.6 (C), 132.5 (CH), 131.9 (C), 126.8 (CH), 124.2 (CH), 68.6 (CH), 61.7 (CH₂), 25.7 (CH₃), 22.9 (CH₃), 14.1 (CH₃); MS (m/z, relative intensity): 209 $(M^+ + 1, 12), 192 (13), 191 (100), 115 (75), 106 (21), 102$ (42); exact mass calculate for $C_{12}H_{16}O_3$ (M⁺): 208.1100; found 208.1091.

4.1.2. 5-(**1**-Dimethylamino-ethylidene)-cyclopenta-1,3dienyl]-hydroxy-acetic acid ethyl ester (2b and 2b'). Prepared from **1b** according to procedure in Section 4.1.1. R_f =0.42 in 85% EtOAc–hexane, 73% yield, yellow oil; IR (neat): 3460, 2926, 1730, 1565, 1453, 1368, 1300, 1192, 1058, 1020, 916, 862, 803, 750 cm⁻¹; ¹H NMR (C₆D₆, 500 MHz): δ 6.90–6.54 (m, 3H), 5.49 (s, 1H), 4.02–3.89 (m, 2H), 2.32 (s, 6H), 1.74 (s, 3H), 0.87 (t, *J*=7.0 Hz, 3H); ¹³C NMR (C₆D₆, 125 MHz, 5:4 ratio, *note minor product): δ 175.0 (C and C*), 157.5 (C and C*), 135.2 (C and C*), 132.3 (C and C*), 122.0 (CH), 121.1 (CH*), 119.0 (CH*), 118.5 (CH), 118.2 (CH*), 115.5 (CH), 71.0 (CH), 70.8 (CH*), 61.04 (CH₂), 61.01 (CH₂*), 42.73 (2CH₃), 42.70 (2CH₃*), 20.2 (CH₃*), 20.0 (CH₃), 14.2 (CH₃*), 14.1 (CH₃); MS (*m/z*, relative intensity): 237 (M⁺, 1), 71 (15), 70 (10), 69 (16), 57 (43), 44 (31), 43 (100), 32 (17); exact mass calculate for $C_{13}H_{19}NO_3$ (M⁺): 237.1366; found: 237.1362.

(5-Dimethylaminomethylene-cyclopenta-1,3-4.1.3. dienvl)-hydroxy-acetic acid ethyl ester (2c). Prepared from 1c according to procedure in Section 4.1.1. $R_{\rm f}$ = 0.47 in 60% EtOAc-hexane, 75% yield, yellow oil; IR (neat): 3433, 2922, 2857, 1728, 1621, 1452, 1387, 1323, 1197, 1101, 799, 735 cm⁻¹; ¹H NMR (C₆D₆, 500 MHz): δ 7.52 (s, 1H), 6.67 (dd, J=4.0, 2.5 Hz, 1H), 6.63 (s, 1H), 6.59 (dd, J=4.5,1.5 Hz, 1H), 5.49 (s, 1H), 4.08–3.84 (m, 1H), 3.96–3.89 (m, 1H), 2.31 (s, 6H), 0.88 (t, J = 7.0 Hz, 3H); ¹³C NMR (CDCl₃, 100 MHz): δ 174.3 (C), 147.8 (CH), 130.7 (C), 121.6 (CH), 119.4 (CH), 116.6 (CH), 113.0 (C), 68.2 (CH), 61.5 (CH₂), 47.5 (CH₃), 40.3 (CH₃), 14.1 (CH₃); MS (m/z, relative intensity): 223 (M⁺, 45), 150 (86), 149 (45), 122 (57), 121 (49), 77 (33), 44 (45) 43 (55), 42 (100), 32 (72); exact mass calculate for $C_{12}H_{17}NO_3$ (M⁺): 223.1209; found: 223.1206.

4.1.4. [5-(1-Ethyl-propylidene)-cyclopenta-1,3-dienyl]hydroxy-acetic acid ethyl ester (2d). Prepared from 1d according to procedure in Section 4.1.1. $R_{\rm f} = 0.51$ in 33% EtOAc-hexane, 72% yield, yellow oil; IR (neat): 3473, 2971, 2936, 1735, 1616, 1368, 1182, 1463, 1371, 1300, 1201, 1063, 862, 764 cm $^{-1};$ $^1\mathrm{H}\,\mathrm{NMR}$ (CDCl₃, 500 MHz): δ 6.55 (d, J=5.0 Hz, 1H), 6.32 (s, 1H), 6.27 (dd, J=5.0, 2.0 Hz, 1H), 5.20 (s, 1H), 4.29-4.21 (m, 2H), 2.90-2.85 (m, 1H), 2.64–2.60 (m, 1H), 2.55–2.49 (m, 2H), 1.27 (t, J =7.0 Hz, 3H), 1.23–1.16 (m, 6H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.9 (C), 163.9 (C), 138.5 (C), 132.5 (CH), 131.9 (C), 127.1 (CH), 124.2 (CH), 68.6 (CH), 61.6 (CH₂), 29.1 (CH₂), 26.4 (CH₂), 15.4 (CH₃), 14.09 (CH₃), 14.06 (CH₃); MS (*m*/*z*, relative intensity): 236 (M⁺, 24), 218 (100), 164 (90), 163 (88), 145 (63), 144 (83), 107 (42), 77 (41), 43 (45), 41 (44); exact mass calculate for $C_{14}H_{20}O_3$ (M⁺): 236.1413; found: 236.1411.

4.1.5. Hydroxy-[5-(1-propyl-butylidene)-cyclopenta-1,3dienyl]-acetic acid ethyl ester (2e). Prepared from 1e according to procedure in Section 4.1.1. $R_{\rm f}$ = 0.67 in 33% EtOAc-hexane, 75% yield, yellow oil; IR (neat): 3503, 2960, 2932, 2871, 1733, 1613, 1456, 1368, 1206, 1058, 785, 758, 668 cm $^{-1};~^1\mathrm{H}$ NMR (CDCl_3, 500 MHz): δ 6.54 (dd, J=5.5, 1.5 Hz, 1H), 6.33 (s, 1H), 6.26 (dd, J=5.5, 2.5 Hz, 1H), 5.19 (s, 1H), 4.27–4.23 (m, 2H), 2.82–2.78 (m, 1H), 2.56-2.48 (m, 1H), 2.47-2.43 (m, 2H), 1.66-1.56 (m, 4H), 1.27 (t, J = 7.0 Hz, 3H), 1.01 (t, J = 7.5 Hz, 3H), 0.96 (t, J = 7.5 Hz, 3Hz), 0.96 (t, J = 7.5 Hz, 3Hz), 0.96 (t, J = 7.5 Hz, 3Hz), 0.96 (t, J = 7.5 Hz), 0.7.5 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.9 (C), 161.2 (C), 139.4 (C), 132.5 (CH), 132.0 (C), 126.9 (CH), 124.4 (CH), 68.6 (CH), 61.6 (CH₂), 38.7 (CH₂), 36.0 (CH₂), 24.2 (CH₂), 23.5 (CH₂), 14.6 (CH₃), 14.3 (CH₃), 14.1 (CH₃); MS (*m*/*z*, relative intensity): 264 (M⁺, 24), 246 (62), 192 (100), 174 (60), 172 (74), 107 (45), 91 (45), 79 (35), 55 (41); exact mass calculate for $C_{16}H_{24}O_3$ (M⁺): 264.1725; found: 264.1716.

4.1.6. Bicyclopentylidene-2,4-dien-2-yl-hydroxy-acetic acid ethyl ester (2f). Prepared from 1f according to procedure in Section 4.1.1. R_f =0.47 in 33% EtOAc-hexane, 60% yield, yellow oil; IR (neat): 3462, 2932, 2857, 1735, 1447, 1372, 1213, 1025, 859, 754 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 6.48 (dd, J=5.5, 1.5 Hz, 1H), 6.39 (s, 1H), 6.28 (dd, J=5.5, 2.5 Hz, 1H), 5.26 (d, J=7.5 Hz, 1H),

4.28–4.23 (m, 2H), 2.78–2.76 (m, 2H), 2.69–2.64 (m, 2H), 1.79–1.74 (m, 4H), 1.68–1.65 (m, 2H), 1.28 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.9 (C), 161.4 (C), 136.4 (C), 133.5 (CH), 131.8 (C), 126.7 (CH), 123.8 (CH), 69.0 (CH), 61.7 (CH₂), 35.6 (CH₂), 33.3 (CH₂), 29.7 (CH₂), 28.8 (CH₂), 26.4 (CH₂), 14.2 (CH₃); MS (*m*/*z*, relative intensity): 248 (M⁺, 26), 230 (36), 176 (93), 174 (100), 158 (47), 145 (57), 107 (33), 91 (48), 79 (39), 67 (31); exact mass calculate for C₁₅H₂₀O₃ (M⁺): 248.1412; found: 248.1411.

4.1.7. (5-Benzhydrylidene-cyclopenta-1,3-dienyl)hydroxy-acetic acid ethyl ester (2g). Prepared from 1g according to procedure in Section 4.1.1. $R_{\rm f} = 0.46$ in 33% EtOAc-hexane, 73% yield, yellow oil; IR (neat): 3502, 3056, 2980, 1713, 1583, 1491, 1443, 1198, 1053, 700 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.43–7.32 (m, 8H), 7.29–7.22 (m, 2H), 6.66 (s, 1H), 6.41 (dd, J=5.0, 2.5 Hz, 1H), 6.17 (dd, J=5.5, 2.0 Hz, 1H), 4.26 (s, 1H), 4.12 (q, J=7.0 Hz, 2H), 2.73 (br s, 1H), 1.21 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.8 (C), 154.0 (C), 142.2 (C), 141.8 (C), 140.4 (C), 133.8 (2CH), 132.3 (2CH), 131.6 (C and CH), 129.2 (CH), 129.03 (CH), 128.96 (CH), 128.1 (CH), 127.9 (2CH), 127.6 (2CH), 66.5 (CH), 61.4 (CH₂), 14.0 (CH₃); MS (m/z, relative intensity): 332 (M⁺, 13), 259 (100), 239 (11), 215 (24), 151 (10); exact mass calculate for C₂₂H₂₀O₃ (M⁺): 332.1413; found: 332.1409.

4.1.8. Representative procedure for the conventional Diels-Alder reaction for the synthesis of hydroxy-(10isopropylidene-3,5-dioxo-4-oxa-tricyclo[5.2.1.0^{2,6}]dec-8en-1-yl)-acetic acid ethyl ester (3a and 3b). A solution of fulvene 2a (30 mg, 0.14 mmol) and maleic anhydride (16 mg, 0.17 mmol) in benzene (2 mL) was heated to reflux for 8 h. The solution was concentrated in vacuo to give the residue as brown oil. The crude product was purified by flash column chromatography with 25% EtOAc-hexane (for **3a**: $R_{\rm f} = 0.38$ in 33% EtOAc-hexane; for **3b**: $R_{\rm f} = 0.37$ in 33% EtOAc-hexane) to give adduct 3a (30 mg; 71%) and **3b** (6 mg; 14%) as the yellow oils. For **3a**: IR (neat): 3448, 2923, 1780, 1731, 1636, 1373, 1227, 1075, 1016, 926, 754 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 6.53 (dd, J=6.0, 3.5 Hz, 1H), 6.16 (d, J = 6.0 Hz, 1H), 5.13 (s, 1H), 4.39-4.36 (m, 2H), 3.92 (dd, J=2.0, 1.0 Hz, 1H), 3.44 (d, J=8.0 Hz, 1H), 3.22 (br s, 1-OH), 3.10 (d, J=8.0 Hz, 1H), 1.74 (s, 3H), 1.64 (s, 3H), 1.35 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.1 (C), 170.6 (C), 169.4 (C), 138.9 (C), 138.3 (CH), 137.6 (CH), 119.5 (C), 67.6 (CH), 62.7 (CH₂), 59.7 (C), 51.0 (CH), 50.5 (CH), 48.5 (CH), 22.0 (CH₃), 18.9 (CH₃), 14.2 (CH₃). For **3b**: ¹H NMR (CDCl₃, 500 MHz): δ 6.42 (d, J=3.0 Hz, 1H), 6.31 (d, J=5.5 Hz, 1H), 5.20 (s, 1H), 4.26-4.22 (m, 2H), 3.92 (s, 1H), 3.38 (d, J = 8.0 Hz, 1H), 3.22 (br s, 1-OH), 3.08 (d, J = 9.0 Hz, 1H), 1.74 (s, 3H), 1.63 (s, 3H), 1.35 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 172.3 (C), 170.1 (C), 169.4 (C), 138.6 (C), 138.0 (CH), 136.5 (CH), 119.3 (C), 68.1 (CH), 62.3 (CH₂), 60.7 (C), 50.5 (CH), 50.1 (CH), 49.1 (CH), 20.6 (CH₃), 18.9 (CH₃), 14.0 (CH₃); MS (*m*/*z*, relative intensity): 306 (M⁺, 2), 191 (17), 190 (21), 136 (56), 135 (100); exact mass calculate for $C_{16}H_{18}O_6$ (M⁺): 306.1103; found: 306.1108.

4.1.9. Representative procedure for the microwave condition for the synthesis of (5-methyl-1,3-dioxo-1.3.3a,4,8a,8b-hexahydro-indeno[4,5-c]furan-6-vlidene)acetic acid ethyl ester (5). A mixture of fulvene 2 (30 mg, 0.14 mol) and maleic anhydride (16 mg, 0.17 mmol) in DMF (1 mL) were placed in a 10 mL quartz vial and subjected to programmed microwave irradiation at 180 W for 30 min. After a period of 2-3 min, the temperature reached a plateau of 130 °C where it remained throughout the reaction. After irradiation for 30 min and cooling, the solution was concentrated and the residue was purified by flash column chromatography with 45% EtOAc-hexane $(R_f=0.16 \text{ in } 33\% \text{ EtOAc-hexane})$ to give adduct 5 as a yellow liquid (33 mg, 77% yield). IR (neat): 3416, 2926, 1777, 1701, 1603, 1378, 1237, 1209, 1150, 1016, 987, 930, 786, 757 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.65 (dd, J=6.0, 1.5 Hz, 1H), 6.61-6.64 (m, 1H), 5.91 (s, 1H),4.20-4.14 (m, 2H), 3.65 (dd, J=10.0, 7.5 Hz, 1H), 3.54-3.49 (m, 1H), 3.35 (d, J=7.5 Hz, 1H), 2.77 (dd, J=15.0, 1.5 Hz, 1H), 2.56 (dd, J = 15.5, 6.0 Hz, 1H), 2.13 (s, 3H), 1.28 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 173.4 (C), 169.7 (C), 166.8 (C), 153.5 (C), 139.3 (CH), 135.7 (C),134.3 (CH), 134.0 (C), 111.2 (CH), 60.00 (CH₂), 44.9 (CH), 42.5 (CH), 41.8 (CH), 34.3 (CH₂), 21.5 (CH₃), 14.3 (CH₃); MS (*m*/*z*, relative intensity): 288 (M⁺, 4), 131 (20), 69 (36), 44 (100), 43 (64); exact mass calculate for $C_{16}H_{16}O_5$ (M⁺): 288.0998; found: 288.1007.

4.1.10. threo-1-(Ethoxycarbonyl-hydroxy-methyl)-7isopropylidene-bicyclo[2.2.1]hepta-2,5-diene-2,3-dicarboxylic acid diethyl ester (6a) and erythro-1-(ethoxycarbonyl-hydroxy-methyl)-7-isopropylidene-bicyclo-[2.2.1]hepta-2,5-diene-2,3-dicarboxylic acid diethyl ester (6b). Prepared from 2a and dimethylacetylenedicarboxylate according to procedure in Section 4.1.8. For **6a**. $R_{\rm f}$ = 0.39 in 33% EtOAc-hexane, 51% yield, yellow oil; IR (neat): 3472, 2952, 2920, 1717, 1626, 1435, 1371, 1285, 1214, 1123, 1097, 745 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.03–7.05 (m, 2H), 4.83 (d, J = 5.0 Hz, 1H), 4.43 (s, 1H), 4.37 (d, J =6.0 Hz, 1H), 4.27 (q, J=7.0 Hz, 2H), 3.76 (s, 3H), 3.72 (s, 3H), 1.65 (s, 3H), 1.52 (s, 3H), 1.25 (t, J = 7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 172.5 (C), 168.0 (C), 163.5 (C), 161.8 (C), 153.9 (C), 148.9 (C), 144.0 (CH), 142.2 (CH), 101.4 (C), 69.3 (C), 68.6 (CH), 62.0 (CH₂), 52.4 (2CH), 51.9 (CH₃), 21.0 (CH₃), 17.9 (CH₃), 13.9 (CH₃); MS (*m/z*, relative intensity): 350 (M⁺, 15), 245 (100), 217 (81), 185 (44), 157 (51), 128 (59), 115 (56), 32 (30); exact mass calculate for $C_{18}H_{22}O_7$ (M⁺): 350.1366; found: 350.1360. For **6b**: $R_f =$ 0.31 in 33% EtOAc-hexane, 32% yield, yellow oil; IR (neat): 3504, 2923, 1723, 1625, 1567, 1435, 1372, 1283, 1120, 1058, 940, 738 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.04–7.05 (m, 1H), 6.98 (d, J = 4.0, 2.0 Hz, 1H), 5.05 (br s, 1H), 4.46 (d, J)J=3.5 Hz, 1H), 4.24 (q, J=7.0 Hz, 2H), 3.78 (s, 3H), 3.75 (s, 3H), 1.53 (s, 3H), 1.44 (s, 3H), 1.27 (t, J = 7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 172.2 (C), 167.1 (C), 164.2 (C), 160.00 (C), 153.4 (C), 150.4 (C), 143.8 (CH), 141.3 (CH), 101.7 (C), 68.6 (CH), 68.1 (C), 61.9 (CH₂), 53.5 (CH), 52.3 (CH₃), 52.0 (CH₃), 20.8 (CH₃), 19.0 (CH₃), 14.0 (CH₃); MS $(m/z, \text{ relative intensity}): 350 (M^+, 15), 245 (100) 217 (81),$ 185 (44), 157 (51), 128 (59), 115 (56), 32 (30); exact mass calculate for C₁₈H₂₂O₇ (M⁺): 350.1366; found: 350.1361.

4.1.11. threo-Hydroxy-(10-isopropylidene-3,5-dioxo-4phenyl-4-aza-tricyclo[5.2.1.02,6]dec-8-en-1-yl)-acetic acid ethyl ester (7a) and erythro-hydroxy-(10-isopropylidene-3,5-dioxo-4-phenyl-4-aza-tricyclo[5.2.1.0^{2,6}]dec-8-en-1-yl)-acetic acid ethyl ester (7b). Prepared from 2a and N-phenyl-maleimide according to procedure in Section 4.1.8. For **7a**: $R_f = 0.53$ in 33% EtOAc-hexane, 60% yield, white solid; mp = 130-134 °C; IR (neat): 3472, 2924, 1709, 1597, 1497, 1455, 1379, 1185, 740, 691 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.32–7.50 (m, 2H), 7.24 (s, 1H), 7.06 (d, J=7.5 Hz, 2H), 6.83 (d, J=6.0 Hz, 1H), 6.43 (s, 1H),4.91 (s, 1H), 4.31-4.33 (m, 2H), 4.23 (s, 1H), 4.00 (s, 1H), 3.84 (d, J=8.0 Hz, 1H), 3.43–3.45 (m, 1H), 1.70 (s, 3H), 1.64 (s, 3H), 1.27 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 100 MHz): δ 176.4 (C), 176.3 (C), 172.9 (C), 146.7 (C), 137.1 (CH), 133.9 (CH), 131.4 (C), 129.2 (2CH), 128.8 (CH), 126.5 (2CH), 113.2 (C), 68.3 (CH), 62.7 (CH₂), 59.9 (C), 46.7 (CH), 44.9 (CH), 44.8 (CH), 22.6 (CH₃), 18.5 (CH₃), 14.0 (CH₃); MS (m/z, relative intensity): 381 (M⁺, 13), 317 (20), 308 (45), 191 (20), 135 (100), 91 (37), 77 (13), 32 (10); exact mass calculate for $C_{22}H_{23}NO_5$ (M⁺): 381.1583; found: 381.1585. For **7b**: $R_f = 0.41$ in 33% EtOAc-hexane, 35% yield, white solid; mp = 132-136; IR (neat): 3491, 2924, 1707, 1497, 1379, 1186, 754 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.45–7.47 (m, 2H), 7.38 (d, J= 7.5 Hz, 1H), 7.13 (d, J=7.5 Hz, 2H), 6.54 (dd, J=6.0, 3.5 Hz, 1H), 6.22 (d, J=6.0 Hz, 1H), 6.21 (d, J=2.0 Hz, 1H), 4.35–4.39 (m, 2H), 3.90 (d, J=3.0 Hz, 1H), 3.30 (d, J=7.5 Hz, 1H), 3.20 (s, 1H), 2.96 (d, J=7.5 Hz, 1H), 1.75 (s, 3H), 1.66 (s, 3H), 1.35 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 176.0 (C), 175.0 (C), 173.5 (C), 139.9 (C), 138.0 (CH), 137.6 (CH), 131.9 (C), 129.2 (CH), 128.9 (CH), 128.6 (CH), 126.9 (CH), 126.3 (CH), 117.9 (C), 68.1 (CH), 62.4 (CH₂), 59.5 (C), 50.0 (CH), 49.1 (CH), 48.1 (CH), 22.2 (CH₃), 19.2 (CH₃), 14.2 (CH₃); MS (*m/z*, relative intensity): 381 (M⁺, 10), 317 (20), 308 (45), 135 (100), 117 (17), 91 (37); exact mass calculate for $C_{22}H_{23}NO_5$ (M⁺): 381.1577; found: 381.1585.

4.1.12. threo-Hydroxy-(10-isopropylidene-3,5-dioxo-4aza-tricyclo[5.2.1.0^{2,6}]dec-8-en-1-yl)- acetic acid ethyl ester (8a) and erythro-hydroxy-(10-isopropylidene-3,5dioxo-4-aza-tricyclo[5.2.1.0^{2,6}]dec-8-en-1-yl)- acetic acid ethyl ester (8b). Prepared from 2a and maleimide according to procedure in Section 4.1.8. For 8a: $R_f = 0.36$ in 60% EtOAc-hexane, 50% yield, yellow oil; IR (neat): 3306, 1709, 1186, 788, 756 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.77 (s, 1H), 6.49 (dd, J=6.0, 3.5 Hz, 1H), 6.13 (d, J=6.0 Hz, 1H), 5.17 (s, 1H), 4.31–4.41 (m, 2H), 3.80 (d, J =2.5 Hz, 1H), 3.20 (br s, 1-OH), 3.17 (d, J = 7.5 Hz, 1H), 2.83 (d, J=7.0 Hz, 1H), 1.75 (s, 3H), 1.58 (s, 3H), 1.27 (t, J=7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 176.4 (C), 175.7 (C), 173.5 (C), 139.5 (C), 138.0 (CH), 137.4 (CH), 118.4 (C), 68.0 (CH), 62.4 (CH₂), 59.2 (C), 51.3 (CH), 50.2 (CH), 47.6 (CH), 22.1 (CH₃), 19.0 (CH₃), 14.2 (CH₃); MS $(m/z, \text{ relative intensity}): 305 (M^+, 2), 43 (100), 117 (16),$ 105 (13), 91 (40), 79 (12); exact mass calculate for $C_{16}H_{19}NO_5$ (M⁺): 305.1264; found: 305.1255. For **8b**: $R_{\rm f}$ =0.33 in 60% EtOAc-hexane, 37% yield, yellow oil; IR (neat): 3208, 1710, 1186, 783, 761 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 7.82 (s, 1H), 6.39 (dd, J=9.0, 3.5 Hz, 1H), 6.32 (d, J = 6.0 Hz, 1H), 5.20 (s, 1H), 4.22–4.26 (m, 2H), 3.82 (d, J=3.0 Hz, 1H), 3.12 (d, J=7.5 Hz, 1H), 2.83 (d,

J=7.5 Hz, 1H), 1.62 (s, 3H), 1.61 (s, 3H), 1.25 (t, J= 7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 177.0 (C), 176.4 (C), 172.1 (C), 139.1 (C), 138.2 (CH), 136.2 (CH), 118.3 (C), 68.8 (CH), 61.8 (CH₂), 60.2 (C), 50.5 (CH), 50.1 (CH), 47.9 (CH), 22.2 (CH₃), 20.4 (CH₃), 14.0 (CH₃); MS (*m*/*z*, relative intensity): 305 (M⁺, 6), 198 (21), 191 (25), 149 (23), 135 (100), 117 (13), 91 (36), 57 (45), 43 (47), 41 (37), 32 (38), 31 (43) exact mass calculate for C₁₆H₁₉NO₅ (M⁺): 305.1264; found: 305.1271.

4.1.13. threo-Hydroxy-(10-isopropylidene-4-methyl-3,5dioxo-4-aza-tricyclo[5.2.1.02,6]dec-8-en-1-yl)-acetic acid ethyl ester (9a) and erythro-hydroxy-(10-isopropylidene-4-methyl-3,5-dioxo-4-aza-tricyclo[5.2.1.0^{2,6}]dec-8-en-1yl)-acetic acid ethyl ester (9b). Prepared from 2a and N-methyl-maleimide according to procedure in Section 4.1.8. For **9a**: $R_f = 0.36$ in 50% EtOAc-hexane, 33% yield, yellow oil; IR (neat): 3460, 2924, 1696, 1343, 1379, 1292, 1132, 783 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 6.48 (dd, J = 6.0, 3.0 Hz, 1H), 6.14 (d, J = 6.0 Hz, 1H), 5.17 (s, 1H), 4.32-4.40 (m, 2H), 3.76 (d, J=3.0 Hz, 1H), 3.20 (br s, 1H), 3.10 (d, J=7.0 Hz, 1H), 2.90 (s, 3H), 2.79 (d, J=7.5 Hz, 1H), 1.65 (s, 3H), 1.54 (s, 3H), 1.35 (t, J = 7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 177.0 (C), 176.2 (C), 173.6 (C), 139.6 (C), 137.8 (CH), 137.2 (CH), 118.1 (C), 68.1 (CH), 62.4 (CH₂), 59.3 (C), 50.00 (CH), 49.00 (CH), 47.7 (CH), 24.5 (CH₃), 21.9 (CH₃), 18.9 (CH₃); 14.2 (CH₃) MS (m/z, relative intensity): 319 (M⁺, 7), 246 (22), 190 (44), 135 (100), 117 (18), 91 (36), 79 (10); exact mass calculate for $C_{17}H_{21}NO_5 (M^+)$: 319.1420; found: 319.1418. For **9b**: $R_f =$ 0.24 in 50% EtOAc-hexane, 63% yield, yellow oil; IR (neat): 3477, 2925, 1691, 1436, 1380, 1290, 1189, 1134, 794, 755 cm⁻¹; δ 6.38 (dd, J=6.0, 3.5 Hz, 1H), 6.32 (d, J= 6.0 Hz, 1H), 5.19 (s, 1H), 4.21–4.25 (m, 2H), 3.79 (dd, J =2.5, 1.0 Hz, 1H), 3.06 (d, J=7.0 Hz, 1H), 2.91 (s, 3H), 2.78 (d, J=7.0 Hz, 1H), 1.56 (s, 3H), 1.54 (s, 3H), 1.24 (t, J = 7.0 Hz, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 177.5 (C), 177.1 (C), 172.1 (C), 139.3 (C), 138.1 (CH), 136.0 (CH), 118.0 (C), 68.8 (CH), 61.7 (CH₂), 60.3 (C), 49.3 (CH), 48.9 (CH), 48.0 (CH), 24.6 (CH₃), 22.1 (CH₃), 20.2 (CH₃), 14.0 (CH₃); MS (m/z, relative intensity): 319 (M⁺, 7), 246 (24), 190 (27), 135 (100), 117 (14), 91 (22), 41 (13) exact mass calculate for $C_{17}H_{21}NO_5$ (M⁺): 319.1420; found: 319.1414.

(10-Isopropylidene-3,5-dioxo-4-oxa-tricy-4.1.14. clo[5.2.1.0^{2,6}]dec-8-en-1-yl)-oxo-acetic acid ethyl ester (4). To a solution of 9a and 9b mixture (4 mg, 0.012 mmol) in CH₂Cl₂ (2 mL) was added Dess-Martin periodinate (6 mg, 0.014 mmol) under Ar. The solution was stirred at room temperature for 20 min and diluted with EtOAc (20 mL). The mixture was washed with aqueous NaHCO₃ (1.0 mL), dried over Na₂SO₄ and concentrated in vacuo to give a yellow oil. The crude product was purified by flash column chromatography with 30% EtOAc-hexane ($R_{\rm f}$ = 0.56 in 50% EtOAc-hexane) to give adduct 4 as a yellow oil (3.6 mg, 95% yield). IR (neat): 2923, 1697, 1434, 1378, 1292, 1292, 1187, 749 cm⁻¹; ¹H NMR (CDCl₃, 500 MHz): δ 6.57 (dd, J=6.0, 3.5 Hz, 1H), 6.28 (d, J=6.0 Hz, 1H), 4.40 (q, J=7.5 Hz, 2H), 3.82 (s, 1H), 3.59 (d, J=7.0 Hz, 1H), 2.86 (s, 3H), 2.85 (s, 1H), 1.58 (s, 1H), 1.54 (s, 3H), 1.41 (t, J=7.5 Hz, 3H), 1.39 (s, 3H); ¹³C NMR (CDCl₃, 125 MHz): δ 193.0 (C), 176.5 (C), 176.5 (C), 162.1 (C),

138.9 (C), 138.5 (CH), 137.1 (CH), 119.6 (C), 64.5 (C), 62.7 (CH₂), 49.4 (CH), 48.3 (CH), 47.6 (CH), 24.6 (CH₃), 21.4 (CH₃), 20.9 (CH₃), 14.0 (CH₃); MS (*m*/*z*, relative intensity): 317 (M⁺, 3), 244 (9), 133 (100), 77 (6), 58 (9), 55 (8), exact mass calculate for $C_{17}H_{19}NO_5$ (M⁺): 317.1264; found: 317.1264.

5. Supplementary material

Experimental procedures and characterization data for the new compounds (2–9); and X-ray crystallographic cif file for compound **7b**. The X-ray cif file of compound **7b** was deposited with Cambridge Crystallographic Data Centre, CCDC No. 288537.

Acknowledgements

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A simple approach for the regioselective synthesis of imidazo[1,2-*a*]pyrimidiones and pyrimido[1,2-*a*]pyrimidinones

David Font,^a Anthony Linden,^b Montserrat Heras^{a,*} and José M. Villalgordo^{a,*,†}

^aDepartment of Chemistry, Faculty of Sciences, University of Girona, Campus de Montilivi, 17071 Girona, Spain ^bInstitute of Organic Chemistry, University of Zürich, Winterthurerstrasse 190, CH-8057 Zürich, Switzerland

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Abstract—Several imidazo and pyrimido[1,2-*a*]pyrimidinones of type **1** and **2** were synthesized through intramolecular cyclization of pyrimidines **9** or pyrimidinones **10** bearing a variety of β and γ -aminoalcohols at the 2-position. Ring closure of the pyrimidinones of type **10** under Mitsunobu conditions lead to mixtures of both bicyclic regioisomers **1** and **2**. Treatment of pyrimidines of type **9** with H₂SO₄ provided an efficient and operationally simple one-pot hydrolysis–cyclization procedure for obtaining imidazo and pyrimido[1,2-*a*]pyrimidinones **1** in good yields as the sole regioisomeric bicyclic product.

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1. Introduction

The imidazo[1,2-*a*]pyrimidinones 1 (n=1) and 2 (n=1) and the pyrimido[1,2-*a*]pyrimidinones 1 (n=2) and 2 (n=2) (Fig. 1) constitute an important class of natural and non-natural products, many of which exhibit useful biological activities. The imidazo[1,2-*a*]pyrimidinone structure 1 (n=1) has been found in the Y base of yeast as a component of phenylalanine transfer ribonucleic acid,¹ and new acyclovir analogs possessing this ring system have exhibited antiherpetic activity on HIV-1,2.² Other members of this family of compounds also have pharmacological interest for their hypnotic,³ anesthetic⁴ and antiallergic⁵ properties. Imidazo[1,2*a*]pyrimidinone derivatives type 2 (n=1) are of considerable interest because of their activities as phosphodiesterase (PDE) inhibitors,⁶ and antihypertensives.⁷ Furthermore, some derivatives of pyrimido[1,2-*a*]pyrimidinones **1** (n=2) and **2** (n=2) have some utility for preventive and/or therapeutic treatment of a neuro-degenerative disease caused by abnormal activity of GSK3 β , such as Alzheimer's disease.⁸

The imidazo and pyrimido[1,2-a]pyrimidinones **1** and **2** incorporate both the guanidine and pyrimidinone functionalities (Fig. 1). It is well known that pyrimidin-4(*3H*)-ones are valuable scaffolds in different areas of research. For example, this class of compounds displays potent and selective activity as non-nucleoside HIV-1 reverse transcriptase inhibitors.⁹ Other members of this family of compounds have found utility as herbicides,¹⁰ antidepressants¹¹ and leishmanicides.¹² In addition, when pyrimidin-4(*3H*)-ones are substituted at the

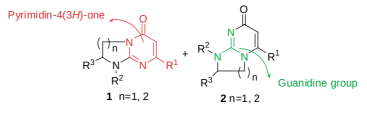


Figure 1. Imidazo and pyrimido[1,2-a]pyrimidinones 1 and 2.

* Corresponding authors. Tel.: +34 972418247; fax: +34 972418150 (M.H.); Tel.: +34 968809089; fax: +34 968890024 (J.M.V.); e-mail addresses: montserrat.heras@udg.es; info@villapharma.com

Keywords: Imidazo[1,2-*a*]pyrimidinone; Pyrimido[1,2-*b*]pyrimidinone; Mitsunobu reaction; *ipso*-Substitution; Acidic hydrolysis.

[†] Present address: Villapharma Research, Polígono Industrial Oeste, c/Paraguay. Parcela 7/5-A, Módulo A, 30169 Murcia, Spain.

2-position by an amino group, it can be considered to be a cyclic guanidine. Due to the hydrogen-bonding acceptor and donor abilities of the guanidine group,¹³ 2-aminopyrimidin-4(3H)-ones have also served as suitable models for studies conducted on self-association¹⁴ and the subsequent application of those studies to supramolecular chemistry.¹⁵

Numerous methods for the synthesis of the imidazo and pyrimido[1,2-*a*]pyrimidinones involve approaches based on either (i) cyclocondensation between 2-substituted pyrimidinone ring systems with appropriate 1,2 or 1,3-difunctionalized synthons, such as α or β -halocarbonyl compounds,^{2,16} 1,2-dihaloalkanes,¹⁷ acrolein,¹⁸ glyoxal,¹⁹ glycidaldehyde,²⁰ and α or β -aminoalcohols,^{6,7,21} or (ii) cyclocondensation between 2-substituted imidazole or pyrimidine ring systems with appropriate 1,3-difunctionalized synthons, such as β -aminoesters²² and α -acetylenic esters.²³ However, both routes can give mixtures of regioisomers. Other useful routes to these type of heterocycles involve the fusion of two heterocycles in one single step²⁴ or the ring contraction of other heterocyclic systems.²⁵

During the last few years, we have been engaged in a research program focused on the development of efficient methodologies that could be adapted readily for combinatorial and/or parallel synthesis of relevant core structures with potential therapeutic interest. We have described the synthesis of novel 2,3-dihydroimidazo[2,1-b][1,3]oxazoles²⁶ and multiple substituted pyrimidines.²⁷ The method has a nucleophilic *ipso*-substitution of the corresponding activated sulfones as one of the key steps, not only for introducing molecular diversity but also as cleavage reaction on solid phase synthesis (Scheme 1). In this way, several purines,²⁸ aminopyridazines²⁹ and pteridines³⁰ have also been prepared using an activatable sulfur linkage. Within this context, we recently reported on the synthesis of novel 2,6-disubstituted 3,4-dihydropyrimidin-4(3H)-ones³¹ 7 starting from 2-alkylsulfanylpyrimidinones 3. The methodology is based on a selective O-alkylation reaction with *i*-PrOH under Mitsunobu conditions, followed by a

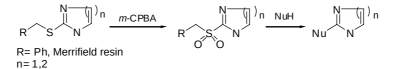
nucleophilic displacement of the corresponding activated sulfones with a wide variety of nucleophiles (phenoxides, Grignard reagents, and primary and secondary amines). Finally, the acidic hydrolysis of the 4-isopropoxy group under standard conditions afforded pyrimidinones 7 in good yields (Scheme 2).

As an extension of this work, an investigation was undertaken to expand the scope of this methodology and its potential application in the synthesis of more elaborate heterocyclic scaffolds based on the pyrimidin-4(3H)-one nucleus. Specifically, we focused our attention on imidazo[1,2-*a*]pyrimidinones and pyrimido[1,2-*a*]pyrimidinones **1** and **2**. The results of this investigation are disclosed herein.

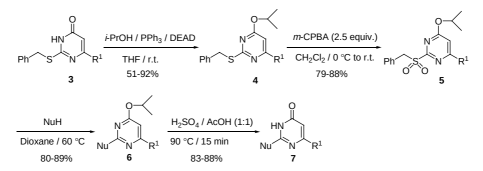
2. Results and discussion

Consistent with the goal of synthesizing more elaborate heterocyclic scaffolds based on the pyrimidin-4(3*H*)-one nucleus, and in complete analogy with the above-mentioned results, we reasoned that nucleophilic *ipso*-substitution of the sulfones **5** with a wide variety of β and γ -aminoal-cohols³² **8**, followed by subsequent acidic hydrolysis and a final cyclization step under Mitsunobu conditions would lead to the formation of a collection of imidazo and pyrimido[1,2-*a*]pyrimidin-5-ones **1** and imidazo and pyrimido[1,2-*a*]pyrimidin-7-ones **2** (Scheme 3). From these intermediates **10** the cyclization could, in principle, take place onto *N*(1) or *N*(3) to afford the regioisomers **2** and **1**, respectively.

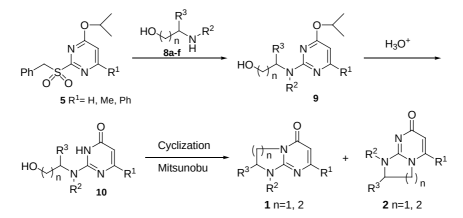
Thus, when pyrimidinyl sulfone derivatives **5** were allowed to react in 1,4-dioxane at reflux with several β and γ -aminoalcohols **8a–f** (Fig. 2), which are readily available from commercial sources and/or from the reduction of the corresponding amino acids,³³ the corresponding pyrimidines **9a–j** were obtained generally in good yields (Scheme 3, Table 1).



Scheme 1. Nucleophilic ipso-substitution of activated sulfones.



Scheme 2. Synthesis of 2,6-disubstituted 3,4-dihydropyrimidin-4(3H)-ones 7.



Scheme 3. Preparation of imidazo and pyrimido[1,2-a]pyrimidinones.

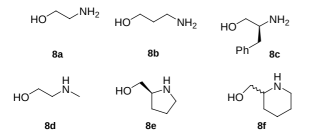


Figure 2. The employed β and γ -aminoalcohols 8.

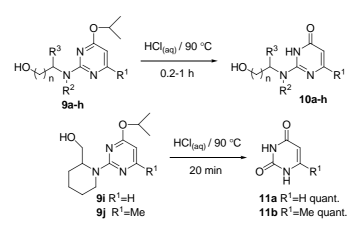
Table 1. Yields of compounds 9a-j

Entry	Compound	\mathbb{R}^1	Aminoalcohol	Yield (%) ^a
1	9a	Н	8a	79
2	9b	Me	8a	76
3	9c	Н	8b	65
4	9d	Me	8b	94
5	9e	Me	8c	79
6	9f	Ph	8d	85
7	9g	Me	8d	80
8	9h	Me	8e	95
9	9i	Н	8f	79
10	9j	Me	8f	90

^a Yields of isolated pure products.

Acidic hydrolysis of the pyrimidines **9** with concd HCl at 90 °C, followed by simple chromatographic filtration, yielded the corresponding target pyrimidinones **10** also in good yields (71–97%), except with the pyrimidines **9i** and **9j**, which led to the formation of the uracils **11** in quantitative yields (Scheme 4, Table 2). These results clearly indicate that the piperidinyl group in the 2-position of the pyrimidine ring is labile under these acidic conditions. We then focused our attention on the search for other acidic hydrolysis conditions that could selectively cleave the 4-alkoxy group in these two compounds, **9i** and **9j**.

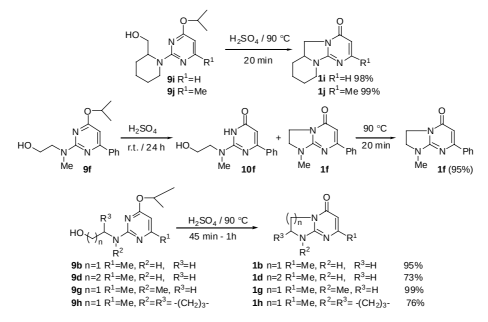
When derivatives **9i** and **9j** were allowed to react with H_2SO_4 at 90 °C, compounds **1i** and **1j** were isolated in near quantitative yields (Scheme 5). The formation of products **1** could be rationalized in terms of a one-pot procedure simply by hydrolysis of the 4-isopropoxy group, followed by complete regioselective cyclization onto N(3) of the pyrimidinone ring to afford the corresponding imidazo-pyrimidinones **1**. In good agreement with this procedure, when pyrimidine **9f** was treated with H_2SO_4 at room temperature for 24 h, a mixture of pyrimidinone **10f** and imidazopyrimidinone **1f** was observed. After heating to 90 °C, the ring closure was completed in only 20 min and compound **1f** was isolated in good yield (Scheme 5). This one-pot hydrolysis–cyclization reaction was successfully extended to other pyrimidines to afford the corresponding



Entry	Compound	\mathbb{R}^1	\mathbb{R}^2	\mathbb{R}^3	n	Yield (%) ^a
1	10a	Н	Н	Н	1	86
2	10b	Me	Н	Н	1	97
3	10c	Н	Н	Н	2	97
4	10d	Me	Н	Н	2	87
5	10e	Н	Н	Bn	1	71
6	10f	Ph	Me	Н	1	95
7	10g	Me	Me	Н	1	85
8	10h	Me	-(Cl	$H_2)_3 -$	1	94
9	10i	Н	-(Cl	$H_2)_4-$	1	_
10	10j	Me		$H_2)_4-$	1	_

Table 2. Yields of compounds 10a-j

^a Yields of isolated pure products.

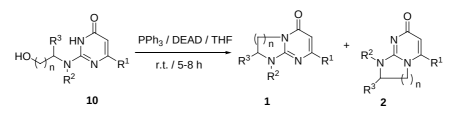


Scheme 5. Regioselective synthesis of imidazo and pyrimido[1,2-a]pyrimidinones 1.

imidazopyrimidinones 1 (n=1) and pyrimidopyrimidinone 1 (n=2) with an absolute regioselectivity and in good yields (Scheme 5).

Following our initial plans we decided to investigate the feasibility of an intramolecular cyclization of the 4(3*H*)pyrimidinones **10**, under Mitsunobu conditions. Thus, when pyrimidinones **10** were treated with PPh₃ and DEAD in anhydrous THF at room temperature for 5–8 h, a separable mixture of the regioisomeric bicyclic compounds **1** and **2** were isolated by flash chromatography in good yields (Scheme 6, Table 3). The cyclization reaction took place predominantly onto N(1) affording compounds of type **2** as the major regioisomer (Table 3, entries 1–4). However, when the *N*-atom at the 2-position on the pyrimidinone ring had an alkyl substituent ($R^2 = Me$), the Mitsunobu reaction proceeded in high yield and with a high degree of selectivity. Only the isomer **1g** from cyclization onto N(3) of the pyrimidinone ring was obtained (Table 3, entry 5).

Generally, the intramolecular cyclization reaction of 2-substituted pyrimidinone ring systems takes place onto N(3), except when N(3) has an alkyl substituent,^{5b,c} which blocks this nitrogen, and cyclization is only possible onto N(1) or when the 2-position on the pyrimidinone ring has a substituent prone to tautomerize, such as an amino^{16b–c} group, with the cyclization then taking place onto both N(1) and N(3) to afford mixtures of the regioisomers **1** and **2**. This last case can explain the results in the Mitsunobu reaction. Thus, when the reaction was carried out by employing pyrimidinones **10** with a secondary amine in the 2-position ($\mathbb{R}^2 = \mathbb{H}$), both regioisomers **1** and **2** were obtained (Table 3,



Scheme 6. Intramolecular cyclization of pyrimidinones 10 under Mitsunobu conditions.

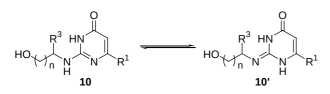
Entry	\mathbb{R}^1	\mathbb{R}^2	R ³	n	Regioisomer 1	Regioisomer 2	1:2 ^a	Yield (%) ^b
1	Н	Н	Н	1	1a	2a	12:88	91
2	Н	Н	Н	2	1c	2c	23:77	95
3	Me	Н	Н	2	1d	2d	27:73	91
4	Me	Н	Bn	1	1e	2e	42:58	78
5	Me	Me	Н	2	1g	—	100:0	96

Table 3. Yields of compounds 1 and 2

^a Ratios were calculated by yields of isolated compounds.

^b Yields of isolated pure products.

entries 1–4). The cyclization reaction onto N(1) probably proceeded via its tautomeric form 10' (Scheme 7). However, when pyrimidinone 10e with a tertiary amine in the 2-position (R²=Me) was employed, the tautomeric form 10' was not possible and the cyclization reaction was completely regioselective in favor of the isomer 1 (Table 3, entry 5).



Scheme 7. Tautomerism of compounds 10.

On the other hand, the absolute regioselectivity obtained during the synthesis of compounds 1, regardless of the R^2 group (H, Me), when intramolecular cyclization was carried out with H₂SO₄, are in good agreement with the literature. To the best of our knowledge, intramolecular cyclization of

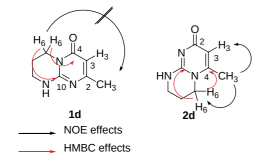


Figure 3. HMBC and NOE experiments of compounds 1d and 2d.

2-subtituted pyrimidin-4(3*H*)-ones always took place exclusively onto N(3) under acid reaction conditions.³⁴

All products and intermediates were characterized by the usual spectroscopic methods, such as ¹H and ¹³C spectroscopy, mass spectrometry, IR, and elemental analysis. The ¹H and ¹³C NMR spectra for 1 and 2 were assigned by means of DEPT and HMQC experiments and the regiochemistry of both of these isomers 1 and 2 was established unequivocally by NOE and heteronuclear multiple bond correlation (HMBC) experiments. A NOE effect was observed between methyl protons and proton H₃, as well as between methyl protons and protons H₆ in isomers 1d, while in isomers 2d, a NOE effect was not observed between protons H₆ and methyl protons (Fig. 3). In addition, the HMBC experiments gave supplementary information: for isomer 1, long range ¹H-¹³C correlations are observed between protons H_6 and both carbons 4 (C=O) and 10 (C=N) (Fig. 3), while isomer 2 presented correlations between protons H_6 and carbon 2 (C=O) and carbon 4 (Fig. 3). Moreover, the structures of compounds 2e and 1h were established unambiguously by X-ray crystallography (Fig. 4).

3. Conclusion

In summary, we have shown that 2-substituted pyrimidinones **10** with a variety of β and γ -aminoalcohols, which are easily available from pyrimidinyl sulfone derivatives **5**, are good synthetic precursors for the preparation of imidazo and pyrimido[1,2-*a*]pyrimidinones **1** and **2** via an intramolecular ring closure. When cyclization was carried out under Mitsunobu conditions by employing pyrimidinones **10** with a secondary amine

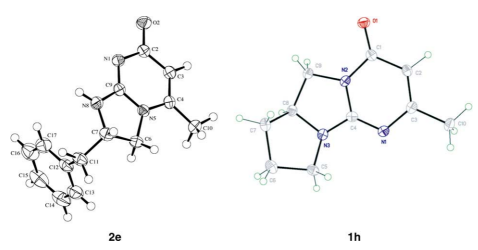


Figure 4. The molecular structures of 2e and 1h (Ortep-plot with ellipsoids at the 50% probability levels).

in the 2-position $(R^2=H)$, both regioisomers 1 and 2 were obtained. In contrast, when the N atom in the 2-position on the pyrimidinone ring has an alkyl substituent ($R^2 = Me$), the Mitsunobu reaction yielded the regioisomer 1 as the only product. On the other hand, treatment of the pyrimidines $\boldsymbol{9}$ with H_2SO_4 afforded, with an absolute regioselectivity, the imidazo and pyrimido[1,2-a]pyrimidinones 1 through a one-pot hydrolysis-cyclization procedure. Considering the easily available starting materials, generality of the reaction, simplicity of the procedure and good yields, this provides a straightforward method to construct a diverse array of imidazo and pyrimido[1,2-a] pyrimidinones 1 and 2. However, we are aware that the orientation of the cyclization reaction could change when pyrimidinone ring would be substituted with strong electron-withdrawing or electron-donating groups. In this way, further investigation on regioselective cyclization of the pyrimidinones substituted at 6-position with nitro, amino, alkylamino or alkoxy groups are currently in progress in our laboratories.

4. Experimental

4.1. General remarks

4-Isopropoxy-2-phenylmethanesulfonyl-pyrimidine 5a, 4-isopropoxy-6-methyl-2-phenylmethanesulfonyl-pyrimidine **5b** and 4-isopropoxy-6-phenyl-2-phenylmethanesulfonyl-pyrimidine 5c, were prepared as previously reported by us.²⁸ All commercially available chemicals were used as purchased without further purification. DMF and dioxane were dried over activated molecular sieves (4 Å). THF was dried over Na/benzophenone prior use. Melting points (capillary tube) were measured with an Electrothermal digital melting point apparatus IA 91000 and are uncorrected. IR spectra were recorded on a Mattson-Galaxy Satellite FT-IR using a single reflection ATR system as a sampling accessory. ¹H and ¹³C NMR spectra were recorded at 200 and 50 MHz, respectively, on a Brucker DPX200 Advance instrument. Spectra recorded in CDCl₃ were referenced to residual CHCl₃ at 7.26 ppm for ¹H or 77.0 ppm for ¹³C. Spectra recorded in DMSO- d_6 were referenced to residual DMSO at 2.49 ppm for ¹H or 39.5 ppm for ¹³C. Coupling constants (J) are given in Hertz (Hz). The terms s, d, t, q, sept, m, dd, refer to singlet, doublet, triplet, quartet, septet, multiplet; double doublet, br implies the signal is broad. Mass spectra were recorded by electron impact (EI, 70 eV) on a Thermo Quest 2000 series apparatus or by fast-atom bombardment (FAB) on a VG Quattro instrument or by electrospray ionitzation (ESI) using a quadrupole mass spectrometer equipped with an electrospray ion source. Elemental analyses were performed on an apparatus from Thermo Instruments, model EA1110-CHNS. Analytical thin-layer chromatography (TLC) was performed on glass plates precoated with silica gel 60 F_{254} (Merck). Visualitzation was accomplished by UV light (254 nm) and potassium permanganate. Flash-chromatography (FC) purifications were performed on silica gel 60 (230-400 mesh, Merck).

4.2. General procedure for the *ipso*-substitution reaction of pyrimidinyl sulfone derivatives 5 with amino alcohols **8.** Synthesis of pyrimidines 9

Under a nitrogen atmosphere, to a solution of pyrimidinyl sulfones 5a-c (1 equiv) in dry dioxane (3 mL/mmol), the corresponding amino alcohol 8a-f (1.5–2 equiv) was added. The resulting mixture was refluxed for 5–24 h until the reaction was completed (TLC monitoring). The solvent was removed under reduced pressure and the resulting residue was purified by flash-chromatography (*n*-hexane/ethyl acetate 4:1 gradually increasing to pure ethyl acetate) to afford pyrimidines **9**.

4.2.1. 2-(4-Isopropoxy-pyrimidin-2-ylamino)-ethanol (**9a).** From 1.61 g (5.51 mmol) of sulfone **5a** and 0.50 mL (8.1 mmol) of 2-amino-ethanol **8a**, 857 mg (79%) of compound **9a** was obtained as a colorless solid. Mp: 100–101 °C. $R_{\rm f}$ 0.23 (dichloromethane/methanol 10:1). IR (neat): 3298, 3259, 1574 cm⁻¹; ¹H NMR (CDCl₃): δ 1.33 (d, 6H, J=6.2 Hz), 3.56 (t, 2H, J=4.4 Hz), 3.81 (t, 2H, J=4.4 Hz), 4.55 (br, 1H), 5.28 (sept, 1H, J=6.2 Hz), 5.75 (br, 1H), 5.95 (d, 1H, J=5.6 Hz), 7.94 (d, 1H, J=5.6 Hz); ¹³C NMR (CDCl₃): δ 21.8 (q, 2 CH₃), 44.4, 62.6 (2t, 2 CH₂), 68.5 (d, CH), 98.1, 157.3 (2d, 2 CH_{pyrim}), 169.4, 162.7 (2s, 2 $C_{\rm pyrim}$); MS (EI) *m/z*: 197 ([M]⁺⁺, 13). Anal. Calcd for C₉H₁₅N₃O₂: C, 54.81; H, 7.67; N, 21.30. Found: C, 54.60; H, 7.78; N, 21.11.

4.2.2. 2-(4-Isopropoxy-6-methyl-pyrimidin-2-ylamino)ethanol (9b). From 1.00 g (3.27 mmol) of sulfone **5b** and 0.30 mL (4.85 mmol) of 2-amino-ethanol **8a**, 524 mg (76%) of compound **9b** was obtained as a colorless solid. Mp: 109–110 °C. R_f 0.79 (dichloromethane/methanol 3:1). IR (neat): 3265, 3190, 1568 cm⁻¹; ¹H NMR (CDCl₃): δ 1.33 (d, 6H, J=6.2 Hz), 2.25 (s, 3H), 3.56 (t, 2H, J=4.4 Hz), 3.83 (t, 2H, J=4.4 Hz), 4.65 (br, 1H), 5.25 (sept, 1H, J= 6.2 Hz), 5.45 (br, 1H), 5.86 (s, 1H); ¹³C NMR (CDCl₃): δ 21.9, 23.4 (2q, 3 CH₃), 44.7, 63.5 (2t, 2 CH₂), 68.4 (d, CH), 96.9 (d, CH_{pyrim}), 162.7, 167.4, 170.0 (3s, 3 C_{pyrim}); MS (EI) m/z: 211 ([M]⁺⁺, 14). Anal. Calcd for C₁₀H₁₇N₃O₂: C, 56.85; H, 8.11; N, 19.89. Found: C, 56.96; H, 8.29; N, 19.70.

4.2.3. 3-(4-Isopropoxy-pyrimidin-2-ylamino)-propan-1ol (9c). From 1.03 g (3.5 mmol) of sulfone **5a** and 0.4 mL (5.42 mmol) of 3-amino-propan-1-ol **8b**, 487 mg (65%) of compound **9c** was obtained as a colorless solid. Mp: 82–83 °C. $R_{\rm f}$ 0.33 (dichloromethane/methanol 10:1). IR (neat): 3280, 3219, 1560 cm⁻¹; ¹H NMR (CDCl₃): δ 1.33 (d, 6H, J=6.2 Hz), 1.77 (m, 2H), 3.57 (q, 2H, J=6.0 Hz), 3.67 (t, 2H, J=5.7 Hz), 4.15 (br, 1H), 5.26 (sept, 1H, J= 6.2 Hz), 5.30 (br, 1H), 5.96 (d, 1H, J=5.8 Hz), 7.96 (d, 1H, J=5.8 Hz); ¹³C NMR (CDCl₃): δ 22.5 (q, 2 CH₃), 33.8, 38.4, 59.4 (3t, 3 CH₂), 69.2 (d, CH), 98.8, 158.1 (2d, CH_{pyrim}), 163.6, 170.1 (2s, 2 C_{pyrim}); MS (EI) m/z: 211 ([M]⁺⁺, 9). Anal. Calcd for C₁₀H₁₇N₃O₂: C, 56.85; H, 8.11; N, 19.89. Found: C, 56.74; H, 8.32; N, 19.86.

4.2.4. 3-(**4**-Isopropoxy-6-methyl-pyrimidin-2-ylamino)propan-1-ol (9d). From 1.23 g (4.02 mmol) of sulfone **5b** 0.46 mL (6.24 mmol) of and 3-amino-propan-1-ol **8b**, 847 mg (94%) of compound **9d** was obtained as a colorless oil. $R_{\rm f}$ 0.12 (*n*-hexane/ethyl acetate 1:1). IR (neat): 3305, 3106, 1578 cm⁻¹; ¹H NMR (CDCl₃): δ 1.33 (d, 6H, J= 6.2 Hz), 1.81 (q, 2H, J=5.8 Hz), 2.33 (s, 3H), 3.56 (q, 2H, J=6.0 Hz), 3.70 (t, 2H, J=5.6 Hz), 5.30 (sept, 1H, J= 6.2 Hz), 5.55 (br, 2H), 5.83 (s, 1H); ¹³C NMR (CDCl₃): δ 21.8, 22.6 (2q, 3 CH₃), 32.5, 38.7, 59.9 (3t, 3 CH₂), 71.4 (d, CH), 98.4 (d, CH_{pyrim}), 158.3, 167.0, 171.3 (3s, 3 C_{pyrim}); MS (EI) *m/z*: 225 ([M]⁺⁺, 64). Anal. Calcd for C₁₁H₁₉N₃O₂: C, 58.64; H, 8.50; N, 18.65. Found: C, 58.87; H, 8.62; N, 18.84.

4.2.5. 2-(4-Isopropoxy-6-methyl-pyrimidin-2-ylamino)-3-phenyl-propan-1-ol (9e). From 785 mg (2.56 mmol) of sulfone 5b and 774 mg (5.13 mmol) of phenylalaninol 8c, 612 mg (79%) of compound 9e was obtained as a colorless oil. R_f 0.12 (chloroform/methanol 6:1). IR (neat): 3400, 3028, 1577 cm⁻¹; ¹H NMR (CDCl₃): δ 1.34 (d, 6H, J= 6.2 Hz), 2.97 (d, 2H, J = 7.4 Hz), 3.67 (dd, 1H, J = 10.8 Hz, J' = 5.6 Hz, 3.82 (dd, 1H, J = 10.8 Hz, J' = 2.8 Hz), 4.20 (m, 1H), 4.30 (br, 1H), 5.27 (sept, 1H, J = 6.2 Hz), 5.40 (br, 1H), 5.86 (s, 1H), 7.20–7.40 (m, 5H); ¹³C NMR (CDCl₃): δ 22.6, 23.8 (2q, 3 CH₃), 38.4 (t, CH₂), 55.6 (d, CH), 65.4 (t, CH₂), 69.3 (d, CH), 97.5 (d, CH_{pvrim}), 127.1, 129.2, 129.9 (3d, 5 CH_{arom}), 139.0 (s, C_{arom}), 162.2, 167.3, 170.8 (3s, 3 C_{pyrim} ; MS (EI) m/z: 301 ([M]⁺, 4). Anal. Calcd for C₁₇H₂₃N₃O₂: C, 67.75; H, 7.69; N, 13.94. Found: C, 67.56; H, 7.52; N, 14.12.

4.2.6. 2-[(4-Isopropoxy-6-phenyl-pyrimidin-2-yl)methyl-amino]-ethanol (9f). From 1.00 g (2.72 mmol) of sulfone **5c** and 0.38 mL (5.4 mmol) of 2-methylaminoethanol **8d**, 667 mg (85%) of compound **9f** was obtained as a colorless oil. $R_{\rm f}$ 0.23 (*n*-hexane/ethyl acetate 1:1). IR (neat): 3400–3200, 1534 cm⁻¹; ¹H NMR (CDCl₃): δ 1.42 (d, 6H, J=6.2 Hz), 3.30 (s, 3H), 3.85–4.00 (m, 4H), 5.40 (sept, 1H, J=6.2 Hz), 6.40 (s, 1H), 7.40, 7.50 (m, 3H), 7.95–8.00 (m, 2H); ¹³C NMR (CDCl₃): δ 22.6, 37.3 (2q, 3 CH₃), 53.5, 63.7 (2t, 2 CH₂), 69.3 (d, CH), 93.7 (d, CH_{pyrim}), 127.5, 129.2, 130.8 (3d, 5 CH_{arom}), 138.4 (s, $C_{\rm arom}$), 163.6, 165.7, 170.9 (3s, 3 $C_{\rm pyrim}$); MS (EI) *m/z*: 287 ([M]⁻⁺, 11). Anal. Calcd for C₁₆H₂₁N₃O₂: C, 66.88; H, 7.37; N, 14.62. Found: C, 67.15; H, 7.45; N, 14.35.

4.2.7. 2-[(4-Isopropoxy-6-methyl-pyrimidin-2-yl)methyl-amino]-ethanol (9g). From 1.23 g (4.02 mmol) of sulfone **5b** and 0.48 mL (6.83 mmol) of 2-methylaminoethanol **8d**, 721 mg (80%) of compound **9g** was obtained as a colorless oil. $R_{\rm f}$ 0.38 (*n*-hexane/ethyl acetate 1:1). IR (neat): 3450–3250, 1576 cm⁻¹; ¹H NMR (CDCl₃): δ 1.34 (d, 6H, J=6.2 Hz), 2.24 (s, 3H), 3.20 (s, 3H), 3.74 (t, 2H, J=4.0 Hz), 3.89 (t, 2H, J=4.0 Hz), 5.30 (sept, 1H, J= 6.2 Hz), 5.81 (s, 1H); ¹³C NMR (CDCl₃): δ 21.9, 23.5, 36.6 (3q, 4 CH₃), 53.2, 63.3 (2t, 2 CH₂), 68.2 (d, CH), 95.7 (d, CH_{pyrim}), 162.6, 167.0, 169.6 (3s, 3 C_{pyrim}); MS (EI) *m/z*: 225 ([M]⁺⁺, 29). Anal. Calcd for C₁₁H₁₉N₃O₂: C, 58.64; H, 8.50; N, 18.65. Found: C, 58.66; H, 8.62; N, 18.56.

4.2.8. [1-(4-Isopropoxy-6-methyl-pyrimidin-2-yl)-pyrrolidin-2-yl]-methanol (9h). From 1.00 g (3.27 mmol) of sulfone **5b** and 0.58 mL (5.6 mmol) of prolinol **8e**, 825 mg (95%) of compound **9h** was obtained as a colorless oil. $R_{\rm f}$ 0.34 (*n*-hexane/ethyl acetate 1:1). IR (neat): 3400–3300, 1574 cm⁻¹; ¹H NMR (CDCl₃): δ 1.33 (d, 6H, J=6.2 Hz), 1.70–2.20 (m, 4H), 2.23 (s, 3H), 3.50–3.80 (m, 4H), 4.25

(m, 1H), 5.30 (sept, 1H, J=6.2 Hz), 5.81 (s, 1H); ¹³C NMR (CDCl₃): δ 21.9, 23.3 (2q, 3 CH₃), 23.9, 29.9, 48.2 (3t, 3 CH₂), 61.1 (d, CH), 68.3 (t, CH₂), 68.8 (d, CH), 95.7 (d, CH_{pyrim}), 161.1, 166.5, 169.6 (3s, 3 C_{pyrim}); MS (EI) *m/z*: 251 ([M]⁺, 3). Anal. Calcd for C₁₃H₂₁N₃O₂: C, 62.13; H, 8.42; N, 16.72. Found: C, 62.19; H, 8.64; N, 16.44.

4.2.9. [1-(4-Isopropoxypyrimidin-2-yl)piperidin-2-yl]methanol (9i). From 875 mg (3.0 mmol) of sulfone 5a and 618 mg (5.37 mmol) of piperidin-2-yl-methanol 8f, 596 mg (79%) of compound 9i was obtanied as a colorless oil. R_f 0.40 (*n*-hexane/ethyl acetate 1:1). IR (neat): 3405, 1587 cm⁻¹; ¹H NMR (CDCl₃): δ 1.37 (d, 6H, *J*=6.2 Hz), 1.65–1.80 (m, 6H), 3.05 (m, 1H), 3.40 (br, 1H), 3.73 (dd, 1H, *J*=10.7 Hz, *J'*=5, 5 Hz), 3.96 (dd, 1H, *J*=10.7 Hz, *J'*=8.8 Hz, CH₂N), 4.60 (m, 1H), 4.90 (m, 1H), 5.28 (sept, 1H, *J*=6.2 Hz), 5.90 (d, 1H, *J*=6.0 Hz), 7.98 (d, 1H, *J*= 6.0 Hz); ¹³C NMR (CDCl₃): δ 19.8 (t, CH₂), 21.8 (q, 2 CH₃), 24.9, 25.6, 39.6 (3t, 3 CH₂), 52.7 (d, CH), 62.7 (t, CH₂), 68.2 (d, CH), 97.1, 157.4 (2d, 2 CH_{pyrim}), 162.6, 169.0 (2s, 2 C_{pyrim}); MS (EI) *m/z*: 251 ([M]⁺⁺, 6). Anal. Calcd for C₁₃H₂₁N₃O₂: C, 62.13; H, 8.42; N, 16.72. Found: C, 62.22; H, 8.34; N, 16.54.

4.2.10. [1-(4-Isopropoxy-6-methylpyrimidin-2-yl)-piperidin-2-yl]-methanol (9j). From 900 mg (2.94 mmol) of sulfone **5b** and 672 mg (5.84 mmol) piperidin-2-ylmethanol **8f**, 716 mg (90%) of compound **9j** was obtained as a colorless oil. R_f 0.44 (*n*-hexane/ethyl acetate 1:1). IR (neat): 3400, 1573 cm⁻¹; ¹H NMR (CDCl₃): δ 1.31 (d, 6H, J=6.2 Hz), 1.50–1.70 (m, 6H), 2.20 (s, 3H), 3.05 (m, 1H), 3.71 (dd, 1H, J=10.7 Hz, J'=4.7 Hz, CH_2 N), 4.00 (dd, 1H, J=10.7 Hz, J'=9.0 Hz), 4.15 (br, 1H), 4.60 (m, 1H), 4.90 (m, 1H), 5.25 (sept, 1H, J=6.2 Hz), 5.77 (s, 1H); ¹³C NMR (CDCl₃): δ 20.6 (t, CH_2), 21.5, 24.4 (2q, 3 CH_3), 25.6, 26.5, 40.3 (3t, 3 CH_2), 53.6 (d, CH), 64.1 (t, CH_2), 68.7 (d, CH), 96.3 (d, CH_{pyrim}), 163.4, 167.9, 170.2 (3s, 3 C_{pyrim}); MS (EI) m/z: 265 ([M]⁺⁺, 3). Anal. Calcd for C₁₄H₂₃N₃O₂: C, 63.37; H, 8.74; N, 15.84. Found: C, 63.09; H, 8.83; N, 15.66.

4.3. General procedure for the hydrolysis of compounds 9 with HCl. Synthesis of pyrimidinones 10

A suspension of the corresponding 4-isopropoxypyrimidine **9** (1 equiv) in concd HCl (2 mL/mmol) was heated at 90 °C for 30 min. After cooling, the mixture was neutralized with aq 5 N NaOH and extracted with CH_2Cl_2 (3×10 mL/mmol). The combined organic layers were washed with brine (1×10 mL/mmol) and the organic layer was dried over MgSO₄, filtered and concentrated under reduced pressure. The resulting crude product was purified by flash-chromatography (ethyl acetate/methanol 10:1) to afford pyrimidinones **10**.

4.3.1. 2-(2-Hydroxy-ethylamino)-3H-pyrimidin-4-one (**10a).** From 336 mg (1.7 mmol) of 4-isopropoxipyrimidine **9a**, 226 mg (86%) of compound **10a** was obtained as a colorless solid. Mp: 176–177 °C. $R_{\rm f}$ 0.17 (dichloromethane/methanol 10:1). IR (neat): 3220–2880, 1681, 1621 cm⁻¹; ¹H NMR (DMSO- d_6): δ 3.30 (m, 2H), 3.51 (t, 2H, J= 5.6 Hz), 4.90 (br, 1H), 5.53 (d, 1H, J=6.6 Hz), 6.55 (br, 1H), 7.57 (d, 1H, J=6.6 Hz), 10.60 (br, 1H); ¹³C NMR (DMSO- d_6): δ 43.4, 59.2 (2t, 2 CH₂), 103.2, 149.2 (2d, 2

 CH_{pyrim}), 153.9, 162.0 (2s, 2 C_{pyrim}); MS (ESI) m/z: 178 $[M+23]^+$, 156 $[M+1]^+$. Anal. Calcd for $C_6H_9N_3O_2$: C, 46.45; H, 5.85; N, 27.08. Found: C, 46.43; H, 5.96; N, 27.10.

4.3.2. 2-(2-Hydroxyethylamino)-6-methylpyrimidin-4(3H)-one (10b). From 446 mg (2.11 mmol) of 4-isopropoxipyrimidine **9b**, 346 mg (97%) of compound **10b** was obtained as a colorless solid. Mp: 192–193 °C. R_f 0.51 (dichloromethane/methanol 3:1). IR (neat): 3250, 1612 cm⁻¹; ¹H NMR (DMSO- d_6): δ 2.10 (s, 3H), 3.30 (m, 2H), 3.40 (t, 2H, J=6.0 Hz), 3.59 (t, 2H, J=6.0 Hz), 4.95 (br, 1H), 5.50 (s, 1H), 6.70 (br, 1H), 10.65 (br, 1H); ¹³C NMR (DMSO- d_6): δ 24.4 (q, CH₃), 43.1, 60.0 (2t, 2 CH₂), 100.8 (d, CH_{pyrim}), 154.7, 163.0, 166.0 (3s, 3 C_{pyrim}); MS (ESI) *m*/*z*: 192 [M+23]⁺, 170 [M+1]⁺. Anal. Calcd for C₇H₁₁N₃O₂: C, 49.70; H, 6.55; N, 24.84. Found: C, 49.91; H, 6.68; N, 24.56.

4.3.3. 2-(3-Hydroxy-propylamino)-3*H***-pyrimidin-4-one (10c). From 344 mg (1.63 mmol) of 4-isopropoxipyrimidine 9c**, 268 mg (97%) of compound **10c** was obtained as a colorless solid. Mp: 141–142 °C. R_f 0.20 (dichloromethane/ methanol 10:1). IR (neat): 3210–2873, 1676, 1609 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 1.73 (m, 2H), 3.39 (m, 2H), 3.54 (m, 2H), 4.65 (br, 1H), 5.62 (d, 1H, *J*=6.6 Hz), 6.62 (br, 1H), 7.66 (d, 1H, *J*=6.6 Hz), 10.85 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 31.9, 37.6, 58.3 (3t, 3 CH₂), 102.7, 154.4 (2d, 2 CH_{pyrim}), 155.2 162.9 (2s, 2 C_{pyrim}); MS (ESI) *m/z*: 192 [M+23]⁺, 170 [M+1]⁺. Anal. Calcd for C₇H₁₁N₃O₂: C, 49.70; H, 6.55; N, 24.84. Found: C, 49.57; H, 6.80; N, 25.05.

4.3.4. 2-(3-Hydroxy-propylamino)-6-methyl-3*H***-pyrimidin-4-one (10d).** From 570 mg (2.53 mmol) of 4-isopropoxipyrimidine **9d**, 403 mg (87%) of compound **10d** was obtained as a colorless solid. Mp: 160–161 °C. $R_{\rm f}$ 0.60 (dichloromethane/methanol 3:1). IR (neat): 3215–2890, 1675, 1611 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 1.71 (m, 2H), 2.10 (s, 3H), 3.40 (t, 2H, *J*=6.0 Hz), 3.54 (t, 2H, *J*= 6.0 Hz), 4.70 (br, 1H), 5.47 (s, 1H), 6.90 (br, 1H), 10.75 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 23.9 (q, CH₃), 32.0, 37.3, 58.3 (3t, 3 CH₂), 100.2 (d, CH_{pyrim}), 154.4, 162.7, 165.5 (3s, 3 *C*_{pyrim}); MS (ESI) *m/z*: 206 [M+23]⁺, 184 [M+1]⁺. Anal. Calcd for C₈H₁₃N₃O₂: C, 52.45; H, 7.15; N, 22.94. Found: C, 52.70; H, 7.13; N, 23.11.

4.3.5. 2-(1-Benzyl-2-hydroxy-ethylamino)-6-methyl-3*H***-pyrimidin-4-one (10e).** From 410 mg (1.36 mmol) of 4-isopropoxipyrimidine **9e**, 251 mg (71%) of compound **10e** was obtained as a colorless solid. Mp: 145–146 °C. *R*_f 0.61 (dichloromethane/methanol 6:1). IR (neat): 3215–2890, 1675, 1611 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 2.10 (s, 3H), 2.90 (m, 2H), 3.51 (d, 2H, *J*=4.2 Hz), 4.15 (m, 1H), 5.48 (s, 1H), 6.57 (d, 1H, *J*=7.6 Hz), 7.25–7.40 (m, 5H); ¹³C NMR (DMSO-*d*₆): δ 24.3 (q, CH₃), 37.2 (t, CH₂), 53.8 (d, CH), 61.9 (t, CH₂), 100.8 (d, CH_{pyrim}), 126.5, 128.6, 129.7 (3d, 5 CH_{arom}), 139.3 (s, *C*_{arom}), 154.4, 163.4, 165.8 (3s, 3 *C*_{pyrim}); MS (FAB⁺) *m*/*z*: 260 ([M+1]⁺, 100). Anal. Calcd for C₁₄H₁₇N₃O₂: C, 64.85; H, 6.61; N, 16.21. Found: C, 65.06; H, 6.74; N, 15.93.

4.3.6. 2-(*N*-(**2-Hydroxyethyl**)-*N*-methylamino)-6-phenylpyrimidin-4(3*H*)-one (10f). From 53 mg (0.18 mmol) of 4-isopropoxipyrimidine 9f, 39 mg (95%) of compound 10f was obtained as a colorless solid. Mp: 197–198 °C. $R_{\rm f}$ 0.40 (dichloromethane/methanol 10:1). IR (neat): 3350, 1639 cm⁻¹; ¹H NMR (DMSO- d_6): δ 3.26 (s, 3H), 3.75 (m, 4H), 5.00 (br, 1H), 6.26 (s, 1H), 7.50–7.60 (m, 3H), 8.05–8.10 (m, 2H); ¹³C NMR (DMSO- d_6): δ 36.5 (q, CH₃), 51.8, 58.9 (2t, 2 CH₂), 95.3 (d, CH_{pyrim}), 126.6, 128.4, 130.0 (3d, 5 CH_{arom}), 137.4 (s, $C_{\rm arom}$), 155.5, 161.9, 165.1 (3s, 3 $C_{\rm pyrim}$); MS (FAB⁺) m/z: 246 ([M+1]⁺, 100). Anal. Calcd for C₁₃H₁₅N₃O₂: C, 63.66; H, 6.16; N, 17.13. Found: C, 63.72; H, 6.03; N, 17.30.

4.3.7. 2-[(2-Hydroxy-ethyl)-methyl-amino]-6-methyl-*3H*-pyrimidin-4-one (10g). From 560 mg (2.49 mmol) of 4-isopropoxipyrimidine **9g**, 387 mg (85%) of compound **10g** was obtained as a colorless solid. Mp: 118–119 °C. $R_{\rm f}$ 0.26 (dichloromethane/methanol 10:1). IR (neat): 3363–2851, 1647, 1565 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 2.13 (s, 3H), 3.15 (s, 3H), 3.65 (m, 4H), 4.90 (br, 1H), 5.55 (s, 1H), 10.85 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 23.0, 35.4 (2q, 2 CH₃), 58.0, 50.6, (2t, 2 CH₂), 97.5 (d, CH_{pyrim}), 154.2, 163.6, 164.8 (3s, 3 C_{pyrim}); MS (FAB⁺) *m/z*: 206 [M+ 23]⁺, 184 [M+1]⁺. Anal. Calcd for C₈H₁₃N₃O₂: C, 52.45; H, 7.15; N, 22.94. Found: C, 52.31; H, 7.23; N, 22.81.

4.3.8. 2-(2-(Hydroxymethyl)pyrrolidin-1-yl)-6-methylpyrimidin-4(3*H***)-one (10h). From 630 mg (2.51 mmol) of 4-isopropoxipyrimidine 9h**, 490 mg (94%) of compound **10h** was obtained as a colorless solid. Mp: 110–111 °C. $R_{\rm f}$ 0.73 (dichloromethane/methanol 6:1). IR (neat): 3380, 1686 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 1.95 (m, 4H), 2.38 (s, 3H), 3.40–3.65 (m, 4H), 4.45 (br, 1H), 5.99 (s, 1H); ¹³C NMR (DMSO-*d*₆): δ 19.6, (q, *C*H₃), 23.2, 27.9, 49.8 (3q, 3 *C*H₂), 61.6 (d, *C*H), 61.9 (t, *C*H₂), 100.6 (d, *C*H_{pyrim}), 151.2, 156.4, 165.6 (3s, 3 C_{pyrim}); MS (FAB⁺) *m/z*: 210 [M+1]⁺. Anal. Calcd for C₁₀H₁₅N₃O₂: C, 57.40; H, 7.23; N, 20.08. Found: C, 57.51; H, 7.04; N, 20.11.

4.4. General procedure for the sequential hydrolysiscyclization of compounds 9. Synthesis of compounds 1

A suspension of the corresponding 4-isopropoxypyrimidine **9** (1 equiv) in concd H_2SO_4 (3 mL/mmol) was heated at 90 °C for 20 min–1 h until the reaction was completed (TLC monitoring). After cooling, the mixture was neutralized with aq 5 N NaOH and extracted with CH_2Cl_2 (3×10 mL/mmol). The combined organic layers were washed with brine (1×10 mL/mmol) and the organic layer was dried over MgSO₄, filtered and evaporated under reduced pressure to afford imidazo[1,2-*a*]pyrimidinones **1** (*n*=1) or pyrimido[1,2-*a*]pyrimidinones **1** (*n*=2).

4.4.1. 5,6,7,8,8a,9-Hexahydro-4,4b,9a-triaza-fluoren-1one (**1i**). From 123 mg (0.49 mmol) of 4-isopropoxipyrimidine **9i**, 92 mg (98%) of compound **1i** was obtained as a colorless solid. Mp: 93–94 °C. $R_{\rm f}$ 0.38 (dichloromethane/ methanol 10:1). IR (neat): 1659, 1579, 1545 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 1.40–1.90 (m, 6H), 3.00 (m, 1H), 3.60–4.25 (m, 4H), 5.68 (d, 1H, *J*=6.4 Hz), 7.66 (d, 1H, *J*= 6.4 Hz); ¹³C NMR (DMSO-*d*₆): δ 22.5, 23.9, 41.0, 46.4 (5t, 5 *C*H₂), 54.6 (d, *C*H), 102.9, 155.3 (2d, 2 *C*H_{pyrim}), 155.6, 160.5 (2s, 2 *C*_{pyrim}); MS (ESI) *m/z*: 192 [M+1]⁺. Anal. Calcd for C₁₀H₁₃N₃O: C, 62.81; H, 6.85; N, 21.97. Found: C, 62.92; H, 6.66; N, 22.00. **4.4.2. 3-Methyl-5,6,7,8,8a,9-hexahydro-4,4b,9a-triaza-fluoren-1-one** (**1j**). From 103 mg (0.39 mmol) of 4-iso-propoxipyrimidine 9j, 79 mg (99%) of compound **1j** was obtained as a colorless solid. Mp: 121–123 °C. $R_{\rm f}$ 0.34 (dichloromethane/methanol 10:1). IR (neat): 1657, 1584, 1553 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 1.40–1.95 (m, 6H), 2.13 (s, 3H), 2.95 (m, 1H), 3.55–4.20 (m, 4H), 5.58 (s, 1H); ¹³C NMR (DMSO-*d*₆): δ 22.6 (t, CH₂), 23.7 (q, CH₃), 23.9, 30.1, 41.0, 46.3 (4t, 4 CH₂), 54.8 (d, CH), 100.6 (d, CH_{pyrim}), 154.7, 160.7, 165.3 (3s, 3 C_{pyrim}); MS (ESI) *m/z*: 206 [M+1]⁺. Anal. Calcd for C₁₁H₁₅N₃O: C, 64.37; H, 7.37; N, 20.47. Found: C, 64.15; H, 7.46; N, 20.32.

4.4.3. 1-Methyl-7-phenyl-2,3-dihydro-1*H***-imidazo**[**1,2-***a*]**pyrimidin-5-one** (**1f**). From 527 mg (1.84 mmol) of 4-isopropoxipyrimidine **9f**, 394 mg (95%) of compound **1f** was obtained as a colorless solid. Mp: 130–131 °C. $R_{\rm f}$ 0.54 (dichloromethane/methanol 10:1). IR (neat): 1665, 1590, 1553 cm⁻¹; ¹H NMR (CDCl₃): δ 3.12 (s, 3H), 3.69 (t, 2H, J=8.8 Hz), 4.16 (t, 2H, J=8.7 Hz), 6.29 (s, 1H), 7.40–7.50 (m, 3H), 7.95–8.00 (m, 2H); ¹³C NMR (CDCl₃): δ 32.3 (q, CH₃), 41.0, 47.8 (2t, 2 CH₂), 99.8 (d, CH_{pyrim}), 127.8, 129.1, 130.7 (3d, 5 CH_{arom}), 138.1 (s, $C_{\rm arom}$), 157.3, 163.2, 164.1 (3s, 3 $C_{\rm pyrim}$); MS (ESI) *m/z*: 228 [M+1]⁺. Anal. Calcd for C₁₃H₁₃N₃O: C, 68.70; H, 5.77; N, 18.49. Found: C, 68.49; H, 5.93; N, 18.54.

4.4.4. 7-Methyl-2,3-dihydro-1*H***-imidazo**[1,2-a]**pyrimidin-5-one** (**1b**). From 50 mg (0.24 mmol) of 4-isopropoxipyrimidine **9b**, 34 mg (95%) of compound **1b** was obtained as a colorless solid. Mp: 233–234 °C. $R_{\rm f}$ 0.54 (dichloromethane/methanol 6:1). IR (neat): 1658, 1617, 1567 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 2.09 (s, 3H), 3.69 (t, 2H, *J*=8.8 Hz), 4.05 (t, 2H, *J*=8.8 Hz), 5.53 (s, 1H), 7.90 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 42.9, 62.1 (2t, 2 *C*H₂), 105.4 (d, *C*H_{pyrim}), 150.1, 160.0, 164.8 (3s, 3 *C*_{pyrim}); MS (ESI) *m/z*: 152 [M+1]⁺. Anal. Calcd for C₇H₉N₃O: C, 55.62; H, 6.00; N, 27.80. Found: C, 55.70; H, 5.77; N, 27.82.

4.4.5. 2-Methyl-6,7,8,9-tetrahydro-pyrimido[**1,2-***a*]**pyrimidin-4-one** (**1d**). From 420 mg (1.87 mmol) of 4-isopropoxipyrimidine **9d**, 223 mg (73%) of compound **1d** was obtained as a colorless solid. Mp: 204–205 °C. $R_{\rm f}$ 0.71 (dichloromethane/methanol 6:1). IR (neat): 3262, 1661, 1600, 1580 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 1.98 (t, 2H, *J*= 5.6 Hz), 2.06 (s, 3H), 3.34 (t, 2H, *J*=5.4 Hz), 3.86 (t, 2H, *J*=5.7 Hz), 5.51 (s, 1H), 7.90 (br, 1H); ¹³C NMR (DMSO*d*₆): δ 19.5 (t, *C*H₂), 23.4 (q, *C*H₃), 38.3, 38.4 (2t, 2 *C*H₂), 98.0 (d, *C*H_{pyrim}), 153.0, 161.5, 164.0 (3s, 3 *C*_{pyrim}); MS (ESI) *m/z*: 166 [M+1]⁺. Anal. Calcd for C₈H₁₁N₃O: C, 58.17; H, 6.71; N, 25.44. Found: C, 58.34; H, 6.60; N, 25.66.

4.4.6. 1,7-Dimethyl-2,3-dihydro-1*H***-imidazo**[**1,2**-*a*]**pyri-midin-5-one** (**1g**). From 55 mg (2.44 mmol) of 4-isopropoxipyrimidine **9g**, 39 mg (99%) of compound **1g** was obtained as a colorless solid. Mp: 132–133 °C. $R_{\rm f}$ 0.40 (dichloromethane/methanol 10:1). IR (neat): 1661, 1591, 1567 cm⁻¹; ¹H NMR (CDCl₃): δ 2.13 (s, 3H), 2.97 (s, 3H), 3.64 (t, 2H, *J*=8.9 Hz), 4.02 (t, 2H, *J*=8.9 Hz), 5.60 (s, 1H); ¹³C NMR (CDCl₃): δ 24.7, 32.2 (2q, 2 CH₃), 40.8, 47.7 (2t, 2 CH₂), 102.3 (d, CH_{pyrim}), 157.0, 162.5, 168.8 (3s, 3 $C_{\rm pyrim}$); MS (ESI) *m/z*: 188 [M+23]⁺, 166 [M+1]⁺. Anal. Calcd for C₈H₁₁N₃O: C, 58.17; H, 6.71; N, 25.44. Found: C, 58.27; H, 6.71; N, 25.23.

4.4.7. 5-Methyl-2,3,8,8a-tetrahydro-1*H*-3a,4,7a-triazacyclopenta[*a*]inden-7-one (1h). From 300 mg (1.19 mmol) of 4-isopropoxipyrimidine 9h, 174 mg (76%) of compound 1h was obtained as a colorless solid. Mp: 77–78 °C. R_f 0.35 (dichloromethane/methanol 10:1). IR (neat): 1661, 1582, 1536 cm⁻¹; ¹H NMR (CDCl₃): δ 1.30-1.50 (m, 1H), 1.85-2.10 (m, 3H), 2.17 (s, 3H), 3.30-3.40 (m, 1H), 3.65-3.75 (m, 1H), 3.90-4.15 (m, 3H), 5.74 (s, 1H); ¹³C NMR (CDCl₃): δ 24.0 (q, CH₃), 25.0, 30.9, 45.6, 47.1 (4t, 4 CH₂), 59.1 (d, CH), 103.4 (d, CH_{pyrim}), 159.0, 161.8, 166.1 (3s, 3 C_{pyrim}); MS (ESI) m/z: $192 [M+1]^+$. Anal. Calcd for $C_{10}H_{13}N_3O$: C, 62.81; H, 6.85; N, 21.97. Found: C, 63.08; H, 6.96; N, 22.01.

4.5. General procedure for the intramolecular Mitsunobu cyclization. Synthesis of compounds 1 and 2

Under nitrogen atmosphere, a solution of DEAD (1.1 equiv) in dry THF (5 mL/mmol) was added dropwise to a solution of Ph₃P (1.1 equiv) and the appropriate pyrimidinone **10** (1 equiv) in dry THF (10 mL/mmol) at 0 °C. The resulting mixture was warmed to room temperature and stirred for 5–8 h until the reaction was completed (TLC monitoring). The solvent was removed under reduced pressure and the crude product was purified by flash chromatography (ethyl acetate/methanol 10:1) to afford compounds **1** and **2**.

4.5.1. Intramolecular cyclization of pyrimdinone 10a. From 197 mg (1.27 mmol) of pyrimidinone **10a**, 19 mg (11%) of compound **1a** and 140 mg (80%) of compound **2a** were obtained.

4.5.1.1. 2,3-Dihydro-1*H***-imidazo**[**1,2***-a*]**pyrimidin-5-one** (**1a**). Isolated as a colorless solid. Mp: 151–153 °C. $R_{\rm f}$ 0.54 (dichloromethane/methanol 6:1). IR (neat): 3216, 1661, 1605, 1530 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 3.80 (t, 2H, *J*=8.9 Hz), 4.27 (t, 2H, *J*=8.8 Hz), 5.81 (d, 1H, *J*= 6.8 Hz), 7.10 (br, 1H), 7.55 (d, 1H, *J*=6.6 Hz); ¹³C NMR (DMSO-*d*₆): δ 42.8, 59.5 (2t, 2 *C*H₂), 102.7, 155.3 (2d, 2 *C*H_{pyrim}), 155.7, 162.6 (2s, 2 *C*_{pyrim}); MS (ESI) *m/z*: 138 [M+1]⁺. Anal. Calcd for C₆H₇N₃O: C, 52.55; H, 5.14; N, 30.64. Found: C, 52.36; H, 5.23; N, 30.71.

4.5.1.2. 2,3-Dihydro-1*H***-imidazo**[**1,2***-a*]**pyrimidin-7-one** (2a). Isolated as a colorless solid. Mp: 195–196 °C. $R_{\rm f}$ 0.14 (dichloromethane/methanol 6:1). IR (neat): 3080, 1649, 1604 cm⁻¹; ¹H NMR (DMSO-*d*₆): δ 3.67 (t, 2H, *J*=8.5 Hz), 4.14 (t, 2H, *J*=8.6 Hz), 5.57 (d, 1H, *J*=7.4 Hz), 7.56 (d, 1H, *J*=7.4 Hz), 7.75 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 40.2, 46.3 (2t, 2 *C*H₂), 105.1, 138.3 (2d, 2 *C*H_{pyrim}), 159.2, 171.2 (2s, 2 *C*_{pyrim}); MS (ESI) *m/z*: 160 [M+23]⁺, 138 [M+1]⁺. Anal. Calcd for C₆H₇N₃O: C, 52.55; H, 5.14; N, 30.64. Found: C, 52.67; H, 5.22; N, 30.76.

4.5.2. Intramolecular cyclization of pyrimdinone 10e. From 130 mg (0.50 mmol) of pyrimidinone **10e**, 39 mg (33%) of compound **1e** and 54 mg (45%) of compound **2e** were obtained.

4.5.2.1. 2-Benzyl-7-methyl-2,3-dihydro-1*H*-imidazo-[1,2-*a*]pyrimidin-5-one (1e). Isolated as a colorless solid. Mp: 182–183 °C. $R_{\rm f}$ 0.51 (dichloromethane/methanol 3:1). IR (neat): 3128, 1659, 1564 cm⁻¹; ¹H NMR (CDCl₃): δ 2.10 (s, 3H), 2.90 (m, 2H), 3.50 (br, 2H), 4.15 (m, 1H), 5.48 (s, 1H), 7.30–7.40 (m, 5H), 7.95 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 19.2 (q, CH₃), 45.6, 49.0 (2t, 2 CH₂), 50.9, (d, CH), 99.1 (d, CH_{pyrim}), 126.9, 128.6, 129.5 (3d, 5 CH_{arom}), 138.1 (s, C_{arom}), 155.4, 160.1, 163.2 (3s, 3 C_{pyrim}); MS (FAB⁺) *m/z*: 242 ([M+1]⁺, 100). Anal. Calcd for C₁₄H₁₅N₃O: C, 69.69; H, 6.27; N, 17.41. Found: C, 69.82; H, 6.12; N, 17.63.

4.5.2.2. 2-Benzyl-5-methyl-2,3-dihydro-1*H***-imidazo[1,2-***a***]pyrimidin-7-one** (**2e**). Isolated as a colorless solid. Mp: 227–228 °C. $R_{\rm f}$ 0.12 (dichloromethane/methanol 3:1). IR (neat): 3104, 1673, 1620, 1552 cm⁻¹; ¹H NMR (CDCl₃): δ 2.03 (s, 3H), 2.97 (dd, 1H, *J*=13.8 Hz, *J'*= 8.2 Hz), 3.35 (dd, 1H, *J*=13.7 Hz, *J'*=4.4 Hz), 3.71 (dd, 1H, *J*=10.1 Hz, *J'*=6.4 Hz), 3.99 (dd, 1H, *J*=10.1 Hz, *J'*=9.4 Hz), 4.50 (m, 1H), 5.51 (s, 1H), 7.25–7.35 (m, 5H), 9.50 (br, 1H); ¹³C NMR (DMSO-*d*₆): δ 17.4 (q, CH₃), 41.0, 48.9 (2t, 2 CH₂), 53.0, (d, CH), 103.7 (d, CH_{pyrim}), 127.1, 128.9, 129.8 (3d, 5 CH_{arom}), 137.0 (s, *C*_{arom}), 148.1, 158.9, 171.9 (3s, 3 *C*_{pyrim}); MS (FAB⁺) *m/z*: 242 ([M+1]⁺, 100). Anal. Calcd for C₁₄H₁₅N₃O: C, 69.69; H, 6.27; N, 17.41. Found: C, 69.84; H, 6.24; N, 17.54.

4.5.3. Intramolecular cyclization of pyrimdinone 10d. From 350 mg (1.91 mmol) of pyrimidinone **10d**, 73 mg (25%) of **1d** and 187 mg (66%) of **2d** were obtained. The spectroscopic features of **1d** was identical to those reported above.

4.5.3.1. Spectroscopic data for 4-methyl-6,7,8,9-tetrahydro-pyrimido[1,2-*a*]pyrimidin-2-one (2d). Isolated as a colorless solid. Mp: > 300 °C. $R_{\rm f}$ 0.10 (dichloromethane/ methanol 10:1). IR (neat): 3099, 1668, 1618, 1555 cm⁻¹; ¹H NMR (DMSO- d_6): δ 2.05 (m, 2H), 2.22 (s, 3H), 3.32 (t, 2H, J=5.6 Hz), 3.88 (t, 2H, J=5.8 Hz), 5.55 (s, 1H), 7.90 (br, 1H); ¹³C NMR (DMSO- d_6): δ 18.1 (q, CH₃), 20.3, 38.0, 43.0 (t, CH₂), 106.2 (d, CH_{pyrim}), 149.5, 153.5, 169.2 (s, $C_{\rm pyrim}$); MS (ESI) *m/z*: 166 [M+1]⁺. Anal. Calcd for C₈H₁₁N₃O: C, 52.55; H, 5.14; N, 30.64. Found: C, 52.74; H, 5.35; N, 30.41.

4.5.4. Intramolecular cyclization of pyrimdinone 10c. From 182 mg (1.08 mmol) of pyrimidinone **10c**, 36 mg (22%) of **1c** and 119 mg (73%) of **2c** were obtained.

4.5.4.1. 6,7,8,9-Tetrahydro-pyrimido[**1,2**-*a*]**pyrimidin-4-one** (**1c**). Isolated as a colorless solid. Mp: 178–179 °C. $R_{\rm f}$ 0.60 (dichloromethane/methanol 3:1). IR (neat): 3219, 1660, 1603, 1555 cm⁻¹; ¹H NMR (DMSO- d_6): δ 1.99 (m, 2H), 3.36 (t, 2H, J= 5.7 Hz), 3.90 (t, 2H, J= 5.9 Hz), 5.62 (d, 1H, J=6.2 Hz), 7.59 (d, 1H, J=6.2 Hz), 7.95 (br, 1H, NH); ¹³C NMR (DMSO- d_6): δ 19.3, 38.3, 38.6 (3t, 3 CH₂), 99.9 (d, CH_{pyrim}), 154.1 (s, C_{pyrim}), 154.6 (d, CH_{pyrim}), 161.3 (s, C_{pyrim}); MS (ESI) m/z: 152 [M+1]⁺. Anal. Calcd for C₇H₉N₃O: C, 55.62; H, 6.00; N, 27.80. Found: C, 55.49; H, 6.22; N, 27.86.

4.5.4.2. 6,7,8,9-Tetrahydro-pyrimido[**1,2**-*a*]**pyrimidin-2-one** (**2c**). Isolated as a colorless solid. Mp:

235–236 °C. $R_{\rm f}$ 0.22 (dichloromethane/methanol 3:1). IR (neat): 3170, 1673, 1621, 1553 cm⁻¹; ¹H NMR (DMSOd₆): δ 2.01 (m, 2H), 3.34 (t, 2H, J=5.3 Hz), 3.88 (t, 2H, J= 5.3 Hz), 5.60 (d, 1H, J=7.4 Hz), 7.36 (d, 1H, J=7.4 Hz), 8.10 (br, 1H); ¹³C NMR (DMSO-d₆): δ 20.0, 37.8, 47.5 (3t, 3 CH₂), 106.2, 142.3 (2d, 2 CH_{pyrim}), 152.8, 169.8 (2s, 2 C_{pyrim}); MS (ESI) m/z: 174 [M+23]⁺, 152 [M+1]⁺. Anal. Calcd for C₇H₉N₃O: C, 55.62; H, 6.00; N, 27.80. Found: C, 55.45; H, 6.03; N, 27.75.

4.5.5. Intramolecular cyclization of pyrimdinone 10g. From 230 mg (1.26 mmol) of pyrimidinone **10g**, 197 mg (96%) of **1g** was obtained. The spectroscopic features of this product were identical to those reported above.

4.6. X-ray crystallographic details

4.6.1. Compound 1h. $C_{10}H_{13}N_3O \cdot 3H_2O$, $M_r = 245.28$, trigonal, space group $P3_1$, a=10.185(3) Å, c=10.217(6) Å, V=917.9(7) Å³, Z=3, $D_x=1.331$ g cm⁻³, T = -173 °C, crystal dimensions: $0.01 \times 0.05 \times 0.30$ mm, BRUKER SMART APEX-CCD area-detector diffractometer, Mo K α radiation, $\lambda = 0.71073$ Å, $\mu =$ 0.103 mm^{-1} , $\theta_{\text{max}} = 28^{\circ}$, 13947 measured reflections, 1489 symmetry-independent reflections, 1459 reflections with $I > 2\sigma(I)$, refinement on F^2 with SHELXL-97,³⁵ 173 parameters, 1 restraint, R(F) [$I > 2\sigma(I)$ reflections]=0.043, $wR(F^2)$ [all reflections]=0.101, $S(F^2)=1.144$, $\Delta \rho_{max}=$ 0.40 e \AA^{-3} . The asymmetric unit contains one molecule of 1h plus three molecules of water. The enantiomer used in the refinement model was chosen arbitrarily.

4.6.2. Compound 2e. $2C_{14}H_{15}N_3O \cdot 3H_2O$, $M_r = 536.63$, monoclinic, space group $P2_1$, a = 11.6063(2) Å, b = 10.0751(2) Å, c = 12.4411(2) Å, $\beta = 108.5550(7)^\circ$, V = 1319.17(4) Å³, Z = 2, $D_x = 1.292$ g cm⁻³, T = -113 °C, crystal dimensions: $0.15 \times 0.25 \times 0.27$ mm, Nonius KappaCCD area-detector diffractometer, Mo K α radiation, $\lambda = 0.71073$ Å, $\mu = 0.0905$ mm⁻¹, $\theta_{max} = 30^\circ$, 36998 measured reflections, 4254 symmetry-independent reflections, 3502 reflections with $I > 2\sigma(I)$, refinement on F^2 with SHELXL-97, 387 parameters, 1 restraint, R(F) [$I > 2\sigma(I)$ reflections] = 0.038, $wR(F^2)$ [all reflections] = 0.092, $S(F^2) = 1.037$, $\Delta \rho_{max} = 0.15$ e Å⁻³. The asymmetric unit contains two molecules of **2e** plus three molecules of water. The chosen enantiomer was based on the assumption that the chiral centre in the molecule has the *S*-configuration as a result of the known configuration of the reagents used in the reaction.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11. 014. CCDC-276097 and CCDC-287052 contain the supplementary crystallographic data for compounds **1h** and **2e**, respectively. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

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New fluorinated 1,3-vinylogous amidines as versatile intermediates: synthesis of fluorinated pyrimidin-2(1*H*)-ones

Santos Fustero,^{a,b,*} Julio Piera,^{a,†} Juan F. Sanz-Cervera,^{a,b,*} Raquel Román,^a Benjamin H. Brodsky,^{a,‡} María Sánchez-Roselló,^a José Luis Aceña^b and Carmen Ramírez de Arellano^{a,§}

^aDepartamento de Química Orgánica, Universidad de Valencia, E-46100 Burjassot, Spain ^bLaboratorio de Moléculas Orgánicas, Centro de Investigación Príncipe Felipe, E-46013 Valencia, Spain

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Abstract—The condensation of the azaenolates derived from readily available ketimines with fluorinated nitriles offers an efficient and straightforward entry to new fluorinated 1,3-vinylogous amidines. These versatile compounds, in turn, react with triphosgene to yield new fluorinated pyrimidin-2(1H)-ones in high yields.

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1. Introduction

The introduction of fluorine atoms into organic molecules leads to important changes in the biological activities of the latter.¹ Particularly appealing as synthetic targets are fluorine-containing pyrimidine derivatives, which have been shown to have a wide variety of biological effects. For example, these derivatives are currently being used as insecticides (e.g., Flufenerim),^{2a} herbicides (e.g., Prim-sulfuron-methyl),^{2b} fungicides (e.g., Diflumetorim),^{2c} and plant growth regulators in crop protection and optimization (e.g., Flurprimidol).^{2d,3} They also have numerous important applications as pharmaceuticals,^{2e} with one example being 5-fluorouracil, 5-(trifluoromethyl)uracil and their analogs, which have been shown to display potent antitumoral activities.^{2f,g} Of the various fluorinated pyrimidine derivatives, however, pyrimidin-2(1H)-ones have perhaps received the greatest amount of attention in the past few years; indeed, several have been patented as

suitable treatments for neurological,^{4a} immunological,^{4b} and viral^{4c} diseases, as well as for cancer,^{4a} CNS,^{4d} metabolic^{4e} disorders, asthma,^{4f} and even as herbicides.^{4g}

Among several possible precursors of pyrimidine derivatives, 1,3-vinylogous amidines are particularly interesting, as they can be used for the preparation of both acyclic and heterocyclic compounds with potential biological activity.⁵ The two main methods for preparing these compounds include either the condensation of amines with 1,3-dicarbonylic compounds,⁶ or the addition of azaenolates to either imidoyl halides,⁷ nitriles,⁸ or, as was very recently shown, imidoyl alkyl thioethers.⁹ However, up until now these methodologies have not been useful for producing the fluorinated counterparts of 1,3-vinylogous amidines. Thus, Butler et al. found that while the condensation of 5-aminolevulinic acid with 1.3-diketones gives pyrroles in non-fluorinated systems, with fluorinated 1,3-diketones the condensation either stops at the intermediate enaminoketone stage when there is only a CF₃ group on the 1,3-diketone, or does not proceed at all when the diketone is fluorinated at both C_{α} positions.¹⁰ Soloshonok et al. experienced similar problems in their attempts to condense fluorinated 1,3-diketones with benzylamine.¹¹ As for the second approach, our group has successfully managed to carry out the condensation of azaenolates with fluorinated imidoyl chlorides, a method that is, to the best of our knowledge, the only available strategy for the preparation of fluorinated N,N'-disubstituted vinylogous amidines.¹² This method allows the use of both cyclic and acyclic ketimines as starting

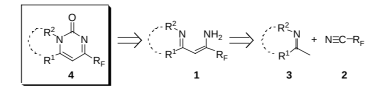
Keywords: Fluorinated compounds; 1,3-Vinylogous amidines; Pyrimidin-2(1*H*)-ones; Imines; Fluorinated nitriles.

^{*} Corresponding authors. Address: Departamento de Química Orgánica, Universidad de Valencia, E-46100 Burjassot, Spain Tel.: +34 963544279; fax: +34 963544938 (S.F.); tel.: +34 963543041; fax: +34 963544228 (J.F.S.-C.); e-mail addresses: santos.fustero@uv.es; juan.f.sanz@uv.es

⁺ Current address: Stockholms universitet, Organisk kemi, Arrheniuslaboratoriet, 106 91 Stockholm, Sweden.

[‡] Current address: Department of Chemistry, Stanford University, Stanford, CA 94305-5080, USA.

[§] Corresponding author for X-ray analysis.



Scheme 1. Retrosynthetic analysis for compounds 1 and 4.

material, and either aliphatic or aromatic substituents on the N atoms. Finally, Barluenga et al. have used the addition of azaenolates to non-fluorinated nitriles to furnish 1,3-vinylogous amidines,⁵ which have subsequently been used in the preparation of acyclic as well as heterocyclic compounds.¹³ Regarding the preparation of pyrimidin-2(1H)-ones, two main strategies have been used thus far: (i) the reaction between 1,3-dicarbonylic compounds or suitable derivatives with a urea,¹⁴ and (ii) the condensation of 1,3-diimines with a carbonic acid derivative.^{13,15,16}

We have now been able to devise an efficient synthesis of new fluorinated 1,3-vinylogous amidines 1 by reacting fluorinated nitriles 2 with ketimines 3 to furnish the corresponding compounds 1 (Scheme 1, retrosynthetic analysis). As an example of the usefulness of these compounds as synthetic intermediates, they were reacted with a suitable carbonic acid derivative to furnish new C4-fluoroalkylated N1,C6-disubstituted pyrimidin-2(1*H*)-ones 4.

2. Results and discussion

In our synthesis, one aliphatic (2c) and two aromatic (2a, 2b) fluorinated nitriles were used as starting materials.

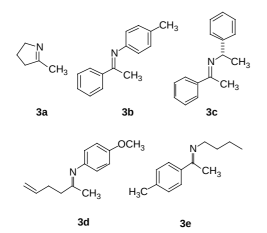


Figure 1. Imines 3 used in the condensation reactions with fluorinated nitriles 2.

While perfluorooctanenitrile (2c) is commercially available, 2a (2,2-difluoro-2-phenylacetonitrile)^{17,18} and 2b (2,2-difluoro-2- α -naphtylacetonitrile)¹⁸ were prepared with slightly modified¹⁹ procedures previously described in the literature.

We chose five representative *N*-substituted imines for our synthetic study: 2-methyl-1-pyrroline (**3a**), which is commercially available, and four acyclic imines derived from acetophenone (**3b** and **3c**), 5-hexen-2-one (**3d**), and *p*-methylacetophenone (**3e**), respectively. Compounds **3a** and **3d** have two different enolizable positions, while **3c** bears a chiral substituent on the N atom (Fig. 1).²⁰

Thus, imines **3** were treated with 1.2 equiv of LDA in THF at -78 °C for 1 h in order to generate their aza-enolates. The temperature was then lowered to -90 °C and a solution of the fluorinated nitrile **2** (1.0 equiv) in THF was added slowly. The reaction was monitored by means of TLC (up to 2 h) and after quenching with aqueous NH₄Cl solution and standard work-up, the resulting crude reaction product was purified through column chromatography on deactivated silica gel (2% Et₃N in hexane) to afford pure fluorinated 1,3-vinylogous amidines **1a–h** in good yields (69–87%, Table 1). The results show that this condensation can be applied to any combination of fluorinated nitrile **2** with an imine **3**, thus allowing for the easy preparation of a variety of fluorinated 1,3-vinylogous amidines **1** (Scheme 2 and Table 1).

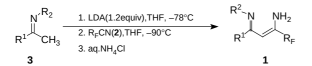
Although in principle compounds **1** might appear in three tautomeric forms (1α - γ in Scheme 3), only the corresponding β -imino enaminic tautomer 1β was present, as confirmed by the ¹H NMR spectra of compounds **1** in CDCl₃ at 300 MHz, which showed the presence of a single tautomer in all cases.¹²

While all attempts to prepare suitable monocrystals of compounds 1a-h failed, it was possible to prepare a complex of compound 1a and ZnI_2 in CH_2Cl_2 , which, in turn, did allow the preparation of suitable monocrystals for X-ray diffraction analysis. The X-ray diffraction structure for the complex $1a \cdot ZnI_2$ clearly shows a 1,3-dimine

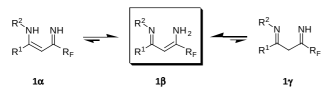
Table 1. Results for the preparation of the fluorinated 1,3-vinylogous amidines 1 (Scheme 2)

Entry	R_F	\mathbb{R}^1		\mathbb{R}^2	1	Yield (%) ^a
1	CF ₂ C ₆ H ₅		-CH2CH2CH2-		1a	87
2	$CF_2C_6H_5$	C ₆ H ₅		$(S)-(+)-C_{6}H_{5}(Me)CH$	1b	81
3	$CF_2C_6H_5$	C_6H_5		p-MeC ₆ H ₄	1c	83
4	$CF_2C_6H_5$	CH2=CH2CH2CH2CH2	,	p-MeOC ₆ H ₄	1d	70
5	$CF_2(\alpha - C_{10}H_7)$		-CH ₂ CH ₂ CH ₂ -		1e	75
6	$CF_3(CF_2)_6$		-CH ₂ CH ₂ CH ₂ -		1f	71
7	$CF_3(CF_2)_6$	C ₆ H ₅		p-MeC ₆ H ₄	1g	79
8	$CF_2C_6H_5$	p-MeC ₆ H ₄		CH ₃ CH ₂ CH ₂ CH ₂ CH ₂	1h	69

^a Yield for purified product.



Scheme 2. The synthesis of 1,3-vinylogous amidines 1 from imines 3 and fluorinated nitriles 2.



Scheme 3. The tautomeric equilibrium in compounds 1.

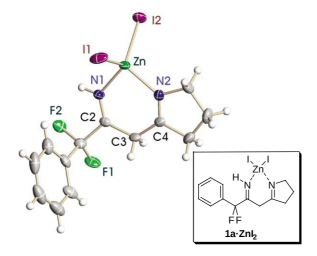


Figure 2. X-ray Structure of complex 1a · ZnI₂. Arbitrary numbering is used in the ORTEP diagram.

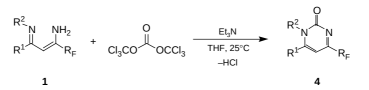
structure that coordinates to the Zn atom with both N atoms (Fig. 2).^{21,22} Thus, the formation of the Zn complex favors the presence of the diiminic form, in contrast with the situation observed in solution (see above). In the crystal

structure, compound **1a** is *N*,*N*-chelated to the Zn atom, which is also coordinated to two iodine atoms. The geometry around the Zn atom is tetrahedral. The *N*,*N*-**1a** chelated ligand forms a quasiplanar six-member metallacycle (mean deviation 0.030 Å), with the metallacycle and the dihydropirrole ring being coplanar [N(1)–Zn–N(2)–C(4)–1.3(5) Å]. In the crystal, intermolecular N–H–I hydrogen bonds form infinite zigzag chains.²¹

The synthetic usefulness of amidines **1** was proven by their transformation into the newly described fluorinated pyrimidin-2(1*H*)-ones **4** through reaction with a suitable carbonic acid derivative. We chose triphosgene [bis(trichloromethyl)carbonate (BTC)]²³ for this purpose because of its ease of handling and high reactivity towards N,N'-binucleophilic compounds. Thus, a solution of triphosgene (1.0 equiv) in THF was added to a solution of compound **1** (1.0 equiv) and Et₃N (2.0 equiv) in THF at room temperature. The reaction mixture was stirred until the starting material was no longer present (0.5–3 h, TLC analysis). Standard work-up furnished crude derivatives **4**, which were then purified by means of flash chromatography to afford fluorinated pyrimidin-2(1*H*)-ones **4a**–**h** in yields that ranged from 70 to 94% (Scheme 4 and Table 2).

3. Conclusion

In conclusion, fluorinated nitriles have once again proven to be versatile starting materials for the preparation of fluorinated heterocycles. In this case, the condensation of the azaenolates derived from readily available ketimines with fluorinated nitriles affords fluorinated 1,3-vinylogous amidines 1 in good yields. These compounds, in turn, easily react with triphosgene to yield fluorinated pyrimidin-2(1*H*)-ones 4 in high yields. Further studies on the reactivity of fluorinated derivatives 1 are currently under way in our laboratories and will be published in due course.



Scheme 4. The reaction of compounds 1 with triphosgene affords compounds 4.

Table 2. Pyrimidin-2(1H)-ones 1 synthesized from derivatives 4 and triphosgene (Scheme 4)

Entry	1	\mathbb{R}^1	R^2	R _F	4	Yield (%) ^a
1	1a	-CH ₂ Cl	H2CH2-	CF ₂ C ₆ H ₅	4 a	94
2	1b	C ₆ H ₅	(S)-(+)-C ₆ H ₅ (Me)CH	$CF_2C_6H_5$	4b	90
3	1c	C ₆ H ₅	p-MeC ₆ H ₄	CF ₂ C ₆ H ₅	4c	87
ł	1d	CH ₂ CH ₂ CH=CH ₂	p-MeOC ₆ H ₄	CF ₂ C ₆ H ₅	4d	84
í	1e	-CH ₂ Cl	H ₂ CH ₂ -	$CF_2(\alpha - C_{10}H_7)$	4 e	77
5	1f	-CH ₂ Cl	H ₂ CH ₂ -	$CF_3(CF_2)_6$	4f	93
,	1g	C ₆ H ₅	$p-MeC_6H_4$	$CF_3(CF_2)_6$	4g	90
3	1h	$p-MeC_6H_4$	CH ₃ CH ₂ CH ₂ CH ₂	CF ₂ C ₆ H ₅	4h	70

^a Yield for purified product.

4. Experimental

4.1. General

All reactions were performed with magnetic stirring in flamedried glassware under an argon atmosphere using dry, distilled solvents. Tetrahydrofuran (THF) was distilled over Na–K alloy while dichloromethane (CH₂Cl₂) was distilled over CaH₂. All other commercially obtained reagents were used as received. All reactions were monitored with thinlayer chromatography (TLC) in which precoated 250 micron softlayer silica gel GF uniplates (Merck) were used. TLC plates were visualized with UV light (254 nm), vanillin, or ammonium molybdate sprays. Flash chromatography was performed with the indicated solvent system on 60 Å (230–400 mesh, particle size 0.040–0.063 mm) normal phase silica gel. In several cases, all of which are clearly identified in the text, the silica gel for column chromatography was deactivated prior to the actual separation through treatment overnight with a 2% solution of triethylamine in hexane, followed by equilibration with the solvent mixture finally employed. 'Concentrated' refers to the removal of solvent with a rotary evaporator at normal water aspirator pressure followed by further evacuation with a two-stage mechanical pump. Yields refer to chromatographically and spectroscopically pure compounds, except where otherwise noted. All new compounds were determined to be at least 95% pure by means of NMR or GC. All melting points were determined with an open capillary. Chemical shifts were reported in δ values relative to tetramethylsilane in ¹H NMR standard, fluorotrichloromethane in ¹⁹F NMR, and the solvent peak in ¹³C NMR. Peak splitting patterns in the NMR are reported as follows: s, singlet; d, doublet; t, triplet; q, quartet; m, multiplet; br, broad.

4.2. Preparation of imines 3b-e

A standard method was used for the preparation of imines 3b-e:^{20,24} the ketone and the primary amine were refluxed with *p*-toluenesulfonic acid catalysis in refluxing toluene in a Dean–Stark apparatus until water formation was no longer observed. After standard work-up, the desired imines were purified through vacuum distillation. Yields for the purified imines were 93% for 3b, 80% for 3c, 65% for 3d, and 72% for 3e.

4.2.1. (4-Methoxyphenyl)-[1-methylpent-4-(*E*)-enylidene]amine (3d). Yellowish oil. ¹H NMR (300 MHz, CDCl₃) δ 1.79 (s, 3H), 2.46 (m, 4H), 3.78 (s, 3H), 5.02 (dd, J_1 = 10.2 Hz, J_2 =1.7 Hz, 1H), 5.09 (dd, J_1 =17.3 Hz, J_2 = 1.7 Hz, 1H), 5.91 (m, 1H), 6.63 (d, J=8.7 Hz, 2H), 6.84 (d, J=8.7 Hz, 2H); ¹³C NMR (100 MHz, CDCl₃) δ 19.6 (q), 30.5 (t), 40.7 (t), 55.4 (q), 114.1 (d), 115.0 (t), 120.6 (d), 137.7 (d), 144.7 (s), 155.7 (s), 171.4 (s). HRMS (EI⁺) calcd for C₁₃H₁₇NO (M⁺): 203.1310, found: 203.1313.

4.2.2. (*E*)-Butyl-[1-*p*-tolylethylidene]-amine (3e). Yellowish oil. ¹H NMR (300 MHz, CDCl₃) δ 1.01 (t, *J* = 7.5 Hz, 3H), 1.49 (m, 2H), 1.76 (m, 2H), 2.23 (s, 3H), 2.39 (s, 3H), 3.50 (t, *J*=7.2 Hz, 2H), 7.20 (d, *J*=8.1 Hz, 2H), 7.70 (d, *J*=8.1 Hz, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 14.5 (q), 15.6 (q), 21.3 (t), 21.6 (q), 33.6 (t), 52.3 (t), 126.9

(d), 129.3 (d), 139.2 (s), 139.6 (s), 164.9 (s). HRMS (EI⁺) calcd for $C_{13}H_{19}N$ (M⁺): 189.1517, found: 189.1522.

4.3. General procedure for the preparation of 1,3-vinylogous amidines 1a-h

n-Butyllithium (3.0 mmol, 2.5 M in hexane) was slowly added to a solution of diisopropylamine (3.0 mmol) in THF (3 mL) at -30 °C. The mixture was stirred for 30 min, after which the temperature was lowered to -90 °C. A solution of the imine **3** (2.5 mmol) in THF (5 mL) was then added dropwise and the reaction mixture was stirred for 1 h at that temperature to allow azaenolate formation, after which a solution of the nitrile **2** (2.5 mmol) in THF (5 mL) was slowly added. The progress of the reaction was monitored with TLC, and after ca. 1–2 h the reaction was quenched with saturated aqueous NH₄Cl solution and extracted with AcOEt (3×10 mL). The organic layers were pooled together, washed with brine, dried over anhydrous Na₂SO₄, and concentrated to give crude product **1**, which was purified as described below in each case.

4.3.1. (*Z*)-1-(Difluorophenylmethyl)-2-(4,5-dihydro-3*H*pyrrol-2-yl)vinylamine (1a). Flash chromatography of the crude reaction product [*n*-hexane/EtOAc (3:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a yellowish solid (87% yield): mp 82–84 °C. ¹H NMR (300 MHz, CDCl₃) δ 1.69–1.79 (m, 2H), 2.48 (t, *J*=8.2 Hz, 2H), 3.85 (t, *J*=7.2 Hz, 2H), 4.96 (s, 1H), 6.60 (br s, 2H), 7.35–7.37 (m, 3H), 7.50–7.53 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.9 (t), 37.0 (t), 59.0 (t), 88.4 (t, ³*J*_{CF}=6.0 Hz), 117.2 (t, ¹*J*_{CF}=243.7 Hz), 124.6 (t, ³*J*_{CF}= 5.7 Hz), 127.4 (d), 129.3 (t, ⁴*J*_{CF}=1.7 Hz), 134.2 (t, ²*J*_{CF}= 27.3 Hz), 148.2 (t, ²*J*_{CF}=28.1 Hz), 172.2 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –98.5 (s). HRMS (EI⁺) calcd for C₁₃H₁₄F₂N₂ (M⁺): 236.1125, found: 236.1136.

4.3.2. (+)-(*Z*)-1-(Diffuorophenylmethyl)-3-[(*Z*)-(*S*)-1phenylethylimino]-3-phenylpropenylamine (1b). Flash chromatography of the crude reaction product [*n*-hexane/ EtOAc (10:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a yellowish oil (81% yield). $[\alpha]_D^{25}$ +233.8 (*c* 1.11, CHCl₃). ¹H NMR (300 MHz, CDCl₃) δ 1.37 (d, *J*=6.6 Hz, 3H), 4.43 (q, *J*=6.6 Hz, 1H), 4.81 (s, 1H), 7.04–7.11 (m, 4H), 7.13–7.22 (m, 6H), 7.23–7.32 (m, 3H), 7.33–7.49 (m, 2H), 8.95 (br, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 25.9 (q), 57.2 (d), 92.9 (t, ³*J*_{CF}=3.5 Hz), 117.9 (t, ¹*J*_{CF}=245.5 Hz), 126.0 (t, ³*J*_{CF}=6.0 Hz), 126.4 (d), 127.1 (d), 127.8 (d), 128.5 (d), 128.8 (d), 128.9 (d), 130.6 (d), 136.0 (t, ²*J*_{CF}=27.6 Hz), 138.2 (s), 146.2 (s), 161.5 (t, ²*J*_{CF}=28.2 Hz), 164.2 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –100.1 (s). HRMS (EI⁺) calcd for C₂₄H₂₂F₂N₂ (M⁺): 376.1751, found: 376.1768.

4.3.3. [(Z)-3-Amino-4,4-difluoro-1,4-diphenyl-but-2-en-(Z)-ylidene]-*p*-tolylamine (1c). Flash chromatography of the crude reaction product [*n*-hexane/EtOAc (10:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a yellow oil (83% yield). ¹H NMR (300 MHz, CDCl₃) δ 2.13 (s, 3H), 5.18 (s, 1H), 6.49 (d, *J*=8.3 Hz, 2H), 6.81 (d, *J*=8.1 Hz, 2H), 7.06–7.13 (m, 5H), 7.35–7.38 (m, 3H), 7.52–7.55 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 21.1 (q), 95.5 (t, ³*J*_{CF}=4.9 Hz), 118.2 (t, ¹*J*_{CF}=245.1 Hz), 122.5 (d), 126.0 (t, ${}^{3}J_{CF}$ =5.7 Hz), 128.4 (d), 128.7 (d), 128.7 (d), 128.9 (d), 129.4 (d), 130.8 (d), 132.6 (s), 135.5 (t, ${}^{2}J_{CF}$ =27.3 Hz), 138.8 (s), 145.6 (s), 155.0 (t, ${}^{2}J_{CF}$ =28.4 Hz), 165.3 (s); ${}^{19}F$ NMR (282.4 MHz, CDCl₃) δ -99.2 (s). HRMS (EI⁺) calcd for C₂₃H₂₀F₂N₂ (M⁺): 362.1594, found: 362.1557.

4.3.4. [1-((*Z*)-2-Amino-3,3-difluoro-3-phenylpropenyl)pent-4-en-(*E*)-ylidene]-(4-methoxyphenyl)amine (1d). Flash chromatography of the crude reaction product [*n*-hexane/EtOAc (4:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a brownish oil (70% yield). ¹H NMR (300 MHz, CDCl₃) δ 2.04–2.12 (m, 2H), 2.18–2.22 (m, 2H), 3.68 (s, 3H), 4.77–4.84 (m, 2H), 5.02 (s, 1H), 5.48–5.61 (m, 1H), 6.62 (d, *J*=8.8 Hz, 2H), 6.75 (d, *J*=8.8 Hz, 2H), 7.34–7.37 (m, 3H), 7.51–7.54 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 31.1 (t), 32.2 (t), 54.3 (c), 92.5 (t, ³*J*_{CF}=5.7 Hz), 113.0 (d), 114.0 (t), 117.1 (t, ¹*J*_{CF}= 243.9 Hz), 120.7 (d), 124.6 (t, ³*J*_{CF}=5.7 Hz), 127.4 (d), 129.4 (d), 134.2 (t, ²*J*_{CF}=27.3 Hz), 136.1 (d), 142.1 (s), 149.1 (t, ²*J*_{CF}=27.9 Hz), 154.6 (s), 168.9 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –98.6 (s). HRMS (EI⁺) calcd for C₂₁H₂₂F₂N₂O (M⁺) 356.1700, found: 356.1661.

4.3.5. (Z)-1-(Difluoronaphthalen-1-yl-methyl)-2-(4, 5-dihydro-3*H*-pyrrol-2-yl)vinylamine (1e). Flash chromatography of the crude reaction product [*n*-hexane/EtOAc (4:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a yellowish solid (75% yield): mp 163–165 °C. ¹H NMR (300, CDCl₃) δ 1.55–1.65 (m, 2H), 2.29 (t, J=8.2 Hz, 2H), 3.77 (t, J=7.2 Hz, 2H), 4.85 (s, 1H), 6.99 (br s, 2H), 7.29–7.37 (m, 3H), 7.69–7.79 (m, 3H), 8.04–8.07 (m, 1H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.8 (t), 36.9 (t), 58.9 (t), 89.0 (t, ³J_{CF}=6.0 Hz), 118.4 (t, ¹J_{CF}= 243.1 Hz), 123.2 (d), 124.1 (t, ⁴J_{CF}=2.6 Hz), 124.5 (t, ³J_{CF}=8.6 Hz), 125.0 (d), 125.8 (d), 127.5 (d), 128.9 (t, ³J_{CF}=1.7 Hz), 129.0 (t, ²J_{CF}=24.9 Hz), 130.5 (d), 132.8 (s), 148.1 (t, ²J_{CF}=27.0 Hz), 172.2 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –93.2 (s). HRMS (EI⁺) calcd for C₁₇H₁₆F₂N₂ (M⁺): 286.1281, found: 286.1311.

1-[1-(4,5-Dihydro-3*H*-pyrrol-2-yl)-meth-(*Z*)-4.3.6. ylidene]-2,2,3,3,4,4,5,5,6,6,7,7,8,8,8-pentadecafluorooctylamine (1f). Flash chromatography of the crude reaction product [n-hexane/EtOAc (10:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a white solid (71%) yield): mp 140–142 °C. ¹H NMR (300 MHz, CDCl₃) δ 1.74–1.85 (m, 2H), 2.55 (t, J=8.2 Hz, 2H), 3.88 (t, J=7.2 Hz, 2H), 5.14 (s, 1H), 7.02 (br s, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.8 (t), 36.8 (t), 58.7 (t), 90.3 (t, ${}^{3}J_{CF}$ = 6.6 Hz), 107–130 (signals for the C₇F₁₅ group were obscured because of their low intensity), 140.9 (t, ${}^{2}J_{CF}$ = 24.1 Hz), 171.5 (s); 19 F NMR (282.4 MHz, CDCl₃) δ -81.1 (m, 3F), -118.4 (m, 2F), -122.0 (m, 2F), -122.4 (m, 2F), -122.8 (m, 2F), -123.1 (m, 2F), -126.5 (m, 2F). HRMS (EI^+) calcd for $C_{13}H_9F_{15}N_2$ (M⁺): 478.0526, found: 478.0527.

4.3.7. [(*Z*)-3-Amino-4,4,5,5,6,6,7,7,8,8,9,9,10,10,10pentadecafluoro-1-phenyldec-2-en-(*Z*)-ylidene]-*p*-tolylamine (1g). Recrystallization from *n*-hexane/CH₂Cl₂ (20:1) gave a yellowish solid (79% yield): mp 63–65 °C. ¹H NMR (300 MHz, CDCl₃) δ 2.16 (s, 3H), 5.34 (s, 1H), 6.56 (d, *J*= 8.1 Hz, 2H), 6.86 (d, J=8.1 Hz, 2H), 7.10–7.20 (m, 3H), 7.49–7.57 (m, 2H), 8.12–8.18 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 21.1 (t), 96.9 (d), 107–130 (signals for the C₇F₁₅ group were obscured because of their low intensity), 138.2 (s), 145.0 (s), 165.1 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –81.2 (m, 3F), –118.7 (m, 2F), –121.9 (m, 2F), –122.3 (m, 2F), –122.3 (m, 2F), –123.1 (m, 2F), –126.6 (m, 2F). HRMS (EI⁺) calcd for C₂₃H₁₅F₁₅N₂ (M⁺): 604.0995, found: 604.0978.

4.3.8. (*Z*)-3-[(*Z*)-Butylimino]-1-(difluorophenylmethyl)-3-*p*-tolylpropenylamine (1h). Flash chromatography of the crude reaction product [*n*-hexane/EtOAc (10:1)] on deactivated silica gel (2% Et₃N in hexane overnight) gave a yellow oil (69% yield). ¹H NMR (300 MHz, CDCl₃) δ 0.78 (t, *J*=7.1 Hz, 3H), 1.19–1.31 (m, 2H), 1.38–1.47 (m, 2H), 2.29 (s, 3H), 3.09 (t, *J*=6.8 Hz, 2H), 4.72 (s, 1H), 7.10 (s, 4H), 7.31–7.35 (m, 3H), 7.44–7.47 (m, 2H), 9.17 (br s, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 14.2 (q), 20.5 (t), 21.7 (q), 33.7 (t), 46.4 (t), 90.6 (d), 125.9 (t, ³*J*_{C-F}=5.5 Hz), 128.1 (d), 128.8 (d), 129.2 (d), 130.5 (d), 134.7 (s), 139.0 (s), 163.8 (s), 165.7 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ – 100.58 (s). HRMS (EI⁺) calcd for C₂₁H₂₄F₂N₂ (M⁺): 342.1907, found: 342.1897.

4.4. General procedure for the preparation of pyrimidin-2(1*H*)-ones 4a-h

A solution of triphosgene (2.0 mmol) in THF (5 mL) was slowly added to a solution of compound **1** (2.0 mmol) and triethylamine (4 mmol) in THF (10 mL) and the resulting mixture was stirred at room temperature. When TLC monitoring indicated that the reaction was complete, it was quenched with aqueous 2 M KOH solution (5 mL) and extracted with AcOEt (3×5 mL). The organic layers were pooled together, washed with brine, dried over anhydrous Na₂SO₄, and the solvent was removed under vacuum to give a solid, which was purified as indicated below in each case.

4.4.1. 3-[Difluoro(phenyl)methyl]-6,7-dihydropyrrolo-[**1,2-f]pyrimidin-1(5H)-one (4a).** Flash chromatography [*n*-hexane/EtOAc (2:1)] gave a colorless oil (94% yield). ¹H NMR (300 MHz, CDCl₃) δ 2.09–2.19 (m, 2H), 3.07 (t, *J*=7.9 Hz, 2H), 4.02 (t, *J*=7.4 Hz, 2H), 6.51 (s, 1H), 7.30–7.34 (m, 3H), 7.52–7.56 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 18.9 (t), 31.6 (t), 49.0 (t), 95.1 (t, ³*J*_{CF}=4.3 Hz), 116.0 (t, ¹*J*_{CF}=247.1 Hz), 124.6 (t, ³*J*_{CF}=6.3 Hz), 127.4 (d), 129.4 (d), 133.8 (t, ²*J*_{CF}=26.7 Hz), 154.4 (s), 163.9 (s), 169.3 (t, ²*J*_{CF}=31.9 Hz); ¹⁹F NMR (282.4 MHz, CDCl₃) δ – 100.7 (s). HRMS (EI⁺) calcd for C₁₄H₁₂F₂N₂O (M⁺): 262.0917, found: 262.0925.

4.4.2. 4-(Diffuoro(phenyl)methyl)-6-phenyl-1-((S)-1-phenylethyl)pyrimidin-2(1*H***)-one (4b). Flash chromatography [***n***-hexane/EtOAc (4:1)] gave a white solid (90% yield): mp 193–195 °C. [\alpha]_{D}^{25} – 5.79 (***c* **0.91, CHCl₃). ¹H NMR (300 MHz, CDCl₃) \delta 1.82 (d,** *J***=7.1 Hz, 3H), 5.46 (q,** *J***=7.1 Hz, 1H), 6.44 (s, 1H), 7.04–7.18 (m, 7H), 7.34–7.41 (m, 6H), 7.59 (dd,** *J***₁=7.5 Hz,** *J***₂=4.9 Hz, 2H); ¹³C NMR (75.5 MHz, CDCl₃) \delta 17.2 (q), 58.9 (q), 102.1 (t, ³***J***_{CF}= 4.0 Hz), 117.2 (s), 126.3 (t, ³***J***_{CF}=6.3 Hz), 127.1 (d), 127.8 (d), 127.9 (d), 128.7 (d), 128.9 (d), 129.4 (d), 130.9 (d), 134.1 (s), 134.9 (s), 139.1 (s), 155.6 (s), 163.4 (s), 169.7 (s);**

¹⁹F NMR (282.4 MHz, CDCl₃) δ -100.5 (d, ³*J*_{HF}= 3.4 Hz). HRMS (EI⁺) calcd for C₂₅H₂₀F₂N₂O (M⁺): 402.1544, found: 402.1546.

4.4.3. 4-[Difluoro(phenyl)methyl]-6-phenyl-1*-p***-tolyl-pyrimidin-2(1***H***)-one** (**4c**). Recrystallization from *n*-hexane/AcOEt (10:1) gave a white solid (87% yield): mp 204–206 °C. ¹H NMR (300 MHz, CDCl₃) δ 2.19 (s, 3H), 6.64 (s, 1H), 6.89 (d, *J*=8.3 Hz, 2H), 6.99 (d, *J*=8.1 Hz, 2H), 7.03–7.16 (m, 2H), 7.13–7.23 (m, 3H), 7.37–7.40 (m, 3H), 7.65–7.68 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.1 (q), 100.0 (t, ³*J*_{CF}=4.0 Hz), 115.8 (t, ¹*J*_{CF}=247.4 Hz), 124.8 (t, ³*J*_{CF}=6.0 Hz), 126.9 (d), 127.3 (d), 127.4 (d), 127.5 (d), 128.7 (d), 129.0 (d), 129.5 (d), 132.1 (s), 133.5 (t, ²*J*_{CF}=26.7 Hz), 133.8 (s), 137.8 (s), 155.3 (s), 161.1 (s), 169.7 (t, ²*J*_{CF}=32.4 Hz); ¹⁹F NMR (282.4 MHz, CDCl₃) δ – 100.9 (s). HRMS (EI⁺) calcd for C₂₄H₁₈F₂N₂O (M⁺): 388.1387, found: 388.1298.

4.4.4. 6-(But-3-enyl)-4-[difluoro(phenyl)methyl]-1-(**4-methoxyphenyl)pyrimidin-2(1***H***)-one (4d).** Flash chromatography [*n*-hexane/EtOAc (4:1)] gave a white solid (84% yield): mp 115–117 °C. ¹H NMR (300 MHz, CDCl₃) δ 2.13–2.20 (m, 2H), 2.30–2.35 (m, 2H), 3.75 (s, 3H), 4.84–4.93 (m, 2H), 5.46–5.60 (m, 1H), 6.53 (s, 1H), 6.93 (d, *J*=9.0 Hz, 2H), 7.01 (d, *J*=9.0 Hz, 2H), 7.35–7.38 (m, 3H), 7.60–7.63 (m, 2H); ¹³C NMR (75.5 MHz, CDCl₃) δ 30.0 (t), 32.1 (t), 54.5 (q), 97.9 (t, ³*J*_{CF}=4.0 Hz), 114.3 (d), 115.8 (t, ¹*J*_{CF}=247.1 Hz), 115.9 (t), 124.7 (t, ³*J*_{CF}= 6.3 Hz), 127.3 (d), 127.5 (d), 128.3 (s), 129.5 (d), 133.6 (t, ²*J*_{CF}=26.7 Hz), 134.0 (d), 155.9 (s), 159.1 (s), 163.3 (s), 169.4 (t, ²*J*_{CF}=32.2 Hz); ¹⁹F NMR (282.4 MHz, CDCl₃) δ – 101.0 (s). HRMS (EI⁺) calcd for C₂₂H₂₀F₂N₂O₂ (M⁺): 382.1492, found: 382.1493.

4.4.5. 3-[Diffuoro(naphthalen-1-yl)methyl]-6,7-dihydropyrrolo[1,2-f]pyrimidin-1(5H)-one (4e). Recrystallization from *n*-hexane/CH₂Cl₂ (20:1) gave a white solid (77% yield): mp 228–230 °C. ¹H NMR (300 MHz, CDCl₃) δ 2.05–2.15 (m, 2H), 3.03 (t, J=7.9 Hz, 2H), 4.01 (t, J= 7.4 Hz, 2H), 6.53 (s, 1H), 7.37–7.46 (m, 3H), 7.75–7.79 (m, 1H), 7.82–7.87 (m, 2H), 8.04–8.07 (m, 1H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.4 (t), 32.9 (t), 50.3 (t), 97.2 (t, ³J_{CF}=3.1 Hz), 118.7 (t, ¹J_{CF}=247.1 Hz), 125.0 (d), 125.3 (t, ⁴J_{CF}=3.1 Hz), 125.5 (t, ³J_{CF}=9.4 Hz), 126.4 (d), 127.4 (d), 129.1 (d), 129.7 (t, ³J_{CF}=2.3 Hz), 130.5 (t, ²J_{CF}=24.4 Hz), 132.0 (d), 134.3 (s), 155.8 (s), 164.8 (s), 170.7 (t, ²J_{CF}=31.3 Hz); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –95.8 (s). HRMS (EI⁺) calcd for C₁₈H₁₄F₂N₂O (M⁺): 312.1074, found: 312.1053.

4.4.6. 6,7-Dihydro-3-(perfluoroheptyl)pyrrolo[**1,2-***f***]-pyrimidin-1(5***H***)-one (4f**). Recrystallization from *n*-hexane/CHCl₃ (20:1) gave a reddish solid (93% yield): mp 129–131 °C. ¹H NMR (300 MHz, CDCl₃) δ 2.21–2.31 (m, 2H), 3.17 (t, *J*=7.8 Hz, 2H), 4.17 (t, *J*=7.5 Hz, 2H), 6.52 (s, 1H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.3 (t), 33.2 (t), 50.7 (t), 97.7 (t, ³*J*_{CF}=4.3 Hz), 104–134 (signals for the C₇F₁₅ group were obscured because of their low intensity), 163.5 (t, ²*J*_{CF}=25.8 Hz), 166.1 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –81.2 (m, 3F), –116.4 (m, 2F), –121.7 (m, 4F), –122.4 (m, 2F), –123.1 (m, 2F), –126.5 (m, 2F). HRMS

(EI⁺) calcd for $C_{14}H_7F_{15}N_2O$ (M⁺): 504.0318, found: 504.0327.

4.4.7. 4-(Perfluoroheptyl)-6-phenyl-1-*p***-tolylpyrimidin-2(1***H***)-one** (**4g**). Flash chromatography [*n*-hexane/EtOAc (4:1)] gave a colorless oil (90% yield). ¹H NMR (300 MHz, CDCl₃) δ 2.21 (s, 3H), 6.59 (s, 1H), 6.94 (d, *J* = 8.3 Hz, 2H), 7.02–7.09 (m, 4H), 7.17–7.27 (m, 3H); ¹³C NMR (75.5 MHz, CDCl₃) δ 21.5 (q), 102.5 (d), 110–135 (signals for the C₇F₁₅ group were obscured because of their low intensity), 128.2 (d), 128.8 (d), 128.9 (d), 130.3 (d), 130.9 (d), 132.9 (s), 134.8 (s), 139.6 (s), 155.8 (s), 163.6 (s), 163.7 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –81.2 (m, 3F), –117.0 (m, 2F), –121.5 (m, 4F), –122.3 (m, 2F), –123.0 (m, 2F), –126.5 (m, 2F). HRMS (EI⁺) calcd for C₂₄H₁₄F₁₅N₂O (M+H⁺): 631.0886, found: 631.0862.

4.4.8. 1-Butyl-4-(difluoro(phenyl)methyl)-6-*p***-tolyl-pyrimidin-2(1***H***)-one (4h).** Flash chromatography [*n*-hexane/ EtOAc (4:1)] gave a colorless oil (70% yield). ¹H NMR (300 MHz, CDCl₃) δ 0.65 (t, *J*=7.2 Hz, 3H), 0.98–1.11 (m, 2H), 1.47–1.57 (m, 2H), 2.35 (s, 3H), 3.80 (t, *J*=7.9 Hz, 2H), 6.42 (s, 1H), 7.13–7.15 (m, 2H), 7.22–7.25 (m, 2H), 7.33–7.35 (m, 3H), 7.58 (dd, *J*₁=6.4 Hz, *J*₂=1.5 Hz, 2H); ¹³C NMR (75.5 Hz, CDCl₃) δ 13.8 (q), 20.2 (t), 21.8 (q), 30.5 (t), 47.8 (t), 101.9 (t, ³*J*_{CF}=4.0 Hz), 117.2 (t, ¹*J*_{CF}=272.6 Hz), 126.2 (t, ³*J*_{CF}=6.3 Hz), 127.9 (d), 128.9 (d), 130.0 (d), 130.6 (d), 130.8 (d), 135.1 (t, ²*J*_{CF}=26.4 Hz), 141.4 (s), 156.7 (s), 163.3 (s), 169.4 (t, ²*J*_{CF}=32.2 Hz); ¹⁹F NMR (282.4 MHz, CDCl₃) δ –100.8 (s). HRMS (EI⁺) calcd for C₂₂H₂₂F₂N₂O (M⁺): 368.1700, found: 368.1677.

4.5. Preparation of compound 1a · ZnI₂

A solution of **1a** (243 mg; 1.03 mmol) in CH₂Cl₂ (3 mL) was slowly added to a solution of ZnI₂ (330 mg; 1.03 mmol) in CH₂Cl₂ (4 mL) at 0 °C. The mixture was stirred for 1 h, after which the solvent was removed under vacuum to give a solid, which was then purified by means of recrystallization from *n*-hexane/CH₂Cl₂. Mp 158–159 °C. ¹H NMR (300 MHz, CDCl₃) δ 2.01 (m, 2H), 2.74 (t, *J*=8.0 Hz, 2H), 3.71 (s, 2H), 4.14 (t, *J*=7.5 Hz, 2H), 7.51 (m, 5H), 10.22 (s, 1H); ¹³C NMR (75.5 MHz, CDCl₃) δ 20.1 (t), 32.0 (t), 41.2 (t), 60.6 (t), 116.3 (s), 125.9 (t, ³*J*_{CF}=5.8 Hz), 129.3 (s), 130.3 (d), 133.0 (d), 174.3 (s), 177.4 (s); ¹⁹F NMR (282.4 MHz, CDCl₃) δ – 104.8 (s, 2F).

The crystal structure of compound $1a \cdot ZnI_2$ was determined through X-ray diffraction of a single crystal obtained by slow evaporation of a dichloromethane/*n*-hexane solution (Fig. 2).

X-ray data for compound **1a**·**ZnI**₂. Colorless lath, 0.18× 0.05×0.02 mm size, monoclinic, $P2_1/c$, a=8.2737(8), b=21.343(2), c=9.6901(14) Å, $\beta=93.965(9)$, V=1707.0(3) Å³, Z=4, $\rho_{calcd}=2.161$ g cm⁻³, $\theta_{max}=25.00$, Mo K α , $\lambda=0.71073$ Å, ω -scan, diffractometer Siemens P4, T=173(2) K, 3254 reflections collected of which 2992 were independent ($R_{int}=0.071$), absorption correction based on Psi-scans, T_{min}/T_{max} 0.174/0.328, direct primary solution and refinement on F² (Sheldrick, G. M. SHELXS-97 and SHELXL-97, University of Göttingen, 1997), 181 refined parameters, hydrogen atoms refined as riding, the largest difference peaks near the iodine atoms are probably due to absorption, $R_1[I > 2\sigma(I)] = 0.0410$, $wR_2(all data) = 0.1024$.

Selected bond lengths (Å) and angles (°) Zn–N2 2.017(5), Zn–N1 2.041(5), Zn–I2 2.5393(7), Zn–I1 2.5601(8), N1–C2 1.265(8), N2–C4 1.272(8), C1–C2 1.527(8), C2–C3 1.510(8), C3–C4 1.493(8), I2–Zn–I1 117.04(3), N2–Zn–N1 92.1(2), N2–Zn–I2 112.13(13), N1–Zn–I2 113.98(13), N2–Zn–I1 111.36(13), N1–Zn–I1 107.46(14), C2–N1–Zn 126.6(4), C4–N2–Zn 127.4(4), N1–C2–C3 125.5(5), C4–C3–C2 121.7(5), N2–C4–C3 125.9(5).

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Zinc perchlorate catalyzed one-pot amination–annulation of α-cyanomethyl-β-ketoesters in water. Regioselective synthesis of 2-aminopyrrole-4-carboxylates

Ayhan S. Demir* and Mustafa Emrullahoglu

Department of Chemistry, Middle East Technical University, 06531 Ankara, Turkey

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Abstract—In this paper, we report the efficient and regioselective synthesis of 2-aminopyrrole-4-carboxylates as derivatives of conformationally restricted analogues of γ -amino butyrates (GABA) via a zinc perchlorate catalyzed amination–annulation of α -cyanomethyl- β -ketoesters under mild reaction conditions in water.

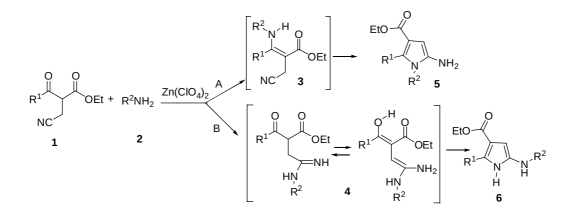
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1. Introduction

Pyrroles are one of the important classes of heterocyclic compounds and are used widely in both synthetic organic chemistry and material science.^{1,2} Pyrroles are often seen as building blocks in naturally occurring and biologically active compounds. Aminopyrroles have been found to show interesting biological properties^{3,4} or have been used as precursors for known drugs⁵ and they are used as synthetic precursors for acyclic nucleoside analogues of the pyrrolo[2,3-*d*]pyrimidine ring system.⁶ Aminopyrroles are not readily available through general pyrrole ring-formation

methods. Many excellent methodologies have been developed for constructing pyrrole rings, in which relatively few examples have been reported for the preparation of simple 2-amino derivatives.⁷

As we have described in previous paper, the condensation reaction of α -cyanomethyl- β -dicarbonyl compounds with amines catalyzed by *p*-TsOH affords the corresponding enamines in good yields. Base catalyzed cyclization via the addition of an amine moiety to the carbon–nitrogen triple bond of nitrile furnished 2-aminopyrroles in high yields (Scheme 1, route A).⁸



Scheme 1.

Keywords: Aminopyrroles; Amination; Hetero annulation; GABA analogs; 1,3-Dicarbonyls.

^{*} Corresponding author. Tel.: +90 312 2103242; fax: +90 312 2101280; e-mail: asdemir@metu.edu.tr

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The addition of nitrogen nucleophiles to CN triple bonds of nitriles is one of the most attractive transformations of nitriles. However, the reported methods are limited because of the low reactivity of nitriles. Development of a catalytic method, which proceeds under neutral and mild conditions is desired in view of the synthetic and environmental aspects. As a line of our study on the development of a synthetic methodology for exploring environmentally friendly processes, and to continue our investigations that are directed towards the synthesis of substituted pyrroles and related compounds,¹⁰ we were especially interested in obtaining 2-amino-4-carboxyl- derivatives of pyrroles, which are conformationally restricted GABA structure analogous. We have found that perchlorate salts are effective catalysts for the activation of both C=O bond and the CN triple bond. These principles have led us to find a novel catalytic one-pot synthesis of 2-aminopyrroles starting from α -cyanomethyl- β -ketoesters.

Herein, we report the novel chemo- and regioselective metalperchlorate-catalyzed amination and annulation of α -cyanomethyl- β -ketoesters.

2. Result and discussion

Metal-coordinated dicarbonyl compound 1 undergoes either C=O activation to react with amines to form enamines 3, or CN triple bond activation of nitriles to have a direct reaction with amines to afford 4 as shown in Scheme 1. This step determines the selective formation of pyrrole isomers 5 and 6 (Scheme 1).

In an initial reaction, we attempted to synthesize the enamine $3a^8$ starting with β -dicarbonyl compound 1a and aniline by using 5 mol% of Zn(ClO₄)₂ in DCM in which the reaction was monitored by TLC. The isolated product was identified as a pyrrole derivative **6a** in excellent yield. The structure of the product showed that pyrrole nitrogen was not from the aniline as expected but from nitrile.

We continued our study by comparing the catalytic activity of zinc perchlorate with other metallic derivatives. Among all of the catalysts tested zinc perchlorate proved to be the most efficient, and 5 mol% of zinc perchlorate showed the highest efficiency. Moreover, the effects of other zinc salts were also tested (Table 1).

This reaction is carried out in different solvents by using 5 mol% Zn(ClO₄)₂ as a catalyst and as shown in Table 2 in which most of the solvents gave comparable yields. The highest yield was obtained with DCM and surprisingly water as a solvent gave comparable yields of DCM. The addition of α -cyanomethyl- β -ketoester and amine into water furnished a heterogen solution, which was heated at 80 °C and the reaction monitored by TLC. The product formation took longer than the DCM but with a comparable yield.

Various α -cyanomethyl- β -ketoesters and amines reacted under the above described conditions and the corresponding pyrroles were obtained in high yields as summarized in Table 2.

Tal	ble	1.	Metal	salts	catal	lyzed	for	mation	of	6a ^a	
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Entry	Catalyst (5 mol%)	Time (h)	Yield (%)
1	$Zn(ClO_4)_2$	3	91
2	$Mg(ClO_4)_2$	4	78
3	$Cu(ClO_4)_2$	4	No product
4	LiClO ₄	5	No product
5	$Co(ClO_4)_2$	5	52
6	NaClO ₄	4	No product
7	$Mn(ClO_4)_2$	4	No product
8	Zinctriflate	3	75
9	$ZnCl_2$	4	27
10	$Zn(OAc)_2$	4	45

^a Commercially available metal salts are used without further purification or drying.

The above described selective amination and annulation reaction shows substrate dependent selectivity by the formation of aminopyrroles. As shown in Table 2 all representative α -cyanomethyl- β -ketoesters furnished with aromatic amines of the pyrrole products **6a–m** gave high yields (Scheme 1, path B). Only one exception was observed when we started with aliphatic α -cyanomethyl- β -ketoesters **2a–c** and aliphatic amines **2f**,g. In this case (Scheme 1, path A) metal-coordinated α -cyanomethyl- β -ketoester undergo C==O activation to give a dehydrative coupling between ketones and amines to form enamines **3**. The addition of a amine moiety to the carbon–nitrogen triple bond furnished 2-aminopyrroles **5a–c** (Table 3).

Many attempts were made for the direct- one-pot synthesis of pyrroles starting with a β -dicarbonyl compound. The reaction of bromoacetonitrile and amine in the presence of catalytical amount of Zn(ClO₄)₂·6H₂O and Mg(ClO₄)₂ by refluxing in DCE and H₂O furnished *N*-substituted β -enamino esters via the condensation of β -ketoesters with amines in a 25–32% yields.⁹

The present reaction can be rationalized by assuming the mechanisms depicted in Scheme 1. The catalytically active species, which would be formed by either the activation of C=O bond or CN bond from α -cyanomethyl- β -ketoesters. Coordination of nitriles to the Zn followed by the addition of the amine into the CN bonds would occur to afford α -cyanomethyl- β -ketoesters metal complex. Coordination of Zn to C=O followed by a dehydrative coupling between ketones and amines would give enamines **3**. Annulation of **3** and **4** would afford product pyrroles **5** and **6** to complete the catalytic cycle. For the chemo- and regioselective processes sterical and electronic factors of the substituents and the enamine–imine tautomeric ratio certainly play an important role. A detailed search for the mechanism of the reaction is under investigation.

3. Conclusions

Typically, when α -cyanomethyl- β -ketoesters was allowed to react with amine in the presence of $Zn(ClO_4)_2$, in addition to the CN triple bond of **1** and subsequent to cyclocondensation took place to afford 2-aminopyrrole. The reactions can be applied to the synthesis of various multifunctionalized

Table 2. Synthesis of aminopyrroles

Entry	Ketoester $1, R_1 =$	Amine 2 , $R_2 =$	Product 6	Solvent	Yield (%) ^a	Reaction time (h)
1	CH ₃ a	C_6H_5 a		DCE H ₂ O Benzene DMF/H ₂ O	91 80 85 40	3 5 4 5
2	а	2,3-(CH ₃) ₂ C ₆ H ₃ b		DCE H ₂ O	78 73	5 7
3	а	2-Cl–C ₆ H ₄ c		DCE H ₂ O	79 77	3 6
4	а	3-Cl-C ₆ H ₄ d		DCE H ₂ O	93 81	3 6
5	a	4-Cl–C ₆ H ₄ e		DCE H ₂ O	94 83	3 7
6	C_2H_5 b	a		DCE H ₂ O	93 77	3 6
7	b	b		DCE H ₂ O	87 75	5 7
8	(CH ₃) ₂ CH c	а		DCE H ₂ O	95 81	4 7
9	C_6H_5 d	а		DCE H ₂ O	89 77	3 6
10	d	e		DCE H ₂ O	85 74	4 7

(continued on next page)

Table 2 (continued)

Entry	Ketoester 1 , $R_1 =$	Amine 2, $R_2 =$	Product 6	Solvent	Yield (%) ^a	Reaction time (h)
11	d	C ₆ H ₅ CH ₂ f		DCE H ₂ O	89 70	5 7
12	2F-C ₆ H ₄ e	a		DCE H ₂ O	87 74	4 6
13	e	f		DCE H ₂ O	85 73	4 6
			г m			

^a Isolated yields.

Table 3.	Synthesis of	f aminopyrroles	with aliph	atic amines in	n DCE and water

Entry	Ketoester 1	Amine 2	Product 5 ^a	Yield (%) DCE/H ₂ O	Reaction time (h) DCE/H ₂ O
14	a	f		76/71	4/7
15	а	C6H5CHCH3 g	eto N NH ₂ b	72/73	5/6
16	b	f		73/76	6/7

^a The compounds are known and have been identified by comparison of spectral data with those reported in the literature.⁸

aminopyrroles. The present $Zn(ClO_4)_2$ catalyzed nitrogencarbon bond formation will provide a wide scope of selective transformations of nitriles and even other substrates under neutral conditions in water. The key point of the present reaction is the selective activation of both C==O bonds of 2-cyanomethyl- β -ketoesters as electrophiles and nitriles as pronucleophiles.

4. Experimental

4.1. Materials and methods

NMR spectra were recorded on a Bruker DPX 400. Chemical shifts δ are reported in ppm relative to CHCl₃ (¹H: δ =7.27), CDCl₃ (¹³C: δ =77.0) and CCl₄ (¹³C: δ =

96.4) as internal standards. IR spectra were recorded on a Perkin Elmer 1600 FTIR series instrument.

Column chromatography was conducted on silica gel 60 (40–63 µm). TLC was carried out on aluminum sheets precoated with silica gel 60F₂₅₄ (Merck), and the spots were visualized with UV light (λ =254 nm). Optical rotations were measured with a Krüss P3002RS automatic polarimeter. Cyanomethylation of β-ketoesters and data for known compounds **5a,b,c**: See Ref.8

4.2. General procedure for the synthesis of pyrroles

a. β -Ketoester (1 mmol) was dissolved in DCE (5 ml). Corresponding amine (1.2 mmol) together with catalytic amount of Zn(ClO₄)₂ (5 mol%) was added to the stirring mixture and refluxed for 3–6 h. Reaction was monitored by TLC. The reaction mixture was extracted with ethyl acetate. The extract was dried over MgSO₄ and the solvent evaporated under reduced pressure and the crude pruduct was purified by column chromotography (hexane–ethyl acetate (4/1)).

b. β -Ketoester (1 mmol) was dissolved in water (5 ml). Corresponding amine (1.2 mmol) together with catalytic amount of Zn(ClO₄)₂ (5 mol%). was added to the stirring mixture. The resulting heterogeneous mixture was heated at 80 °C for 5–7 h. Reaction was monitored by TLC. The reaction mixture was extracted with ethyl acetate. The extract was dried over MgSO₄ and the solvent evaporated under reduced pressure and the crude pruduct was purified by column chromotography (hexane–ethyl acetate (4/1)).

4.2.1. Ethyl 2-methyl-5-(phenylamino)-1*H*-**pyrrole-3carboxylate (6a).** Yield: (222 mg, 91%), white solid (mp=123-125 °C), IR (CHCl₃): 3426, 3295, 3043, 2982, 2930, 2369, 2326, 1682, 1595 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.24 (3H, t, *J*=7.1 Hz), 2.40 (3H, s), 4.16 (2H, q, *J*=7.1 Hz), 5.06 (1H, br s, N*H*), 6.14 (1H, d, *J*=2.6 Hz), 6.52–7.12 (5H, m), 7.98 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 13.1, 14.5, 59.1, 104.1, 111.4, 113.8, 119.3, 127.5, 129.2, 131.9, 146.4, 165.0. Anal. Calcd for C₁₄H₁₆N₂O₂ (244.12): C, 68.83; H, 6.60; N, 11.47. Found: C, 68.61; H, 6.42; N, 11.28.

4.2.2. Ethyl 5-(2,3-dimethylphenylamino)-2-methyl-1*H*-**pyrrole-3-carboxylate (6b).** Yield: (212 mg, 78%), white solid (mp=130–132 °C), IR (CHCl₃): 3414, 3217, 2991, 2978, 2856, 2356, 2330, 1721, 1669 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.36 (3H, t, *J*=7.1 Hz), 2.15 (3H, s), 2.32 (3H, s), 2.51 (3H, s), 4.29 (2H, q, *J*=7.1 Hz), 5.02 (1H, br s, N*H*), 6.21 (1H, d, *J*=2.6 Hz), 6.49–6.93 (3H, m), 8.04 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 13.4, 14.0, 15.4, 21.4, 60.0, 97.0, 104.6, 112.3, 121.7, 122.4, 127.1, 129.1, 132.8, 137.7, 145.4, 166.0. Anal. Calcd for C₁₆H₂₀N₂O₂ (272.15): C, 70.56; H, 7.40; N, 10.29. Found: C, 70.38; H, 7.21; N, 10.06.

4.2.3. Ethyl **5-(2-chlorophenylamino)-2-methyl-1H-pyrrole-3-carboxylate (6c).** Yield: (219 mg, 79%), white solid (mp=149–150 °C), IR (CHCl₃): 3673, 3304, 3052, 2930, 2895, 2353, 2326, 1682, 1591 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.24 (3H, t, *J*=7.1 Hz), 2.44 (3H, s), 4.11 (2H, q, J=7.1 Hz), 5.63 (1H, br s, NH), 6.24 (1H, d, J=2.9 Hz), 6.62–7.21 (4H, m), 7.84 (1H, br s, NH); ¹³C NMR (100 MHz, CDCl₃): 13.1, 14.5, 59.2, 105.6, 111.8, 113.6, 118.8, 119.4, 125.7, 127.7, 129.2, 132.4, 142.7, 164.8. Anal. Calcd for C₁₄H₁₅ClN₂O₂ (278.7): C, 60.33; H, 5.42; N, 10.05. Found: C, 60.12; H, 5.22; N, 9.83.

4.2.4. Ethyl **5-(3-chlorophenylamino)-2-methyl-***1H*-pyrrole-3-carboxylate (6d). Yield: (258 mg, 93%), white solid (mp=137–139 °C), IR (CHCl₃): 3523, 3387, 3022, 2894, 2336, 2229, 2136, 1732, 1681 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.20 (3H, t, J=7.1 Hz), 2.44 (3H, s), 4.18 (2H, q, J=7.1 Hz), 5.15 (1H, br s, NH), 6.15 (1H, d, J=2.6 Hz), 6.46–7.01 (4H, m); ¹³C NMR (100 MHz, CDCl₃): 13.2, 14.4, 59.4, 105.1, 111.6, 111.9, 113.6, 119.2, 126.3, 130.3, 132.6, 135.1, 147.9, 165.2. Anal. Calcd for C₁₄H₁₅ClN₂O₂ (278.7): C, 60.33; H, 5.42; N, 10.05. Found: C, 60.15; H, 5.21; N, 9.84.

4.2.5. Ethyl **5-(4-chlorophenylamino)-2-methyl-***1H*-pyrrole-3-carboxylate (6e). Yield: (261 mg, 94%), white solid (mp=136–137 °C), IR (CHCl₃): 3443, 3391, 3052, 2943, 2336, 2356, 2326, 1739, 1682 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.27 (3H, t, *J*=7.1 Hz), 2.40 (3H, s), 4.16 (2H, q, *J*=7.1 Hz), 5.13 (1H, br s, *NH*), 6.13 (1H, d, *J*=2.6 Hz), 6.50 (2H, d, *J*=8.7 Hz, Ph-H), 7.01 (2H, d, *J*= 8.7 Hz), 8.12 (1H, br s, *NH*); ¹³C NMR (100 MHz, CDCl₃): 13.1, 14.5, 59.3, 104.3, 111.4, 114.9, 124.0, 127.1, 129.1, 132.2, 145.1, 165.2. Anal. Calcd for C₁₄H₁₅ClN₂O₂ (278.7): C, 60.33; H, 5.42; N, 10.05. Found: C, 60.18; H, 5.33; N, 10.22.

4.2.6. Ethyl 2-ethyl-5-(phenylamino)-1*H*-pyrrole-3-carboxylate (6f). Yield: (239 mg, 93%), white solid (mp = 125 °C), IR (CHCl₃): 3316, 3275, 3047, 2962, 2839, 2329, 2316, 1785, 1621 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.14 (3H, t, *J*=7.5 Hz), 1.23 (3H, t, *J*= 7.2 Hz), 2.89 (2H, q, *J*=7.5 Hz), 4.19 (2H, q, *J*=7.2 Hz), 5.13 (1H, br s, N*H*), 6.21 (1H, d, *J*=2.9 Hz), 6.61–7.13 (5H, m), 7.98 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 13.4, 14.4, 20.5, 59.3, 104.1, 110.4, 113.8, 119.3, 127.6, 129.3, 138.1, 146.5, 165.3. Anal. Calcd for C₁₅H₁₈N₂O₂ (258.32): C, 69.74; H, 7.02; N, 10.84. Found: C, 69.53; H, 7.11; N, 10.63.

4.2.7. Ethyl 5-(2,3-dimethylphenylamino)-2-ethyl-1*H*-**pyrrole-3-carboxylate (6g).** Yield: (248 mg, 87%), semisolid, IR (CHCl₃): 3512, 3312, 2871, 2934, 2867, 2312, 2336, 1678, 1678 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.14 (3H, t, *J*=7.5 Hz), 1.24 (3H, t, *J*=7.1 Hz), 2.03 (3H, s), 2.23 (3H, s), 2.85 (2H, q, *J*=7.5 Hz), 4.13 (2H, q, *J*=7.1 Hz), 4.94 (1H, br s, NH), 6.09 (1H, d, *J*=2.8 Hz), 6.39–6.71 (3H, m), 8.06 (1H, br s, NH); ¹³C NMR (100 MHz, CDCl₃): 12.5, 13.6, 14.5, 20.3, 20.5, 59.1, 103.5, 110.4, 111.3, 120.8, 121.4, 126.2, 128.3, 136.8, 137.8, 144.5, 165.0. Anal. Calcd for C₁₇H₂₂N₂O₂ (286.37): C, 71.30; H, 7.74; N, 9.78. Found: C, 71.12; H, 7.51; N, 9.48.

4.2.8. Ethyl 2-isopropyl-5-(phenylamino)-1*H***-pyrrole-3-carboxylate (6h).** Yield: (187 mg, 95%), semisolid, IR (CHCl₃): 3523, 3285, 3061, 2979, 2922, 2345, 2313, 1672, 1612 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.26 (6H, d, *J* =

7.0 Hz), 1.34 (3H, t, J=7.1 Hz), 3.79–3.83 (1H, m), 4.27 (2H, q, J=7.1 Hz), 5.21 (1H, br s, NH), 6.24 (1H, d, J= 2.8 Hz), 6.63–7.31 (5H, m), 8.07 (1H, br s, NH); ¹³C NMR (100 MHz, CDCl₃): 14.5, 22.1, 25.8, 59.2, 104.1, 109.9, 113.7, 119.3, 127.4, 129.3, 142.0, 146.5, 164.9. Anal. Calcd for C₁₆H₂₀N₂O₂ (272.15): C, 70.56; H, 7.40; N, 10.29. Found: C, 70.41; H, 7.38; N, 10.15.

4.2.9. Ethyl 2-phenyl-5-(phenylamino)-1*H***-pyrrole-3carboxylate (6i). Yield: (272 mg, 89%), white solid (mp=141–142 °C), IR (CHCl₃): 3421, 3065, 2982, 2934, 2904, 1695, 1591 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): \delta 1.17 (3H, t,** *J***=7.1 Hz), 4.10 (2H, q,** *J***=7.1 Hz), 5.2 (1H, br s, N***H***), 6.31 (1H, d,** *J***=2.3 Hz), 6.53–7.55 (10H, m), 8.15 (1H, br s, N***H***); ¹³C NMR (100 MHz, CDCl₃): 14.3, 30.7, 59.4, 105.4, 111.9, 114.1, 119.7, 128.0, 128.8, 129.4, 129.6, 131.8, 133.6, 145.9, 164.1. Anal. Calcd for C₁₉H₁₈N₂O₂ (306.14): C, 74.49; H, 5.92; N, 9.14. Found: C, 74.28; H, 5.83; N, 8.91.**

4.2.10. Ethyl 5-(4-chlorophenylamino)-2-phenyl-1H-pyrrole-3-carboxylate (6j). Yield: (289 mg, 85%), white solid (mp=132–133 °C), IR (CHCl₃): 3513, 3372, 3153, 2897, 2239, 2450, 2389, 1732, 1679 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.14 (3H, t, *J*=7.1 Hz), 4.04 (2H, q, *J*=7.1 Hz), 5.27 (1H, br s, N*H*), 6.25 (1H, s), 6.53–7.44 (9H, m), 8.29 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 15.1, 60.6, 105.4, 112.5, 116.1, 125.2, 128.9, 128.9, 129.7, 130.1, 130.5, 132.6, 145.2, 165.6. Anal. Calcd for C₁₉H₁₇ClN₂O₂ (340.8): C, 66.96; H, 5.03; N, 8.22. Found: C, 66.85; H, 5.13; N, 8.02.

4.2.11. Ethyl 5-(benzylamino)-2-phenyl-1*H*-pyrrole-3carboxylate (6k). Yield: (284 mg, 89%), yellow oil, IR (neat): 3413, 3272, 3142, 2787, 2199, 2421, 2298, 1752, 1685 cm^{-1.} ¹H NMR (400 MHz, CDCl₃): δ 1.10 (3H, t, *J*= 7.1 Hz), 4.04 (3H, q, *J*=7.1 Hz), 4.07 (2H, br s), 5.68 (1H, d, *J*=2.8 Hz), 7.11–7.39 (10H, m), 8.04 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 14.3, 50.6, 59.3, 92.4, 111.1, 127.3, 127.4, 127.7, 127.8, 128.1, 128.5, 128.6, 128.9, 130.8, 132.3, 139.0, 139.3, 165.1. Anal. Calcd for C₂₀H₂₀N₂O₂ (320.38): C, 74.98; H, 6.29; N, 8.74. Found: C, 74.81; H, 6.14; N, 8.51.

4.2.12. Ethyl 2-(2-fluorophenyl)-5-(phenylamino)-1*H*-pyrrole-3-carboxylate (6l). Yield: (281 mg, 87%), white solid (mp=113 °C), IR (CHCl₃): 3525, 3266, 2987, 2921, 2964, 1699, 1601 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.16 (3H, t, *J*=7.2 Hz), 4.15 (2H, q, *J*=7.2 Hz), 5.25 (1H, br s, N*H*), 6.39 (1H, s), 6.71–7.61 (9H, m), 8.31 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 11.1, 56.6, 99.9, 110.3, 111.1, 112.3 (*J*=22 Hz), 116.4, 120.4, 122.8, 126.2, 126.5, 126.7, 127.6, 128.8, 142.2, 156.9 (d, *J*=246 Hz), 161.6. Anal. Calcd for C₁₉H₁₇FN₂O₂ (324.35): C, 70.36; H, 5.28; N, 8.64. Found: C, 70.32; H, 5.22; N, 8.41.

4.2.13. Ethyl 5-(benzylamino)-2-(2-fluorophenyl)-1*H*pyrrole-3-carboxylate (6m). Yield: (287 mg, 85%), yellow oil, IR (neat): 3523, 3472, 3347, 2687, 2231, 2521, 2342, 1749, 1699 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.12 (3H, t, *J*=7.0 Hz), 4.04 (2H, q, *J*=7.0 Hz), 4.07 (2H, br s), 5.73 (1H, s), 6.91–7.48 (9H, m), 8.14 (1H, br s, N*H*); ¹³C NMR (100 MHz, CDCl₃): 14.2, 50.5, 59.5, 92.0, 113.2, 115.4 (d, J=22 Hz), 120.3 (d, J=13 Hz), 123.6 (d, J=8 Hz), 127.4, 127.7, 128.1, 128.6, 129.0, 131.9, 138.9, 139.6, 159.1 (d, J=245 Hz), 164.9. Anal. Calcd for C₂₀H₁₉FN₂O₂ (338.38): C, 70.99; H, 5.66; N, 8.28. Found: C, 70.82; H, 5.45; N, 8.02.

4.2.14. Ethyl 2-(cyanomethyl)-4-methyl-3-oxopentanoate (1c). Yield: (155 mg, 79%), yellow oil, IR (neat): 2978, 2249, 1736, 1689 cm⁻¹. ¹H NMR (400 MHz, CDCl₃): δ 1.14 (3H, d, *J*=6.7 Hz), 1.20 (3H, d, *J*=7.1 Hz), 1.31 (3H, t, *J*=7.2 Hz), 2.83 (2H, d, *J*=7.3 Hz), 2.91 (1H, m), 3.98 (1H, t, *J*=7.3 Hz), 4.27 (2H, q, *J*=7.2 Hz); ¹³C NMR (100 MHz, CDCl₃): 13.9, 15.9, 18.4, 19.2, 40.5, 52.5, 62.3, 116.9, 166.3, 204.7. Anal. Calcd for C₁₀H₁₅NO₃ (197.23): C, 60.90; H, 7.67; N, 7.10. Found: C, 60.81; H, 7.48; N, 7.33.

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Tetrahedron

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Stable chiral spirocyclic [5,5]-ammonium ylides using a metallo carbenoid approach

Daniele Muroni,^a Antonio Saba^{a,*} and Nicola Culeddu^b

^aDipartimento di Chimica, Facoltà di Scienze, Università di Sassari, Via Vienna 2, 07100 Sassari, Italy ^bCNR Istituto di Chimica Biomolecolare Sez. di Sassari, Via La Crucca, Baldinca- Li Punti (Sassari) 07040, Italy

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Abstract—Enantiomerically pure spiro[5,5]-ammonium ylides were obtained by Rh(II)-catalyzed decomposition of α -diazo- β -carbonylesters. In the crude decomposition mixtures, variable quantities of enamino- α , β -keto esters were detected as secondary products. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Recently, we prepared enantiopure indolizidinone alkaloids 2 by a carbenoid/spiro[5,5]-ammonium ylide/Stevens [1,2]shift with ring-expansion tandem sequence¹ (Scheme 1). In this contest, isolation and characterization of vlide intermediates 1 in a stable and enantiopure form, enabled us, by studying their reactivity, to probe unambiguously the complete reaction cascade pathway previously proposed for similar processes, including the stereochemical reaction course.² We proved high chirality transfer from the original proline template to a second temporary spirocyclic ammonium nitrogen through its stereoselective quaternarization. The second one permitted the preservation of the original stereocenter during its [1,2]-shift to the newly formed neighbour quaternary indolizidinone stereocenter, a synthetic application of the SRS (self-regeneration of stereocenters) Seebach principle.³

In some related studies involving metallo carbenoid generation and Stevens [1,2]-rearrangement of similar

non-isolable ylide intermediates, a high degree of chirality transfer from the ammonium N atom was previously observed.²

This stereoselectivity is in line with the extensive investigations of Ollis that demonstrate a solvent caged radical pair Stevens [1,2]-rearrangement reaction mechanism⁴ or with an alternative recently proposed ion pair mediated mechanism.⁵

Concerning the Stevens [1,2]-rearrangement, notwithstanding the considerable examples performed in related studies, there are few synthetic applications. One limitation of this methodology is the need for activating groups on the migrating carbons, such as allyl, aryl, carbonyl⁶ and silyl;^{2c} consequently, no examples of a primary carbon migration have been reported, except for a few cases.⁷

 $\begin{array}{c} & & & \\ &$

Scheme 1.

Keywords: Diazocompounds; Catalytic decomposition; Cascade process; Rearrangement.

* Corresponding author. Tel.: +39 079 229538; fax: +39 079 229559; e-mail: saba@uniss.it

In view of the mild conditions, the carbenoid route seems most suitable for intramolecular ylide trapping, and given the rarity of stable spirocyclic ammonium ylides obtained

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by this protocol,⁸ we were interested in increasing their number and in studying their reactivity, including rearrangement and cycloaddition reactions.

For our purposes, the synthesis of diazoketones **4a**, **4b**, **4c**, **4d** and **4e** was planned (Figs. 1 and 2). Due to the proper diazo group position on the chain tethered to pyrrolidine or isoindoline nitrogen atom, [5,5]-spirocyclic ammonium ylides **5a**, **5b**, **5c**, **5d** and **5e** (Fig. 3) were expected to form predominantly as intermediates, by nitrogen trapping of the metallocarbene precursors, over competitive C–H insertion processes.

Moreover, for evaluating the enantioselectivity in generating quaternary ammonium stereocenters, the chiral diazo-substrates **4b**, **4c** and **4e** were prepared as single enantiomers.

Finally, the diazocompound **4d** synthesis was suggested by our interest in evaluating the possibility of the Stevens [1,2]-

rearrangement of the corresponding ylide **5d**; to our knowledge, no examples of tertiary carbon Stevens-type migration are reported.

2. Results and discussion

All the diazoketoesters **4a**–**e**, were conveniently prepared in two steps. First step was the conjugate addition of isoindoline **3a** to (-)-menthyl-3-keto-pent-4-enoate,⁹ and 3,4-bis-methoxymethoxy-pyrrolidine **3b**,¹⁰ 1,4-dideoxy-2,3,5,6-*O*-isopropylidene-1,4-imino-D-talitol **3c**,¹¹ 2-methoxymethyl-2-methyl-pyrrolidine **3d**,¹² and 1,4-dideoxy-2,3di-*O*-isopropylidene-1,4-imino-5-*O*-trityl-L-lyxitol **3e**¹³ to ethyl-3-keto-pent-4-enoate.¹⁴ Then the *N*-alkyl-isoindoline and the *N*-alkyl-pyrrolidines obtained were submitted to diazo-transfer reaction with tosylazide.

¹H NMR analysis of diazoketoesters 4b and 4c and 4e indicated their enantiomeric purity.

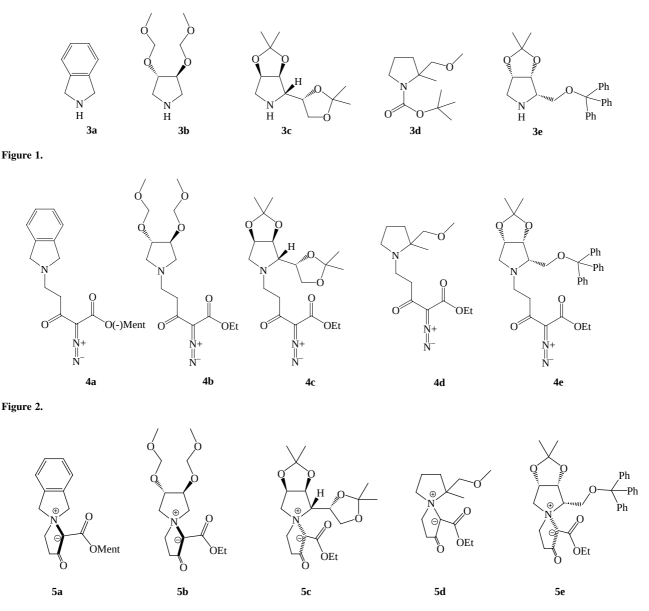
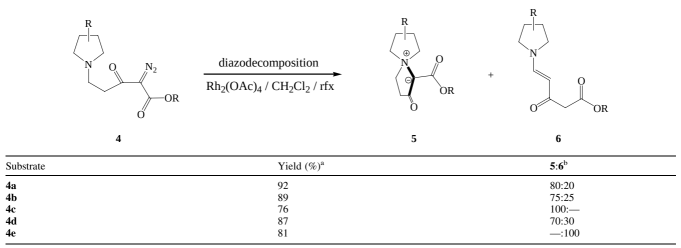


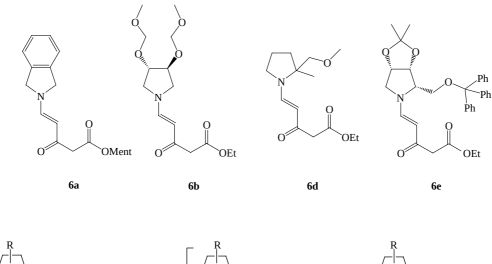
Figure 3.



^a Isolated yield after column chromatography.

^b Reflects percent conversion (measured by ¹H NMR of crude mixture after removal of solvents).

When the diazocompounds **4a**, **4b**, **4c** and **4d** were refluxed in CH_2Cl_2 solution, in the presence of rhodium(II) acetate, the corresponding ylides **5a**, **5b**, **5c** and **5d** were obtained¹⁵ (Fig. 3 and Table 1). Ylide **5d** was obtained as a 44:56 diastereomeric mixture. In the crude decomposition mixtures, variable quantities of enamino- α , β -keto esters **6a**, **6b** and **6c** were detected as secondary products (Fig. 4). No detectable amount of ylide



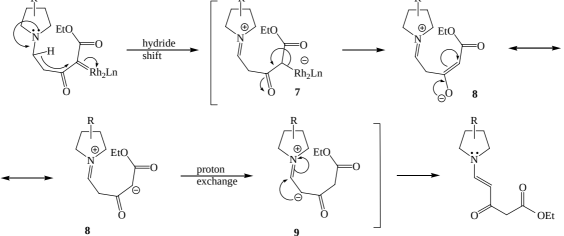


Figure 4.

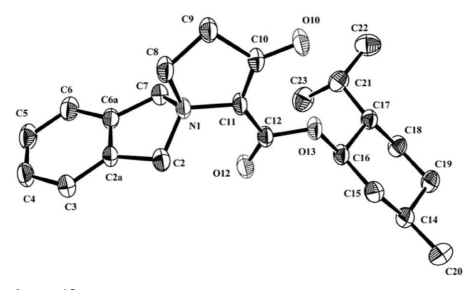
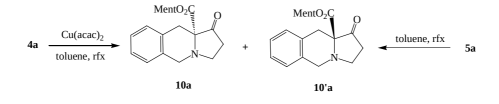


Figure 5. ORTEP view of compound 5a.



Scheme 3.

was found in the crude reaction mixture of the dirhodium tetraacetate decomposition of diazoketone **4e** in boiling CH_2Cl_2 solution. In this case, ketopentenoic ester **6e** was obtained as the exclusive product in 81% isolated yield.

The formation of unsaturated compounds can be rationalized by a mechanism (Scheme 2), which involves an initial intramolecular metal–carbene hydride-abstraction, ¹⁶ to give a transient zwitterion intermediate **7**, which then produces the zwitterion enolate/ketone **8**. This seemingly undergoes proton exchange¹⁷ with an adjacent α -hydrogen to give a new zwitterionic intermediate **9**. A subsequent dissipation of the charges then occurs to furnish enamine **6** with trans configuration (¹H NMR spectroscopy).

Isolation of ylide 5a crystals enabled the single crystal X-ray analysis¹⁸ to be performed (Fig. 5).

The absolute configuration of ylide **5c** was deduced from 2D NOESY correlation studies. Thus, the NOESY trace of ethyl ester methylene protons shows a positive NOE effect for the acetonide methylene protons.

The ¹H NMR analysis of ylides **5b** and **5c** indicated their enantiomeric purity.

Decomposition of the diazocompound 4a in boiling toluene in the presence of Cu(acac)₂ provided an inseparable 1:1 diastereomeric mixture of indolizidinone alkaloids **10a** and **10'a**; surprisingly no enantioselection was obtained. By heating of ylide **5a** in toluene at reflux,

without catalyst, the same diastereomeric mixture was obtained (Scheme 3).

No Stevens rearrangement products but undetectable decomposition mixtures were obtained by heating of ylides **5b** and **5c** and **5d** under the above reaction conditions.

3. Conclusion

These preliminary studies have shown that stable ammonium ylides may also be obtained using the mild and concise metallo carbenoid protocol. The ready isolation of these intermediates could be attributed to the stabilizing effect exerted on the charged ylide carbon atom by the presence of both carbonyl and ester groups. The lack of ylide formation in the catalytic decomposition of diazocompound **4e** can be rationalized in terms of steric factors.

Moreover, the thermal decomposition of ylide **5a**, affording the same mixture of alkaloids **10a** and **10'a** as obtained by catalytic decomposition of the starting diazoketone **4a**, provides a further clear confirmation of a carbenoid/ spiro[5,5]-ammonium ylide/Stevens [1,2]-shift with ringexpansion tandem sequence. In this case, the [1,2]-shift is assisted by the presence of the aryl group on the migrating carbon.

No Stevens-type rearrangement products were detected in the crude reaction mixtures of the attempted thermal decomposition of ylides **5b** and **5c**: this is presumably due to the absence of activating groups on the potential migrating carbons.

No rearrangement products were also observed by heating of ylide **5d**; in our opinion, this result rules out the radical pair reaction mechanism. Due to its stability, a putative migrating tertiary radical intermediate would favour the rearrangement process.

4. Experimental

4.1. General

¹H (300 MHz) and ¹³C NMR (75 MHz) spectra were recorded on a Varian VXR-300 spectrometer with TMS as internal standard. COSY, NOESY, HSQC and USQC-TOSCY spectra were recorded with a Bruker Avance 600 NMR spectrometer equipped with inverse detection probe. Infrared (IR) spectra were performed on a FT/IR-480plus JASKO spectrophotometer. The optical rotations were measured by a polarimeter P-1010 JASKO in a 1 dm tube. All reagents and solvents employed were reagent grade materials purified by standard methods and redistilled before use.

4.1.1. 2-Diazo-3-oxo-5-[1,3-dihydroisoindol-yl]-pentanoic acid (-)-menthyl ester 4a. To a stirred solution of isoindoline **3a** (0.5 g, 0.0042 mol) and 3-oxo-pent-4-enoic acid (-)-menthyl ester (1.1 g, 0.004 mol) in CH_2Cl_2 (15 mL), a solution of Et₃N (0.76 mL, 0.0055 mol) in CH₂Cl₂ (10 mL) was added dropwise and stirred for 30 min. To the reaction mixture, tosyl azide (1.1 g, 0.0055 mol) and a solution of Et₃N (1.1 mL, 0.015 mol) in CH₂Cl₂ (10 mL) were added dropwise at 0 °C. After the addition was complete, the solution was stirred at room temperature overnight. The solvent was evaporated and the residue was purified by flash chromatography (petroleum ether/Et₂O, 8:2) affording the title compound 4a (0.995 g, 60% yield) as a yellow-brown oil: $\left[\alpha\right]_{D}^{25}$ – 53 (c 0.7, CHCl₃); ¹H NMR: δ 0.80 (d, 3H, J=6.9 Hz), 0.92 (two overlapping doublets, 6H, J = 6.9 Hz), 0.99–1.12 (m, 2H), 1.39–1.53 (m, 2H), 1.69–1.73 (m, 3H), 1.80–1.90 (m, 1H), 2.04–2.17 (m, 1H), 3.07-3.20 (m, 4H), 3.98 (s, 4H), 4.84 (dt, 1H, d: J=4.5 Hz, t: J=10.8 Hz), 7.18 (s, 4H); ¹³C NMR (CDCl₃): δ 16.4, 20.6, 21.9, 23.5, 26.5, 31.4, 34.0, 39.1, 41.0, 46.9, 50.6, 58.9, 75.7, 76.5, 122.2, 126.7, 139.8, 160.9, 191.4; IR (neat): 2955, 2870, 2765, 2368, 2131, 1710, 1656, 1455, 1368, 1209, 1039, 1014, 913, 873 and 742 cm^{-1} . Anal. Calcd for C₂₃H₃₁N₃O₃: C, 69.49; H, 7.86; N, 10.57. Found: C, 69.61; H, 7.58; N, 10.75.

4.1.2. Diazo-decomposition of 4a. A solution of diazoketoester **4a** (0.5 g, 0.00126 mol) in CH₂Cl₂ (5 mL) was refluxed for 30 min in the presence of Rh₂(OAc)₄ (0.007 g, 2 mol%). The solvent was evaporated and the residue gave a 80:20 mixture of **5a** and **6a**. After flash chromatography (Et₂O/MeOH/Et₃N, 5:5:0.1) **5a** was obtained as a white solid (0.336 g, 74% yield) and **6a** as impure brown oil (0.081 g, 18% yield).

2'-[(-)-Menthoxycarbonyl]-3'-oxo-1,3-dihydrospiro [isoindole-2,1'-pyrrolidine]-2'-ylide **5a** colourless needles 1463

(ethyl acetate/Et₂O, 10:1), mp 185–187 °C; $[\alpha]_{D}^{25}$ –45 (*c* 0.4, CHCl₃); ¹H NMR: δ 0.76 (d, 3H, *J*=6.9 Hz), 0.88 (two overlapping doublets, 6H, *J*=6.9 Hz), 1.02–1.06 (m, 1H), 1.07–1.25 (m, 1H), 1.36–1.65 (m. 4H), 2.06–2.24 (m, 3H), 2.45–2.73 (m, 2H), 3.65–3.69 (m, 2H), 4.27–4.50 (m, 2H), 4.74 (dt, 1H, d: *J*=4.5 Hz, t: *J*=10.8 Hz), 5.74–5.86 (m, 2H), 7.26–7.41 (m, 4H); ¹³C NMR (CDCl₃): δ 16.1, 20.9, 22.1, 23.3, 25.7, 32.6, 32.8, 34.3, 41.5, 46.8, 63.3, 68.6, 72.9, 123.0, 128.9, 133.8, 163.2, 178.3; IR (Nujol): 2924, 2854, 1640, 1606, 1460, 1413, 1376, 1200, 1112, 1013, 973, 767and 723 cm⁻¹. Anal. Calcd for C₂₃H₃₁NO₃: C, 74.46; H, 8.46; N, 3.79. Found: C, 74.20; H, 8.76; N, 3.55.

3-Oxo-5-[1,3dihydroisoindol-yl]-pent-3,4-enoic acid (-)menthyl ester **6a**, brown oil; $[\alpha]_{25}^{25} - 34$ (c 0.11, CHCl₃); ¹H NMR: δ 0.76 (d, 3H, J=6.9 Hz), 0.86–0.91 (m, 6H), 0.98–1.10 (m, 2H), 1.25 (s, 2H), 1.32–1.51 (m, 2H), 1.63–1.70 (m, 3H), 1.83–2.06 (m, 2H), 3.42 (s, 2H), 4.75 (dt, 1H, d: J=4.5 Hz, t: J=10.8 Hz), 4.87 (s, 2H), 5.21 (d, 1H, J=12.6 Hz), 7.26–7.34 (m, 4H), 7.90 (d, 1H, J= 12.6 Hz); ¹³C NMR (CDCl₃): δ 16.2, 20.7, 22.0, 23.2, 26.0, 29.7, 31.4, 34.2, 40.7, 46.8, 53.3, 57.1, 74.9, 122.5, 122.7, 127.9, 128.0, 135.7, 149.1, 168.5, 188.7; IR (neat): 2955, 2869, 1715, 1650, 1571, 1463, 1361, 1271, 1143, 1097, 989 and 741 cm⁻¹. Anal. Calcd for C₂₃H₃₁NO₃: C, 74.46; H, 8.46; N, 3.79. Found: C, 74.54; H, 8.31; N, 3.40.

4.1.3. (10aR,S)-1-Oxo-2,3,4,5,10-tetrahydro-1H-pyrrolo[1,2-b]-isoquinoline-10a-carboxylic acid (-)menthyl esters 10a and 10'a. A solution of diazoketoester 4a (0.44 g, 0.00110 mol) in toluene (5 mL) was refluxed for 30 min in the presence of $Cu(acac)_2$ (0.005 g, 2 mol%). The solvent was evaporated and the residue gave, after flash chromatography (petroleum ether/ethyl acetate, 9:1) the title compounds 10a and 10'a as a 1:1 mixture of diastereoisomers (0.21 g, 52% yield). ¹H NMR: δ 0.62 (0.41) (d, 3H, J = 6.9 Hz), 0.76–0.92 (m, 9H), 1.19–1.37 (m, 5H), 1.56–1.74 (m, 4H), 2.48–2.67 (m, 2H), 2.87 (2.82) (d, 1H, J = 15.6 Hz), 3.24–3.38 (m, 2H), 3.52 (q, 1H, J = 7.8 Hz), 4.02-4.23 (m, 2H), 4.60 (4.56) (dt, 1H, d: J=4.5 Hz, t: J=10.8 Hz), 7.07–7.17 (m, 4H). Anal. Calcd for C₂₃H₃₁NO₃: C, 74.46; H, 8.46; N, 3.79. Found: C, 74.31; H, 8.18; N, 3.98.

4.1.4. 5-[(3S,4S)-3,4-Bis(methoxymethoxy)pyrrolidin-1yl]-2-diazo-3-oxo-pentanoic acid ethyl ester 4b. To a stirred solution of 3,4-bis-methoxymethoxy-pyrrolidine 3b (1.6 g, 0.008 mol) in CH₂Cl₂ (15 mL) a solution of 3-oxopent-4-enoic acid ethyl ester (1.1 g, 0.008 mol) in CH₂Cl₂ (10 mL) was added dropwise and stirred for 30 min. To the reaction mixture tosyl azide (1.6 g, 0.008 mol) and a solution of Et₃N (2.8 mL, 0.02 mol) in CH₂Cl₂ (10 mL) were added dropwise at 0 °C. After the addition was complete, the solution was warmed to room temperature and stirred overnight. The solvent was evaporated and the residue was purified by flash chromatography (Et₂O/ petroleum ether/Et₃N, 8:2:0.1) to give the title compound **4b** (1.8 g, 70% yield) as a yellow oil; $[\alpha]_D^{25} + 3$ (c 0.6, CHCl₃); ¹H NMR (CDCl₃): δ 1.33 (t, 3H, J=7.2 Hz), 2.58 (dd, 2H, J=4.2, 10.5 Hz); 2.70-2.90 (m, 2H), (dd, 2H, J=6.0, 9.9 Hz), 3.09 (dd, 2H, J=6.9, 8.1 Hz), 3.37 (s, 6H), 4.13 (dd, 2H, J=4.2, 5.1 Hz), 4.29 (q, 2H, J=7.2 Hz), 4.67 (AB system, 4H); ¹³C NMR (CDCl₃): δ 14.2, 38.6. 50.4,

55.3, 58.6, 61.2, 76.0, 81.1, 95.4, 161.0, 191.1; IR (neat): 2945, 2822, 2136, 1718 and 1655 cm⁻¹. Anal. Calcd for C₁₅H₂₅N₃O₇: C, 50.13; H, 7.01; N, 11.69. Found C, 50.24; H, 7.03; N, 11.65.

4.1.5. Diazo-decomposition of 4b. To a refluxing solution of $Rh_2(OAc)_4$ (0.013 g, 3 mol%) in 30 mL of dry CH_2Cl_2 , a solution of **4b** (0.360 g, 0.001 mol) in dry CH_2Cl_2 (20 mL) was added dropwise over 30 min. After stirring for another 30 min at reflux, the reaction mixture was cooled and evaporated to give a 75:25 mixture of **5b** and **6b**. Purification by flash chromatography (Et₂O/ethyl acetate/Et₃N, 5:5:0.1; Et₂O/MeOH/Et₃N, 5:5:0.1) gave 0.22 g (67% yield) of **5b** as a yellow amorphous solid and 0.074 g (22% yield) of **6b** as yellow oil.

(75,85)-1-Ethoxycarbonyl-7,8-bis-methoxymethoxy-2-oxo-5-azonia-spiro[4.4] nonane-1-ylide **5b** $[\alpha]_D^{25} - 3$ (*c* 0.15, CHCl₃); ¹H NMR: δ 1.35 (t, 3H, J=7.2 Hz), 2.60 (t, 2H, J= 7.8 Hz), 3.25 (d, 1H, J=12.9 Hz), 3.39 (d, 6H, J=6.6 Hz), 3.55–3.64 (m, 2H), 3.70–3.90 (m, 1H), 4.26 (q, 2H, J= 7.2 Hz), 4.37–4.58 (m, 3H), 4.68 (dd, 2H, J=6.9, 10.2 Hz), 4.75 (dd, 2H, J=3.3, 6.9 Hz), 4.88 (dd, 1H, J=6.9, 12.9 Hz); ¹³C NMR (CDCl₃): δ 14.7, 32.8, 55.9, 59.0, 63.3, 66.0, 80.0, 80.2, 96.3, 96.4, 102.3, 162.3, 178.3; IR (neat): 2933, 2826, 1735, 1653, 1609 and 1567 cm⁻¹. Anal. Calcd for C₁₅H₂₅NO₇: C, 54.37; H, 7.60; N, 4.23. Found: C, 54.17; H, 7.58; N, 4.21.

(4*E*)-5-[(3*S*,4*S*)-3,4-bis(Methoxymethoxy)pyrrolidin-1-yl]-3-oxo-pentenoic acid ethyl ester **6b** $[\alpha]_{25}^{15}$ +5 (*c* 0.10, CHCl₃); ¹H NMR: δ 1.27 (t, 3H, *J*=7.1 Hz), 3.23 (d, 1H, *J*= 12.6 Hz), 3.36 (s, 6H), 3.38 (s, 2H), 3.44 (d, 1H, *J*=11.8 Hz), 3.53 (d, 1H, *J*=11.7 Hz), 4.14–4.21 (m, 1H), 4.17 (q, 2H, *J*= 7.1 Hz), 4.26 (broad s, 1H), 4.60–4.75 (m, 4H), 5.09 (d, 1H, *J*=12.6 Hz), 7.74 (d, 1H, *J*=12.6 Hz); ¹³C NMR (CDCl₃): δ 14.13, 48.2, 51.5, 55.6, 55.7, 60.9, 78.3, 78.7, 95.8, 96.4, 149.5, 168.9, 188.3; IR (neat): 3031, 2948, 2853, 2829, 1730, 1655 and 1593 cm⁻¹. Anal. Calcd for C₁₅H₂₅NO₇: C, 54.37; H, 7.60; N, 4.23. Found: C, 54.25; H, 7.63; N, 4.24.

4.1.6. 2-Diazo-5-[(3aS,4aS,6aR)-4-[(4S)-2,2-dimethyl-1.3-dioxolan-4-vl]-2,2-dimethyltetrahydro-3aH-[1,3]dioxolo[4,5-c]pyrrol-5-yl]-3-oxo-pentanoic acid ethyl ester 4c. To a stirred solution of 1,4-dideoxy-2,3,5, 6-O-isopropylidene-1,4-imino-D-talitol 3c (0.679 g, 0.00279 mol) and 3-oxo-pent-4-enoic acid ethyl ester (0.397 g, 0.00279 mol), in CH₂Cl₂ (15 mL), a solution of Et₃N (0.60 mL, 0.0042 mol) in CH₂Cl₂ (10 mL) was added dropwise and stirred for 30 min. Tosyl azide (0.840 g, 0.0042 mol) and a solution of Et₃N (0.60 mL, 0.0042 mol) in CH₂Cl₂ (10 mL) were then added dropwise at 0 °C. After the addition was complete, the solution was stirred at room temperature overnight. The solvent was evaporated and the residue gave, after flash chromatography (petroleum ether/ Et₂O, 8:2), the title compound 4c (0.580 g, 84% yield) as a yellow-brown oil: $[\alpha]_{D}^{25} + 81.6 (c \ 0.32, \text{CHCl}_{3}); ^{1}\text{H NMR}: \delta$ 1.29 (s, 3H); 1.33 (t, 3H, J = 6.6 Hz), 1.34 (s, 3H), 1.42 (s, 3H), 1.50 (s, 3H), 2.58 (dd, 1H, J = 5.7, 10.2 Hz), 2.83–2.89 (m, 2H), 2.95-3.13 (m, 2H), 3.30-3.39 (m, 2H), 3.76 (dd, 1H, J = 1.2, 8.1 Hz), 4.01 (dd, 1H, J = 1.8, 8.1 Hz), 4.20 (q, 1H, J=6.6 Hz), 4.31 (AB system, 2H), 4.36 (dd, 1H, J=3.9, 6.9 Hz), 4.57–4.63 (m, 1H); ¹³C NMR (CDCl₃): δ 24.8,

25.1, 26.3, 27.1, 38.5, 49.5, 58.9, 61.2, 66.1, 70.6, 76.0, 78.6, 81.8, 109.3, 112.7, 161.1, 191.5; IR (neat): 2985, 2936, 2134, 1715, 1654, 1455, 1372, 1300, 1211, 1078, 1159, 1053, 856 and 747 cm⁻¹. Anal. Calcd for $C_{19}H_{29}N_3O_7$: C, 55.46; H, 7.10; N, 10.21. Found: C, 55.71; H, 6.96; N, 10.44.

4.1.7. (6R,7S,8S)-1-Ethoxy carbonyl-6a-[(6aS)-2,2dimethyl-1,3-dioxolan-6a-yl]-2,2-dimethyl tetrahydro-7H-[1,3]-dioxolo 2-oxo-5-azonia-spiro[4,4]nonane-1ylide 5c. A solution of the diazoketoester 4c (0.517 g, 0.00126 mol) in CH₂Cl₂ (10 mL) was refluxed for 30' in the presence of Rh₂(OAc)₄ (0.015 g, 3 mol%). The solvent was evaporated and the residue gave, after flash chromatography $(Et_2O/MeOH/Et_3N, 5:5:0.1)$ the title compound 5c as amorphous grey solid (0.365 g, 76% yield). $[\alpha]_D^{25} - 47$ (c 0.37, CHCl₃); ¹H NMR: δ 1.32 (s, 3H), 1.34 (t, 3H, J= 6.9 Hz), 1.35 (s, 3H), 1.42 (s, 3H), 1.56 (s, 3H), 2.05–2.60 (m, 1H), 2.86 (quintet, 1H, J=4 Hz), 3.61 (t, 1H, J=7.8 Hz), 3.83 (dt, 1H, J = 2.4, 8.7 Hz), 3.94–4.37 (m, 7H), 4.41 (dd, 1H, J=7.2, 12.8 Hz), 4.96 (t, 1H, J=6.6 Hz), 5.09 (dt, 1H, J=0.6, 6.6 Hz); ¹³C NMR (CDCl₃): δ 14.6, 24.8, 26.6, 27.3, 33.4, 59.3, 65.2, 68.6, 76.7, 77.9, 80.8, 86.5, 105.8, 110.4, 113.7, 163.4, 179.4; IR (neat): 2926, 2854, 1652, 1607, 1460, 1419, 1376, 1263, 1210, 1078, 1055, 861and 844 cm⁻¹. Anal. Calcd for C₁₉H₂₉NO₇: C, 59.52; H, 7.62; N, 3.65. Found: C, 59.40; H, 7.83; N, 3.43.

4.1.8. rac-2-Diazo-5-(2-methoxymethyl-2-methyl-pyrrolidin-1-yl)-3-oxo-pentanoic acid ethyl ester 4d. TFA (0.48 mL, 0.006 mol) was added dropwise under nitrogen atmosphere to rac-N-Boc-2-methoxymethyl-2-methyl-pyrrolidine 3d (0.31 g, 0.0016 mol) at 0 °C. The mixture was stirred at room temperature until the disappearance of the substrate (¹H NMR) and formation of trifluoroacetate of the amine (about 1 h) and then the excess of TFA was evaporated in vacuum at room temperature. To a stirred solution of the residue and 3-oxo-pent-4-enoic acid ethyl ester (0.35 g, 0.0025 mol) in CH₂Cl₂ (10 mL), a solution of Et₃N (0.35 mL, 0.0025 mol) in CH₂Cl₂ (10 mL) was added dropwise and stirred for 30 min. To the reaction mixture, tosyl azide (0.36 g, 0.0025 mol) and a solution of Et₃N (1.0 mL, 0.0075 mol) in CH_2Cl_2 (10 mL) were added dropwise at 0 °C. After the addition was complete, the solution was stirred at room temperature overnight. The solvent was evaporated and the residue was purified by flash chromatography (Et₂O/petroleum ether/Et₃N, 6:4:0.1) to give the title compound 4d (0.30 g, 70% yield) as a yellow oil: ¹H NMR (CDCl₃): δ 0.96 (s, 3H), 1.33 (t, 3H, J= 7.2 Hz), 1.48–1.59 (m, 1H), 1.74 (quintet, 2H, J=7.2 Hz), 1.84–1.96 (m, 2H), 2.61 (q, 1H, J=7.8 Hz), 2.67–2.77 (m, 1H), 2.93-3.07 (m, 2H), 3.20 (s, 3H), 3.33 (s, 3H), 4.30 (q, 2H, J = 7.2 Hz); ¹³C NMR (CDCl₃): δ 14.3, 17.9, 21.2, 35.9, 40.3, 44.3, 51.6, 59.2, 61.3, 62.8, 76.1, 78.7, 161.3, 192.0; IR (neat): 2965, 2928, 2132, 1718 and 1656 cm⁻¹. Anal. Calcd for C₁₄H₂₃N₃O₄: C, 56.55; H, 7.80; N, 14.13. Found: C, 56.73; H, 7.77; N, 14.05.

4.1.9. Diazo-decomposition of 4d. To a refluxing solution of $Rh_2(OAc)_4$ (0.013 g, 3 mol%) in 30 mL of dry CH_2Cl_2 , a solution of **4d** (0.297 g, 0.001 mol) in dry CH_2Cl_2 (20 mL) was added dropwise over 30 min. After stirring for another 30 min at reflux, the reaction mixture was cooled and concentrated to give a 69:31 mixture of **5d** and **6d** (¹H

NMR). Purification by flash chromatography (Et₂O/ethyl acetate/Et₃N, 5:5:0.1; MeOH) gave 0.081 g of **6d** as a yellow oil and 0.180 g of inseparable mixture of ylides **5d** in the ratio of 44:56 (¹H NMR).

(*4E*)-5-(2-Methoxymethyl-2-methyl-pyrrolidin-1-yl)-3oxo-pent-4-enoic acid ethyl ester **6d** ¹H NMR (CDCl₃): δ 1.27 (t, 3H, *J*=7.2 Hz), 1.29 (s, 3H), 1.66–1.78 (m, 1H), 1.87–2.01 (m, 2H), 2.01–2.15 (m, 1H), 3.22–3.34 (m, 4H), 3.34 (s, 3H), 3.39 (s, 2H), 2.61 (q, 1H, *J*=7.8 Hz), 2.67–2.77 (m, 1H), 2.93–3.07 (m, 2H), 3.20 (s, 3H), 3.33 (s, 3H), 4.18 (q, 2H, *J*=7.2 Hz) 5.06 (d, 1H, *J*= 12.6 Hz), 7.78 (d, 1H, *J*=12.6 Hz); ¹³C NMR (CDCl₃): δ 14.2, 21.7. 23.7, 35.9, 48.6, 48.7, 59.4, 60.9, 65.1, 78.3, 98.0, 146.9, 169.1, 188.0; IR (neat): 2977, 2934, 2874, 1735, 1651, 1600 and 1556 cm⁻¹. Anal. Calcd for C₁₄H₂₃NO₄: C, 62.43; H, 8.61; N, 5.20. Found: C, 62.51; H, 8.60; N, 5.18.

4.1.10. 2-Diazo-5-[(3aR,4S,6aS)-(2,2-dimethyl-4-trityloxymethyl-tetrahydro[1,3] dioxolo[4,5-c]-pyrrol-5-yl)]-3-oxopentanoic acid ethyl ester 4e. To a stirred solution of 1,4dideoxy-2,3-di-O-isopropylidene-1,4-imino-5-O-trityl-L-lyxitol 3e (2.57 g, 0.0062 mol), and 3-oxo-pent-4-enoic acid ethyl ester (0.88 g, 0.0062 mol), in CH₂Cl₂ (15 mL), a solution of Et₃N (1.35 mL, 0.0096 mol) in CH₂Cl₂ (10 mL) was added dropwise and stirred for 30 min. The solvent was evaporated and to the residue a solution tosyl azide (1.21 g, 0.0062 mol) and Et₃N (1.35. mL, 0.0096 mol) in CH₃CN (15 mL) were added dropwise at 0 °C. After the addition was complete, the solution was stirred at room temperature overnight. The solvent was evaporated and the residue gave, after flash chromatography (petroleum ether/Et₂O, 8:2), the title compound 4e (1.83 g, 50% yield) as a yellow oil: $[\alpha]_D^{22}$ +52.2 (c 0.52, CHCl₃); ¹H NMR: δ 1.27 (s, 3H), 1.28 (t, 3H, J=7.2 Hz), 1.29 (s, 3H), 2.07 (dd, 1H, J=4.5, 10.8 Hz), 2.30 (q, 1H, J = 5.4 Hz), 2.40 (AB system, 1H), 2.92-2.99 (m, 2H),3.13-3.22 (m, 2H), 3.26 (dd, 1H, J=3.9, 9.9 Hz), 3.56 (dd, 1H, J=3.9, 9.9 Hz), 4.22 (dq, 2H, J=1.2, 7.2 Hz), 4.58 (dt, 2H, J=9.1, 10.8 Hz), 7.21–7.48 (m, 15H); ¹³C NMR (CDCl₃): δ 14.2, 25.5, 25.9, 37.9, 48.4, 59.4, 61.3, 62.3, 67.5, 75.9, 78.0, 80.8, 86.9, 110.9, 126.8, 127.6, 128.8, 144.2, 161.2, 191.5; IR (neat): 3085, 3057, 2982, 2936, 2800, 2132, 1714, 1654, 1490, 1448, 1371, 1299, 1209, 1172, 1152, 1117, 1068, 862, 763, 747, 706 and 633 cm⁻¹. Anal. Calcd for C₃₄H₃₇N₃O₆: C, 69.96; H, 6.39; N, 7.20. Found: C, 70.11; H, 6.25; N, 7.17.

4.1.11. Diazo-decomposition of 4e. A solution of the diazoketoester 4e (1.105 g, 0.00110 mol) in CH₂Cl₂ (10 mL) was refluxed for 30 min in the presence of Rh₂(OAc)₄ (0.015 g, 3 mol%). The solvent was evaporated and the residue gave, after flash chromatography (petroleum ether/Et₂O, 8:2), the unsaturated compound (4*E*)-5-[(3a*R*,4*S*,6a*S*)-(2,2-dimethyl-4-trityloxymethyl-tetrahydro-[1,3]dioxolo[4,5-*c*]pyrrol-5-yl)]-3-oxo-pent-4-enoic acid ethyl ester 6e (0.852 g, 81% yield), brown oil. $[\alpha]_D^{25}$ +47.2 (*c* 0.25, CHCl₃); ¹H NMR: δ 1.19 (s, 3H), 1.22 (t, 3H, *J*=7.2 Hz), 1.24 (s, 3H), 3.23–3.42 (m, 5H), (dd, 1H, *J*=3.9, 10.8 Hz), 3.70–3.78 (m, 1H), 4.13 (q, 1H, *J*=7.2 Hz), 4.69 (t, 1H, *J*=6.3 Hz), 4.75 (dt, 1H, *J*=2.4, 6.3 Hz), 5.01 (d, 1H, *J*=13.2 Hz), 7.20–7.45 (m, 15H), 7.91 (d, 1H, *J*=13.2 Hz); ¹³C NMR (CDCl₃): δ 14.0, 24.7, 25.8,

1465

60.8, 62.6, 63.9, 77.6, 79.0, 87.5, 112.7, 127.2, 127.8, 128.5, 143.4, 149.7, 168.7, 188.7; IR (neat): 3057, 3032, 2984, 2938, 1731, 1659, 1556, 1490, 1448, 1372, 1211, 1155, 1082, 982, 861, 764, 748, 706 and 648 cm⁻¹. Anal. Calcd for $C_{34}H_{37}NO_6$: C, 73.49; H, 6.71; N, 2.52. Found: C, 73.22; H, 6.50; N, 2.51.

Acknowledgements

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- Padwa, A.; Hasegawa, T.; Liu, B.; Zhang, Z. J. Org. Chem. 2000, 65, 7124.
- 18. Crystal data and structure refinement for ylide **5a**: single crystals were obtained by recrystallization from ethyl acetate/Et₂O, 10:1. The substance (C₂₃H₃₁NO₃, M_r =369.49) crystallized in the monoclinic space group $P2_1$, a=11.022(1) Å, b=7.793(1) Å,

c = 12.104(1) Å; $\alpha = 90^{\circ}$, $\beta = 90.46(1)^{\circ}$, $\gamma = 90^{\circ}$; V =1039.63(18) Å³, Z=2, $D_{calcd}=1.180 \text{ Mg/m}^3$, F (000)=400, T = 295(2) K, crystal size $0.33 \times 0.27 \times 0.010$ mm³, $\theta =$ $1.85-25.36^{\circ}$, limiting indices = -12 < =h < =13, -9 < =k < =9, -15 < =l < =14, reflections collected = 11503, independent reflections = 3602 [R_{int} = 0.0244], completness to θ = 25.36°: 97%, max. and min. transmission: 0.9923 and 0.9750, refinement method: full-matrix least-squares on F^2 , data = 3602, restrains = 32, parameters = 369, goodness-of-fit on $F^2 = 1.075$, final R indices $[I > 2\sigma(I)]$: $R^1 = 0.0348$, $wR^2 = 0.0774$, R indices (all data): $R^1 = 0.0504$, $wR^2 = 0.0846$, absolute structure parameter = -1.0(3), extinction coefficient = 0.018(3), largest diff. peak and hole=0.109 and $-0.116 \text{ e} \text{ Å}^{-3}$. Crystallografic data have been deposited with the Cambridge Crystallografic Data Center (Deposition number: CCDC 279550).



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Photooxygenations of 1-naphthols: an environmentally friendly access to 1,4-naphthoquinones

Oliver Suchard,^a Ronan Kane,^b Bernard J. Roe,^b Elmar Zimmermann,^c Christian Jung,^d Prashant A. Waske,^a Jochen Mattay^{a,*} and Michael Oelgemöller^{b,*}

^aOrganische Chemie I, Fakultät für Chemie, Universität Bielefeld, Postfach 10 01 31, D-33501 Bielefeld, Germany

^bSchool of Chemical Sciences and National Institute for Cellular Biotechnology, Dublin City University, Glasnevin, Dublin 9, Ireland

^cInstitut für Organische Chemie, Universität zu Köln, Greinstr. 4, D-50939 Köln, Germany

^dDeutsches Zentrum für Luft- und Raumfahrt e.V. (DLR), Solarforschung, Linder Höhe, D-51147 Köln, Germany

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Abstract—Dye sensitized photooxygenations of 1-naphthols were investigated with soluble and solid-supported sensitizers and moderate to excellent yields of the corresponding 1,4-naphthoquinones were achieved in relatively short irradiation times. The mild and environmentally friendly reaction conditions made this application particularly attractive for 'Green Photochemistry'. Consequently, the photooxygenation of 1,5-dihydroxynaphthalene was studied with non-concentrated and moderately concentrated sunlight and 5-hydroxy-1,4-naphthoquinone (Juglone) was obtained in yields up to 71%.

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1. Introduction

Naphthoquinone derivatives based on 5-hydroxy-1, 4-naphthoquinone (Juglone, **1**) represent an important class of natural products.¹ Additionally, Juglone serves as a valuable building block for the synthesis of biologically active quinonoid compounds (Chart 1),² and was thus selected by us as starting material for our ongoing photoacylation study.³ Most commonly, Juglone is synthesized from the cheap and commercially available 1,5-dihydroxynaphthalene by oxidation, but many of these thermal pathways suffer from severe disadvantages concerning yield, selectivity, sustainability, scale-up or reproducibility, respectively.⁴ Dye sensitized photooxygenations can serve as a versatile alternative and various examples involving 1-naphthols have been reported in the literature. $^{5-7}$

Due to the favorable absorption of most dyes within the visible spectrum, photooxygenation reactions have been subjected to concentrated sunlight and served as model systems for environmentally friendly and benign 'Green Photochemistry'.^{8–10} Recently, we have briefly reported on solar photooxygenations to Juglone using novel holographic mirror elements.^{8b} In this publication, we would like to present a comprehensive study on solar and artificial light induced photooxygenations of 1-naphthols and 1,5-dihydroxynaphthalene in particular.

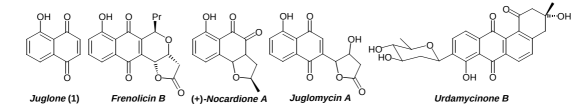


Chart 1. Quinonoid natural products synthesized from Juglone 1 (for more examples see Ref. 2).

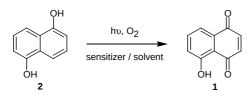
Keywords: Quinones; Photochemistry; Photooxygenations; Green chemistry.

^{*} Corresponding authors. Tel.: +49 521 106 2072; fax: +49 521 106 6417 (J.M.); tel.: +353 1 700 5312; fax: +353 1 700 5503 (M.O.); e-mail addresses: mattay@uni-bielefeld.de; michael.oelgemoeller@dcu.ie

2. Results and discussion

2.1. Experiments with artificial light

To find suitable reaction conditions for the solar chemical campaign, we have launched a detailed laboratory study with artificial light and selected the photooxygenation of 1,5-dihydroxynaphthalene **2** to Juglone **1** as model system (Scheme 1). A major disadvantage of the literature procedures was the usage of the hazardous solvents dichloromethane, acetonitrile or methanol, respectively,⁵ which needed to be replaced for a solar 'outdoor' application. In order to simplify the work-up procedure, we furthermore examined the usage of solid-supported sensitizers, in particular Sensitox[®] (rose bengal on Merry-field resin; RB_{MF})¹¹ and methylene blue on ion exchange resin (MB_{IE}).¹² Both materials can be easily removed by filtration and are, in principal, reusable.



Scheme 1. Photooxygenation of 1,5-dihydroxynaphthalene 2.

Following a standardized procedure, a 0.01 M solution of 1,5-dihydroxynaphthalene **2** was irradiated with a 150 W medium-pressure mercury lamp in the presence of a sensitizer while a gentle stream of oxygen was passed through the solution (Table 1). The progress of the reaction

 Table 1. Experimental data for the photooxygenations of 2 with artificial light (150 W medium-pressure mercury lamp)

Entry	Sens. ^a	Solvent	Time (h)	1 (%)
1	MB	MeOH	5	51
2	RB	MeOH	5	34
3	MB_{IE}	MeOH	5	43
4	RB _{MF}	MeOH	5	32
5	b	MeOH	5	2
6	MB	EtOH	5	54
7	RB	EtOH	5	37
8	MB_{IE}	EtOH	5	47
9	RB _{MF}	EtOH	5	32
10	b	EtOH	5	2
11	MB	<i>i</i> -PrOH	5	58
12	RB	<i>i</i> -PrOH	5	38
13	MB_{IE}	<i>i</i> -PrOH	5	46
14	RB _{MF}	i-PrOH	5	33
15	b	<i>i</i> -PrOH	5	2
16	MB	Acetone	5	48
17	RB	Acetone	5	71
18	MB_{IE}	Acetone	5	41
19	RB _{MF}	Acetone	5	68
20	b	Acetone	5	8
21	MB	CH ₂ Cl ₂ /MeOH ^c	5	51
22	RB	CH ₂ Cl ₂ /MeOH ^c	5	34
23	MB_{IE}	CH ₂ Cl ₂ /MeOH ^c	5	43
24	RB _{MF}	CH ₂ Cl ₂ /MeOH ^c	5	28
25	b	CH ₂ Cl ₂ /MeOH ^c	5	3

^a Sensitizers: methylene blue (MB), rose bengal (RB), methylene blue on ion exchange resin (MB_{IE}), rose bengal on Merryfield resin (Sensitox[®], RB_{MF}).

^b Without sensitizer.

^c CH₂Cl₂/MeOH (9:1).

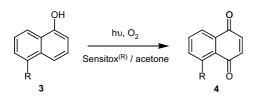
was followed by GC or TLC analysis. After 5 h, Juglone was isolated via column chromatography using chloroform as eluent or, more conveniently, via continuous extraction with *n*-hexane in a Soxhlet extractor.

In methanol, methylene blue was found to be the most effective sensitizer and the desired 1 was isolated in a reasonable yield of 51%. As would be expected for heterogeneous conditions, the yields for the solid-supported sensitizers were slightly lower with 43% (MB_{IE}) and 32% (RB_{MF}), respectively. With Sensitox[®], the characteristic orange color of Juglone became clearly visible after a relatively short irradiation time. After 5 h, TLC analysis showed no sign of sensitizer leaching. In contrast to the literature,¹² significant leaching was, however, observed for methylene blue on ion exchange resin as noticeable from the green color of the final reaction mixture. Similar preferences and yields were obtained when ethanol was used as solvent. Likewise, the photooxygenation proceeded satisfactory in isopropanol and 2 was isolated in yields of 33-58%. The best Juglone yields of 71 and 68% were obtained using acetone as solvent and rose bengal or Sensitox[®] as sensitizer (entries 17 and 19), respectively. In comparison to the irradiations in alcohols, methylene blue gave a somewhat lower yield of 48%. Due to the limited solubility of the diol 2 in pure dichloromethane, photosensitized oxygenations were conducted alternatively in a 9:1 mixture with methanol, and Juglone was formed in yields of 28-51%. Surprisingly, the product yields did not improve as would be expected from the longer ¹O₂ lifetime in this solvent mixture.¹³ Enhanced photobleaching of the dye and photodecomposition of 1 due to the rather harsh radiation emitted from the medium-pressure mercury lamp,^{5b} in combination with the formation of acid from the halogenated solvent, might explain this unexpected drop.

In the absence of sensitizer, **1** was formed in only small amounts of 2-3% in all alcoholic solvents and in the dichloromethane/methanol mixture. Solely the irradiation of **2** in pure acetone furnished Juglone in a significant yield of 8% (entry 20),¹⁴ and we tentatively postulate a type-I photooxidation for its formation as known for phenols.^{5d}

A scale-up of the photooxygenation to **1** was furthermore examined in acetone with Sensitox[®], and the concentration of **2** was stepwise increased in 0.01 mol/l intervals. Up to a concentration of **2** of 0.05 mol/l, complete conversions were achieved within 5 h and **1** was isolated in yields of 65–70%. At higher concentrations, prolonged irradiation times up to 10 h were required but **1** was still isolated in good yields of 63–68%. At a diol concentration of 0.1 mol/l, the reaction was stopped after 10 h. At this stage, GC analysis showed a conversion of ca. 80%. After work-up, Juglone was obtained in 55% yield (69% based on conversion).

The photooxygenation protocol was additionally applied to 1-naphthol **3a**, 1-acetoxy-5-hydroxynaphthalene **3b** and 5-acetamido-1-hydroxynaphthalene **3c** (Scheme 2; Table 2), respectively. 1-Naphthol **3a** readily gave 53% of 1,4-naphthoquinone **4a** when irradiated with a mediumpressure mercury lamp in acetone and in the presence of Sensitox[®]. In line with the literature, ^{15b} irradiation of **3b** under identical conditions furnished Juglone **1** in 68% yield. Obviously, the acetate-group is cleaved during the course of the reaction. In contrast, the related amide-linked compound **3c** readily gave the corresponding quinone **4c** in a good yield of 61%.^{15a}



Scheme 2. Photooxygenations of 3.

 Table 2. Experimental data for the photooxygenations of 3 with artificial light (150 W medium-pressure mercury lamp)

Entry	R	Sens. ^a	Solvent	Time (h)	4 (%)
1	H (3a)	RB _{MF}	Acetone	5	53 (4a)
2	OAc (3b)	RB _{MF}	Acetone	5	68 (1)
3	NHAc (3c)	RB _{MF}	Acetone	5	61 (4c)

^a Sensitizer: rose bengal on Merryfield resin (Sensitox[®], RB_{MF}).

Due to the absorption of Juglone within the emission spectra of the medium-pressure mercury lamp,¹⁶ we have conducted a series of experiments using a pair of 500 W halogen lamps (Table 3). Since solution purging with pure oxygen is furthermore problematic for industrial applications, we have examined its replacement with compressed air. Almost all experiments were run in non-hazardous isopropanol. Irradiations with pure oxygen readily furnished Juglone in yields of 25–70% after 5 h. Since the given set-up did not allow an even distribution of the solid-supported sensitizers within the reaction mixture, the experiments involving Sensitox[®] and methylene blue on ion exchanger resin (MB_{IF}) showed significantly lower conversions and yields. With compressed air, prolonged irradiation times of 10 h were required but the desired 1 was still obtained in fair to high yields of 21-71%. For laboratory purposes, we have furthermore modified the conditions reported by Cossy and Belotti for photooxygenations of 8-hydroxyquinolines.^{7b} Irradiation in a 9:1 mixture of dichloromethane and methanol for 2 h and in the presence of TTP as sensitizer yielded 1 in an excellent yield of 88% (entry 5). The yield

Table 3. Experimental data for the photooxygenations of 2 with artificial light (2×500 W halogen lamps)

Entry	Sens. ^a	Solvent	Gas	Time (h)	1 (%)
1	MB	<i>i</i> -PrOH	O ₂	5	69
2	RB	i-PrOH	O_2	5	70
3	MB_{IE}	i-PrOH	O_2	5	34
4	RB _{MF}	<i>i</i> -PrOH	O_2	5	25
5	TPP^{b}	CH ₂ Cl ₂ /MeOH ^c	O_2	2	88
6	MB	<i>i</i> -PrOH	Air	10	71
7	RB	i-PrOH	Air	10	55
8	MB_{IE}	<i>i</i> -PrOH	Air	10	45
9	RB_{MF}	i-PrOH	Air	10	21
10	TPP ^b	CH ₂ Cl ₂ /MeOH ^c	Air	3	78

^a Sensitizers: methylene blue (MB), rose bengal (RB), methylene blue on ion exchange resin (MB_{IE}), rose bengal on Merryfield resin (Sensitox[®], RB_{MF}), tetraphenylporphine (TPP).

^b TPP insoluble in *i*-PrOH.

^c CH₂Cl₂/MeOH (9:1).

was somewhat lower with 78% when air was used as oxygen source (entry 10). Noteworthy, this procedure represents the so far best synthetic pathway to Juglone.⁴

2.2. Solar chemical experiments

In July and August 2005, we have conducted a series of solar chemical experiments at Dublin City University (latitude 53°23'N, 6°15'W, 50 m above sea level). Due to the volatility and flammability of the solvent acetone, we have selected the less hazardous isopropanol for our campaign. Following this strategy, various solutions of 2 were exposed in a Schlenck-flask equipped with a cold finger and a reflux condenser to direct sunlight while the solution was purged with a gentle stream of air. All experiments went smoothly and gave satisfactory results in reasonable periods of time without any noticeable sideproducts (Table 4). The first run (I) was performed with soluble rose bengal under ideal solar conditions and Juglone was isolated in a moderate yield of 39% after 3.5 h of illumination. A somewhat lower yield of 30% of 1 was obtained when the reaction was repeated for 6.5 h during a partly sunny period (II). With soluble methylene blue as sensitizer (III), Juglone became available in 44% yield after 5.5 h of partly sunny weather. Likewise, Sensitox[®] was tested as a heterogeneous sensitizer (IV). Within $\frac{1}{2}$ h, the orange color of 1 became clearly visible and further intensified with progressing illumination. Due to the limited distribution of the solid sensitizer within the reaction mixture, 1 was obtained in just 19% yield after 6.5 h of perfect weather conditions. Noteworthy, all reactions described above could have been driven easily to high conversions with longer illumination times. Thus, the preliminary results obtained clearly indicate that the solar photosensitized oxygenation of 1,5-dihydroxynaphthalene opens a promising and environmentally friendly pathway to Juglone.

 Table 4. Experimental data for the solar photooxygenation reactions of 2 with non-concentrated sunlight

		Experiment				
	I	II	III	IV		
Date	12.07.2005	13.07.2005	25.07.2005	08.08.2005		
Scale						
2 (g)	0.56	0.56	0.56	0.56		
Sens. (g) ^a	0.05 (RB)	0.05 (RB)	0.05 (MB)	0.4 (RB _{MF})		
Solvent	i-PrOH	<i>i</i> -PrOH	<i>i</i> -PrOH	<i>i</i> -PrOH		
V(ml)	350	350	350	350		
Time						
IST ^b	14:15-	11:45-	10:45-	10:15-16:45		
	17:45	18:15	16:15			
Total (h)	3.5	6.5	5.5	6.5		
Weather	Sunny	Partly sunny	Partly sunny	Sunny		
Yield 1 (%)	39	30	44	19		

^a Sensitizers: rose bengal (RB), methylene blue (MB), rose bengal on Merryfield resin (Sensitox[®], RB_{MF}).

^b Irish summer time.

Further solar chemical experiments were performed with moderately concentrated sunlight at the solar chemical facility of the German Aerospace Center (DLR) close to Cologne/Germany (latitude $50^{\circ}51'N$, $7^{\circ}07'E$, 70 m above sea level).¹⁷ A parabolic trough collector designed for laboratory-scale (<500 ml) applications and equipped with

an aluminum mirror (aperture: 41×36 cm) was selected (Fig. 1).¹⁸ The reactor offers a geometric concentration factor of about 18 suns, but its efficiency is reduced in practice due to optical losses. Tracking of the sun is performed manually for the elevation and the azimuth every 15 min. The reaction mixture is pumped through the jacket of a Liebig condenser (diameter: 2.4 cm), which is placed in the focal line of the concentrator. Cooling water is passed through the inner tube of the condenser. Oxygen is added via a simple Y-connector, which limited its homogeneous distribution within the absorber tube.



Figure 1. Laboratory-scale parabolic trough reactor during the solar photooxygenation of 1,5-dihydroxynaphthalene 2 (the red color of the sensitizer rose bengal can be clearly seen).

In August and September 2003, three laboratory-scale experiments were conducted, and the progress of each reaction was followed by GC analysis versus tetradecane as internal standard. Due to its favorable solar sensitization efficiency and overall stability,^{8d} rose bengal was chosen as sensitizer. The experimental details and results from the solar chemical studies are summarized in Table 5.

The first run (V) was performed during a sunny period with 0.5 g of diol 2 and 0.05 g of rose bengal in 100 ml of isopropanol. The starting material was readily consumed and already after 40 min, GC analysis revealed complete conversion (>95%). During that time the reactor collected 0.07 mol of photons in the important absorption range of rose bengal between 500-600 nm.¹⁹ After work-up, the desired product 1 was obtained in 71% yield. For the second experiment (VI) under mostly sunny conditions, the amount of diol 2 was doubled to 1.0 g and after 2.5 h, GC analysis showed a constant value for Juglone 1 of 74% (Fig. 2). The collector received 0.16 mol of photons in the range of 500-600 nm,¹⁹ slightly more than double the amount as during the first experiment. After a total illumination period of 3.5 h, Juglone was obtained in a good yield of 69% (78% based on conversion). For the final experiment (VII), the solvent was replaced by methanol. Due to the less

 Table 5. Experimental data for the solar photooxygenation reactions of 2

 with moderately concentrated sunlight

	Experiment			
	V	VI	VII	
Date	15.08.2003	09.09.2003	11.09.2003	
Scale				
2 (g)	0.5	1.0	1.0	
Rose bengal (g)	0.05	0.01	0.1	
Solvent	<i>i</i> -PrOH	<i>i</i> -PrOH	MeOH	
V(ml)	100	100	100	
Time				
CEST ^a	14:20-16:50	13:45-17:15	10:15-14:45	
Total (h)	2.5	3.5	4.5 ^b	
Effective (h) ^c	2/3	2.5	_	
Weather	Sunny	Mostly sunny	Partly sunny	
Photons (mol) ^d	•			
Total	0.26	0.21	0.16	
Effective ^e	0.07	0.16		
Conversion (%) ^f	>95	88	86	
Yield 1 (%)	71 (75) ^g	69 (78) ^g	46 (54) ^g	

Central European summer time.

^b Stopped due to rainfall.

^c Time until conversion reaches an almost constant value.

^d Estimated amount of photons collected between 500-600 nm.¹⁸

^e Estimated amount of photons (500-600 nm) for effective illumination time.

^f Conversion of **2** as determined by GC analysis (vs tetradecane).

^g Yield based on conversion of **2**.

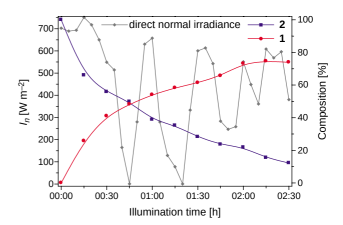


Figure 2. Direct normal irradiance (I_n) and product composition versus illumination time for the photooxygenation of 1,5-dihydroxynaphthalene 2 (Experiment VI).

favorable weather, the illumination time needed to be extended. After 4.5 h of partly sunny conditions, the reaction was stopped at 86% conversion due to beginning rainfall. At this stage the reactor had collected 0.16 mol of photons between 500–600 nm.¹⁹ After work-up, Juglone was isolated in a moderate yield of 46% (54% based on conversion).

3. Conclusion

In conclusion, photooxygenations of 1-naphthols to the corresponding 1,4-naphthoquinones can serve as a useful and environmentally friendly alternative to existing thermal processes. The solar chemical reaction of the cheap and commercially available 1,5-dihydroxynaphthalene can be easily performed with non-concentrated or concentrated

sunlight, and yields the valuable intermediate Juglone. Thus, a realization of Giacomo Ciamician's spectacular vision of 'the photochemistry of the future' (presented at the International Congress of Applied Chemistry in New York in 1912)²⁰ seems feasible.

4. Experimental

4.1. General methods

Melting points were measured on a Büchi B-540 apparatus and are uncorrected. ¹H and ¹³C NMR spectra were recorded on a Bruker Avance 400 spectrometer (400 and 100 MHz, respectively) using the solvent residual peak as internal standard. Chemical shifts (δ) are given in ppm, coupling constants (J) in Hz. MS spectra were recorded on a Finnigan MAT 8230 (EI) spectrometer. IR spectra were recorded as KBr discs on a Perkin-Elmer 298 infrared spectrophotometer, UV/vis spectra on a Perkin-Elmer Lambda 7 spectrophotometer using n-hexane (Janssen Chimica, spectrophotometric grade) as solvent. For combustion analysis a Heraeus CHN-O-Rapid Elemental Analyzer was used. GC analysis was performed on a Shimadzu GC-14A or a Hewlett-Packard GC 5890 Series II. A Hanau TQ-150 medium-pressure mercury lamp (150 W) or Armley 500 W halogen lamps $(2 \times 500 \text{ W})$ and immersion well reactors ($\lambda > 280$ nm) were used for irradiation experiments. TLC was carried out on Merck Kieselgel 60 F₂₅₄, column chromatography on silica gel (Macherey and Nagel) 230-240 mesh using chloroform or a 19:1 mixture of chloroform and methanol. 1,5-Dihydroxynaphthalene 2 was purified according to a modified procedure of Johnson and co-workers.²¹ 1-Acetoxy-5-hydroxynaphthalene **3b** was synthesized as reported by Becher et al.,²² 5-acetamido-1hydroxynaphthalene 3c via a method described by Jindal and co-workers.²³ Sensitox[®] was prepared with chloromethylated styrene-divinylbenzene copolymer (50-100 mesh, 1% cross-linked) according to Schapp et al.,11 methylene blue on ion exchange resin (Lewatit SC 104 or MonoPlus SP 112) according to Williams and co-workers.¹⁷ Solvents and reagents were commercially available and were used without further purification.

4.2. Irradiation and illumination experiments

4.2.1. Irradiations with artificial light. General procedure (medium-pressure mercury lamp). The naphthol (1.5 mmol) was dissolved in 150 ml of solvent. The sensitizer was added (MB: 10 mg; RB: 20 mg; MB_{IE}: 400 mg; RB_{MF}: 100 mg) and the solution was irradiated with a Hanau TQ-150 medium-pressure mercury lamp (150 W) for 5 h at room temperature while purging with a gentle stream of oxygen. Evaporated solvent was frequently refilled. The progress of the reaction was monitored by TLC or GC analysis. The reaction mixture was filtrated, the solvent removed in vacuum, and the residue was purified by column chromatography on silica or by extraction in a Soxhlet extractor with *n*-hexane. Experimental details are given in Tables 1 and 2.

4.2.1.1. 5-Hydroxy-1,4-naphthoquinone (Juglone, 1). Isolated by Soxhlet extraction with *n*-hexane. Orange solid,

mp: 152 °C (lit.:^{5a} 151–154 °C). ¹H NMR (400 MHz, CDCl₃): δ =6.94 (s, 2H), 7.27 (dd, 1H, *J*=2.2, 7.5 Hz), 7.60–7.65 (m, 2H), 11.90 ppm (s, 1H, OH). ¹³C NMR (100 MHz, CDCl₃): δ =114.0, 118.1, 123.5, 130.8, 135.5, 137.6, 138.6, 160.6, 183.2, 189.3 ppm. MS (EI, 70 eV): *m*/*z*=174 (M⁺, 100%), 146, 118, 90, 63, 39. IR (KBr): *ν*= 3400, 3058, 1662, 1641, 1590, 1448, 1289, 1225, 1151, 1098, 1081, 863, 827, 762, 703 cm⁻¹. UV/vis (*n*-hexane): λ_{max} =247.8, 318.0, 427.8 nm. Anal. Calcd for C₁₀H₆O₃: C 68.97, H 3.47. Found: C 68.25, H 3.70.

4.2.1.2. 1,4-Naphthoquinone (**4a**). Isolated by column chromatography using chloroform as eluent. Yellow-brownish solid, mp: 128 °C (lit.:²⁴ 128.5 °C). ¹H NMR (400 MHz, CDCl₃): δ =6.96 (s, 2H), 7.74 (m, 2H), 8.06 ppm (m, 2H). ¹³C NMR (100 MHz, CDCl₃): δ = 126.5, 132.0, 134.1, 138.8, 185.2 ppm. MS (EI, 70 eV): m/z=158 (M⁺, 100%), 130, 102, 76, 62, 50, 40. IR (KBr): ν =1660, 1587, 1331, 1302, 1146, 1115, 1059, 863, 771 cm⁻¹. UV/vis (*n*-hexane): λ_{max} =240.2, 245.3, 328.3 nm. Anal. Calcd for C₁₀H₆O₂: C 75.94, H 3.82. Found: C 75.55, H 3.91.

4.2.1.3. 5-Acetamido-1,4-naphthoquinone (4c). Isolated by column chromatography using chloroform as eluent. Yellow solid, mp: 170 °C (lit.^{14a} 172 °C). ¹H NMR (400 MHz, CDCl₃): δ =2.29 (s, 3H), 6.90 (d, 1H, *J*=10 Hz), 6.94 (d, 1H, *J*=10 Hz), 7.72 (dd, 1H, *J*= 8.4 Hz), 7.81 (dd, 1H, *J*=1.2, 8.4 Hz), 9.07 (dd, 1H, *J*=1.2, 8.4 Hz), 11.85 (s, 1H, NH). ¹³C NMR (100 MHz, CDCl₃): δ =25.8, 116.1, 122.1, 126.2, 132.3, 135.9, 138.1, 140.1, 141.5, 170.1, 184.7, 189.3 ppm. MS (EI, 70 eV): *m/z*=215 (M⁺), 173 (100%), 145, 117, 101, 91, 63, 43. IR (KBr): ν = 3477, 3211, 1707, 1666, 1646, 1609, 1580, 1496, 1408, 1302, 1264, 1159, 833, 766, 723 cm⁻¹. Anal. Calcd for C₁₂H₉N₁O₃: C 66.97, H 4.22, N 6.51. Found: C 66.51, H 4.39, N 6.70.

4.2.2. General procedure (halogen lamps). Five hundred and forty milligrams (3.5 mmol) of 1,5-dihydroxynaphthalene **2** were dissolved in 350 ml of solvent. The sensitizer was added (MB: 50 mg; RB: 50 mg; MB_{IE}: 400 mg; RB_{MF}: 400 mg; TPP: 20 mg) and the solution was irradiated (2×500 W halogen lamps) in a Schlenck-flask equipped with a cold finger and a reflux condenser for 2–10 h at room temperature while purging with a gentle stream of oxygen or air. Evaporated solvent was frequently refilled. The progress of the reaction was monitored by TLC analysis. The reaction mixture was filtrated the solvent removed in vacuum, and the residue was purified by column chromatography (SiO₂, CHCl₃) or by extraction in a Soxhlet extractor with *n*-hexane. Experimental details are given in Table 3.

4.2.3. Illuminations with sunlight. General procedure (non-concentrated sunlight). Five hundred and forty milligrams (3.5 mmol) of 1,5-dihydroxynaphthalene **2** were dissolved in 350 ml of isopropanol. The sensitizer was added (RB: 50 mg; MB: 50 mg; RB_{MF}: 400 mg) and the solution was exposed to direct sunlight in a Schlenck-flask equipped with a cold finger and a reflux condenser for 3.5–6.5 h while purging with a gentle stream of air. Evaporated isopropanol was frequently refilled and

the progress of the reaction was monitored by TLC analysis. The reaction mixture was filtrated, the solvent removed in vacuum, and the residue was purified by column chromatography (SiO₂, CHCl₃) or by extraction in a Soxhlet extractor with *n*-hexane. Experimental details are given in Table 4.

4.2.4. General procedure (concentrated sunlight). 1,5-Dihydroxynaphthalene **2** was dissolved in 100 ml of solvent. Rose bengal was added and the solution was exposed to moderately concentrated sunlight in a parabolic trough reactor for 2.5–4.5 h while purging with a gentle stream of oxygen. The progress of each reaction was monitored by GC analysis versus tetradecane as internal standard. The solvent was removed in vacuum, and the residue was purified by column chromatography (SiO₂, CHCl₃/MeOH=19:1). Experimental details are given in Table 5.

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1473

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Investigation of the active species in a Michael addition promoted by chirally modified tetrahydroborate

Susan Abraham and G. Sundararajan*

Department of Chemistry, Indian Institute of Technology Madras, Chennai 600036, India

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Dedicated to Professor Robert Moss on the occasion of his 65th birthday

Abstract—For the first time, asymmetric 1,4-addition of various malonates to enones has been carried out using tetrabutylammoniumtetrahydroborate (TBATB) in the presence of a chiral ligand. The Michael adducts were formed in reasonably good yields (61–67%) with moderate ee's at 0 °C. ¹¹B NMR spectroscopic studies explain this unexpected reactivity through the predominant formation of an aminodiol modified borate complex in the presence of a hydride acceptor.

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1. Introduction

The Michael addition, being one of the most important C–C bond-forming reactions, has attracted much attention toward the development of enantioselective catalytic procedures in recent years.¹ The current literature abounds with many reports on enantioselective Michael addition catalyzed by chiral complexes of Ru,^{2a} Co,^{2b} Rh,^{2c} Ni,^{2d} Cu,^{2e} Zn,^{2f} Cd,^{2g} Al^{2h} and other heterobimetallics.³ Thus far, however, there are not many reports on boron catalyzed asymmetric Michael reactions.⁴

We have earlier shown that chiral aminodiol, (R,R)-1, in combination with LiAlH₄ or lanthanum–sodium, can be effectively used for asymmetric Michael additions.⁵ As an extrapolation of these findings, we decided to investigate the application of chirally modified borohydrides in promoting the Michael reaction of α , β -unsaturated ketones. Although chirally modified boron has been employed to promote many asymmetric processes^{6a} such as Diels–Alder,^{6b} allylation^{6c} and aldol^{6d} reactions, little has been reported on the chirally modified tetrabutylammoniumtetrahydroborate (TBATB) system in such reactions. However, it is known that chirally modified borohydrides are effective in asymmetric reduction processes⁷ but, in contrast, to chiral auxiliaries of lithium aluminum hydrides that promote asymmetric Michael addition,^{3c,5} chirally modified borohydrides are not known to assist such reactions.^{3c}

Herein, we give a brief report on the results of Michael additions promoted by a mixture of TBATB/(R,R)-1 in THF and attempts to rationalize our observations.

2. Results and discussion

The required ligand (R,R)-1 was prepared from the reaction of (R)-styrene oxide with benzylamine.^{5c} First, a control reaction was performed to study the reduction pattern of cyclic enone with TBATB in the presence of (R,R)-1. As expected the products were alcohol and ketone resulting from an initial 1,4-addition of hydride across the enone to give the enolate, that converts into the ketone (via the enol) and gets reduced further. These findings are in agreement with other literature reports.⁸

Subsequently, (R,R)-1 in combination with TBATB was used as a promoter in the Michael addition of cyclic enones with diethyl malonate⁹ (Eq. 1). The corresponding Michael adducts from cyclohexenone and cyclopentenone were formed in good yields and with moderate enantioselectivities. The reduced products of cyclic enone were also obtained in minor amounts along with the Michael adduct. In all these cases the yields of Michael adducts remained fairly constant. The results are summarized in Table 1.

Keywords: Borohydride; Michael addition; Malonates.

^{*} Corresponding author. Tel.: +91 44 2257 4213; fax: +91 44 2257 4202; e-mail: gsundar@iitm.ac.in

Table 1. Michael addition of various malonates to cyclic enones

$$\begin{array}{c} O \\ (n) \\ ($$

Entry	Enone	Michael donor	Time (h)	Product distribution (%) ^a			% ee of 4^{b}
				4	5	6	
1	2a	3a	7	4a =62	5a =24	6a =14	4a = 35
2	2a	3b	7	4b =64	5a =22	6a =14	4b=40
3	2a	3c	7	4c = 61	5a =25	6a =12	4c=31
5	2b	3a	7	4d=67	5b =22	6b =11	4d=42
6	2b	3b	7	4e = 65	5b =25	6b =10	4e = 45
7	2b	3c	7	4f =63	5b =22	6b =15	4f=39

^a Determined by HPLC.

^b %ee was determined by HPLC connected to a Chiracel OD. The absolute configuration in all cases were determined by comparison of optical rotation and was found to be *R*.

Table 2. Michael addition of benzylidineacetophenone with malonates

	O Ph + Ph	$\begin{array}{c} \text{CO}_2\text{R} \\ \text{CO}_2\text{R} \end{array} \begin{array}{c} 1 & 20 \ \% \left[(R,R) \\ 2 & 2 \end{array} \right] \\ \text{CO}_2\text{R} \end{array} \begin{array}{c} 2 & \text{aq. H}_2\text{O}_2, 1 \end{array}$	1), TBATB]	Ph CO ₂ R CO ₂ R	OH + Ph Ph	+ Ph Ph	
	-	3a : R = Et 3b : R = ⊁Bu 3c : R = Bn		8a : R = Et 8b : R = <i>t</i> -Bu 8c : R = Bn	9	10	
Entry	Enone	Michael donor	Time (h)	Product of	distribution (%) ^a		
				8	9		10
1 2 3	7	3a 3b 3c	7 7 7	8a =62 8b =64 8c =61	22 22 23		16 14 16

^a Isolated yields.

In a similar manner benzylidineacetophenone reacts with malonates to give 1,4-adducts with moderate enantioselectivity, along with minor amounts of reduced products. The results are summarized in Table 2.

Thus, in the presence of (R,R)-1 and TBATB a mixture of enone and malonate gives reasonable yields of the Michael adducts in moderate enantiomeric excess, suggesting the formation of a chirally modified borohydride, an observation that warranted further scrutiny.

To gain better insight into these findings, we chose to study the reaction by ¹¹B NMR spectroscopy. The ¹¹B NMR spectrum of a solution containing (*R*,*R*)-1 and TBATB in a 2:1 ratio gave a quintet centered at -57.4 ppm indicating the presence of free borohydride.¹⁰ To this mixture, the addition of cyclohexenone in portions of 0.5 equiv, promoted the formation of a singlet centered at -15.7 ppm alongside the quintet that could be attributed to a free tetraborate anion having a tetrahedral structure, ^{11,6b} and with 2.1 equiv of cyclohexenone the quintet disappeared completely leaving a sharp singlet at -15.7 ppm (Fig. 1). In the absence of cyclohexenone, a mixture of 1 and TBATB, showed the quintet persisting in the ¹¹B NMR spectrum even after an overnight reflux. Thus, the need for a hydride acceptor to initiate the formation of the tetraalkoxyborate becomes clear.

When the same experiment was performed with cyclohexanone, the quintet did not disappear completely, even after addition of many equivalents of the ketone, indicative of a relatively slow hydride transfer to cyclohexanone. Nevertheless, the appearance of a sharp singlet at -15.9 ppm could be seen here as well. Also as expected, the ¹¹B NMR spectrum of a solution containing (*R*,*R*)-1, TBATB and diethyl malonate in the absence of cyclohexenone gave no other signal than the quintet. Not so surprising also was the sudden appearance of the singlet at -15.5 ppm beside the quintet when a small amount of cyclohexenone was added to this solution at

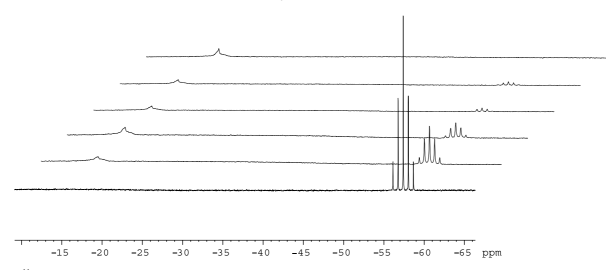
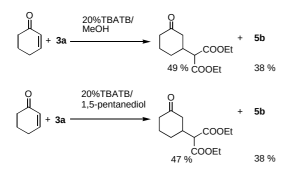


Figure 1. ¹¹B NMR spectra of 1-TBATB and varying equivalents of cyclohexenone (a) 0 equiv (b) 0.5 equiv (c) 1.0 equiv (d) 1.5 equiv (e) 2.0 equiv (f) 2.1 equiv.

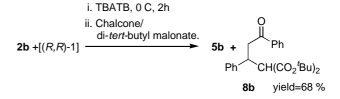
ambient conditions. Thus, the combined role of 1 and cyclohexenone in the generation of the singlet around -15 ppm in ¹¹B NMR needs to be appreciated.

To probe the effect of any interaction of the nitrogen atom in the backbone of (R,R)-1 with the boron, the corresponding borate complex was generated from methanol or pentanediol by reacting with TBATB in the presence of the cyclic enone (Scheme 1). The borate complexes generated here, were effective in the Michael addition with product yields hovering around 47–49%, comparable to the earlier observations with (R,R)-1 as the chelating ligand, pointing to an unlikely role for the nitrogen atom in the scaffold of 1. Predictably, the ¹¹B NMR spectral studies of these systems were highly reminiscent of the earlier results.



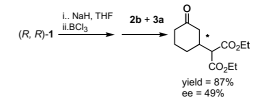
Scheme 1. Michael addition in the presence of achiral alcohols without any ligating atom in the backbone.

In order to confirm the need for a hydride acceptor in the formation of the active catalyst, we deliberately added cyclohexenone as a sacrificial hydride acceptor to the (R,R)-1-TBATB mixture prior to the addition of chalcone as the actual Michael acceptor. Thus, a solution of TBATB, (R,R)-1 and cyclohexenone in the ratio 1:2:2 was stirred for a period of 2 h, to which a mixture of chalcone and malonate was added. As expected, we could get the Michael adduct corresponding to chalcone and di-*tert*-butyl malonate as the major product along with the reduction products of cyclohexenone (Scheme 2).



Scheme 2. Use of cyclohexenone as a sacrificial hydride acceptor.

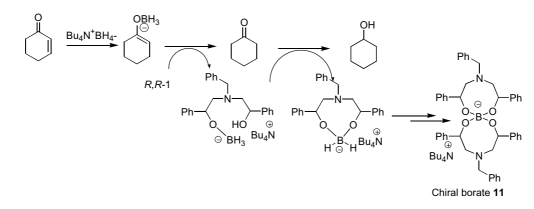
We also examined an alternate possibility for generating the borate, by reacting the disodiated (R,R)-1 with BCl₃, to promote the Michael reaction involving cyclohexenone and diethyl malonate which, as expected, gave the Michael adduct in 87% yield with 49% ee (Scheme 3). It was also not surprising that the ¹¹B NMR spectrum of sodium aminodiolate and BCl₃ gave a peak at -16 ppm, implicating strongly the formation of a tetraborate species as in earlier cases.



Scheme 3. Asymmetric Michael addition with chiral borate generated from *R*,*R*-1 and BCl₃.

2.1. Suggested mechanism for the chirally modified borate promoted asymmetric Michael addition

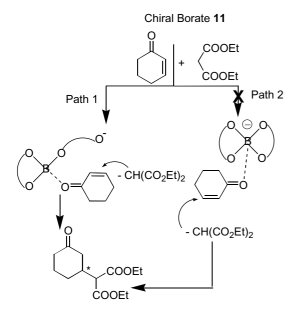
On viewing the above observations collectively, a plausible mechanism for the enantioselective Michael addition emerges (Scheme 4). The less acidic (R,R)-1 does not react with TBATB to form the borate complex upon simple



Scheme 4. Suggested mechanism for the formation of the chirally modified borate in the asymmetric Michael addition.

mixing. However, when the enone is added, an initial hydride transfer from TBATB takes place; the enolate so generated undergoes a protic quench with (R,R)-1 that converts it to the ketone. Stepwise mediation of boron leads to the eventual formation of the bischelate complex, the catalytically active species in the Michael reaction.

Clearly, the moderate (but tangible!) enantioselectivities observed in all these cases suggest probable coordination of cyclohexenone to a chirally modified borate complex. The possibilities could then be, either a tetracoordinate boron with one arm of the aminodiol acting as a detachable tether or a pentacoordinate hypervalent boron, the half life of which is very short on the NMR timescale¹² (vide Scheme 5). Further NMR spectroscopic investigations performed to detect the catalytically active species involved did not offer positive clues even at low temperatures $(-60 \,^{\circ}\text{C})$ when only signals at -57 and -15 ppm could be observed. Since we have no clear proof by boron NMR spectroscopy or otherwise for the occurrence of pentacoordinate boron, we tend to support the former mechanism. The mechanism also explains the fact that the combined yields of the reduced products in the reaction equal a stoichiometric transfer of four hydrides from the borate (Table 1).



Scheme 5. Possible modes of activation of enone.

3. Conclusion

In conclusion, we have shown for the first time that chirally modified TBATB–aminodiol is effective in the Michael addition of α , β -unsaturated ketones with various Michael donors with moderate enantioselectivity. Evidence from ¹¹B NMR spectroscopic studies and other experiments support the formation of chiral tetrahedral borate from aminodiol and borohydride in the presence of a hydride acceptor.

4. Experimental

4.1. General experimental procedures

All operations were carried out under an atmosphere of dry, oxygen-free nitrogen employing vacuum or Schlenk line techniques, unless otherwise noted. Nitrogen was purified by passage through columns of MnO anchored on silica gel catalyst and 4 Å molecular sieves. Solid organometallic compounds were transferred in an argonfilled glove bag. All glassware, syringes and needles were oven dried at 140 °C and cooled to room temperature under nitrogen before use. Tetrahydrofuran was freshly distilled from sodium/benzophenone ketyl under nitrogen atmosphere. Cyclohexenone, di-tert-butylmalonate, di-ethylmalonate, di-benzylmalonate and (R)styreneoxide were purchased from Lancaster synthesis and cyclopentenone was purchased from Aldrich and used as received. Tetrabutylammoniumtetrahydroborate (TBATB) was prepared from tetrabutylammoniumhydrogensulphate according to the literature procedure. ¹H NMR (400 MHz) and ¹³C NMR (100 MHz) spectra were recorded in CDCl₃ at ambient temperature with TMS as the internal standard and ¹¹B NMR (135 MHz) spectra were recorded with boric acid as an external standard using AV400 Bruker spectrometer (BF₃·Et₂O signal appeared at -19.38 ppm). Analytical HPLC was performed with Shimadzu LC-8A HPLC instrument equipped with RI detector and chiralcel OD column. Optical rotations were measured on a JASCO DIP-370 Polarimeter. Melting points were determined in a capillary and are uncorrected. Mass spectra were recorded on a Q-TOF mass spectrometer.

4.2. General reaction procedure of malonate addition on conjugate alkenones

To a solution of TBATB (56 mg, 0.214 mmol) in dry THF (3 mL) was added a solution of aminodiol (150 mg, 0.432 mmol) in THF (3 mL). The mixture was stirred under moisture free nitrogen atmosphere for 30 min at 0 °C, then a mixture of α , β -unsaturated ketone (1.06 mmol) and Michael donor (1.06 mmol) were added. The mixture was stirred for 7 h. The reaction was then quenched by the addition of 3% aqueous hydrogen peroxide (2 mL) and 10% aqueous sodium hydroxide (1 mL). The mixture was stirred for 2 h, the layers were separated and the aqueous phase was extracted with CH₂Cl₂ (2×20 mL). The combined organic layer was dried over anhydrous sodium sulfate, concentrated and the crude product was purified by column chromatography (silica gel 60–120, acetone/hexane 10:90). NMR spectra are identical to those previously reported.⁵

%ee's were determined by HPLC (Daicel Chiralcel OD, 2.0:98.0, 2-propanol/hexane, flow rate = 0.5 mL/min, 254 nm; For example, **4e** had retention times of t_1 =28.6 (*S*), t_2 =36.5 (*R*)). The absolute configuration was established by comparison to the literature.¹³

Acknowledgements

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Tetrahedron

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Fluorinated alcohol directed formation of dispiro-1,2,4,5-tetraoxanes by hydrogen peroxide under acid conditions

Katja Žmitek,^a Stojan Stavber,^a Marko Zupan,^a Danièle Bonnet-Delpon^b and Jernej Iskra^{a,*}

^aLaboratory of Organic and Bioorganic Chemistry, 'Jožef Stefan' Institute and Faculty of Chemistry and Chemical Technology of University of Ljubljana, Jamova 39, 1000 Ljubljana, Slovenia

^bBIOCIS UMR 8076 CNRS, Faculté de Pharmacie, Université Paris-Sud, rue J. B. Clément, Châtenay-Malabry, 92296 Cedex, France

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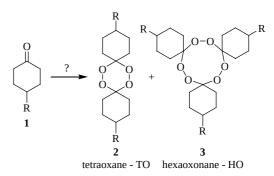
Abstract—The oxidative system MTO/30%H₂O₂/HBF₄/fluorous alcohol is promising for the selective synthesis of biologically important antimalarial dispiro-1,2,4,5-tetraoxanes by direct acid-catalysed cyclisation of various 4-substituted cyclohexanones (1, R = Me, Et, tBu, Ph, $COOEt, CF_3$). The role of the substitutent at the 4-position was important in the selectivity of formation of tetraoxane (2, TO) with respect to hexaoxonane (3, HO). By the use of fluorinated alcohols and under the right reaction conditions, tetraoxanes 2 were selectively formed and synthesised in 46-86% isolated yield from 4-substituted cyclohexanones 1.

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1. Introduction

Malaria is a major communicable disease that causes or contributes to approximately 3 million deaths per year.¹ Malaria mortality has increased in recent years, mainly due to the parasites growing resistance to classical alkaloidal drugs.² Artemisinin and its semi synthetic derivatives as well as synthetic endoperoxides have emerged as potent non-alkaloid compounds that are active against chloroquine-resistant seves of Plasmodium.³ Dispiro-1,2,4,5tetraoxanes, having two endoperoxide bonds, are effective and inexpensive antimalarial agents.^{4,5} The easiest path for their synthesis is the acid-catalysed peroxidation of a carbonyl compound with hydrogen peroxide. However, this method can be employed only for selected substrates, since it results in a mixture of compounds with dispiro-1,2,4,5-tetraoxane (TO) being only one among many.⁶ For example, acid-catalysed cyclisation of 4-methylcyclohexanone with H₂O₂ gave only 1,2,4,5,7,8-hexaoxonane (HO), while the 4-tert-butyl analogue gave a mixture of both cyclic peroxides (Scheme 1).⁷ An alternative route for the selective synthesis of tetraoxanes is ozonolysis of cycloalkylidenecycloalkanes,8 enol ethers,9 and O-methyl oximes,^{7,10} but yields are generally low.

Improving the selectivity of the cyclisation by using H_2O_2 and acid would be particularly advantageous due to its simplicity. Fluorinated alcohols (hexafluoro-2-propanol-HFIP and trifluoroethanol—TFE)¹¹ are known activators of hydrogen peroxide in various oxidation reactions: oxidations of sulfides,¹² epoxidations¹³ and Baeyer–Villiger oxidations.¹⁴ This activation is attributed to the high hydrogen bond donor strength of TFE and HFIP.¹⁵ We have already applied fluorinated alcohols in the methyltrioxorhenium-catalyzed oxidation of 4-methylcyclohexanone with 30% H₂O₂ and the reaction led to the formation of gem-dihydroperoxide. Based on this result, we were able to report the first one-pot synthesis of non-symmetric TO from simple ketones and 30% H₂O₂.¹⁶ During this study, we selectively transformed 4-methylcyclohexanone into the corresponding TO without the formation of



Scheme 1.

Keywords: Hydrogen peroxide; Fluorinated alcohols; Cyclic peroxides; Tetraoxane; Antimalarials; Oxidation.

^{*} Corresponding author. Tel.: +386 14773631; fax: +386 12519385; e-mail: jernej.iskra@ijs.si

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HO. It was already known from the literature that the selectivity of the cyclisation is very dependent on the structure of the ketone and on reaction conditions.

Presented herein is our investigation into the factors that govern the formation of the two cyclic peroxides—tetraoxanes and hexaoxonanes from 4-substituted cyclo-hexanones 1 (Fig. 1) and our results on the selective direct formation of dispiro-1,2,4,5-tetraoxanes 2.

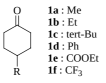
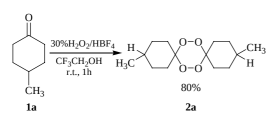


Figure 1.

2. Results and discussion

Initially, we studied the role of reaction conditions (fluorinated alcohol, acid...) on the conversion of 4-methylcyclohexanone 1a to TO 2 and HO 3. In the first experiment, we mixed equimolar amounts of 1a, 30% H₂O₂ and HBF₄ in TFE for 1 h at room temperature. A white precipitate formed immediately and after a filtration, the main product formed was dispiro-1,2,4,5-tetraoxane 2a, which was further isolated in 80% yield (Scheme 2). We did not observe the formation of hexaoxonane 3a but we were able to show the presence of trace amounts of ε-caprolactone. Selective formation of TO 2a in TFE was surprising, since in previous study using acetonitrile as a solvent, only HO was isolated. The NMR spectrum of 2a shows dynamic properties with a characteristic broad signal at 3.05 ppm for two protons (C1-H and C9-H), which agrees with previously published data.¹⁷ Although two possible signals were expected for the CH₃ group there was only one sharp doublet at 0.92 ppm, indicating that the methyl groups are either in the equatorial or axial position.





To gain greater insight into the role of acid in acid-catalysed peroxidation, different acids were tested. In acid-catalysed reactions of **1a** with H_2O_2 in TFE, strong acids (H_2SO_4 , HCl, *p*-toluenesulfonic acid) acted as HBF₄ and the only cyclic peroxide formed was TO **2a**. Peroxidation catalysed with a weaker acid (acetic acid) did not yield any cyclic peroxide, whereas with intermediate acids, trifluoroacetic and phosphoric acid, a mixture of both cyclic peroxides was formed (**2a**:**3a**=3:1 and 1:3.7, respectively).

Next, we took 4-ethyl- **1b** and 4-*tert*-butyl cyclohexanone **1c**. Using reaction conditions with HBF_4 in TFE as for **1a**,

the 4-ethyl derivative 1b was transformed to TO 2b, (isolated in 69% yield, Table 1, entry 2). Surprisingly, with the *tert*-butyl derivative 1c selectivity was lost and cyclisation afforded both cyclic peroxides-TO 2c and HO 3c (Table 1, entry 3). Due to the similar solubility of the two cyclic peroxides, we could only separate a mixture of both 2c and 3c in a ratio of 55/23 by column chromatography (63% yield). To obtain only tetraoxane 2c, we looked for selective conditions that avoid the formation of the trimeric peroxide. First, we applied methyltrioxorhenium-MTO,¹⁸ one of the catalysts with the broadest range of action and the one that has already been applied in the synthesis of non-symmetric TOs.¹⁶ The use of MTO (0.1 mol%) brought some advantage (Table 1, entry 4), but further improvement was obtained by the use of a more diluted solution (0.5 M solution of 1c instead of 1 M) that gave TO 2c as the only cyclic peroxide formed (entry 6), isolated by column chromatography in 64% yield. Reaction at 0 °C produced a complex reaction mixture with HO 3c being the major reaction product (entry 7). Hexafluoro-2propanol (HFIP) is a better hydrogen bond donor than TFE and is therefore a more activating solvent for oxidation reactions.¹¹ The reaction of **2c** in HFIP at 0 °C was stopped after 5 min and TO 2c was the only cyclic peroxide formed and was accompanied by ε -caprolactone 4c (entry 8). Column chromatography gave 40% isolated yield of TO 2c. This reaction performed at room temperature yielded only ε-caprolactone **4c** (96% isolated yield, Scheme 3).

4-Phenylcyclohexanone 1d was the next substrate on which we investigated the effect of reaction conditions on the selectivity of acid-catalysed peroxidation. As in the case of 1c, phenyl-derivative 1d gave mixture of TO 2d and HO 3d in the MTO-catalysed reaction in TFE at standard room temperature. The isolated reaction mixture was separated by column chromatography and resulted in a mixture of a TO 2d and HO 3d (25% yield) in a ratio 1:1.8 as determined by NMR spectroscopy (Table 2, entry 10). HFIP had a beneficial effect on the selective formation of TO as exclusive formation of TO 2d was observed at room temperature (46% isolated yield, entry 12). We also found a similar reactivity pattern for ethyl 4-oxocyclohexanecarboxylate **1e** (entries 13–15) where again the use of more activated conditions (HFIP, room temperature) was needed to achieve selective cyclisation to TO 2e (50% isolated yield; entry 15). The important role of the hydrogen bond donor strength of the solvent is evident in the case of 4-trifluoromethylcyclohexanone 1f. In TFE, the reaction was directed towards the formation of the trimeric-HO 3f (entry 16), while the use of HFIP as a solvent resulted in the exclusive formation of TO 2f (entry 17). After work-up, the isolated yield was 54% for HO and 86% for TO. We could conclude from the preceding experiments that the structure of ketone had a profound effect on the formation of cyclic ketones (TO vs HO). However, only TO can be selectively formed by tuning of the reaction conditions, where the presence of MTO, room temperature and more activating solvent (HFIP) plays an important role.

The sensitivity of this reaction on the type of acid and the substituent on the position 4 of the cyclohexanone ring pose the question whether TO and HO are formed independently or whether they could be inter-converted during the reaction

Table 1. The effect of reaction conditions on the MTO-catalysed oxidative cyclisation of 4-substituted cyclohexanones 1c-f in fluorinated alcohols^a

Entry	Substrate	Reaction conditions	Relative ratio (%) ^b					
			1	2	3	Other	2/3	Yield of 2/3 (%) ^c
1	1a (4-Me)	TFE, 1 M, rt, 1 h ^d	6	85		9 ^e	>100	80
2	1b (4-Et)	TFE, 1 M, rt, 1 h ^d	4	87		9 ^e	>100	69
3	1c (4- <i>t</i> Bu)	TFE, 1 M, rt, 1 h ^d	13	55	23	9 ^e	2.39	63
4		TFE, 1 M, rt, 1 h	11	73	16	_	4.56	63
5		TFE, 1 M,, rt, 1 h ^f	44	Trace	47	9	< 0.05	30
6		TFE, 0.5 M, rt, 1 h	9	77	_	14 ^e	>100	64
7		TFE, 0.5 M, 0 °C, 15 min	16	23	35	26 ^g	0.66	48
8		HFIP, 0.5 M, 0 °C, 5 min	_	52	_	48 ^e	> 100	40
9		HFIP, 0.5 M, rt, 5 min	_	_	_	100 ^e	_	
10	1d (4-Ph)	TFE, 1 M, rt, 1 h	_	12	23	65 ^g	0.52	25
11		HFIP, 0.5 M, 0 °C, 1 h	12	25	13	50 ^g	1.92	26
12		HFIP, 0.5 M, rt, 5 min	_	69	_	31 ^e	>100	46
13	1e (4-COOEt)	TFE, 1 M, rt, 1 h	_	32	38	30 ^g	0.84	17
14		HFIP, 1 M, 0 °C, 1 h	_	41	43	16 ^g	0.95	14
15		HFIP, 0.5 M, rt, 5 min	5	86	_	9 ^e	>100	50
16	1f (4-CF ₃)	TFE, 1 M, rt, 1 h	23	_	61	16 ^g	< 0.01	54
17		HFIP, 0.5 M, rt, 5 min	3	90	—	7 ^g	>100	86

^a Reaction conditions: 1c, 1 equiv 30% H₂O₂, 1 equiv HBF₄, 0.1 mol% MTO.

^b Ratio determined by NMR spectroscopy.

^c Isolated yield of the mixture 2/3 or pure compounds 2 or 3 after column chromatography.

^d Reaction without MTO.

^e 4-Substituted-ε-caprolactone 4.

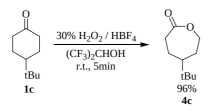
^f H₃PO₄ used as acid.

^g Mixture of hydroperoxides.

process. This subject was already debated in the seventies where it was argued that HOs are kinetically controlled products that can be converted to TOs.¹⁹ Further studies show that TOs are thermodynamically more stable, but which of them is the kinetically preferred product depends on several reaction rates.²⁰

Using both cyclic peroxides 2c and 3c, we made crosscheck experiments to determine their stability under the reaction conditions imposed on the synthesis of TO (with HBF₄ in TFE) and HO (with H₃PO₄ in TFE). The isolated TO 2c was stable in the presence of weak and strong acid in TFE, while HO 3c was stable only in the presence of a weaker acid (Table 2). In contrast, in the presence of a strong acid (HBF₄), HO 3c decomposes into a mixture of hydroperoxide products with a small amount of ketone 1c and lactone 4c as determined by ¹H NMR spectra of the crude product mixture.

Further, we made experiments under conditions as reported in the literature for conversion of HO into TO with perchloric acid in acetic acid.¹⁴ The result was that only 5% of TO **2c** together with 10% of lactone **4c** were formed. Similarly, when using CF₃-substituted HO **3f** we did not detect any conversion of HO to TO; instead **3f** was decomposed into different products. These results indicate that HO is not directly converted in high yield into TO with



Scheme 3.

acid in fluorous alcohol and furthermore, it decomposes under the reported reaction conditions. This implies that the point of decision on the reaction path for the formation of tetraoxane versus hexaoxonane should be made prior to the cyclisation step and as noted by McCullough et al., is dependent on the relative rates of several equilibria between the ketone and precursors of cyclic peroxides.²⁰

3. Conclusion

We have investigated what effect the reaction parameters have on the selectivity of the cyclisation of ketones to tetraoxanes with H_2O_2 under acid conditions. By choosing the appropriate reaction conditions we found that it is possible to select for the formation of dispiro-1,2,4,5tetraoxane over its trimer analogue—hexaoxonane. In particular, we found the role of fluorinated alcohols (TFE and HFIP) to be essential. We conclude that for the preferred formation of dispiro-1,2,4,5-tetraoxane, a higher temperature, a more activating solvent (HFIP>TFE), the presence of a catalyst (MTO) and substrate concentration have a profound influence. Also the effect of the structure of the ketone should not be overlooked. Table 3 gives a summary of the results and reaction conditions needed for selective formation of tetraoxanes **2**.

Furthermore, formation of TO and HO is competitive and it is unlikely that they could be directly inter-converted under

Table 2. Stability of TO and HO under reaction conditions

Substrate	Acid	Reaction conditions	
TO 2c	$\begin{array}{c} HBF_4\\ H_3PO_4\\ HBF_4\\ H_3PO_4\end{array}$	H ₂ O ₂ , MTO, TFE, rt, 15 min	No conversion
TO 2c		H ₂ O ₂ , MTO, TFE, rt, 1 h	No conversion
HO 3c		H ₂ O ₂ , MTO, TFE, rt, 15 min	Decomposition
HO 3c		H ₂ O ₂ , MTO, TFE, rt, 1 h	No conversion

 $\begin{array}{l} \textbf{Table 3. Overview of reaction conditions needed for selective formation of tetraoxane 2 by acid-catalysed peroxidation of cyclohexanones 1 with 30\% H_2O_2 in fluorinated alcohols \\ \end{array}$

Cyclohexanone	Reaction conditions ^a	Yield of $2 (\%)^{b}$
4-Me 1a	TFE, rt, 1 h	80
4-Et 1b	TFE, rt, 1 h	69
4- <i>t</i> -Bu 1c	0.1 mol% MTO, TFE (0.5 M), rt,	64
4-Ph 1d	15 min 0.1 mol% MTO, HFIP, rt, 5 min	46
4-COOEt 1e	0.1 mol% MTO, HFIP, rt, 5 min	50
4-CF ₃ 1f	0.1 mol% MTO, HFIP, rt, 5 min	86

^a Reaction conditions: 1: H_2O_2 :HB8₄=1:1:1(1M).

^b Isolated yield of TO 2 after column chromatography.

the reaction conditions employed in this synthesis. This study opens the way for the selective preparation of TOs by acid-catalysed oxidative cyclisation directly from ketones and 30% hydrogen peroxide. TOs thus obtained will be evaluated as antimalarials.

4. Experimental

4.1. General

Cyclohexanones 1 and methyltrioxorhenium were obtained from commercial sources and were used as received. Trifluoroethanol (TFE), 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP) and other solvents were distilled before use. ¹H and ¹³C spectra were obtained using a Varian Unity-300 spectrometer with TMS and CDCl₃ as standards. Melting points were determined on a Büchi 535 apparatus and were not corrected. Electron-spray mass spectra (8 kV spray needles, CsI or NH₄OAc) were acquired on an AutoSpec hybrid spectrometer. Elemental analyses were performed at the Microanalytisches Labor Pascher (Germany).

Caution! Although we have encountered no difficulties in working with these tetraoxanes, we advise routine precautions (shields, fume hoods, avoidance of transition metal salts) since organic peroxides are potentially hazardous.

4.1.1. Reaction procedure for synthesis of tetraoxanes 2a and 2b. First, $30\% H_2O_2$ (0.23 mL) and $54\% HBF_4$ solution in Et₂O (0.28 mL) were dissolved in TFE (2 mL). Substrate 1a or 1b (2 mmol) was then added and stirred at room temperature for 1 h. Reaction mixture was filtered and precipitate purified by column chromatography (SiO₂, CH₂Cl₂/petroleum ether=4:1) and obtained:

3,12-Dimethyl-7,8,15,16-tetraoxadispiro[5.2.5.2]-hexadecane (**2a**) (205 mg, 80%); white solid, mp 70–71 °C (lit.,⁷ 71–72 °C); ν_{max} (KBr) 1433, 1253, 1095, 1040, 970, 890 cm⁻¹; $\delta_{\rm H}$ (CDCl₃) 0.92 (6H, d, J=6.3 Hz), 1.06–1.34 (4H, m), 1.38–1.85 (12H, m), 3.05 (2H, br s); $\delta_{\rm C}$ (CDCl₃) 21.35, 21.44, 29.02 (br), 30.00 (br), 30.41 (br), 31.43 (br), 31.64, 31.73, 108.12, 108.16; *m*/*z* (ESI) 389 (M+Cs⁺).

3,12-Diethyl-7,8,15,16-tetraoxadispiro[5.2.5.2]-hexadecane (**2b**) (197 mg, 69%) (Found: C, 67.59; H, 9.89. $C_{16}H_{28}O_4$ requires C, 67.57; H, 9.92%); white solid, mp 111–113 °C; ν_{max} (KBr) 1420, 1325, 1145, 1030, 910, 880 cm⁻¹; $\delta_{\rm H}$ (CDCl₃) 0.88 (6H, t, *J*=7.1 Hz), 1.05–1.35 (10H, m), 1.35–1.90 (10H, m), 3.05 (2H, br s); $\delta_{\rm C}$ (CDCl₃) **4.1.2. Reaction procedure for the synthesis of tetraoxane 2c.** MTO (0.5 mg), 30% H_2O_2 (0.23 mL) and 54% HBF_4 solution in Et_2O (0.28 mL) were dissolved in TFE (2 mL). Substrate **1c** (2 mmol) was added and stirred at room temperature for 15 min. A typical isolation procedure followed by purification by column chromatography (SiO₂, CH₂Cl₂/petroleum ether=4:1) yielded:

3,12-Di-tert-butyl-7,8,15,16-tetraoxadispiro[5.2.5.2]-hexadecane (**2c**) (219 mg, 64%); white solid, mp 190–192 °C decomp. (lit.,⁷ 196–198 °C); ν_{max} (KBr) 1435, 1365, 1190, 1063, 1055, 935, 905 cm⁻¹; $\delta_{\rm H}$ (CDCl₃) 0.86 (18H, s), 1.03–1.52 (10H, m), 1.58–1.95 (6H, m), 3.17 (2H, br s); $\delta_{\rm C}$ (CDCl₃) 22.81 (br), 23.09 (br), 27.57, 27.61, 29.66 (br), 31.95 (br), 32.32, 47.40, 47.54, 108.11; *m*/*z* (ESI) 358 (M+NH₄⁺).

4.1.3. Reaction procedure for the synthesis of hexaoxonane 3c. MTO (0.5 mg) and 30% H₂O₂ (0.23 mL) and H₃PO₄ (0.14 mL) were dissolved in TFE (2 mL). Substrate **1c** (2 mmol) was added and stirred at 23 °C for 1 h. After typical isolation procedure and purification by column chromatography (SiO₂, CH₂Cl₂) was obtained 3,12,20-tritert-butyl-7,8,15,16,23,24-hexaoxatrispiro[5.2.5.2.5.2]tetracosane (**3c**) (102 mg, 30%); white solid, mp 194– 196 °C decomp. (lit.,⁷ 195–196 °C); ν_{max} (KBr) 1440, 1365, 1195, 1125, 1080, 1065, 930, 910 cm⁻¹; δ_{H} (CDCl₃) 0.86 (27H, 2 s), 1.05–1.80 (21H, m), 2.18–2.42 (6H, m); δ_{C} (CDCl₃) 23.23, 23.52, 23.64, 23.67, 23.70, 27.57, 27.60, 27.65, 29.07, 32.32, 32.52, 32.55, 47.23, 47.29, 47.52, 107.48, 107.55, 107.84; *m*/z ESI 528 (M+NH⁴₄).

4.1.4. Reaction procedure for the synthesis of lactone 4c. MTO (0.5 mg), 30% H₂O₂ (0.23 mL) and 54% HBF₄ solution in Et₂O (0.28 mL) were dissolved in HFIP (4 mL). Substrate **1c** (2 mmol) was added and stirred at room temperature for 5 min. After a typical isolation procedure pure *4-tert-butyl-ε-caprolactone* **4c** was obtained as colorless oil (326 mg, 96%) and identified by comparison with literature data:²¹: $\delta_{\rm H}$ (CDCl₃) 0.89 (9H, s), 1.26–1.56 (3H, m), 1.96–2.12 (2H, m), 2.51–2.62 (1H, m), 2.70 (1H, ddd, J=1.3, 7.3, 14.0 Hz), 4.17 (1H, dd, J=10.1, 12.8 Hz), 4.34 (1H, ddd, J=1.9, 5.9, 12.8 Hz); $\delta_{\rm C}$ (CDCl₃) 23.59, 27.29, 30.08, 32.88, 33.26, 50.60, 69.02, 177.76.

4.1.5. Reaction procedure for the synthesis of tetraoxanes 2d, 2e and 2f. MTO (0.5 mg), $30\% \text{ H}_2\text{O}_2$ (0.23 mL) and 54% HBF₄ solution in Et₂O (0.28 mL) were dissolved in HFIP (4 mL). Substrate 1d or 1e (2 mmol) was added and stirred at room temperature for 5 min. After typical isolation procedure, tetraoxane 2 was purified by column chromatography (SiO₂, CH₂Cl₂/petroleum ether=9:1) and obtained:

3,12-Diphenyl-7,8,15,16-tetraoxadispiro[5.2.5.2]-hexadecane (2d) (176 mg, 46%) (Found: C, 74.51; H, 7.30. $C_{24}H_{28}O_4 \times 1/4H_2O$ requires C, 74.59; H, 7.48%); white solid, mp 209 210.5 °C decomp.; ν_{max} (KBr) 1470, 1420, 1230, 1050, 1040, 930, 920, 910 cm⁻¹; δ_{H} (CDCl₃) 1.46– 2.02 (14H, m), 2.63 (2H, m), 3.31 (2H, br s), 7.30 (10H, m); $\delta_{\rm C}$ (CDCl₃) 29.66 (br), 31.88 (br), 43.54, 107.85, 126.28, 126.81, 128.45, 145.78; *m*/*z* ESI 398 (M+NH₄⁺).

Diethyl 7,8,15,16-tetraoxadispiro[5.2.5.2]-hexadecane-3,12-dicarboxylate (**2e**) (185 mg, 50%) (Found: C, 57.76; H, 7.64. $C_{18}H_{28}O_8$ requires C 58.05, H 7.58%); white solid, mp 125–129 °C; ν_{max} (KBr) 1705, 1430, 1305, 1245, 1180, 1165, 1120, 1050, 930, 910 cm⁻¹; $\delta_{\rm H}$ (CDCl₃) 1.26 (6H, t, J=7.1 Hz), 1.40–2.05 (14H, m), 2.35–2.48 (2H, m), 2.85 (2H, br s), 4.1 (4H, q, J=7.1 Hz); $\delta_{\rm C}$ (CDCl₃) 14.16, 23.71 (br), 24.60 (br), 28.06 (br), 30.28 (br), 41.43, 41.58, 60.39, 107.48, 174.49; *m*/z ESI 390 (M+NH⁴₄).

3,12-Bis(trifluoromethyl)-7,8,15,16-tetraoxadispiro[5.2.5.2]hexadecane (**2f**) (245 mg, 86%) (Found: C, 45.88; H, 5.18. C₁₄H₁₈F₆O₄ requires C, 46.16; H, 4.98); white solid, mp 102.5–105 °C; ν_{max} (KBr) 1440, 1350, 1330, 1265, 1240, 1190, 1155, 1120, 1080, 1055, 1020, 975, 930, 920, 880 cm⁻¹; $\delta_{\rm H}$ (CDCl₃) 1.43–1.75 (8H, m), 1.75–2.02 (6H, m), 2.02–2.21 (2H, m), 3.20 (2H, br s); $\delta_{\rm C}$ (CDCl₃) 20.48 (br), 20.97 (br), 27.72 (br), 30.02 (br), 40.93 (q, J(C,F)=27 Hz), 41.00 (q, J(C,F)=27 Hz), 107.30, 127.23 (q, J(C,F)=278 Hz).

4.1.6. Reaction procedure for the synthesis of hexaoxonane **3f.** MTO (0.5 mg), 30% H₂O₂ (0.23 mL) and 54% HBF₄ solution in Et₂O (0.28 mL) were dissolved in TFE (2 mL). Substrate **1c** (2 mmol) was added and stirred at room temperature for 1 h. After a typical isolation procedure and column chromatography (SiO₂, CH₂Cl₂/petroleum ether=4:1) was obtained *3*,*12*,20-*tris*(*trifluoromethyl*)-*7*,8,*15*,*16*,23,24-*hexaoxatrispiro*[5.2.5.2.5.2]-*tetracosane* (**3f**) (196 mg, 54%) (Found: C, 45.75; H, 4.79. C₂₁H₂₇F₉O₆ requires C, 46.16; H, 4.98%); white solid, mp 128–132 °C; ν_{max} (KBr) 1460, 1400, 1370, 1340, 1290, 1270, 1195, 1165, 1130, 1100, 1070, 960, 940, 890 cm⁻¹; $\delta_{\rm H}$ (CDCl₃) 1.3–1.5 (3H, m), 1.5–1.8 (9H, m), 1.8–2.2 (9H, m), 2.2–2.5 (6H, m); $\delta_{\rm C}$ (CDCl₃) 21.21 (br), 21.54 (br), 27.19 (br), 27.27 (br), 30.42 (br), 40.55 (q, J(C,F)=27 Hz), 106.84, 106.87, 106.91, 127.45 (q, J(C,F)=277 Hz).

4.1.7. Stability of peroxide 2c. Compound 2c (51 mg, 0.15 mmol), 54% HBF₄ solution in Et₂O (41 μ L, 0.3 mmol) (85% H₃PO₄ (20 μ L, 0.3 mmol), respectively), 30% H₂O₂ (34 μ L, 0.3 mmol) and MTO (0.08 mg, 0.3 μ mol) were mixed with TFE (0.6 mL) and stirred for 15 min (1 h, respectively). CH₂Cl₂ (10 mL) was added and washed with H₂O (10 mL) and saturated solution of NaHCO₃ (10 mL). Organic phase was dried with Na₂SO₄, solvent was evaporated and crude product was obtained (49 mg, 48 mg, respectively). The product was analyzed by ¹H and ¹³C NMR spectroscopy with PhCF₃ as internal standard and only signals of TO **2c** were observed.

4.1.8. Stability of peroxide 3c in the presence of H_3PO_4 .

Compound **3c** (51 mg, 0.1 mmol), 85% H_3PO_4 (20 µL, 0.3 mmol), 30% H_2O_2 (34 µL, 0.3 mmol), MTO (0.08 mg, 0.3 µmol) were mixed with TFE (0.6 mL) and stirred for 1 h. CH₂Cl₂ (10 mL) was added and washed with H₂O (10 mL) and saturated solution of NaHCO₃ (10 mL). Organic phase was dried with Na₂SO₄, solvent evaporated and 46 mg of product was obtained. The product was analyzed by ¹H and ¹³C NMR spectroscopy with PhCF₃ as internal standard and only signals of HO **3c** were observed.

4.1.9. Stability of peroxide 3c in the presence of HBF₄. Compound 3c (51 mg, 0.1 mmol), 54% HBF₄ solution in Et₂O (41 μ L, 0.3 mmol), 30% H₂O₂ (34 μ L, 0.3 mmol), MTO (0.08 mg, 0.3 μ mol) were mixed with TFE (0.6 mL) and stirred for 15 min. CH₂Cl₂ (10 mL) was added and washed with H₂O (10 mL) and saturated solution of NaHCO₃ (10 mL). The organic phase was dried with Na₂SO₄, solvent evaporated and 51 mg of product was isolated. A crude product was analyzed by ¹H and ¹³C NMR spectroscopy with PhCF₃ as internal standard and by comparison with spectra of known compounds. The product was composed of HO 3c (27%), ketone 1c (3%), lactone 4c (7%) and mixture of hydroperoxides (69%) with singlets at 9.4 ppm in ¹H NMR spectra.

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The synthesis of bis(oligophenyleneethynylenes): novel potential nonlinear optical materials

David J. Armitt* and Geoffrey T. Crisp

School of Chemistry and Physics, University of Adelaide, Adelaide, SA 5005, Australia

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Abstract—Various functionalised phenyleneethynylene dimers 10 and trimers 12 were synthesised by palladium-catalyzed Sonogashira methodology. These dimers and trimers were coupled to 1,8-diido-10-methoxyanthracene to generate bis(oligophenyleneethynylenes) 17 and 18. Preliminary results towards the construction of both phenyleneethynylene and phenylenevinylene hybrid motifs are presented. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

The effects produced by a nonlinear optical (NLO) response in a bulk material have applications in optical switching, modulation, amplification, beam steering, wavelength filters and image processing.¹ The most important contributions to this activity come from the second and third order optical susceptibilities, $\chi^{(2)}$ and $\chi^{(3)}$, and can be described on the molecular level by the first and second order hyperpolarizabilities β and γ . In order to achieve large $\chi^{(2)}$ and $\chi^{(3)}$, high values of β and γ and a high density of nonlinear optical chromophores is desired, and since the polarization of a molecule is a vector quantity, the alignment of molecular dipoles which reinforce each other is also important.

The design of NLO materials continues to present a significant challenge to organic chemists.² It is recognized that attributes which enhance β include high polarizability, large anharmonicity and extensive electron delocalization.¹ In order to exhibit high $\chi^{(2)}$, the molecules must also be noncentrosymmetric, and generally, the organic compounds which have shown the most promise are donor–acceptor molecules possessing conjugated spacers with a low HOMO–LUMO band-gap. However, the structural requirements for materials having significant γ and $\chi^{(3)}$ are less well understood; although a high degree of conjugation is again desirable, a non-centrosymmetric geometry is unimportant.³

Oligo- and poly(phenylenevinylene) molecules (OPVs and PPVs) are a well-known class of organic NLO materials, and have some of the highest $\chi^{(2)}$ and $\chi^{(3)}$ values recorded.⁴ However, their alkynyl analogues, oligo(phenyleneethynylene)s (OPEs), have not been as thoroughly assessed for NLO activity, partly because of their poor solubility. Due to the absence of *E*–*Z* photoisomerization, OPEs offer the potential advantage of durability over their double bond counterparts,⁵ but those previously studied have generally shown a lower NLO response in comparison,⁶ which has usually been ascribed to a less effective electron delocalization along the OPE chain.⁵

Since the extent of conjugation is dependant on orbital overlap, coplanarity of the aryl moieties of the chain is an important factor for high β and γ .⁷ Arylacetylenes are known to have a very low barrier (<1 kcal mol⁻¹) for rotation around their sp–sp² bonds,⁸ hence OPEs exist in a conformational equilibrium of planar and various twisted forms.⁹ It has been shown that properties such as fluorescence emission and $\chi^{(3)}$ can be altered or enhanced through coplanarity enforced by π -stacking¹⁰ and hydrophobic/hydrophilic interactions¹¹ in Langmuir films, and by metal–metal bridging in related systems.¹² It is our aim to explore this theme by restricting the rotation of OPEs by covalent linking,¹³ hydrogen bonding or steric bulk.

Earlier we reported the synthesis of the thiacyclophane 1 (Fig. 1) as well as some cyclophane precursors¹⁴ and compared their absorption and emission spectra. However, the X-ray crystal structure of 1 and subsequent molecular modelling revealed that a consequence of the length of the 'arms' of the cyclophane was to induce some twisting of the OPE chains and this made the synthesis of

Keywords: Nonlinear optics; Sonogashira coupling; Alkynes.

^{*} Corresponding author. Tel.: +61 8 8303 5712; fax: +61 8 8303 4358; e-mail: david.armitt@adelaide.edu.au

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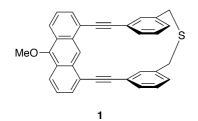


Figure 1. A previously studied thiacyclophane.

dithiacyclophanes extremely challenging. We wished to prepare extended analogues of the precursors to thiacyclophane **1** with a view to exploring the effects of hydrogen bonding and steric bulk in the aryl spacer 'arms' on inducing coplanarity of the arylacetylene spacers. Herein we report the preparation of two bis(OPEs), after investigating various approaches for extension of OPE chains.

2. Results and discussion

The arylacetylene building blocks **2** (Fig. 2) required to generate the OPEs have been reported previously, or are commercially available.¹⁵ In general, they were synthesized by palladium-catalyzed coupling of trimethylsilylacetylene (TMSA) with the corresponding aryl iodides, followed by protodesilylation under mild conditions.¹⁶ The basic repeating unit of the OPE chains, the iodide **3a**,¹⁷ was obtained in 65% overall yield from methyl *o*-anthranilate¹⁸ by following established procedures for similar monomers¹⁹ (Scheme 1). Alternatively, the triflate **3b** could be used.²⁰

The OPE segments could be constructed in one of two ways: (a) the suitably functionalized monomers could be sequentially added to the anthracene template, extending the bis(OPE) one unit at a time; or (b) OPEs could be synthesized independently, then attached to the anthracene template. The ester side-arm acts as a handle that allows for further elaboration to groups, which may restrict the rotation of the aryl rings through hydrogen bonding or steric bulk, as well as to increase the solubility of the bis(OPE) products.²¹ Monomer units containing only one ester side-arm were utilized for the development of OPE and bis(OPE) synthesis since the precursors were readily available.

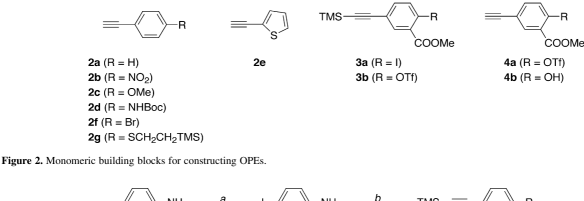
Based on previous reports, route (a) was initially investigated.¹⁴ Protodesilylation of **3b** with potassium fluoride in methanol gave the terminal alkyne **4a** in quantitative yield; these conditions proved more efficient than either potassium carbonate–methanol or TBAF. Coupling of **4a** with 5^{22} using piperidine as a base, even when diluted with DMF, afforded the unexpected product **7**, which represents an example of an unusually facile Pd-catalyzed aryl amination reaction²³ (Scheme 2). It was believed that the formation of **7** was a consequence of the extended conjugation of the desired intermediate **6**. However, further experiments with different primary and secondary amines revealed that this reaction was specific for piperidine. The ditriflate **6** was isolated in 47% yield when the base was changed to triethylamine.

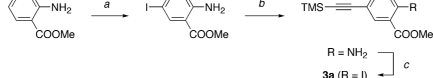
Disappointingly, the reaction of **5** with the phenolic alkyne $4b^{24}$ gave a complex mixture of products, which were not separable by chromatography, either directly or after attempted derivatization of the phenolic groups.²⁵

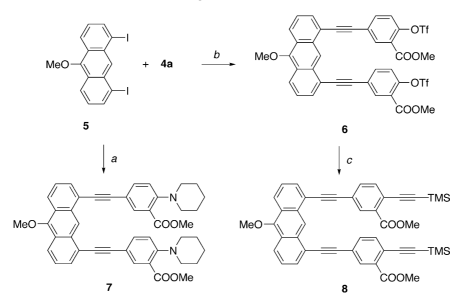
A longer route for chain extension involving alternating addition of the alkynyl and aryl moieties was then explored. Reaction of **6** with an excess of TMSA afforded the disilane **8** in 74% yield; however, deprotection of **8** gave low yields of the corresponding unstable terminal bis-alkyne (Scheme 2).

The alternative route (b) to bis(OPE) formation proved to be more efficient. Sonogashira coupling of the arylacetylenes 2a-2g with 3a afforded the dimers 9a-9g in 58–100% yields (Scheme 3).²⁶ Removal of the TMS groups was readily achieved with potassium carbonate–methanol to yield the terminal alkynes 10a-10g without need for purification.²⁷

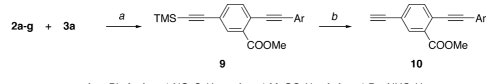
Further coupling of **10a–10e** with **3a** gave the trimers **11a–11e** in good yields (Scheme 4), which again were





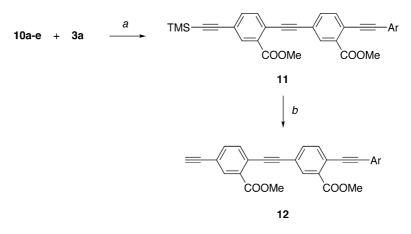


Scheme 2. Conditions: (a) Pd(PPh₃)₄, CuI, piperidine, DMF, 83% (7); (b) Pd(PPh₃)₄, CuI, Et₃N, DMF, 47% (6); (c) TMSA, Pd(PPh₃)₄, CuI, 74%.



a: Ar = Ph, **b**: Ar = 4-NO₂C₆H₄, **c**: Ar = 4-MeOC₆H₄, **d**: Ar = 4-BocNHC₆H₄ **e**: Ar = 2-thienyl, f: Ar = 4-BrC₆H₄, **g**: Ar = 4-(TMSCH₂CH₂S)C₆H₄

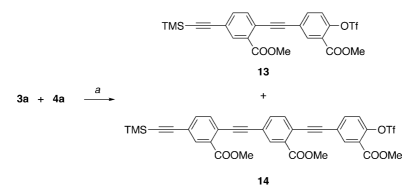
Scheme 3. Conditions: (a) Pd(PPh₃)₂Cl₂, CuI, 89% (9a); 77% (9b); 99% (9c); 58% (9d); 80% (9e); 92% (9f); 100% (9g); 54% (9h); (b) K₂CO₃, MeOH, 100% (10a); 78% (9b); 86% (10c); 95% (10d); 97% (10e); 91% (10f); 93% (10g).



Scheme 4. Conditions: (a) Pd(PPh₃)₂Cl₂, CuI, 74% (11a); 76% (11b); 59% (11c); 63% (11d); 61% (11e); (b) K₂CO₃, MeOH, 100% (12a); 76% (12b); 95% (12c); 87% (12d); 87% (12e).

deprotected under mild conditions. In principle, this iterative method could be employed to generate even longer OPEs. It was found that this approach to OPE synthesis, where the chains were elaborated from the functionalized end group,^{18,26a} was more facile than when the direction of chain growth was reversed.

The dimer 13 was formed by the coupling of monomeric units 3a and 4a (Scheme 5), which also generated a significant amount of trimer 14. The dimer was treated with potassium carbonate-methanol, but partial hydrolysis of the triflate was observed along with deprotection of the alkyne. Dimer 13 and trimer 14 are potentially useful building



Scheme 5. Conditions: (a) Pd(PPh₃)₂Cl₂, CuI, 32% (13); 18% (14).

9h (R = TMS) 10h (R = H)

Scheme 6. Conditions: (a) 3a, Pd(PPh₃)₂Cl₂, CuI, 54%; (b) KF, MeOH, 76%.

blocks for the rapid construction of soluble long chain OPEs.

It was believed that the presence of an ester in the *ortho*position may have contributed to the unusually facile hydrolysis of the triflate.²⁸ This hypothesis was tested by the synthesis and deprotection of the dimer **9h** (Scheme 6), which lacks the ester in close proximity. Removal of the TMS group could be performed with potassium fluoridemethanol without concomitant sulfonate cleavage.

The synthesis of hybrid compounds containing both OPE and OPV motifs, such as **15** (Fig. 3), would provide an interesting new class of potential NLO materials. However, many of the procedures widely used for the preparation of stilbenes²⁹ require strongly basic conditions, which could cause unwanted silyl deprotection of the arylalkyne building blocks. Thus, mild methods for the selective preparation of *E*-stilbenes were explored.

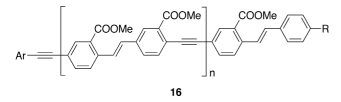


Figure 3. A hypothetical hybrid OPE-OPV structure.

The Heck reaction³⁰ has been utilized for the formation of a new bond between two sp²-carbon atoms under mild conditions. The triflate **3b** was reacted with methyl acrylate under the conditions described by Cabri³¹ to afford the alkynylcinnamate **16a** (Scheme 7); only the *E*-isomer was detected by ¹H NMR. Next, **3b** was reacted with styrene under identical conditions to give a mixture of compounds from which the *E*-stilbene **16b** was isolated in 29% yield. Finally, 4-nitrostyrene³² was treated with **3b**; however, in

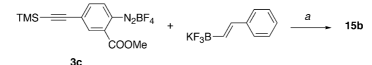
this case, an inseparable 3:1 mixture of regioisomers was obtained where the major component was the desired E-stilbene, and the minor component was the corresponding 1,1-disubstituted alkene.

OTI

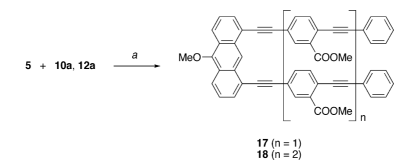
Scheme 7. Conditions: (a) Pd(OAc)₂, dppp, Et₃N, 47% (15a); 29% (15b).

The Suzuki–Miyaura coupling of organic halides and triflates with boronic acids or esters³³ usually requires the presence of a base. Recently Genêt et al. have demonstrated that aryldiazonium salts undergo a rapid palladium-catalyzed coupling reaction with potassium alkenyltrifluoroborates under neutral conditions.³⁴ Hence, the aryldiazonium tetrafluoroborate **3c**, an intermediate in the synthesis of **3a**, was treated with potassium *E*-styryl-trifluoroborate^{34b,35} to give the stilbene **16b** in 58% yield (Scheme 8). Attempts to extend this methodology to synthesize nitro- and methoxy-substituted stilbenes were unsuccessful, although it was suspected the problems lay with the conversion of the corresponding styrylboronic acids into their trifluoroborate salts.

The parent bis(dimer) (17) and bis(trimer) (18) were generated in 35 and 75% yields, respectively, through the palladium-catalyzed coupling of 5 with 10a and 12a (Scheme 9). Dimer 10a reacted at room temperature, but the reaction of 12a was warmed to 60 °C to increase the solubility of the intermediate monoalkynylated anthracene and ensure completion. The preparation of the more functionalized bis(OPEs), and an examination of their properties, will be discussed in a separate article.³⁶



Scheme 8. Conditions: (a) Pd(OAc)₂, 58%.



Scheme 9. Conditions: (a) Pd(PPh₃)₂Cl₂, CuI, 35% (17); 75% (18).

3. Conclusions

In summary, an efficient method for the construction of trimethylsilylethynyl-terminated OPEs bearing different functional groups was elucidated, which allows access to a wide range of potential new NLO materials. Removal of the silyl protecting groups and subsequent attachment to the anthracene scaffold **5** generated bis(OPEs) **17** and **18** in which the OPE strands are held in close proximity, which could encourage coplanarity of the aryl rings and thereby increase the effective NLO response of the materials relative to their parent compounds. Preliminary results aimed at possibly further fine tuning the NLO properties by the combination of OPE and OPV elements were also discussed.

4. Experimental

4.1. General

4.1.1. Methyl 2-iodo-5-(trimethylsilylethynyl)benzoate **3a.** $BF_3 \cdot OEt_2$ (6.15 mL, 48.5 mmol) was cooled to -20 °C under N₂, then a solution of methyl 5-(trimethylsilylethynyl)anthranilate²³ (3.00 g, 12.1 mmol) in dry ether (30 mL) was added dropwise over 5 min. The dropping funnel was rinsed with dry ether (5 mL), then a solution of tert-butyl nitrite (5.05 mL, 42.4 mmol) in dry ether (15 mL) was added dropwise over 0.5 h. The resulting suspension was stirred at -20 °C for a further 10 min, then allowed to warm to 0 °C over 20 min. Dry ether (150 mL) was added and the suspension was kept at 0 °C for 15 min, filtered, and the solid collected was rinsed with ice-cold dry ether (20 mL) and briefly air dried. The crude diazonium salt 3c (3.65 g, 10.5 mmol) was dissolved in dry MeCN (39 mL) and added dropwise to a solution of NaI (1.90 g, 12.6 mmol) and I₂ (0.27 g, 1.05 mmol) in dry MeCN (80 mL) at room temperature under N₂. The dark solution was stirred at room temperature for 1 h, 20% Na₂S₂O₃ (90 mL) was added and the mixture was stirred vigorously for 5 min. The mixture was extracted with CH_2Cl_2 (3×120 mL), the combined extracts were washed with brine (120 mL), dried (Na₂SO₄), filtered and concentrated under vacuum. Squat column chromatography of the residue (1:2 CH₂Cl₂/hexane) afforded **3a**¹⁷ (3.20 g, 74% from anthranilate) as a pale yellow oil ($R_{\rm f}$ 0.28); C₁₃H₁₅IO₂SiNa requires 380.9784, found (M+Na)⁺ 380.9773; $\delta_{\rm H}$ (300 MHz, CDCl₃) 7.93 (1H, d, J=8.2 Hz), 7.88 (1H, d, J=1.9 Hz), 7.20 (1H, dd, J=2.2, 8.2 Hz), 3.93 (3H, s), 0.25 (9H, s); EI-MS *m/z* 358 (M⁺⁻), 343.

4.1.2. Methyl 5-ethynyl-2-*O*-(trifluoromethanesulfonyl)-salicylate **4a.** A mixture of **3b** (0.79 g, 2.08 mmol), anhydrous KF (116 mg, 2.00 mmol) and MeOH (20 mL) was stirred at room temperature for 18 h, then concentrated under vacuum to afford **4a** (0.64 g, 100%) as a colourless oil;²⁷ $\delta_{\rm H}$ (200 MHz, CDCl₃) 8.19 (1H, d, *J*=2.2 Hz), 7.70 (1H, dd, *J*=2.4, 8.6 Hz), 7.27 (1H, d, *J*=8.6 Hz), 3.97 (3H, s), 3.21 (1H, s).

Sonogashira coupling of terminal alkynes to 5. A mixture of 5^{22} (1.0 mmol), alkyne (2.2 mmol) and 1:2 Et₃N/DMF (6 mL) was degassed with a stream of N₂. Pd(PPh₃)₄ or Pd(PPh₃)₂Cl₂ (0.1 mmol) and CuI (0.1 mmol) were added and the mixture was stirred at room temperature under N₂ for 18 h, then poured into saturated NH₄Cl (40 mL). The resulting suspension was extracted with a suitable solvent (3×30 mL), and the combined extracts were washed with water (5×30 mL) and brine (30 mL), dried (Na₂SO₄), filtered and concentrated under vacuum.

4.1.3. 1,8-Bis[4-(trifluoromethanesulfonyloxy)-3-(methoxycarbonyl)phenylethynyl]-10-methoxyanthracene 6. The general procedure was followed using **5** (1.00 g, 2.18 mmol), **4a** (1.48 g, 4.79 mmol), CuI (41 mg, 0.22 mmol) and Pd(PPh_3)_4 (0.25 g, 0.22 mmol). After extraction with CH_2Cl_2, flash chromatography of the residue (1:4 EtOAc/hexane) afforded **6** (0.84 g, 47%) as a yellow solid (R_f 0.18), mp 140–145 °C (dec); C₃₇H₂₂F₆O₁₁S₂Na requires 843.0405, found (M+Na)⁺ 843.0412; δ_H (200 MHz, CDCl_3) 9.20 (1H, s), 8.36 (2H, dt, J=0.9, 8.8 Hz), 8.30 (2H, d, J=2.2 Hz), 7.89 (2H, dd, J=0.9, 6.8 Hz), 7.68 (2H, dd, J=2.2, 8.4 Hz), 7.51 (2H, dd, J=7.0, 8.8 Hz), 7.13 (2H, d, J=8.6 Hz), 4.17 (3H, s), 3.95 (6H, s); δ_C (50 MHz, CDCl_3) 163.5, 153.8, 147.6, 136.7, 136.5, 135.8, 132.0, 131.9, 125.1, 125.0, 124.9, 124.6, 124.4, 124.2, 123.3, 121.4, 120.7, 119.3, 118.8 (q, J_{CF} = 320.5 Hz), 92.1, 90.8, 63.9, 52.9, ν_{max} 1731, 1643, 1429, 1212, 1186, 1062, 1036, 987 cm⁻¹; EI-MS *m/z* 820 (M⁺⁺), 73; UV λ_{max} (log ε) 435 (3.91), 411 (3.98), 390 (3.81), 287 (4.49), 268 (4.82) nm; fluorescence λ_{em} 455, 481 nm.

4.1.4. 1,8-Bis[3-(methoxycarbonyl)-4-(piperidin-1-yl)phenylethynyl]-10-methoxyanthracene 7. A mixture of 5 (0.24 g, 0.53 mmol), **4a** (0.36 g, 1.17 mmol) and 1:1 piperidine/DMF (4 mL) was degassed with a stream of N_2 . Pd(PPh₃)₄ (62 mg, 0.05 mmol) and CuI (10 mg, 0.05 mmol) were added and the mixture was stirred at room temperature under N2 for 23 h, then poured into saturated NH₄Cl (20 mL). The mixture was extracted with CH_2Cl_2 (3×15 mL), the combined extracts were dried (Na₂SO₄), filtered and concentrated under vacuum. Flash chromatography of the residue (3:7 EtOAc/hexane) afforded 7 (0.30 g, 83%) as a yellow solid ($R_{\rm f}$ 0.37), mp 142–144 °C; $\delta_{\rm H}$ (200 MHz, CDCl₃) 9.41 (1H, s), 8.28 (2H, d, J=8.8 Hz), 8.01 (2H, d, J=2.0 Hz), 7.76 (2H, d, J=6.6 Hz), 7.44–7.52 (4H, m), 6.70 (2H, d, *J*=8.6 Hz), 4.14 (3H, s), 3.85 (6H, s), 3.00–3.05 (8H, m), 1.59–1.73 (12H, m); $\delta_{\rm C}$ (75 MHz, CDCl₃) 167.8, 153.1, 152.8, 135.8, 135.1, 131.9, 130.3, 124.9, 124.4, 122.7, 121.9, 119.8, 118.3, 114.5, 94.7, 87.0, 63.5, 53.4, 52.0, 25.9, 24.1; ν_{max} 2201, 1725, 1599, 1497, 1450, 1435, 1246, 1231, 1208, 1129, 1079, 1017, 920, 822, 732 cm⁻¹; EI-MS *m/z* 690 (M⁺⁺), 45.

4.1.5. 1,8-Bis[3-(methoxycarbonyl)-4-(trimethylsilylethynyl)phenylethynyl]-10-methoxyanthracene 8. A mixture of 6 (0.44 g, 0.53 mmol), CuI (10 mg, 0.05 mmol) and 1:2 Et₃N/NMP (6 mL) was degassed with a stream of N₂. Pd(PPh₃)₄ (62 mg, 0.05 mmol) and TMSA (0.30 mL, 2.13 mmol) were added and the mixture was stirred at room temperature for 19 h. The mixture was poured into saturated NH₄Cl (30 mL) and extracted with CH₂Cl₂ (3×20 mL). The combined extracts were washed with water (20 mL) and brine (20 mL), dried (Na₂SO₄), filtered and concentrated under vacuum. The residue was applied to a short column of silica, eluted with 1:1 EtOAc/hexane (100 mL) and the eluant was concentrated under vacuum. Flash chromatography of the residue (1:4 EtOAc/hexane) afforded **8** (0.28 g, 74%) as a yellow oil ($R_{\rm f}$ 0.40); $C_{45}H_{40}O_5Si_2Na$ requires 739.2312, found $(M+Na)^+$ 739.2316; δ_H $(300 \text{ MHz}, \text{CDCl}_3)$ 9.29 (1H, s), 8.33 (2H, d, J=8.5 Hz), 8.17 (2H, d, J=1.6 Hz), 7.83 (2H, dd, J=0.8, 6.9 Hz), 7.61 (2H, dd, J=1.6, 8.0 Hz), 7.51 (2H, dd, J=6.7, 8.8 Hz), 7.45 (2H, d, J=8.0 Hz), 4.16 (3H, s), 3.89 (6H, s), 0.31 (18H, s); δ_C (75 MHz, CDCl₃) 165.9, 153.4, 134.5, 134.4, 134.0, 133.3, 132.9, 131.7, 131.6, 124.9, 124.4, 123.6, 122.9, 121.1, 119.4, 103.0, 101.9, 93.7, 90.6, 63.6, 52.1, -0.1; $\nu_{\rm max}$ 2157, 1732, 1677, 1601, 1491, 1437, 1286, 1249, 1079, 845, 760, 739 cm⁻¹; EI-MS *m*/*z* 716 (M⁺⁻); UV $\lambda_{\rm max}$ (log ε) 308 (4.85), 289 (4.88), 231 (5.10) nm; fluorescence λ_{em} 377 nm.

4.1.6. Methyl 5-(trimethylsilylethynyl)-2-(phenylethynyl)benzoate 9a. A mixture of 3b (1.00 g, 2.79 mmol), 2a (0.33 mL, 3.07 mmol) and 1:2 Et₃N/DMF (12 mL) was degassed with a stream of N₂. CuI (27 mg, 0.14 mmol) and Pd(PPh₃)₂Cl₂ (98 mg, 0.14 mmol) were added and the mixture was stirred at room temperature under N₂ for 18 h. The mixture was poured into saturated NH₄Cl (50 mL) and extracted with ether (3×50 mL). The extracts were washed with water (5×50 mL) and brine (50 mL), dried (MgSO₄), filtered and concentrated under vacuum. Flash chromatography of the residue (3:7 CH₂Cl₂/hexane) afforded **9a** (0.83 g, 89%) as a yellow oil ($R_{\rm f}$ 0.28); C₂₁H₂₁O₂Si requires 333.1311, found (M+H)⁺ 333.1307; $\delta_{\rm H}$ (200 MHz, CDCl₃) 8.08 (1H, dd, *J*=0.9, 1.3 Hz), 7.55–7.60 (4H, m), 7.34–7.39 (3H, m), 3.96 (3H, s), 0.26 (9H, s); $\delta_{\rm C}$ (50 MHz, CDCl₃) 165.9, 134.6, 134.1, 133.8, 131.9, 131.8, 128.7, 128.4, 123.6, 123.1, 122.9, 103.6, 97.3, 96.2, 88.0, 52.3, -0.2; $\nu_{\rm max}$ 2157, 1729, 1250, 1185, 1071, 1036, 848 cm⁻¹; EI-MS *m*/*z* 332 (M⁺⁺), 317; UV $\lambda_{\rm max}$ (log ε) 340 (4.61), 330 (4.62), 306 (4.64), 296 (4.54), 288 (4.50), 267 (4.45), 247 (4.57), 236 (4.63) nm; fluorescence $\lambda_{\rm em}$ 367 nm.

4.1.7. Methyl 5-ethynyl-2-(phenylethynyl)benzoate 10a. A mixture of 9a (0.26 g, 0.78 mmol), anhydrous K₂CO₃ (65 mg, 0.47 mmol) and 1:1 MeOH/CH₂Cl₂ (8 mL) was stirred at room temperature for 3 h, then diluted with CH₂Cl₂ (20 mL), washed with water (15 mL) and brine (15 mL), dried (Na₂SO₄), filtered and concentrated under vacuum to afford **10a** (0.21 g, 100%) as a pale yellow oil;²⁷ $\delta_{\rm H}$ (200 MHz, CDCl₃) 8.10 (1H, t, *J*=1.1 Hz), 7.55–7.60 (4H, m), 7.37 (3H, m), 3.97 (3H, s), 3.21 (1H, s).

4.1.8. Methyl 5-ethynyl-2-[4-(trifluoromethanesulfonyloxy)phenylethynyl]benzoate 10h. A mixture of 9h (0.53 g, 1.10 mmol), anhydrous KF (64 mg, 1.10 mmol) and MeOH (10 mL) was stirred at room temperature for 18 h, then diluted with CH₂Cl₂ (25 mL), washed with water (15 mL) and brine (15 mL), dried (Na₂SO₄), filtered and concentrated under vacuum to afford 10h (0.34 g, 76%) as a yellow oil;²⁷ $\delta_{\rm H}$ (200 MHz, CDCl₃) 8.12 (1H, t, *J*=1.1 Hz), 7.65 (2H, m), 7.60 (2H, d, *J*=1.1 Hz), 7.28 (2H, m), 3.96 (3H, s), 3.23 (1H, s).

4.1.9. Methyl 2-[3-(methoxycarbonyl)-4-(phenylethynyl)phenylethynyl]-5-(trimethylsilylethynyl)benzoate 11a. The procedure for 9a was followed using 3a (0.34 g, 0.94 mmol), **10a** (0.27 g, 1.03 mmol), CuI (9 mg, 0.05 mmol) and Pd(PPh₃)₂Cl₂ (33 mg, 0.05 mmol). After extraction with CH₂Cl₂, flash chromatography of the residue (3:17 EtOAc/hexane) afforded **11a** (0.34 g, 74%) as a pale yellow solid (*R*_f 0.12, 1:1 CH₂Cl₂/hexane), mp 118–121 °C; $C_{31}H_{26}O_4SiNa$ requires 513.1498, found $(M+Na)^+$ 513.1485; $\delta_{\rm H}$ (300 MHz, CDCl₃) 8.17 (1H, dd, J=0.6, 1.6 Hz), 8.10 (1H, t, J=0.8 Hz), 7.66 (1H, d, J=1.6 Hz), 7.64 (1H, br s), 7.57-7.61 (4H, m), 7.35-7.39 (3H, m), 3.98 (3H, s), 3.97 (3H, s), 0.27 (9H, s); $\delta_{\rm C}$ (75 MHz, CDCl₃) 165.9, 165.7, 134.6, 134.4, 134.1, 134.0, 133.9, 133.8, 132.1, 131.9, 131.8, 128.7, 128.4, 123.8, 123.4, 123.1, 122.9, 103.4, 97.7, 97.3, 96.5, 94.8, 90.6, 88.1, 52.32, 52.28, -0.2; ν_{max} 1731, 1647, 1503, 1293, 1248, 1189, 1072, 1036, 847 cm⁻¹; EI-MS *m/z* 490 (M⁺⁺), 475; UV λ_{max} (log ε) 351 (4.78), 249 (4.34) nm; fluorescence λ_{em} 388 nm.

4.1.10. Methyl 5-ethynyl-2-[3-(methoxycarbonyl)-4-(phenylethynyl)phenylethynyl]benzoate 12a. The procedure for 10a was followed using 11a (0.41 g, 0.83 mmol) and anhydrous K₂CO₃ (0.11 g, 0.83 mmol) to afford 12a (0.34 g, 100%) as a yellow-orange solid;²⁷ $\delta_{\rm H}$ (300 MHz, CDCl₃) 8.18 (1H, dd, J=0.4, 1.5 Hz), 8.13 (1H, t, J=1.1 Hz), 7.68 (1H, dd, J=1.6, 8.0 Hz), 7.65 (1H, d, *J*=0.4 Hz), 7.57–7.61 (4H, m), 7.37 (3H, m), 3.984 (3H, s), 3.981 (3H, s), 3.23 (1H, s).

4.1.11. Methyl 2-O-trifluoromethanesulfonyl-5-[2-(methoxycarbonyl)-4-(trimethylsilylethynyl)phenylethynyl]-salicylate 13, and methyl 2-{4-[4-(trifluoromethanesulfonyloxy)-3-(methoxycarbonyl)phenylethynyl]-3-[methoxycarbonyl]phenylethynyl}-5-(trimethylsilylethynyl)benzoate 14. The procedure for 9a was followed using **3a** (0.40 g, 1.10 mmol), **4a** (0.37 g, 1.21 mmol), CuI (11 mg, 0.06 mmol) and Pd(PPh₃)₂Cl₂ (39 mg, 0.06 mmol). After extraction with CH₂Cl₂, flash chromatography of the residue (1:1 CH₂Cl₂/hexane, then 2:1) afforded **13** (0.36 g, 32%) as a pale yellow oil ($R_{\rm f}$ 0.54, 2:1 CH₂Cl₂/hexane); C₂₄H₂₂F₃O₇SSi requires 539.0808, found $(M+H)^+$ 539.0808; δ_H (200 MHz, CDCl₃) 8.25 (1H, d, J=2.2 Hz), 8.10 (1H, t, J=1.3 Hz), 7.79 (1H, dd, J=2.2, 8.4 Hz), 7.58 (2H, d, J=1.1 Hz), 7.30 (1H, d, J=8.6 Hz), 3.98 (3H, s), 3.96 (3H, s), 0.27 (9H, s); δ_C (50 MHz, CDCl₃) 165.5, 163.5, 147.7, 136.9, 135.8, 134.7, 134.2, 134.0, 132.1, 124.8, 124.2, 123.9, 123.1, 122.4, 118.7 (q, $J_{CF} =$ 319.1 Hz), 103.3, 98.1, 92.8, 90.8, 52.8, 52.4, -0.2; EI-MS m/z 538 (M⁺), 523, 405, 375, 332. Further elution afforded 14 (0.14 g, 18%) as a vellow solid ($R_{\rm f}$ 0.22), mp 129– 132 °C; $C_{34}H_{27}F_3O_9SSiNa$ requires 719.0995, found (M+ Na)⁺ 719.1000; $\delta_{\rm H}$ (200 MHz, CDCl₃) 8.26 (1H, d, J= 2.2 Hz), 8.20 (1H, d, J=1.1 Hz), 8.10 (1H, t, J=1.1 Hz), 7.80 (1H, dd, J = 2.2, 8.4 Hz), 7.70 (1H, dd, J = 1.6, 8.1 Hz), 7.63 (1H, d, J=7.9 Hz), 7.58 (2H, d, J=1.3 Hz), 7.30 (1H, d, J=8.4 Hz), 3.99 (3H, s), 3.98 (3H, s), 3.97 (3H, s), 0.26 (9H, s); $\delta_{\rm C}$ (50 MHz, CDCl₃) 165.6, 165.5, 163.5, 147.7, 136.9, 135.8, 134.7, 134.6, 134.2, 134.0, 133.9, 132.3, 131.9, 128.6, 124.8, 124.2, 123.9, 123.6, 123.1, 122.8, 122.6, 125.1 (q, J_{CF}=322.7 Hz), 103.4, 97.9, 94.5, 93.0, 91.1, 90.9, 52.8, 52.44, 52.37, -0.2; ν_{max} 2157, 1734, 1501, 1291, 1249, 1210, 1140, 1072, 985, 845 cm⁻¹; EI-MS m/z696 (M⁺), 563, 274, 259, 237, 77; UV λ_{max} (log ε) 348 (4.52), 318 (4.32); fluorescence λ_{em} 386, 405 nm.

4.1.12. Methyl 2-methoxycarbonyl-4-(trimethylsilylethynyl)cinnamate 15a. To a solution of 3b (0.50 g, 1.31 mmol) in DMF (3 mL) were added Et₃N (0.20 mL, 1.45 mmol), methyl acrylate (0.24 mL, 2.63 mmol), dppp (15 mg, 0.04 mmol) and $Pd(OAc)_2$ (7 mg, 0.03 mmol), and the mixture was stirred at 80 °C under N2 for 6.5 h. After cooling, the mixture was diluted with CH_2Cl_2 (30 mL), washed with 5% HCl (2×20 mL), water (3×20 mL) and brine (20 mL), dried (Na₂SO₄), filtered and concentrated under vacuum. Flash chromatography of the residue (7:3 CH₂Cl₂/hexane) afforded **15a** (0.20 g, 47%) as a colourless oil (R_f 0.38); $C_{17}H_{20}O_4$ Si requires 316.1131, found (M⁺) 316.1134; $\delta_{\rm H}$ (300 MHz, CDCl₃) 8.41 (1H, d, J = 16.2 Hz), 8.04 (1H, d, J=1.6 Hz), 7.59 (1H, dd, J=1.6, 8.2 Hz), 7.54 (1H, d, J=8.0 Hz), 6.31 (1H, d, J=15.9 Hz), 3.93 (3H, s), 3.81 (3H, s), 0.26 (9H, s); $\delta_{\rm C}$ (75 MHz, CDCl₃) 166.8, 166.5, 142.9, 136.0, 135.2, 134.3, 129.9, 129.6, 127.8, 121.2, 103.3, 97.5, 52.5, 51.8, -0.2; EI-MS m/z 316 (M⁺⁺), 301.

4.1.13. Methyl 5-(trimethylsilylethynyl)-2-*E*-(2-phenylethenyl)benzoate 15b. 1,4-Dioxane (4 mL) was degassed with a stream of N₂. Crude **3c** (0.35 g, 1.00 mmol), potassium *E*-styryltrifluoroborate^{34a,35} (0.25 g, 1.20 mmol) and $Pd(OAc)_2$ (11 mg, 0.05 mmol) were added and the mixture was stirred at room temperature under N_2 in the dark for 19 h. The solution was diluted with CH₂Cl₂ (40 mL), washed with water (5×25 mL) and brine (25 mL), dried (Na₂SO₄), filtered and concentrated under vacuum. Flash chromatography of the residue (2:3 CH₂Cl₂/hexane) afforded **15b** (0.19 g, 58%) as a colourless oil (R_f 0.22, 1:2 CH₂Cl₂/hexane); C₂₁H₂₂O₂Si requires 334.1389, found (M^{+}) 334.1385; δ_{H} (200 MHz, CDCl₃) 8.04 (1H, dd, J=0.5, 1.8 Hz), 7.99 (1H, d, J=16.3 Hz), 7.68 (1H, d, J = 8.2 Hz, 7.51–7.59 (3H, m), 7.27–7.41 (3H, m), 7.04 $(1H, d, J = 16.3 \text{ Hz}), 3.93 (3H, s), 0.27 (9H, s); \delta_C (50 \text{ MHz}), \delta_C (50 \text{$ CDCl₃) 167.1, 139.0, 137.2, 135.0, 134.3, 132.3, 128.7, 128.3, 128.1, 126.9, 126.7, 126.5, 122.0, 104.0, 96.0, 52.2, -0.1; ν_{max} 2159, 1722, 1492, 1295, 1248, 1208, 1073, 845 cm⁻¹; EI-MS *m*/*z* 334 (M⁺⁻), 319; UV λ_{max} (log ε) 327 (4.15), 269 (4.14), 259 (4.21) nm; fluorescence $\lambda_{\rm em}$ 398 nm.

4.1.14. 1,8-Bis[3-(methoxycarbonyl)-4-(phenylethynyl)phenylethynyl]-10-methoxyanthracene 17. The general procedure was followed using 5 (0.17 g, 0.37 mmol), 10a (0.21 g, 0.82 mmol), CuI (7 mg, 0.04 mmol) and Pd(PPh₃)₂Cl₂ (26 mg, 0.04 mmol). After extraction with CH₂Cl₂, flash chromatography of the residue (gradient elution 1:9 EtOAc/ hexane to 1:3) afforded 17 (94 mg, 35%) as a yellow solid $(R_{\rm f} 0.29, 1:4 \text{ EtOAc/hexane}), \text{ mp } 95-105 \,^{\circ}\text{C} \, (\text{dec});$ $C_{51}H_{32}O_5$ requires 724.2250, found (M⁺⁺) 724.2246; δ_H $(200 \text{ MHz}, \text{ CDCl}_3) 9.37 (1\text{H}, \text{ br s}), 8.35 (2\text{H}, \text{ dt}, J=1.1,$ 8.8 Hz), 8.24 (2H, dd, J = 0.5, 1.8 Hz), 7.85 (2H, dd, J = 0.7, 6.8 Hz), 7.49–7.62 (8H, m), 7.42 (2H, dd, J=0.5, 8.1 Hz), 7.18–7.31 (6H, m), 4.18 (3H, s), 3.93 (6H, s); δ_C (75 MHz) 165.8, 153.6, 134.3, 134.1, 133.6, 132.1, 132.0, 131.9, 131.2, 128.6, 128.4, 128.3, 125.0, 124.5, 123.6, 123.2, 122.9, 121.3, 119.6, 96.5, 93.9, 90.3, 88.1, 63.7, 52.3; v_{max} 1723, 1500, 1281, 1184, 1074, 1036, 992 cm⁻¹; EI-MS *m/z* 724 (M⁺, 709, 694, 634, 588; UV λ_{max} (log ε) 438 (4.26), 414 (4.31), 391 (4.19), 331 (4.66), 307 (4.72), 267 (4.87), 247 (4.70) nm; fluorescence λ_{em} 457, 484 nm.

4.1.15. 1,8-Bis{3-[methoxycarbonyl]-4-[3-(methoxycarbonyl)-4-(phenylethynyl)phenylethynyl]phenylethynyl}-10-methoxyanthracene 18. The general procedure was followed using 5 (0.17 g, 0.38 mmol), 12a (0.35 g, 0.83 mmol), CuI (7 mg, 0.04 mmol) and Pd(PPh₃)₂Cl₂ (26 mg, 0.04 mmol), except that the reaction was stirred at 60 °C. After extraction with CH₂Cl₂, flash chromatography of the residue (9:1 CH₂Cl₂/hexane then CH₂Cl₂) afforded 18 (0.30 g, 75%) as a dark yellow solid ($R_{\rm f} 0.18$, CH₂Cl₂), mp 158-165 °C (dec); C71H44O9Na requires 1063.2883, found $(M+Na)^+$ 1063.2866; δ_H (300 MHz, CDCl₃) 9.36 (1H, br s), 8.36 (2H, dt, J=1.0, 8.8 Hz), 8.22 (2H, dd, J=0.3, 1.5 Hz), 8.07 (2H, dd, J = 0.4, 1.8 Hz), 7.83 (2H, dd, J = 1.0, 7.0 Hz), 7.50–7.59 (10H, m), 7.43 (2H, dd, J=0.4, 8.0 Hz), 7.37 (2H, dd, J=0.4, 7.8 Hz), 7.22–7.34 (6H, m), 4.18 $(3H, s), 3.95 (6H, s), 3.94 (6H, s); \delta_C (75 \text{ MHz}, \text{CDCl}_3) 96.5,$ 95.1, 93.9, 90.63, 90.60, 88.2, 63.7, 52.3, 52.2; ν_{max} 1771, 1734, 1505, 1287, 1244, 1184, 1074, 1037, 991 cm⁻¹; ESI-MS $m/z \ 1063 \ (M + Na)^+$; UV $\lambda_{max} \ (\log \varepsilon) \ 440 \ (4.48)$, 416 (4.57), 396 (4.55), 351 (5.01), 321 (4.85), 266 (4.95), 248 (4.85); fluorescence $\lambda_{\rm em}$ 465, 488.

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11. 015. General experimental methods and compound characterization data for **9b–g**, **10b–g**, **11b–e**, and **12b–e**.

References and notes

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1,3-Dipolar cycloaddition approach to isoxazole, isoxazoline and isoxazolidine analogues of *C***-nucleosides related to pseudouridine**

Evdoxia Coutouli-Argyropoulou,^{a,*} Pygmalion Lianis,^a Marigoula Mitakou,^a Anestis Giannoulis^a and Joanna Nowak^b

^aDepartment of chemistry, Aristotle University of Thessaloniki, 54124 Thessaloniki, Greece ^bErasmus student from A. Mickiewicz University, Poznan, Poland

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Abstract—Isoxazole, isoxazoline and isoxazolidine analogues of *C*-nucleosides related to pseudouridine have been synthesized by 1,3-dipolar cycloaddition reactions of nitrile oxides and nitrones derived from mono and disubstituted uracil-5-carbaldehydes and 2,4-dimethoxypyrimidine-5-carbaldehyde. The dimethoxy derivatives have been easily deprotected to the corresponding uracils bearing the heterocyclic ring instead of a sugar moiety. The regio and stereoselectivity of the reactions are discussed. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Since the latter part of the 1980s unnatural nucleoside analogues have played an important role as anticancer and antiviral agents.¹ Consequently, several variations have been made to both the heterocyclic base and the sugar moiety in the search for effective and selective derivatives. Due to the need for the base moiety to preserve the basepairing functionalities, only minor modifications of the base are usually found in biologically active nucleosides analogues. The C-5 position is usually the position of choice for the introduction of substituents in pyrimidine nucleosides since it is not involved in the Watson-Crick base-pairing.² On the contrary, a lot of variations have been made in the sugar part replacing it by acyclic moieties or carbo or other heterocyclic rings. Among them, isoxazoline and isoxazolidine nucleosides have emerged as an important class of nucleoside analogues and several approaches for their synthesis have been reported.³

Besides the variations in the sugar and base moieties a crucial modification results from varying their connection, as in the *C*-nucleosides, which have a carbon–carbon linkage instead of an hydrolyzable carbon–nitrogen bond between the sugar and the aglycon. The most abundant natural *C*-nucleoside is pseudouridine a C-5 linked uridine. Pseudouridine is the first *C*-nucleoside found in nature

and has attracted the interest of organic chemists and biochemists since its discovery in 1957.⁴ The occurrence of pseudouridine in highly conserved regions of RNA indicates that certain physicochemical properties of pseudouridine are critical to the biological function of RNA molecules.

Thus the biological significance of pseudouridine has resulted in studies aimed at the incorporation of synthetic pseudouridine analogues with modified sugar moieties.⁵ Recently, the synthesis of isoxazoline analogues of pseudouridine by 1,3-dipolar cycloaddition reactions of 5-uracil nitrones has been described.⁶

During recent years and in connection with our interests to induce nucleoside modifications,⁷ we have also attempted to apply the convenience and diversity of 1,3-dipolar cycloaddition reactions to the synthesis of pseudouridine analogues. However, our initial attempts to isolate cycloaddition products via the in situ formation of nitrile oxides from 5-uracilcarbaldehyde oxime or 1-monosubstituted 5-uracilcarbaldehyde oximes were unsuccessful even in the presence of very active dipolarophiles such as methyl acrylate. On the contrary these oximes gave mixtures of isoxazolidines from the reaction of intermediate nitrones.^{8a} Nitrone generation from oximes via a 1,2-prototropic process or an 1,3-azaprotiocyclotransfer is a well known reaction established by Grigg,^{8b,c} and it has been also described by us for other oximes.^{8d} The last findings indicated that nitrone cycloaddition can work with uracil nitrones barrying free NH bonds. This has been also shown recently by the work of Chiacchio et al.⁶

Keywords: Pyrimidine; Pseudouridine; Cycloaddition.

^{*} Corresponding author. Tel.: +30 2310 997733; fax: +30 2310 997679; e-mail: evd@chem.auth.gr

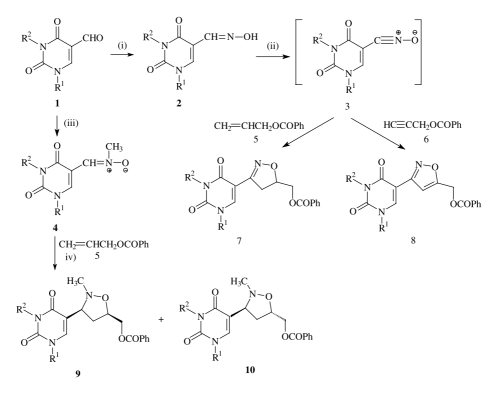
However, nucleoside analogues with restricted conformational flexibility induced by a second ring or by unsaturation are the target compounds in many cases, since they are potent inhibitors of HIV-1 reverse transcriptase.^{3k,o,9} Thus, in order to expand the use of 5-uracil dipoles for the formation of both saturated and unsaturated rings, we report in this paper, the application of cycloaddition reactions of both nitrile oxides and nitrone uracil dipoles by applying monosubsituted, disubstituted and protected uracil derivatives.

2. Results and discussion

As starting materials for the formation of the dipoles we have chosen the mono and disubstituted aldehydes 1a and 1b and the dimethoxy-5-formyl pyrimidine 11 (Schemes 1 and 2). The octyl derivatives 1a and 1b have been chosen for purposes of higher solubility and enhanced hydrophobicity, whereas aldehyde 11 is a protected form of 5-formyluracil. The above aldehydes were prepared according to the procedures we have previously described.¹⁰ The oximes 2a, 2b, 12 as well as the nitrones 4a, 4b and 14 were prepared from the corresponding aldehydes applying conventional procedures. As dipolarophiles, we have used allylic or propargylic alcohol derivatives in order to ensure the presence of a 5'-hydroxymethyl group in the final product, which potentially allows enzymatic phosphorylation for antiviral expression or incorporation into automatic solid phase synthesis.

Nitrile oxide 3b was generated in situ from the corresponding oxime in the presence of the dipolarophile in a biphasic methylene chloride/aqueous bleach system. Generation of nitrile oxide 3a following the same procedure was unsuccessful. As we have already mentioned, in our initial attempts we failed to isolate nitrile oxide cycloaddition products from 1-substituted uracil aldoximes. Thus, as well as the above standard procedure for the generation of the nitrile oxide 3a from the oxime 2a, several other alternative procedures using N-chlorosuccinimide, and several variations in the reaction time, temperatures and work up were also tested without success. Nitrile oxide 3b reacted with both allylic benzoate (5) and propargylic benzoate (6) to give the isoxazoline **7b** and the isoxazole **8b**, respectively, in good yields (70-80%). The reactions were regioselective and only 5-substituted cycloadducts were isolated. The reactions of nitrones 4 with the alkene 5 took place under reflux in xylene to give isoxazolidines 9 as the main products in satisfactory yields (70-72%). The reactions were regio and stereoselective. In both cases only 5-substituted derivatives with a cis arrangement of the 3' and 5' substituents (structure 9) were isolated, although in the crude reaction mixture, traces of compounds with structure 10 were also detected on the basis of some ¹H NMR chemical shifts (Table 1).

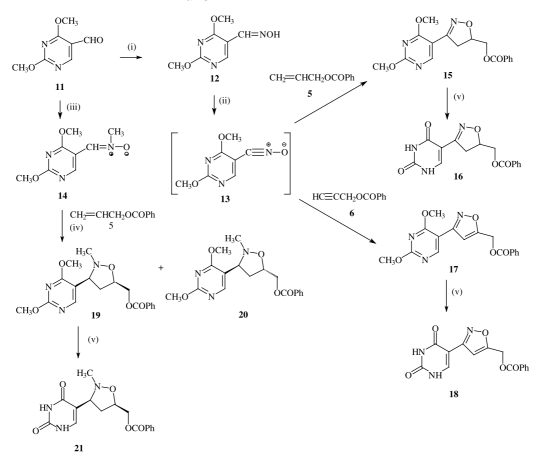
Dimethoxypyridine dipoles 13 and 14 showed analogous behaviour. Nitrile oxide 13 generated in situ from the oxime 12 reacted regioselectively with 5 to give the isoxazoline derivative 15. The reaction of 13 with the alkyne derivative



1a, **2a**, **4a**, **9a**, **10a** $R^1 = CH_3(CH_2)_6CH_2$, $R^2 = H$

1b, **2b**, **3b**, **4b**, **7b**, **8b**, **9b**, **10b** $R^1 = R^2 = CH_3(CH_2)_6CH_2$

Scheme 1. Reagents and conditions: (i) NH₂OH·HCl, Na₂CO₃, EtOH/H₂O, 20 °C, 24 h; (ii) NaOCl, CH₂Cl₂/H₂O, 0–20 °C, 24 h; (iii) CH₃NHOH·HCl, Na₂CO₃, EtOH/H₂O, 20 °C, 24 h; (iv) Xylene, reflux, 48 h.



Scheme 2. Reagents and conditions: (i) NH₂OH·HCl, Na₂CO₃, EtOH/H₂O, 20 °C, 24 h; (ii) NaOCl, CH₂Cl₂/H₂O, 0–20 °C, 24 h; (iii) CH₃NHOH·HCl, Na₂CO₃, EtOH/H₂O, 20 °C, 24 h; (iv) Xylene, reflux, 48 h; (v) CH₃COOH, NaI, 90 °C, 1 h.

Table 1. Selected values for proton chemical shifts and coupling constants of compounds 9, 10, 19, 20

Compound	4′-Ha	4'-Hb	3′-Н
9a	2.10 (dt, $J_{4'a,4'b} = 12.2$ Hz,	3.02 (ddd, $J_{4'a,4'b} = 12.2$ Hz, $J_{3',4'b} = 7.3$ Hz,	4.03 (dd, $J_{3',4'a}$ =5.1 Hz, $J_{3',4'b}$ =7.3 Hz)
9b	$J_{3',4'a} = J_{4'a,5'} = 5.1 \text{ Hz})$ 2.05 (dt, $J_{4'a,4'b} = 13.6 \text{ Hz},$	$J_{4'b,5'} = 8.4 \text{ Hz}$) 3.02 (ddd, $J_{4'a,4'b} = 13.6 \text{ Hz}$, $J_{3',4'b} = 7.8 \text{ Hz}$,	3.99 (dd, $J_{3',4'a}$ =5.1 Hz, $J_{3',4'b}$ =7.8 Hz)
19	$J_{3',4'a} = J_{4'a,5'} = 5.1 \text{ Hz})$ 2.05 (dt, $J_{4'a,4'b} = 12.8 \text{ Hz},$	$J_{4'b,5'} = 8.4 \text{ Hz})$ 2.89 (ddd, $J_{4'a,4'b} = 12.8 \text{ Hz}, J_{3',4'b} = 8.4 \text{ Hz},$	3.85 (dd, $J_{3',4'a} = 6.4$ Hz, $J_{3',4'b} = 8.4$ Hz)
20	$J_{3',4'a} = J_{4'a,5'} = 6.4 \text{ Hz})$ 2.41 (ddd, $J_{4'a,4'b} = 14.2 \text{ Hz},$	$J_{4'b,5'} = 7.7$ Hz) 2.55 (ddd, $J_{4'a,4'b} = 14.2$ Hz, $J_{3',4'b} = 3.9$ Hz,	4.24 (dd, $J_{3',4'a}$ =5.7 Hz, $J_{3',4'b}$ =3.9 Hz)
10a	$J_{3',4'a} = 5.7 \text{ Hz}, J_{4'a,5'} = 7.7 \text{ Hz})$ 2.29–2.37 (m)	$J_{4'b,5'} = 8.9 \text{ Hz}$ 2.60–2.69 (m)	4.22 (dd, $J_{3',4'a} = 6.0$ Hz, $J_{3',4'b} = 4.1$ Hz)
10b	2.25–2.35 (m)	2.60–2.69 (m)	4.22 (dd, $J_{3',4'a} = 5.2$ Hz, $J_{3',4'b} = 3.8$ Hz)

6 was also regioselective affording the 5'-substituted isomer **17**. The reaction of the nitrone **14** with the alkene **5** was also regioselective, but less stereospecific than that of nitrones **4** resulting in the formation of the two 5'-substituted stereoisomes **19** and **20** in a ratio 1.5:1.

The structure elucidation of the obtained cycloadducts was made on the basis of their elemental analysis and their spectral data. All the compounds give molecular ion peaks in the mass spectra and the expected chemical shifts in the ¹H and ¹³C NMR spectra. The differentiation between stereoisomers **9** and **10** and between **19** and **20** was less obvious and was based on observed coupling constants and NOE measurements carried out on compound **9b**. The protons assignment was confirmed by decoupling experiments and selected chemical shifts and coupling constants of diagnostic value for compounds 9, 10, 19 and 20 are given in Table 1.

In 9a, 9b, and 19, the one of 4'-H (4a'-H) appears at a higher field, and exhibits smaller coupling constants with both 3'-H and 5'-H than the other 4'-H (4b'-H), indicating a trans topological relationship with both of them. On the contrary, in compound 20 each of the 4'-H exhibits one large and one small coupling constant indicative that is trans to one and cis to the other. An interesting feature also in the ¹H NMR spectra is the difference in the chemical shifts of the two 4'-H protons, which is remarkably larger in the stereoisomers 9a, 9b, and 19 with a cis arrangement of the 3' and 5' substituents than in 20 with a trans arrangement probably as a result of the shielding effect of both substituents to the same proton. Also, the chemical shift of 3'-H is higher in the trans isomer than in the cis. Thus the presence of multipets in the ¹H NMR of the crude reaction mixtures at the regions 2.25–2.37 and 2.60–269 as well as a dd at δ 4.22 are indicative for isomers **10a** and **10b**.

The proposed stereochemistry for the isolated cycloadducts was further supported by NOE measurements carried out on compound **9b**. As depicted in Figure 1, the mutual large NOE enhancements observed upon saturation of 3'-H, 5'-H and 4b'-H are in accordance with their cis arrangement.

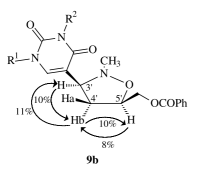


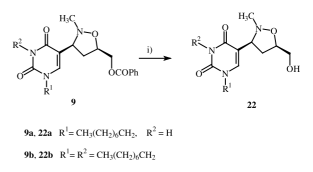
Figure 1.

It should be mentioned that the stereoselectivity of the reactions leading preferentially to cycloadducts with a cis arrangement of 3' and 5' substituents is favorable, since cis cycloaaducts match more the natural analogues. On the contrary, trans cycloadducts were referred as the main products of the reactions of unsubstituted uracil nitrones.⁶ The observed stereoselectivity of the reactions can be explained via an endo approach of the dipolarophile assuming Z-configuration of the nitrone as it has been proved for aldonitrones.^{6,11} Secondary interactions that favor an *endo* approach obviously prevail in the reactions of octyl and dioctyl substituted nitrones 4, leading almost exclusively to the formation of cis cycloadducts 9. In the reaction of the dimethoxy nitrone 14, competition between steric factors and secondary interactions leads to the formation of a substantial amount of the minor trans isomer 20 as a product of the *exo* approach of the dipolarophile.

The dimethoxy derivatives **15**, **17** and **19** were readily transformed to uracil derivatives **16**, **18** and **21**, respectively, in satisfactory yields (67–72%) and without loss of the heterocyclic ring moiety, by heating in acetic acid in the presence of sodium iodide. The obtained uracils, besides the disappearance of the methoxy chemical shifts and the presence of NH resonances, exhibits in their NMR almost the same characteristics with their precursors.

The removal of the benzoyl group from the obtained cycloadducts can be also done easily by alkaline hydrolysis. In a representative experiment compounds **9a** and **9b** were transformed quantitatively to the corresponding hydroxy derivatives **22a** and **22b** with potassium hydroxide in aqueous methanol solution (Scheme 3).

In conclusion, cycloaddition reactions of nitrones or nitrile oxides derived from suitably substituted uracils or dimethoxy pyrimidines can be used as versatile procedures for the synthesis of modified pseudouridine analogues



Scheme 3. Reagents and conditions: (i) KOH, MeOH/H₂O, 20 °C, 24 h.

bearing isoxazole, isoxazoline or isoxazolidine rings instead of a sugar unit. The case of dimethoxy pyrimidine derivatives is significant in the sense that they could be deprotected without affecting the heterocyclic ring moiety. The presence of substituents differentiates the stereoselectivity of the reactions favoring those more close related to the natural products (cis cycloadducts) as a result of enhanced secondary interactions.

3. Experimental

3.1. General

Mps are uncorrected and were determined on a Kofler hot-stage microscope. IR spectra were recorded on a Perkin-Elmer 297 spectrometer. ¹Ĥ NMR spectra were recorded at 300 MHz on a Bruker 300 AM spectrometer and ¹³C NMR spectra at 75.5 MHz on the same spectrometer, and are quoted relative to tetramethylsilane as internal reference, in deuteriochloroform solutions, unless otherwise stated. Mass spectra (EI) were performed on a VG-250 spectrometer with ionization energy maintained at 70 eV. High resolution mass spectra (HRESI) were obtained with a 7 T APEX II spectrometer. Microanalyses were performed on a Perkin-Elmer 2400-II element analyser. Column chromatography was carried out on Merck Kieselgel (particle size 0.063-0.200 mm) and solvents were distilled before use. The preparation of the aldehydes 1 and 11 was made according to previously described procedures.¹⁰

3.2. Synthesis of oximes 2 and 12

General procedure. An aqueous solution (2.5 ml) of hydroxylamine hydrochloride (2.25 mmol) and sodium carbonate (1.5 mmol) were added to an ethanolic solution (5 ml) of the aldehyde **1** or **11** (1 mmol) and the reaction mixture was stirred at room temperature for 24 h. After that the ethanol was evaporated, water was added and the mixture was dried over sodium sulfate and after evaporation of the solvent the oximes were obtained as white solids and they were used without further purification.

3.2.1. 1-Octyl-5-uracilcarbaldehyde oxime (2a). This compound was obtained in 90% yield as a white solid, mp 173–176 °C; IR (Nujol): ν_{max} 3300, 3150, 3040, 1680, 1600 cm⁻¹; ¹H NMR (DMSO- d_6 + CDCl₃)): δ 0.87 (t, J= 7.2 Hz, 3H, CH₃), 1.27–1.31 (m, 10H, CH₂CH₂(CH₂)₅CH₃), 1.68 (br t, 2H, CH₂CH₂(CH₂)₅CH₃), 3.74 (t, J=7.2 Hz, 2H,

CH₂CH₂(CH₂)₅CH₃), 7.82 and 7.95 (two s, 2H, CH=N and 6-H), 10.79 and 11.45 (two br s, 2H, NH and OH); ¹³C NMR (DMSO- d_6 + CDCl₃): δ 12.9 (CH₃), 21.2, 25.0, 27.7, 27.8 and 30.3 (CH₂(CH₂)₆CH₃), 47.6 (CH₂(CH₂)₆CH₃), 105.8 (C-5), 139.6 and 140.1 (C=N and C-6), 149.3 (C-2), 161.2 (C-4); MS (EI): m/z (%) 267 (M⁺, 84). Anal. Calcd for C₁₃H₂₁N₃O₃: C, 58.41; H, 7.92; N, 15.72. Found: C, 58.41; H, 7.86; N, 15.35.

3.2.2. 1,3-Dioctyl-5-uracilcarbaldehyde oxime (2b). This compound was obtained in 87% yield as a white solid, mp 128–130 °C; IR (Nujol): ν_{max} 3290, 3040, 1685, 1620, 1590 cm⁻¹; ¹H NMR: δ 0.83–0.87 (m, 6H, CH₃), 1.27–1.32 (m, 20H, CH₂CH₂(CH₂)₅CH₃), 1.62–1.71 (m, 4H, CH₂CH₂(CH₂)₅CH₃), 3.78 (t, *J*=7.3 Hz, 2H, CH₂CH₂-(CH₂)₅CH₃), 3.95 (t, *J*=7.1 Hz, 2H, CH₂CH₂(CH₂)₅CH₃), 7.66 and 8.13 (two s, 2H, CH=N and 6-H), 8.80 (br s, 1H, OH); ¹³C NMR: δ 14.0 (CH₃), 22.6, 26.4, 26.9, 27.5, 29.0, 29.2, 31.7 and 31.8 (CH₂(CH₂)₆CH₃), 41.8 and 50.4 (CH₂(CH₂)₆CH₃), 106.1 (C-5), 139.6 (C=N), 143.8 (C-6), 150.7 (C-2), 161.2 (C-4); MS (EI): *m/z* (%) 379 (M⁺, 11). Anal. Calcd for C₂₁H₃₇N₃O₃: C, 66.46; H, 9.83; N, 11.07. Found: C, 66.45; H, 9.42; N, 10.79.

3.2.3. 2,4-Dimethoxy-5-pyrimidinecarbaldehyde oxime (12). This compound was obtained in 87% yield as a white solid, mp 150–154 °C; IR (Nujol): ν_{max} 3200, 3010, 1580, 1550 cm⁻¹; ¹H NMR: δ 4.04 (s, 3H, OCH₃), 4.06 (s, 3H, OCH₃), 8.19 and 8.56 (two s, 2H, 6-H and CH=N), 8.85 (br s, 1H, OH); ¹³C NMR: δ 54.3 and 55.1 (OCH₃), 107.7 (C-5), 143.2 (C=N), 156.9 (C-6), 161.7 and 168.3 (C-2 and C-4); HRESIMS for C₇H₉N₃O₃ (M+Na)⁺: calcd 206.0536, found 206.0536.

3.3. Synthesis of nitrones 4 and 14

General procedure. An aqueous solution (2.5 ml) of methylhydroxylamine hydrochloride (2 mmol) and sodium carbonate (1.5 mmol) were added to an ethanolic solution (5 ml) of the aldehyde **1** or **11** (1 mmol) and the reaction mixture was stirred at room temperature for 24 h. After that the ethanol was evaporated, water was added and the mixture was extracted with methylene chloride. After drying and evaporation of the solvent from the organic layer the residue nitrones were used without further purification.

3.3.1. *N*-Methyl-*C*-(1-octyl-5-uracil) nitrone (4a). This compound was obtained in 84% yield as a white solid, mp 190–193 °C; IR (Nujol): ν_{max} 3180, 3110, 3040, 1670, 1590 cm⁻¹; ¹H NMR (45 °C): δ 0.89 (br, 3H, CH₃), 1.29–1.34 (m, 10H, CH₂CH₂(CH₂)₅CH₃), 1.75 (br t, 2H, CH₂CH₂(CH₂)₅CH₃), 3.76–3.80 (m, 5H, CH₂CH₂(CH₂)₅CH₃ and N–CH₃), 7.56 (s, 1H, 6-H), 8.81 (br s, 1H, NH), 9.90 (s, 1H, CH=N(O)); ¹³C NMR (45 °C): δ 13.8 (CH₃), 22.5, 26.5, 29.0 and 31.7 (CH₂(CH₂)₆CH₃), 47.6 (CH₂ (CH₂)₆CH₃), 53.5 (*N*–CH₃), 106.5 (C-5), 127.6 (CH=N(O)), 144.3 (C-6), 149.8 (C-2), 161.8 (C-4); MS (EI): *m/z* (%) 281 (M⁺, 86). Anal. Calcd for C₁₄H₂₃N₃O₃: C, 59.77; H, 8.24; N, 14.93. Found: C, 59.87; H, 8.07; N, 14.89.

3.3.2. *N*-Methyl-*C*-(1,3-dioctyl-5-uracil) nitrone (4b). This compound was obtained in 87% yield as a white

solid, mp 70–72 °C; IR (Nujol): ν_{max} 3070, 3030, 1695, 1630, 1590 cm⁻¹; ¹H NMR: δ 0.85–0.89 (m, 6H, CH₃), 1.26–1.32 (m, 20H, CH₂CH₂(CH₂)₅CH₃), 1.58–1.67 (m, 4H, CH₂CH₂(CH₂)₅CH₃), 3.79 (t, *J*=7.3 Hz, 2H, CH₂CH₂(CH₂)₅CH₃), 3.81 (s, 3H, *N*–CH₃), 3.96 (t, *J*=7.4 Hz, 2H, CH₂CH₂(CH₂)₅CH₃), 7.62 (s, 1H, 6-H), 9.83 (s, 1H, CH=N(O)); ¹³C NMR: δ 14.0 (CH₃), 22.5, 26.4, 26.9, 27.5, 29.0, 29.1, 29.2, 31.7 and 31.8 (CH₂(CH₂)₆CH₃), 41.8, 50.6 and 53.5 (CH₂(CH₂)₆CH₃ and *N*–CH₃), 105.6 (C-5), 128.5 (CH=N(O)), 142.5 (C-6), 150.1 (C-2), 161.4 (C-4); MS (EI): *m/z* (%) 393 (M⁺, 26). Anal. Calcd for C₂₂H₃₉N₃O₃: C, 67.14; H, 9.99; N, 10.68. Found: C, 67.50; H, 9.58; N, 10.53.

3.3.3. *N*-Methyl-*C*-(1,3-dimethoxy-5-pyrimidine) nitrone (14). This compound was obtained in 75% yield as a white solid, mp 168–170 °C; IR (Nujol): ν_{max} 3040, 1585, 1570, 1540 cm⁻¹; ¹H NMR: δ 4.04, 4.05 and 4.12 (three s, 9H, OCH₃ and *N*-CH₃), 7.57 (s, 1H, 6-H), 10.19 (s, 1H, CH=N(O)); ¹³C NMR: δ 53.9, 54.2 and 54.9 (OCH₃ and *N*-CH₃), 107.2 (C-5), 126.4 (CH=N(O)), 157.5 (C-6), 164.8 and 167.6 (C-2 and C-4); MS (EI): *m/z* (%) 197 (M⁺, 100). Anal. Calcd for C₈H₁₁N₃O₃: C, 48.73; H, 5.62; N, 21.31. Found: C, 48.62; H, 5.53; N, 21.71.

3.4. Formation of nitrile oxides 3 and 13 and reactions with the dipolarophiles 5 and 6

General procedure. A solution of the aldoxime 2 or 12 (0.5 mmol) and the dipolarophile 5 or 6 (1 mmol) in methylene chloride (5 ml) was cooled to 0 $^{\circ}$ C and commercial bleach (4 ml) was added. The reaction mixture was warmed to room temperature and allowed to react overnight with stirring. The reaction mixture was extracted with methylene chloride and the organic layer was dried over sodium sulfate. After evaporation of the solvent the residue was chromatographed on a silica gel column with hexane–ethyl acetate (3/1 for the reactions of 2b, 2/1 for reactions of 12) as the eluent.

3.4.1. 5-(5'-Benzoyloxymethyl-isoxazolin-3'-yl)-1,3dioctyluracil (7b). This compound was obtained in 70% yield as an oil; IR (liquid film): v_{max} 3060, 1710–1640, 1595, 1575 cm⁻¹; ¹H NMR: δ 0.87–0.89 (m, 6H, CH₃), 1.26-1.31 (m, 20H, CH₂CH₂(CH₂)₅CH₃), 1.61-1.70 (m, 4H, $CH_2CH_2(CH_2)_5CH_3$), 3.44 (dd, J=18.0, 7.1 Hz, 1H, 4'-H), 3.65 (dd, J = 18.0, 10.9 Hz, 1H, 4'-H), 3.78 (t, J =7.4 Hz, 2H, $CH_2CH_2(CH_2)_5CH_3$), 3.93 (t, J=7.4 Hz, 2H, CH₂CH₂(CH₂)₅CH₃), 4.38–4.49 (m, 2H, CH₂OCOPh), 4.98–5.10 (m, 1H, 5'-H), 7.42 (t, J=7.6 Hz, 2H, Ph-H), 7.56 (t, J = 7.6 Hz, 1H, Ph-H), 7.88 (s, 1H, 6-H), 8.04 (d, J =7.6 Hz, 2H, Ph-H); ¹³C NMR: δ 14.1 (CH₃), 22.7, 26.5, 26.9, 27.1, 27.2, 27.3, 27.5, 29.1, 29.2, 29.3, 31.7 and 31.8 (CH₂(CH₂)₆CH₃), 39.0 (C-4'), 41.9 and 50.5 (CH₂(CH₂)₆CH₃), 65.4 (CH₂OCOPh), 78.5 (C-5[']), 103.5 (C-5), 128.4, 129.7, 129.8 and 133.2 (C-Ph), 141.7 (C-6), 150.6 and 152.8 (C-2 and C=N), 161.0 (C-4), 166.3 (C=O); MS (EI): m/z (%) 539 (M⁺, 8). Anal. Calcd for C₃₁H₄₅N₃O₅: C, 68.99; H, 8.40; N, 7.79. Found: C, 69.27; H, 8.57; N, 7.99.

3.4.2. 5-(5'-Benzoyloxymethyl-isoxazol-3'-yl)-1,3-dioctyluracil (8b). This compound was obtained in 80% yield as a white solid, mp 47–49 °C; IR (Nujol): ν_{max} 3050, 1710, 1650, 1595 cm⁻¹; ¹H NMR: δ 0.85–0.89 (m, 6H, CH₃), 1.26–1.32 (m, 20H, CH₂CH₂(CH₂)₅CH₃), 1.62–1.78 (m, 4H, CH₂CH₂(CH₂)₅CH₃), 3.83 (t, J= 7.4 Hz, 2H, CH₂CH₂(CH₂)₅CH₃), 3.99 (t, J=7.4 Hz, 2H, CH₂CH₂(CH₂)₅CH₃), 5.46 (s, 2H, CH₂OCOPh), 7.13 (s, 1H, 4'-H), 7.46 (t, J=7.4 Hz, 2H, Ph-H), 7.60 (t, J=7.4 Hz, 1H, Ph-H), 8.05 (s, 1H, 6-H), 8.07 (d, J=7.4 Hz, 2H, Ph-H); ¹³C NMR: δ 14.0 (CH₃), 22.5, 26.4, 26.8, 26.9, 27.5, 29.0, 29.1, 29.2, 31.7 and 31.8 (CH₂(CH₂)₆CH₃), 41.9 and 50.5 (CH₂(CH₂)₆CH₃), 56.9 (CH₂OCOPh), 102.4 (C-5), 104.5 (C-4'), 128.4, 129.1, 129.9 and 133.5 (C-Ph), 141.6 (C-6), 150.6 (C-2), 156.8 (C=N), 161.9 (C-4), 165.7 and 166.3 (C=O and C-5'); HRESIMS for C₃₁H₄₃N₃O₅ (M+Na)⁺: calcd 560.3095, found 560.3098.

3.4.3. 5-(5'-Benzoyloxymethyl-isoxazolin-3'-yl)-2,4dimethoxypyrimidine (15). This compound was obtained in 65% yield as a white solid, mp 132–133 °C; IR (Nujol): ν_{max} 3030, 1710, 1595, 1535 cm⁻¹; ¹H NMR: δ 3.32 (dd, J=17.4, 6.8 Hz, 1H, 4'-H), 3.57 (dd, J=17.4, 10.9 Hz, 1H, 4'-H), 4.01 (s, 3H, OCH₃), 4.04 (s, 3H, OCH₃), 4.43–4.56 (m, 2H, CH₂OCOPh), 5.08–5.17 (m, 1H, 5'-H), 7.41 (t, J=7.6 Hz, 2H, Ph-H), 7.55 (t, J=7.6 Hz, 1H, Ph-H), 8.02 (d, J=7.6 Hz, 2H, Ph-H), 8.65 (s, 1H, 6-H); ¹³C NMR: δ 39.0 (C-4'), 54.2 and 54.4 (OCH₃), 65.4 (CH₂OCOPh), 78.0 (C-5'), 105.2 (C-5), 128.3, 129.4, 129.5 and 133.1 (C-Ph), 151.5 (C=N), 158.1 (C-6), 165.7, 166.1 and 167.9 (C-2, C-4 and C=O); MS (EI): m/z (%) 343 (M⁺, 9%). Anal. Calcd for C₁₇H₁₇N₃O₅: C, 59.47; H, 4.99; N, 12.44. Found: C, 59.34; H, 4.89; N, 12.39.

3.4.4. 5-(5'-**Benzoyloxymethyl-isoxazol-3**'-**yl**)-**2,4dimethoxypyrimidine (17).** This compound was obtained in 90% yield as a white solid, mp 98–100 °C; IR (Nujol): ν_{max} 3050, 1715, 1590, 1550 cm⁻¹; ¹H NMR: δ 4.07 (s, 3H, OCH₃), 4.11 (s, 3H, OCH₃), 5.48 (s, 2H, CH₂OCOPh), 6.82 (s, 1H, 4'-H), 7.47 (t, *J*=7.4 Hz, 2H, Ph-H), 7.58 (t, *J*= 7.4 Hz, 1H, Ph-H), 8.09 (d, *J*=7.4 Hz, 2H, Ph-H), 8.85 (s, 1H, 6-H); ¹³C NMR: δ 54.4 and 55.2 (OCH₃), 56.7 (CH₂OCOPh), 104.7 (C-5), 104.9 (C-4'), 128.5, 129.5, 129.9 and 133.6 (C-Ph), 156.6 (C=N), 158.1 (C-6), 163.5, 165.8, 166.7 and 168.1 (C-2, C-4, C-5' and C=O); MS (EI): *m/z* (%) 341 (M⁺, 23). Anal. Calcd for C₁₇H₁₅N₃O₅: C, 59.82; H, 4.43; N, 12.31. Found: C, 59.60; H, 4.53; N, 12.51.

3.5. Reactions of nitrones 4 and 14 with the dipolarophile 5

General procedure. A solution of the nitrone 4 or 14 (0.5 mmol) and the dipolarophile 5 (1 mmol) in xylene (5 ml) was heated to reflux and the reaction was monitored by TLC until the consumption of the nitrone. After 2 days only traces of the nitrone were detected in the TLC. The heating was stopped and after evaporation of the solvent the residue was chromatographed on a silica gel column with hexane–ethyl acetate (1/1 for the reaction of 4a, 3/1 for the reaction of 4b, 2/1 for the reaction of 14) as the eluent.

3.5.1. (3'*R*S,5'*SR*)-5-(5'-Benzoyloxymethyl-isoxazolidin-3'-yl)-1-octyluracil (9a). This compound was obtained in 72% yield as an oil; IR (liquid film): ν_{max} 3190, 3060, 1715–1650, 1595, 1575 cm⁻¹; ¹H NMR: δ 0.87 (t, *J*= 8.5 Hz, 3H, CH₃), 1.15–1.40 (m, 10H, CH₂CH₂(CH₂)₅CH₃), 1.50–1.65 (m, 2H, CH₂CH₂(CH₂)₅CH₃), 2.10 (dt, *J*=12.2, 5.1 Hz, 1H, 4'-H), 2.73 (s, 3H, *N*–CH₃), 3.02 (ddd, J=12.2, 8.4, 7.3 Hz, 1H, 4'-H), 3.48–3.76 (m, 2H, CH₂CH₂(CH₂)₅CH₃), 4.03 (dd, J=7.3, 5.1 Hz, 1H, 3'-H), 4.31 (dd, J=12.0, 6.0 Hz, 1H, CH₂OCOPh), 4.47 (dd, J=12.0, 3.3 Hz, 1H, CH₂OCOPh), 4.67 (dddd, J=8.4, 6.0, 5.1, 3.3 Hz, 1H, 5'-H), 7.41 (t, J=7.4 Hz, 2H, Ph-H), 7.43 (s, 1H, 6-H) 7.55 (t, J=7.4 Hz, 1H, Ph-H), 7.98 (d, J=7.4 Hz, 2H, Ph-H), 9.81 (s, 1H, NH); ¹³C NMR: δ 13.9 (CH₃), 22.5, 26.3, 28.9, 29.0 and 31.6 (CH₂(CH₂)₆CH₃), 37.5 (C-4'), 44.1 and 48.7 (CH₂(CH₂)₆CH₃ and *N*–CH₃), 63.1 and 64.9 (C-3' and CH₂OCOPh), 74.6 (C-5'), 113.6 (C-5), 128.3, 129.4, 129.6 and 133.1 (C-Ph), 141.7 (C-6), 150.5 (C-2), 163.5 (C-4), 166.1 (C=O); MS (EI): *m*/*z* (%) 443 (M⁺, 10). Anal. Calcd for C₂₄H₃₃N₃O₅: C, 64.99; H, 7.50; N, 9.57. Found: C, 65.11; H, 7.50; N, 9.24.

3.5.2. (3'RS,5'SR)-5-(5'-Benzoyloxymethyl-isoxazolidin-3'-yl)-1,3-dioctyluracil (9b). This compound was obtained in 70% yield as an oil; IR (liquid film): v_{max} 3060, 1720–1690, 1660–1630, 1590 cm⁻¹; ¹H NMR: δ 0.85–0.92 (m, 6H, CH₃), 1.15–1.40 (m, 20H, CH₂CH₂(CH₂)₅CH₃), 1.50-1.70 (m, 4H, CH₂CH₂(CH₂)₅CH₃), 2.05 (dt, $J = 13.6, 5.1 \text{ Hz}, 1\text{H}, 4'-\text{H}), 2.73 \text{ (s, 3H, } N-\text{CH}_3),$ 3.02 (ddd, J=13.6, 8.4, 7.8 Hz, 1H, 4'-H), 3.49–3.75 (m, 2H, $CH_2CH_2(CH_2)_5CH_3$), 3.90 (t, J=9.3 Hz, 2H, $CH_2CH_2(CH_2)_5CH_3$, 3.99 (dd, J=7.8, 5.1 Hz, 1H, 3'-H), 4.32 (dd, J = 11.9, 6.1 Hz, 1H, CH_2OCOPh), 4.43 (dd, J =11.9, 3.1 Hz, 1H, CH₂OCOPh), 4.67 (dddd, J=8.4, 6.1, 5.1, 3.1 Hz, 1H, 5'-H), 7.37 (s, 1H, 6-H), 7.41 (t, J=7.6 Hz, 2H, Ph-H), 7.56 (t, J=7.6 Hz, 1H, Ph-H), 7.97 (d, J=7.6 Hz, 2H, Ph-H); ¹³C NMR: δ 14.0 (CH₃), 22.5, 26.4, 26.9, 27.5, 28.9, 29.0, 29.1, 31.6 and 31.7 (CH₂(CH₂)₆CH₃), 37.7 (C-4'), 41.4, 44.2 and 49.7 (CH₂(CH₂)₆CH₃ and N-CH₃), 63.7 and 65.1 (C-3' and CH₂OCOPh), 74.5 (C-5'), 112.9 (C-5), 128.3, 129.5, 129.7 and 133.1 (C-Ph), 139.3 (C-6), 150.8 (C-2), 162.6 (C-4), 166.2 (C=O); MS (EI): m/z (%) 555 (M⁺ , 16). Anal. Calcd for C₃₂H₄₉N₃O₅: C, 69.16; H, 8.89; N, 7.56. Found: C, 69.46; H, 8.81; N, 7.25.

3.5.3. (3'RS,5'SR)-5-(5'-Benzoyloxymethyl-isoxazolidin-3'-yl)-2,4-dimethoxypyrimidine (19). This compound was obtained in 52% yield as an oil; IR (liquid film): v_{max} 3060, 1715, 1595, 1565 cm⁻¹; ¹H NMR: δ 2.05 (dt, J=12.8, 6.4 Hz, 1H, 4'-H), 2.68 (s, 3H, N-CH₃), 2.89 (ddd, J=12.8, 8.4, 7.7 Hz, 1H, 4'-H), 3.85 (dd, J=8.4, 6.4 Hz, 1H, 3'-H), 3.97 (s, 3H, OCH₃), 4.00 (s, 3H, OCH₃), 4.37 (dd, J = 11.5, 3.9 Hz, 1H, CH₂OCOPh), 4.45 (dd, J=11.5, 7.1 Hz, 1H, CH_2OCOPh), 4.61 (dddd, J=7.7, 7.1, 6.4, 3.9 Hz, 1H, 5'-H), 7.37 (s, 1H, 6-H), 7.42 (t, J=7.6 Hz, 2H, Ph-H), 7.54 (t, J = 7.6 Hz, 1H, Ph-H), 8.01 (d, J = 7.6 Hz, 2H, Ph-H); ¹³C NMR: δ 38.9 (C-4′), 43.5 (*N*–CH₃), 53.9 (OCH₃), 54.7 (OCH₃), 64.1 and 66.0 (C-3['] and CH₂OCOPh), 74.3 (C-5[']), 113.1 (C-5), 128.2, 129.6, 129.8 and 132.9 (C-Ph), 156.4 (C-6), 164.6, 166.4 and 168.7 (C-2, C-4 and C=O); MS (EI): m/z (%) 359 (M⁺, 17). Anal. Calcd for C₁₈H₂₁N₃O₅: C, 60.16; H, 5.89; N, 11.69. Found: C, 60.28; H, 6.10; N, 11.39.

3.5.4. (3'*R***R**,5'*S***S**)-5-(5'-Benzoyloxymethyl-isoxazolidin-3'-yl)-2,4-dimethoxy-pyrimidine (20). This compound was obtained in 26% yield as an oil; IR (liquid film): ν_{max} 3060, 1710, 1660, 1600–1560 cm⁻¹; ¹H NMR: δ 2.41 (ddd, J=14.2, 7.7, 5.7 Hz, 1H, 4'-H), 2.55 (ddd, J=14.2, 8.9, 3.9 Hz, 1H, 4'-H), 2.69 (s, 3H, *N*–CH₃), 4.00 (s, 3H, OCH₃), 4.02 (s, 3H, OCH₃), 4.24 (dd, J=5.7, 3.9 Hz, 1H, 3'-H), 4.41 (dd, J=12.2, 6.4 Hz, 1H, CH₂OCOPh), 4.50–4.61 (m, 2H, CH₂OCOPh and 5'-H), 7.47 (t, J=7.4 Hz, 2H, Ph-H), 7.57 (t, J=7.4 Hz, 1H, Ph-H), 8.11 (d, J=7.4 Hz, 2H, Ph-H), 8.35 (s, 1H, 6-H); ¹³C NMR: δ 38.7 (C-4'), 43.8 (*N*–CH₃), 54.0 (OCH₃), 54.7 (OCH₃), 64.0 and 65.4 (C-3' and CH₂OCOPh), 74.8 (C-5'), 112.6 (C-5), 128.3, 129.6, 129.9 and 133.0 (C-Ph), 156.6 (C-6), 164.6, 167.6 and 169.3 (C-2, C-4 and C=O); MS (EI): *m*/*z* (%) 359 (M⁺, 14). Anal. Calcd for C₁₈H₂₁N₃O₅: C, 60.16; H, 5.89; N, 11.69. Found: C, 60.26; H, 5.99; N, 11.30.

3.6. Demethylation of compounds 15, 17 and 19

General procedure. The dimethoxy derivative **15** or **17** or **19** (0.2 mmol) was heated with sodium iodide (0.1 g) in glacial acetic acid (3 ml) at 90 °C for 1 h. The solvent was removed under reduced pressure and the residue was chromatographed on a silica gel column with 3% methanol in methylene chloride as the eluent.

3.6.1. 5-(**5**'-**Benzoyloxymethyl-isoxazolin-3**'-**yl**)-**uracil** (**16**). This compound was obtained in 72% yield as a white solid, mp 253–257 °C; IR (Nujol): ν_{max} 3210, 3080, 3040, 1710, 1640 cm⁻¹; ¹H NMR (CDCl₃/DMSO-*d*₆): δ 3.35 (dd, *J*=17.8, 6.9 Hz, 1H, 4'-H), 3.52 (dd, *J*= 17.8, 11.0 Hz, 1H, 4'-H), 4.33 (dd, *J*=12.3, 5.5 Hz, 1H, CH₂OCOPh), 4.42 (dd, *J*=12.3, 3.5 Hz, 1H, CH₂OCOPh), 4.42 (dd, *J*=17.7 Hz, 1H, Ph-H), 7.48 (t, *J*= 7.7 Hz, 2H, Ph-H), 7.63 (t, *J*=7.7 Hz, 1H, Ph-H), 7.77 (s, 1H, 6-H), 7.96 (d, *J*=7.7 Hz, 2H, Ph-H), 10.12 (br s, 2H, NH); ¹³C NMR (CDCl₃/DMSO-*d*₆): δ 36.5 (C-4'), 63.7 (CH₂OCOPh), 75.5 (C-5'), 101.1 (C-5), 126.7, 127.4, 127.7 and 131.4 (C-Ph), 139.6 (C-6), 148.9 (C-2), 150.4 (C=N), 160.1 (C-4), 163.4 (C=O); HRESIMS for C₁₅H₁₃N₃O₅ (M+Na)⁺: calcd 338.0747, found 338.0748.

3.6.2. 5-(**5**'-**Benzoyloxymethyl-isoxazol-3**'-**yl**)-**uracil** (**18**). This compound was obtained in 67% yield as a white solid, mp 217–220 °C; IR (Nujol): ν_{max} 3210, 3080, 3050, 1715, 1590 cm⁻¹; ¹H NMR (CDCl₃/DMSO-*d*₆): δ 5.47 (s, 2H, C*H*₂OCOPh), 7.02 (s, 1H, 4'-H), 7.51 (t, *J*=7.4 Hz, 2H, Ph-H), 7.64 (t, *J*=7.4 Hz, 1H, Ph-H), 8.01–8.07 (overlapped d and s, 3H, Ph-H and 6-H), 11.34–11.52 (overlapped br s, 2H, NH); ¹³C NMR (CDCl₃/DMSO-*d*₆): δ 55.1 (*C*H₂OCOPh), 100.0 (C-5), 102.7 (C-4'), 126.9, 127.7, 128.0 and 131.9 (C-Ph), 139.5 (C-6), 149.3, 155.1, 160.6 163.5 and 164.6 (C-2, C-4, C=N C-5' and C=O); HRESIMS for C₁₅H₁₁N₃O₅ (M+Na)⁺: calcd 336.0591, found 336.0591.

3.6.3. (*J*'*RS*,5'*SR*)-**5**-(5'-Benzoyloxymethyl-isoxazolidin-*J*'-yl)-uracil (21). This compound was obtained in 76% yield as a white solid, mp 210–212 °C; IR (Nujol): ν_{max} 3190, 3150, 3060, 1710, 1660 cm⁻¹; ¹H NMR (CDCl₃/CD₃OD): δ 2.03 (dt, *J*=12.9, 6.1 Hz, 1H, 4'-H), 2.72 (s, 3H, *N*–CH₃), 2.98 (ddd, *J*=12.9, 9.2, 7.4 Hz, 1H, 4'-H), 3.85 (dd, *J*=7.4, 6.1 Hz, 1H, 3'-H), 4.33–4.42 (m, 2H, CH₂OCOPh), 4.61–4.70 (m, 1H, 5'-H), 7.42 (s, 1H, 6-H), 7.43 (t, *J*=7.4 Hz, 2H, Ph-H), 7.56 (t, *J*=7.4 Hz, 1H, Ph-H), 8.00 (d, *J*=7.4 Hz, 2H, Ph-H); ¹³C NMR (CDCl₃/CD₃OD): δ 37.7 (C-4'), 43.9 (*N*–CH₃), 63.2 and 65.3 (C-3' and CH₂OCOPh), 74.6 (C-5'), 113.0 (C-5), 128.3, 129.5 and 133.2 (C-Ph), 138.2 (C-6), 151.6 (C-2), 163.9 (C-4), 166.5 (C=O); MS: m/z (%) 331 (M⁺, 10). Anal. Calcd for C₁₆H₁₇N₃O₅: C, 58.00; H, 5.17; N, 12.68. Found: C, 57.98; H, 5.01; N, 12.82.

3.7. Hydrolysis of compounds 9

General procedure. An aqueous solution (1 ml) of KOH (10%) was added to a methanolic solution (5 ml) of the compound **9a** or **9b** (0.1 mmol) and the reaction mixture was stirred at room temperature for 24 h. After that the methanol was evaporated, water was added, neutralized with ammonium chloride and the mixture was extracted with methylene chloride. The organic layer was dried over sodium sulfate and after evaporation of the solvent compounds **22** were obtained quantitatively as oils. For analytical purposes, they were further purified by column chromatography on a silica gel column with ethyl acetate as the eluent.

3.7.1. (3'RS,5'SR)-5-(5'-Hydroxymethyl-isoxazolidin-3'-yl)-1-octyluracil (22a). This compound was obtained in 100% yield as an oil; IR (liquid film): ν_{max} 3400, 3180, 3050, 1690–1640 cm⁻¹; ¹H NMR: δ 0.87 (t, J=6.6 Hz, 3H, CH₃), 1.19–1.40 (m, 10H, CH₂CH₂(CH₂)₅CH₃), 1.59–1.73 $(m, 2H, CH_2CH_2(CH_2)_5CH_3), 2.01 (dt, J = 12.9, 5.9 Hz, 1H,$ 4'-H), 2.50 (br s, 1H, OH), 2.69 (s, 3H, N-CH₃), 2.91 (dt, J = 12.9, 7.9 Hz, 1H, 4'-H), 3.59 (dd, J = 12.8, 5.3 Hz, 1H,CH₂OH), 3.64–3.79 (m, 3H, CH₂CH₂(CH₂)₅CH₃ and CH₂OH), 3.90 (dd, J=7.9, 5.9 Hz, 1H, 3'-H), 4.36–4.46 (m, 1H, 5'-H), 7.48 (s, 1H, 6-H), 9.62 (br s, 1H, NH); 13 C NMR: δ 14.0 (CH₃), 22.5, 26.4, 29.1, 29.7 and 31.6 (CH₂(CH₂)₆CH₃), 37.7 (C-4'), 44.0 and 49.1 (CH₂(CH₂)₆CH₃ and N-CH₃), 63.6 and 64.5 (C-3' and CH₂OH), 76.6 (C-5'), 112.9 (C-5), 141.7 (C-6), 150.5 (C-2), 163.5 (C-4); MS (EI): *m/z* (%) 339 (M⁺ 8). Anal. Calcd for C₁₇H₂₉N₃O₄: C, 60.15; H, 8.61; N,12.38. Found: C, 60.01; H, 8.90; N, 12.14.

3.7.2. (3'RS,5'SR)-5-(5'-Hydroxymethyl-isoxazolidin-3'-yl)-1,3-dioctyluracil (22b). This compound was obtained in 100% yield as an oil; IR (liquid film): ν_{max} 3400, 3060, 1690, 1660–1630 cm⁻¹; ¹H NMR: δ 0.85–0.92 (m, 6H, CH₃), 1.14-1.41 (m, 20H, CH₂CH₂(CH₂)₅CH₃), 1.50-1.75 (m, 4H, $CH_2CH_2(CH_2)_5CH_3$, 1.97 (dt, J=12.6, 6.4 Hz, 1H, 4'-H), 2.30 (br s, 1H, OH), 2.68 (s, 3H, N-CH₃), 2.87 (dt, J=12.6, 8.3 Hz, 1H, 4'-H), 3.59 (dd, J=11.9, 5.2 Hz, 1H, CH₂OH), 3.69-3.80 (m, 3H, CH₂CH₂(CH₂)₅CH₃ and CH₂OH), 3.85–3.96 (m, 3H, $CH_2CH_2(CH_2)_5CH_3$ and 3'-H), 4.34–4.43 (m, 1H, 5'-H), 7.35 (s, 1H, 6-H); ¹³C NMR: δ 14.0 (CH₃), 22.6, 26.5, 27.0, 27.5, 29.1, 29.2, 29.7, 31.7 and 31.8 (CH₂(CH₂)₆CH₃), 38.0 (C-4'), 41.6, 44.0 and 50.0 (CH₂(CH₂)₆CH₃ and N-CH₃), 64.1 and 64.9 (C-3' and CH₂OH), 76.9 (C-5[']), 112.1 (C-5), 139.2 (C-6), 150.8 (C-2), 162.7 (C-4); MS (EI): *m/z* (%) 451 (M⁺, 9). Anal. Calcd for C₂₅H₄₅N₃O₄: C, 66.48; H, 10.04; N, 9.30. Found: C, 66.26; H, 10.21; N, 9.25.

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Optical sensor arrays: one-pot, multiparallel synthesis and cellulose immobilization of pH and metal ion sensitive azo-dyes

Tommaso Carofiglio,^{a,*} Carlo Fregonese,^a Gerhard J. Mohr,^b Federico Rastrelli^a and Umberto Tonellato^a

^aDepartimento di Scienze Chimiche, Università di Padova, Via Marzolo 1, 35131 Padova, Italy ^bInstitute of Phys. Chem., Friedrich-Schiller University Jena, Lessing St. 10, D-07743 Jena, Germany

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Abstract—A sixteen component array of acid–base and metal ion sensors was synthesized and covalently immobilized onto a transparent cellulose-based membrane. Dye synthesis and cellulose dyeing were carried out in one-pot, parallel, microscale reactions not requiring any isolation or purification steps. In addition, pH and metal ion optical sensing properties of the sixteen component array have been tested using a high-throughput screening based on digital imaging analysis.

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1. Introduction

Chemical sensors are now recognized as a valid alternative to conventional instruments. This is especially true for analytical problems requiring on-site and realtime acquisition of data such as process control, environmental and biomedical monitoring.¹ Research in chemical sensors has traditionally been dominated by the quest for substance-specific detectors. This makes the development of new chemical sensors a formidable challenge. However, an emerging strategy that obviates the need for absolute selectivity is based on the use of broadly-tuned, spatially addressable sensing elements arranged in array configuration.² With this approach, inspired by mammalian senses of smell and taste, identification of the analyte is achieved via deconvolution of the multiplexed array response using smart data processing methods. This has produced some outstanding applications such as the so-called artificial noses and tongues.³

Recently, we focussed our research into developing arrays of optical sensors covalently attached onto transparent supports, especially for metal ion detection in aqueous solution. Covalent bonding is the most reliable method for immobilizing probe molecules since it excludes the problem of leaching. However, many grafting procedures described so far involve several chemical steps for modification of both the indicator and the support matrix before performing the actual coupling reaction. Therefore, we surveyed the literature in order to find simpler means for manufacturing chemical sensors in array format. Ideally, such methods should involve a straightforward sequence of high yield reactions so that purification steps can be reduced or totally eliminated. In addition, parallel synthesis, miniaturization, and a combinatorial analysis should also become possible. We reasoned that azo-dyes could address most of the above issues for the following reasons: (i) azo-dye synthesis is based only on two modular reactions⁴ (i.e., diazotization and azo-coupling); (ii) an almost unlimited choice of commercially available starting compounds allows for accessing the structural diversity required for combinatorial chemistry; (iii) the well-established reactive-dye technology can be exploited for their covalent grafting to a given substrate;⁴ (iv) the intense color of azo-dyes enables the use of digital image analysis as the highthroughput method for screening the libraries.⁵

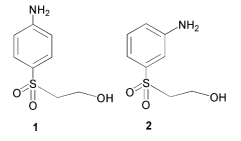
Within this context, we describe a procedure for arraying pH and metal ion sensitive azo-dyes onto transparent cellulose-based membranes.

Keywords: Sensor arrays; Azo-dyes.

^{*} Corresponding author. Tel.: +39 049 827 5670; fax: +39 049 827 5239; e-mail: tommaso.carofiglio@unipd.it

2. Results and discussion

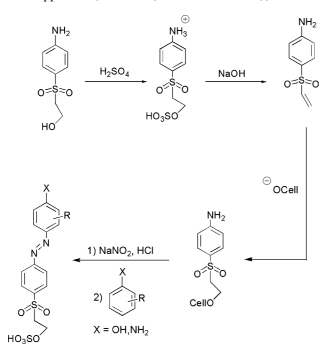
The method reported here exploits the hetero-bifunctional scaffold **1** (or **2**, Scheme 1) formerly reported in the literature for the production of nonleaching pH-sensors supported onto surface hydrolysed cellulose acetate overhead transparencies⁶ or polyvinyl alcohol based materials.⁷



Scheme 1. Hetero-bifunctional scaffolds

The aniline group allows for the synthesis of the azochromophore (via diazotization followed by coupling with aromatic amines or phenols) whereas the 2-hydroxyethyl sulfonyl moiety provides a handle for attaching the dye onto supports bearing nucleophilic hydroxyls. The procedure for making single-sensor based systems implies three multi-step stages: (1) azo-dye synthesis and isolation, (2) formation of the vinylsulfone reactive group from the 2-hydroxyethyl sulfonyl moiety (hereafter referred to as 'dye activation'), and (3) cellulose dyeing. In principle, this procedure could also be applied for preparing arrays of chemical sensors. However, all array members must be separately synthesized, purified, and characterised before array fabrication.

In an initial attempt to optimize this procedure for array production we envisaged a direct and parallel synthesis of dye molecules on the hydrolysed cellulose acetate membrane. Such approach (Scheme 2) would involve: (i) covalent



Scheme 2. Direct parallel synthesis onto cellulose.

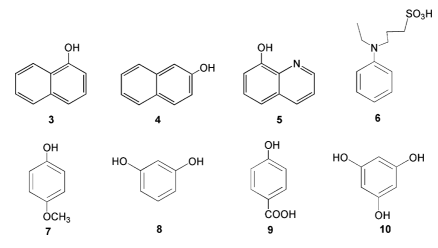
attachment of the scaffold molecules 1 or 2 to cellulose membrane (using the same chemistries involved in dye activation) followed by (ii) the in situ synthesis of dyes in spatially indexed spots of the activated polymer surface. Unfortunately, the strongly acidic conditions required for the diazotization reaction proved to be detrimental for mechanical and transparency properties of the cellulose acetate support.

As an alternative, we then considered the multiparallel synthesis and activation of dye molecules followed by simultaneous printing into the cellulose membrane. According to this approach, dye isolation would be necessary both for obtaining the dye in analytically pure form and for its activation process in concentrated sulfuric acid (vide infra). However, dye isolation is troublesome, especially to perform in parallel, since it frequently requires salting-out, slow filtrations, and reiterative chromatographic purification steps. Furthermore, dye activation must be replicated for all the dyes that need to be incorporated into the array.

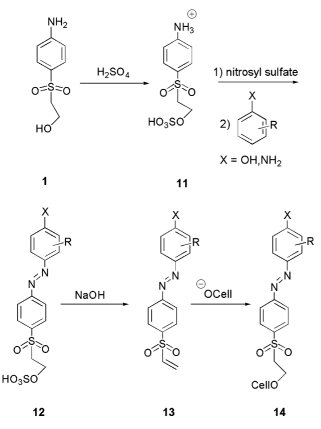
To address these issues we devised the following strategy: (i) synthesis of the common precursor 2-sulfatoethylsulfone of 1 (or 2) and its one-pot diazotization; (ii) parallel azocoupling to generate the dye library and (iii) array formation by parallel dyeing of cellulose. In order to demonstrate the validity of this approach, a library of 16 azo-dyes was synthesized by combining compounds 1 or 2 with aromatic derivatives 3-10 (Scheme 3) according to the procedure depicted in Scheme 4 for compound 1.

Treating 1 with concentrated sulfuric acid gave the sulfate ester (11). The subsequent diazotisation step was accomplished using nitrosyl sulfate (instead of the more common NaNO₂/HCl) to preserve the reaction medium necessary for a one-pot synthesis. As soon as the diazonium salt was formed, the synthesis of the azo-dyes advanced in a parallel fashion inside polypropylene micro centrifuge tubes. Compounds 3-10 (Scheme 1) were mixed with a substoichiometric amount of the diazonium salt of 11 to encourage quantitative conversion in the azo-coupling reaction. At this stage, dye isolation could be avoided because target dyes are the only species in the reaction mixture able to react with cellulose (i.e., containing the 2-hydroxyethylsulfonyl moiety). In this way, the subsequent cellulose dyeing (after activation) also performs the in situ purification of the target molecules.

The solution pH is the experimental factor requiring most careful control in azo-coupling reactions.⁴ Indeed, the ideal pH range depends on the particular coupling component used. Phenols typically require alkaline conditions whereas aromatic amines react in acidic or neutral solution. Although the coupling components chosen for this study were, apart from compound 6, all phenol derivatives, we wished to find a protocol of general use. Thus, we decided to gradually increase the pH by adding two subsequent portions of sodium hydroxide solution to each vial making the pH first almost neutral (for coupling of amines), then slightly alkaline (for coupling of phenols). It should be noted that stronger alkaline conditions are disadvantageous at this point not only since they would promote diazonium salt decomposition but also because they would cause premature formation of the vinylsulfone reactive group.



Scheme 3. Coupling molecules for dye formation.



Scheme 4. One-pot synthetic protocol for array manufacturing.

Before parallel dyeing, cellulose acetate membranes were hydrolysed with 0.1 M NaOH for 30 min in order to form a layer of cellulose. Then, dyeing was performed by transferring a suitable amount of the above reaction mixture in a homemade teflon reactor containing a 4×2 array of 8 mm diameter wells (see electronic Supplementary information for details) and adding a concentrated solution of NaOH to promote the formation of vinyl sulfone and cellulosate anion.

The reactor allowed immobilization of the sensor molecules only on one side of the cellulose substrate with minimum volume of the reaction mixture. The entire procedure was repeated for scaffold **2** and the resulting dyes were patterned in the same membrane thus obtaining a 16 member array. As an additional benefit, since isolation of dyes is not required, the reactions can be performed on a small scale. Indeed, microarrays containing spots of about 1.5 mm diameter were prepared using adhesive membranes in PVC (see electronic Supplementary information). It has been possible to fit 16 dyes in about 1 cm² and only about 2 μ L of dying solution was necessary to obtain each spot. Figure 1 reports two images of the same membrane after exposition to an acidic (pH=2.06) and to a basic solution (pH=12.03). Simple visual inspection shows, which dyes are most applicable, for example because of a more intense dyeing and/or a marked chromogenic difference between acid and base forms.

In order to prove that the one-pot approach yielded the desired indicator dyes in pure form, some of the spots were cut out from the array and glued to disposable plastic cuvettes. The UV–vis spectra were collected in acidic and basic conditions and compared with those obtained upon dying cellulose with the same dyes synthesized according to the published procedures.⁶ The matching of the UV–vis spectra (Fig. 2) confirms the cleanness of our procedure.[†]

In order to evaluate our sensor arrays under different experimental conditions (i.e., pH or metal ion concentration) within a reasonable timeframe, we decided to use digital image analysis in a high-throughput approach.⁸ To this end, images of the membranes were acquired using a plane scanner. Then, image post-processing was carried out using a script written in Matlab. The program first located the coordinates of the spots, next it randomly sampled a circular area of fixed radius around each centroid and it calculated the average color coded in the CIE L^*a^*b color space.

Figure 3 shows a titration plot obtained by this method for

[†] UV-vis matching was confirmed also for spots 2A, 3A, 4A, and 1B (data not reported here). Authentic dyes were synthesized according to the procedure in Ref. 6 The same test was considered unessential for spots 1C, 2C, 3C, 4C, and 1D since they derived from scaffold 2, which is a positional isomer of 1. On the other hand, the remaining spots (2B, 3B, 4B, 2D, 3D, 4D) displayed a low dyeing of cellulose membrane and a minimal pH/metal ion response thus were not further investigated.

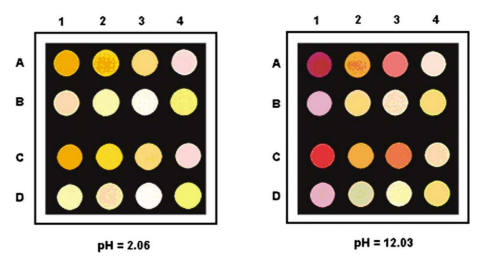


Figure 1. Membrane images in acidic (pH=2.06) and basic (pH=12.03) conditions. Spot identification (scaffold-coupling agent): 1A(1-3), 2A(1-4), 3A(1-5), 4A(1-6), 1B(1-7), 2B(1-8), 3B(1-9), 4B(1-10), 1C(2-3), 2C(2-4), 3C(2-5), 4C(2-6), 1D(2-7), 2D(2-8), 3D(2-9), 4D(2-10). Black background results from superimposing a black mask using a photo-editing software.

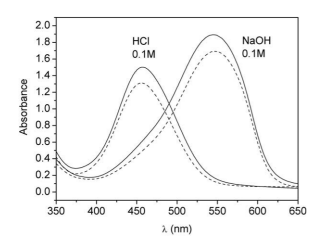


Figure 2. Spectral comparison of layers prepared by the one-pot approach from scaffolds 1 and 3 (dotted lines) and layers prepared by the conventional method (continuous lines).

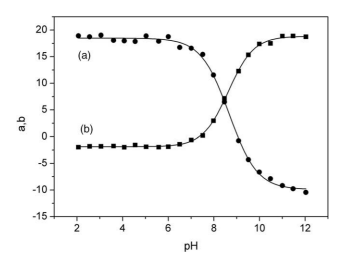


Figure 3. Effect of pH variation on red/green (a) and yellow/blue (b) coordinates for spot 1B.

the dye at the position 1B (corresponding to the 1–7 sensor) of the array. The plot reports only the red/green coordinate (*a*) and the yellow/blue coordinate (*b*) since the lightness coordinate (*L*) was almost unaffected by the variation of pH. Fitting of the two curves gave a pK_a of 8.6, which is in good agreement with the value found using the conventional UV–vis spectroscopic method, thus validating the present procedure.

As a further application, we used our sensor array for the detection of metal ions in aqueous solution, stimulated by the metal chelating properties of azo-dyes.⁹ In a preliminary study, we detected the color changes (ΔE^{\ddagger}) of the different dyes on the array upon exposure to 12 different metal ions (0.1 mM in 0.1 M MOPS buffer, pH=7.0). We obtained a set of data, which is plotted in Figure 4. From these data it is evident that coupling agents **3**, **4**, and **5**, when reacted with both scaffold **1** or **2**, gave azo-dyes, which respond to all the metal ions examined, albeit with different sensitivity. Thus, such compounds are good candidates for the construction of an array of cross-reactive sensors. On the other hand, if the goal were to find an ion-specific sensor, a careful examination of the array response showed that coupling agent **5**, especially in conjunction with scaffold **2**, gave a sensor, which is selective for cobalt(II) ions.

3. Conclusion

In conclusion, we have reported a simple procedure for the construction of arrays of azo-dyes covalently immobilized onto a transparent cellulose membrane. Both the syntheses of the dyes and the chemistries involved in the immobilization stage have been carried out in parallel, in microscale amount, in a one-pot fashion, and without any isolation

^{*} $\Delta E = ((a-a_0)^2 + (b-b_0)^2 + (c-c_0)^2)^{1/2}$ where a_0 , b_0 , L_0 and a, b, L are the coordinates in the L^*a^*b CIELAB space of an initial color taken as a reference (i.e., the color of an indicator spot under acidic conditions or before the addition of a metal ion) and a generic color (i.e., after increasing the pH or after addition of a metal ion) respectively. In other words, ΔE represents the Euclidean distance between two colors in the L^*a^*b CIELAB space.

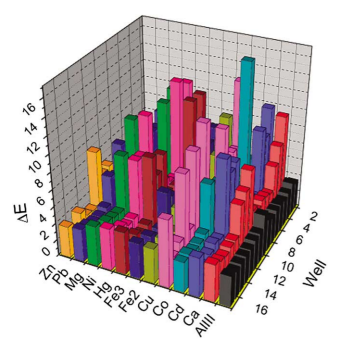


Figure 4. Color changes (ΔE) upon membrane exposition to metal ions (0.1 mM in 0.1 M MOPS buffer, pH=7.0. Spot identification (scaffold-coupling agent): 1(1–3), 2(1–4), 3(1–5), 4(1–6), 5(1–7), 6(1–8), 7(1–9), 8(1–10), 9(2–3), 10(2–4), 11(2–5), 12(2–6), 13(2–7), 14(2–8), 15(2–9), 16(2–10).

steps. Furthermore, the array response to several metal ions in aqueous solution was studied. A detailed chemometric evaluation of these data, also in terms of reproducibility and metal ion selectivity at different pH values, will be the subject of a future report.

4. Experimental

4.1. General

Cellulose triacetate membranes were purchased from Folex Film Systems (22.5×30 cm, no. 35302). Before dyeing, membranes were immersed in a solution of NaOH 0.1 M for 30 min then carefully washed with milli-Q water. UV–vis spectra were collected using a Perkin-Elmer Lambda 45 Spectrophotometer. Images of the membranes were acquired (300 dpi resolution, 24 bit color) using a plane scanner (Epson Perfection 1240U).

4.2. Array preparation

4.2.1. Formation of the sulfate ester of 1 (or 2). In a 10 mL glass vial, 100 mg of compound **1** (or **2**) (as the hydrochloride salt, 0.42 mmol) were dissolved in 96% sulfuric acid (1.0 mL) and stirred for 30 min at room temperature.

4.2.2. Diazotization. The sulfate ester solution was cooled down to 0 °C. Then a solution of NaNO₂ (35 mg, 0.51 mmol) in 96% H₂SO₄ (0.5 mL) was added drop-wise. After 15 min the solution was carefully diluted with 2.0 mL of cold water (hereafter referred to as DIAZO solution).

4.2.3. Azo-coupling. Coupling was carried out in a parallel fashion in 1 mL vials. Solutions of coupling compounds **3–10** (200 μ L of 25 mM solution in NaOH 0.1 M, 5.0 μ mol)) were mixed with DIAZO solution (20 μ L, 2.4 μ mol). After 5 min a portion of NaOH was added (2 M, 100 μ L, 400 μ mol) to each vial in order to render the solution neutral/slightly alkaline.

4.2.4. Membrane dyeing. After 15-30 min another portion of NaOH was added (2 M, 200 µL, 200 µmol) to each vial to render the solution strongly alkaline and promote the formation of the vinylsulfone reactive species. A portion of each reaction mixture (150μ L) was introduced into the teflon reactor (see Supplementary data). The teflon array was covered with the membrane, sealed and turned up-side-down so that the solution came into contact with the cellulose membrane. After 1 h the membrane was removed from the reactor, washed abundantly with water and left to dry in air.

4.3. Image processing

Image post-processing has been carried out by means of a custom software, which exploits the interoperability among graphic user interfaces (GUIs), scripts and functions within the Matlab [Matlab 6.5 R13, The MathWorks, Inc.1984-2002] programming environment. In order to isolate the relevant color areas, the $m \times n$ source image is first imported into Matlab workspace as an $m \times n \times 3$ RGB tensor, which is subsequently downgraded to an $m \times n$ grayscaled matrix. Such $m \times n$ grayscaled matrix is then converted into an $m \times n$ binary matrix representing a mask of the original image. To compensate for slight misalignments in the arrayed cells, a down-sampled version of the binary mask is fed into a k-means clustering algorithm, with the purpose of computing the centroids of the original color spots. As the next step, the software randomly samples a number of points in a disk of fixed radius around each centroid coordinates and it calculates the average color in the RGB space. Finally, RGB values are converted to L^*a^*b CIELAB space, which is a uniform color space.¹⁰

Acknowledgements

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Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.tet.2005.11. 016.

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Synthesis of alkoxymethyl derivatives of resorcinarene via the Mannich reaction catalysed with iminodiacetic acid

Mariusz Urbaniak and Waldemar Iwanek*

Institute of Chemistry, Pedagogical University, Checińska 5, 25-020 Kielce, Poland

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Abstract—The conditions of a simple synthesis of tetra(alkoxymethyl) derivatives of resorcinarene via the Mannich reaction catalysed with iminodiacetic acid are described. A possibility of high yield synthesis of such derivatives for the selected straight-chain alcohols is shown. A possible mechanism of this reaction is suggested.

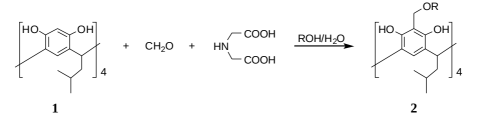
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1. Introduction

The Mannich reaction is one of the most important multicomponent reactions in the field of organic chemistry.¹ It is often also employed in the chemistry of resorcinarenes to prepare, inter alia, the derivatives of resorcinarenes such as: aminomethyl-, oxazine-, oxazolidine-, bridged and partially functionalised resorcinar-enes. $^{2-6}$

NMR spectra demonstrate unequivocally that this product is a tetra(methoxymethyl) derivative 2a of resorcinarene 1 (Scheme 1). Higher straight-chain alkyl alcohols also undergo this reaction easily and very efficiently.

Table 1 reports the yields of formation of tetra(alkoxymethyl) derivatives 2a-f from various alcohols in the presence of stoichiometric amount of iminodiacetic acid. The reaction was carried out under reflux for 12 h.



Scheme 1.

This paper presents the synthesis of alkoxymethyl derivatives of resorcinarene via the Mannich reaction catalysed using iminodiacetic acid. Recently, Rissanen et al. described the synthesis of such derivatives using trihydroxymethylmethylamine.⁷ Our method of synthesis of these derivatives is efficient and uses readily available and inexpensive substrates. This reaction can be applied for synthesis of other derivatives of resorcinarene, which is currently a subject of further studies.

Heating of resorcinarene 1 with 6 equiv of formaldehyde and 4 equiv of iminodiacetic acid in methanol-water (10/1) causes precipitation of a product, which is not a result of aminomethylation in the Mannich reaction. The ¹H and ¹³C

Table 1. Yields of formation of the tetra(alkoxymethyl) derivatives 2a-f of various alcohols in the presence of stoichiometric amount of iminodiacetic acid

Product	R	Stoichiometric amounts of iminodiacetic acid (%	
2a	CH ₃	88.2	
2b	C_2H_5	75.4	
2c	C_3H_7	72.6	
2d	C_4H_9	56.6	
2e	$C_{5}H_{11}$	49.9	
2f	C ₆ H ₁₃	36.5	

Keywords: Resorcinarene; Mannich reaction; Catalyse; Alcohols.

^{*} Corresponding author. Tel.: +48 41 349 70 16; e-mail: iwanek@pu.kielce.pl

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Table 2. The yield of formation of the tetra(ethoxymethyl) derivative 2b versus the amount of iminodiacetic acid with respect to the amount of resorcinarene 1

Relative amount of	of iminodiacetic acid	Yield of 2b (%)
(weight%)	(mole%)	
10	13.4	45.0
7	9.4	44.9
4	5.4	8.8
2	2.7	0.0

Table 3. Yields of formation of the tetra(alkoxymethyl) derivatives 2a–f of various alcohols in the presence of catalytic amounts of iminodiacetic acid

Product	R	Catalytic amounts of iminodiacetic acid (%)
2a	CH ₃	79.3
2b	C_2H_5	65.8
2c	C_3H_7	64.1
2d	C_4H_9	45.8
2e	$C_{5}H_{11}$	40.2
2f	C ₆ H ₁₃	27.6

It was found that the stoichiometric amount of iminodiacetic acid was not necessary for maintaining a high yield of this reaction. We conducted a number of trial runs in order to optimise the amount of iminodiacetic acid and determined chromatographically the yields of the tetra(ethoxymethyl) derivative **2b** obtained for various amounts of iminodiacetic acid (reflux, 4 h). According to the experiments, the optimal amount of iminodiacetic acid was 7% by weight with

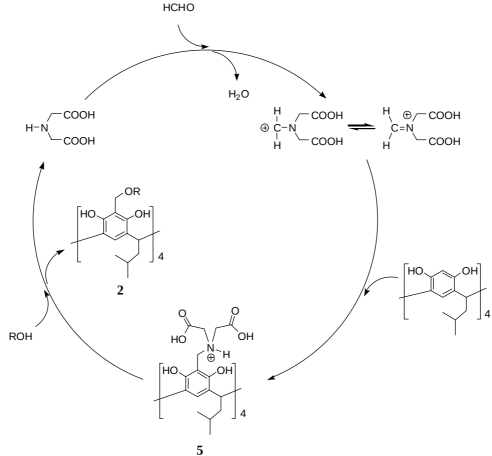
respect to resorcinarene 1 (i.e., 9.4 mol%). These results are shown in Table 2.

Table 3 shows the yields of formation of tetra(alkoxymethyl) derivatives 2a-f from various alcohols in the presence of catalytic amount of iminodiacetic acid. The reaction was carried out under reflux for 12 h. When the reflux in lower alcohols was continued for 12 h, the reaction yields rose to about 80%. The trial reaction carried out in the absence of iminodiacetic acid gave no corresponding product **2**.

We also investigated the alkoxymethylation of other resorcinarenes. Refluxing of the reaction mixture in the presence of catalytic amounts (9.4 mol%) of iminodiacetic acid for 4 h led to formation of the tetra(ethoxymethyl) derivatives **3** and **4**. The reaction yields were satisfactory and amounted to 42 and 65%, respectively.



The alkoxymethylation of resorcinarenes does not occur without iminodiacetic acid. The possible mechanism of formation of the alkoxymethyl derivatives 2 in the presence of iminodiacetic acid is shown in Scheme 2. First, the



formaldehyde and iminodiacetic acid molecules form the iminium cation, which then adds at the 2-position to the resorcinarene 1 to form the aminomethylene derivative 5.

The presence of two carboxy groups in the adduct **5** favours an intramolecular transfer of a proton from the carboxy group to the amino group. The thus-formed quaternary aminomethyl salt of resorcinarene is susceptible to elimination followed by addition of the alcohol. The displacement of tertiary and quaternary nitrogen atoms by nucleophilic reagents is rather well known reaction.^{8–11} As a result, the 2-position of resorcinarene **1** is substituted with a simultaneous elimination of iminodiacetic acid. The latter molecule can then form the next reactive iminium cation with formaldehyde, and the reaction cycle repeats, to form finally the tetra(alkoxymethyl) derivative **2** of resorcinarene **1**.

2. Conclusion

Refluxing of an alcoholic solution of a resorcinarene, formaldehyde and a catalytic amount of iminodiacetic acid affords the tetra(alkoxymethyl) derivatives of resorcinarene in high yields. The optimal amount of iminodiacetic acid is 10 mol% with respect to the resorcinarene.

3. Experimental

The ¹H and ¹³C NMR spectra were recorded with a Bruker AC 400 (400 MHz) spectrometer. The FAB-mass spectra were obtained with a Finnigan MAT90 mass spectrometer using *m*-nitrobenzoyl alcohol (NBA) as a matrix. The melting points were determined with a Büchi B-540 melting point apparatus and are uncorrected. The reagents and solvents were obtained from Fluka and Merck and were used without purification. The chromatographic analysis was performed using a Varian Polaris 201 HPLC chromatograph equipped with a C-18 column and an UV–vis detector. The chromatographic separations were performed on silica gel 60 (SiO₂, Merck, particle size 0.040–0.063 mm, 230–240 mesh).

3.1. Preparation of the alkoxymethyl derivatives of resorcinarene

Iminodiacetic acid (stoichiometric amount: 0.37 g, 2.8 mmol; or catalytic amount: 0.05 g, 0.37 mmol) was dissolved in hot water (1 mL). An alcohol (10 mL) was added to resorcinarene 1 (0.5 g, 0.7 mmol) and warmed until complete dissolution of resorcinarene. Aqueous solvent of formaldehyde (37%, 0.26 mL, 3.5 mmol) was added, followed after 5 min by the aqueous solution of iminodiacetic acid. The reaction mixture was refluxed for 12 h. After about half an hour, the iminodiacetic acid dissolved again. The solution became then cloudy and a solid started to precipitate. Finally, the solid was filtered off. Chloroform (20 mL) was added to the filtrate, and the undissolved part of iminodiacetic acid was filtered off. The chloroform solution of the product 2 was concentrated by evaporation and purified by column chromatography.

3.1.1. Tetramethoxyisopentylresorcinarene 2a. The solid was purified by flash chromatography (CH₃COOC₂H₅/ *n*-hexane, 2:5) to give the title compound **2a** as a white solid, $R_{\rm f}$ 0.38; mp>300 °C; $\nu_{\rm max}$ (KBr) 3344, 2952, 1608, 1472, 1296, 1088 cm⁻¹; $\delta_{\rm H}$ NMR (400 MHz, CDCl₃) 8.69 (8H, s, OH), 7.14 (4H, s, ArH), 4.75 (8H, s, ArCH₂O), 4.40 (4H, t, J=8.0 Hz, ArCHRAr), 3.43 (12H, s, CH₃O), 2.05 (8H, t, J=7.4 Hz, CHCH₂CH(CH₃)₂), 1.54–1.39 (4H, m, CH₂CH(CH₃)₂), 0.96 (24H, d, J=6.6 Hz, CH₂CH(CH₃)₂); $\delta_{\rm C}$ (100 MHz, CDCl₃) 22.8, 26.0, 30.5, 42.7, 58.8, 70.2, 76.7, 77.0, 77.3, 109.0, 122.8, 124.1, 149.7; *m*/z (MALDI): [M⁺ + Na⁺] found 910.4856, requires 911.1270.

3.1.2. Tetraethoxyisopentylresorcinarene 2b. The solid was purified by flash chromatography (CH₃COOC₂H₅/*n*-hexane, 2:5) to give the title compound 2b as a white solid, $R_{\rm f}$ 0.66; mp > 300 °C; $\nu_{\rm max}$ (KBr) 3376, 2952, 1608, 1472, 1296, 1088 cm⁻¹; $\delta_{\rm H}$ (400 MHz, CDCl₃) 8.83 (8H, s, OH), 7.14 (4H, s, ArH), 4.80 (8H, s, ArCH₂O), 4.41 (4H, t, J=8.6 Hz, ArCHRAr), 3.59 (8H, q, J=7.0 Hz, CH₃CH₂O), 2.05 (8H, t, J=7.6 Hz, CHCH₂CH(CH₃)₂), 1.52–1.42 (4H, m, CH₂CH(CH₃)₂), 1.25 (12H, t, J=6.8 Hz, CH₃CH₂O), 0.97 (24H, d, J=7.1 Hz, CH₂CH(CH₃)₂); $\delta_{\rm C}$ (100 MHz, CDCl₃) 14.9, 22.8, 26.1, 30.6, 42.7, 66.8, 68.0, 76.7, 77.0, 77.3, 109.3, 122.7, 124.1, 149.7; *m*/z (MALDI): [M⁺ + Na⁺] found 966.6733, requires 967.2353.

3.1.3. Tetrapropoxyisopentylresorcinarene 2c. The solid was purified by flash chromatography (CH₃COOC₂H₅/*n*-hexane, 2:5) to give the title compound **2c** as a pale yellow solid, R_f 0.88; mp > 300 °C; ν_{max} (KBr) 3344, 2952, 1616, 1472, 1296, 1088 cm⁻¹; δ_H (400 MHz, CDCl₃) 8.83 (8H, s, OH), 7.13 (4H, s, ArH), 4.79 (8H, s, ArCH₂O), 4.41 (4H, t, *J*=8.9 Hz, ArCHRAr), 3.48 (8H, t, *J*=6.7 Hz, C₂H₅CH₂O), 2.05 (8H, t, *J*=7.4 Hz, CHCH₂CH(CH₃)₂), 1.69–1.59 (8H, m, CH₃CH₂CH₂O), 1.52–1.42 (4H, m, CH₂CH(CH₃)₂), 0.97 (24H, d, *J*=6.6 Hz, CH₂CH(CH₃)₂), 0.93 (12H, t, *J*=7.4 Hz, CH₃C₂H₄O); δ_C (100 MHz, CDCl₃) 10.4, 22.6, 22.8, 26.0, 26.9, 30.6, 42.7, 68.2, 73.0, 109.3, 122.7, 124.1, 149.8; *m*/z (MALDI): [M⁺ + Na⁺] found 1022.7309, requires 1023.3436.

3.1.4. Tetrabutoxyisopentylresorcinarene 2d. The solid was purified by flash chromatography (CH₃COOC₂H₅/*n*-hexane, 1:4) to give the title compound **2d** as a pale yellow solid, $R_f 0.48$; mp>300 °C; ν_{max} (KBr) 3352, 2960, 1624, 1472, 1296, 1088 cm⁻¹; δ_H (400 MHz, CDCl₃) 8.82 (8H, s, OH), 7.13 (4H, s, ArH), 4.78 (8H, s, ArCH₂O), 4.41 (4H, t, J=8.0 Hz, ArCHRAr), 3.52 (8H, t, J=6.6 Hz, C₃H₇CH₂O), 2.05 (8H, t, J=7.9 Hz, CHCH₂CH(CH₃)₂), 1.63–1.56 (8H, m, C₂H₅CH₂CH₂O), 1.51–1.42 (4H, m, CH₂CH(CH₃)₂), 1.41–1.31 (8H, m, CH₃CH₂C₂H₄O), 0.96 (24H, d, J=6.6 Hz, CH₂CH(CH₃)₂), 0.92 (12H, t, J=7.5 Hz, CH₃C₃H₆O); δ_C (100 MHz, CDCl₃) 13.8, 19.2, 19.4, 22.8, 26.0, 26.9, 30.6, 31.4, 32.0, 42.7, 65.1, 68.3, 71.2, 109.3, 122.7, 124.1, 149.8; *m*/z (MALDI): [M⁺ + Na⁺] found 1079.008, requires 1079.4520.

3.1.5. Tetrapentoxyisopentylresorcinarene 2e. The residue was purified by flash chromatography (CH₃COOC₂H₅/*n*-hexane, 1:10) to give the title compound **2e** as a brown oil, $R_{\rm f}$ 0.46; mp>300 °C; $\nu_{\rm max}$ (KBr) 3344, 2952, 1632, 1472, 1296, 1088 cm⁻¹; $\delta_{\rm H}$ (400 MHz, DMSO- d_6) 8.72 (8H, s,

OH), 7.38 (4H, s, ArH), 4.47 (8H, s, ArCH₂O), 4.35 (4H, t, J=7.9 Hz, ArCHRAr), 3.37 (8H, t, J=7.6 Hz, $C_4H_9CH_2O$), 2.12 (8H, t, J=7.0 Hz, CHCH₂CH(CH₃)₂), 1.53–1.43 (8H, m, C₃H₇CH₂CH₂O), 1.40–1.32 (4H, m, CH₂CH(CH₃)₂), 1.29–1.25 (8H, m, C₂H₅CH₂C₂H₄O), 1.25–1.21 (8H, m, CH₃CH₂C₃H₆O), 0.93 (24H, d, J=6.6 Hz, CH₂CH(CH₃)₂), 0.83 (12H, t, J=7.0 Hz, $CH_3C_4H_8O$); δ_C (100 MHz, DMSO- d_6) 13.9, 21.9, 22.8, 25.9, 27.9, 28.7, 38.9, 39.1, 39.3, 39.5, 39.7, 39.9, 40.1, 63.3, 66.9, 69.7, 94.4, 111.4, 124.0; m/z (MALDI): [M⁺ + Na⁺] found 1135.9865, requires 1135.5603.

3.1.6. Tetrahexoxyisopentylresorcinarene 2f. The residue was purified by flash chromatography (CH₃COOC₂H₅/ *n*-hexane, 1:10) to give the title compound 2f as a brown oil, $R_{\rm f}$ 0.58; mp 147–148 °C; $\nu_{\rm max}$ (KBr) 3352, 2952, 1640, 1472, 1296, 1088 cm⁻¹; $\delta_{\rm H}$ (400 MHz, DMSO- d_6) 8.75 (8H, s, OH), 7.38 (4H, s, ArH), 4.46 (8H, s, ArCH₂O), 4.35 (4H, t, J=7.6 Hz, ArCHRAr), 3.36 (8H, t, J=6.8 Hz, $C_5H_{11}CH_2O$, 2.12 (8H, t, J=7.0 Hz, $CHCH_2CH(CH_3)_2$), 1.48-1.41 (8H, m, C₄H₉CH₂CH₂O), 1.38-1.32 (4H, m, CH₂CH(CH₃)₂), 1.26–1.18 (24H, m, CH₃(CH₂)₃(CH₂)₂O), 0.92 (24H, d, J=6.5 Hz, CH₂CH(CH₃)₂), 0.83 (12H, t, J=7.5 Hz, $CH_3(CH_2)_5O$; δ_C (100 MHz, DMSO- d_6) 13.9, 22.1, 22.7, 25.3, 25.9, 29.0, 31.1, 31.4, 38.9, 39.1, 39.3, 39.5, 39.7, 39.9, 40.1, 42.1, 63.3, 69.7, 111.4, 124.5, 149.8; m/z (MALDI): $[M^+ + Na^+]$ found 1192.1146, requires 1191.6687.

3.1.7. Tetraethoxydecylresorcinarene 3. The solid was purified by flash chromatography (CH₃COOC₂H₅/*n*-hexane, 1:6) to give the title compound **3** as a white solid, $R_{\rm f}$ 0.41; mp > 300 °C; $\nu_{\rm max}$ (KBr) 3296, 2968, 1608, 1464, 1296, 1104 cm⁻¹; $\delta_{\rm H}$ (400 MHz, CDCl₃) 8.82 (8H, s, OH), 7.15 (4H, s, ArH), 4.81 (8H, s, ArCH₂O), 4.28 (4H, t, *J*=7.9 Hz, ArCHRAr), 3.61 (8H, q, *J*=7.0 Hz, CH₃CH₂O), 2.17 (8H, q, *J*=7.6 Hz, CHCH₂C₁₀H₂₁), 1.37 (12H, t, *J*=4.0 Hz, CH₃CH₂O), 1.27 (72H, m, CH₂(C₉H₁₈)CH₃), 0.89 (12H, t, *J*=6.9 Hz, C₁₀H₂₀CH₃); $\delta_{\rm C}$ (100 MHz, CDCl₃) 14.1, 14.9, 22.7, 28.1, 29.4, 29.7, 31.9, 33.0, 33.7, 66.8, 68.0, 76.7, 77.0, 77.3, 109.2, 122.4, 124.1, 149.7; MS (MALDI): *m*/*z* [M+Na⁺] calcd 1359.9684, found 1359.9937.

3.1.8. Tetraethoxymethylresorcinarene 4. The solid was purified by flash chromatography (CH₃COOC₂H₅/*n*-hexane, ?:?) to give the title compound **4** as a white solid, $R_{\rm f}$ 0.55; mp > 300 °C; $\nu_{\rm max}$ (liquid film) 3416, 2928, 1712, 1368,

1224, 1096 cm⁻¹; $\delta_{\rm H}$ (400 MHz, CDCl₃) 8.85 (8H, s, OH), 7.28 (4H, s, ArH), 4.79 (8H, s, ArCH₂O), 4.53 (4H, q, J= 7.9 Hz, ArCHRAr), 3.58 (8H, q, J=7.0 Hz, CH₃CH₂O), 1.72 (12H, d, J=7.6 Hz, CHCH₃), 1.23 (12H, t, J=4.0 Hz, CH₃CH₂O); $\delta_{\rm C}$ (100 MHz, CDCl₃) 14.9, 19.9, 27.3, 66.8, 67.9, 76.8, 77.0, 77.3, 109.3, 122.1, 125.1, 149.4; m/z(MALDI): [M⁺ + Na⁺] found 799.0133, requires 798.9103.

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Tetrahedron

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Tricalysiolides A–F, new rearranged *ent*-kaurane diterpenes from *Tricalysia dubia*

Koichi Nishimura,^a Yukio Hitotsuyanagi,^a Noriko Sugeta,^a Kei-ichi Sakakura,^a Kazuya Fujita,^a Haruhiko Fukaya,^a Yutaka Aoyagi,^a Tomoyo Hasuda,^a Takeshi Kinoshita,^b Dong-Hui He,^c Hideaki Otsuka,^c Yoshio Takeda^d and Koichi Takeya^{a,*}

^aSchool of Pharmacy, Tokyo University of Pharmacy and Life Science, 1432-1 Horinouchi, Hachioji, Tokyo 192-0392, Japan ^bFaculty of Pharmaceutical Sciences, Teikyo University, 1091-1 Suarashi, Sagamiko-machi, Tsukui-gun, Kanagawa 199-0195, Japan ^cGraduate School of Biomedical Sciences, Hiroshima University, 1-2-3 Kasumi, Minami-ku, Hiroshima 734-8551, Japan ^dFaculty of Integrated Arts and Sciences, The University of Tokushima, 1-1 Minamijosanjima-cho, Tokushima 770-8502, Japan

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Abstract—Six rearranged *ent*-kaurane diterpenes, tricalysiolides A–F, having the cafestol-type carbon framework were isolated from the wood of *Tricalysia dubia* (Rubiaceae). Their absolute structures were determined on the basis of the 2D NMR spectroscopy, X-ray crystallographic analysis, and chemical methods.

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1. Introduction

Tricalysia dubia (Lindl.) Ohwi (Rubiaceae) is an evergreen shrub or tree that is wildly distributed in Taiwan and the southern parts of China and Japan. From the leaves of this plant, unique rearranged *ent*-kaurane glycosides, tricalysio-sides A–G, have been isolated.¹ In the present study, from the wood of this plant, we isolated six new rearranged *ent*-kaurane diterpenes, tricalysiolides A–F (**1–6**), and determined their structures.

2. Results and discussion

By a series of column chromatography on highly porous synthetic resin (Diaion HP-20), silica gel, aminopropylbonded silica gel, and ODS HPLC, a hot MeOH extract of air-dried wood of *T. dubia* afforded six diterpenes named tricalysiolides A–F (**1–6**) (Fig. 1).

Tricalysiolide A (1) was isolated as an amorphous solid, whose molecular formula was determined to be $C_{20}H_{28}O_4$ from the $[M+H]^+$ peak at m/z 333.2074 (calcd for $C_{20}H_{29}O_4$, 333.2066) in the HRESIMS. The IR spectrum indicated that **1** possessed hydroxyl

Keywords: Tricalysiolide; Diterpene; Tricalysia dubia.

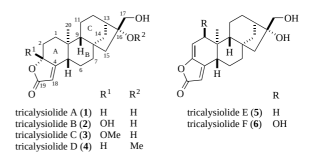


Figure 1. Structures of tricalysiolides A–F (1–6).

(3365 cm⁻¹) and lactone (1742 cm⁻¹) groups. The ¹H NMR spectrum showed the presence of one tertiary methyl (δ 0.67) and one olefinic proton (δ 5.73) (Table 1). The ¹³C NMR spectrum suggested that **1** was a diterpenoid derivative with a total of 20 carbons, consisting of one methyl, nine methylenes, four methines, one trisubstituted double bond giving highly shielded (δ 111.4) and deshielded (δ 175.1) signals, one carbonyl, and three quaternary carbons (Table 2). On the basis of detailed analysis of the ¹H–¹H COSY, HMQC, and HMBC spectra, **1** was shown to be a cafestol-type rearranged kaurane diterpenoid with two hydroxyls at C-16 and C-17. In the HMBC spectrum, H-18 was correlated with C-3, C-4, C-5, and C-19, which implied that **1** had an α , β -unsaturated- γ -lactone group attached to ring A (Fig. 2). From the NOESY correlations between

^{*} Corresponding author. Tel.: +81 426 76 3007; fax: +81 426 77 1436; e-mail: takeyak@ps.toyaku.ac.jp

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Table 1. ¹H NMR data for tricalysiolides A-F (1-6) in C₅D₅N^a

Proton	1 ^b	2 ^b	3 ^b	4 ^b	5 °	6 ^b
1a	1.67 (m)	1.78 (m)	1.68 (m)	1.66 (m)	2.32 (dd, 17.6, 6.7)	4.40 (t, 6.1)
1b	0.97 (td, 13.8, 3.7)	1.52 (m)	1.27 (m)	0.94 (td, 13.8, 3.6)	1.80 (m)	
2a	2.21 (m)	2.52 (d, 13.7)	2.40 (ddd, 14.0, 4.1, 2.3)	2.20 (m)	5.62 (dt, 6.7, 2.1)	6.09 (dd, 6.4, 1.6)
2b	1.52 (m)	2.00 (m)	1.87 (dd, 14.0, 4.7)	1.49 (m)		
3	4.75 (t, 8.9)			4.75 (t, 9.3)		
5	1.90 (m)	2.57 (d, 9.7)	2.12 (m)	1.87 (m)	2.17 (dd, 11.7, 2.4)	3.05 (dt, 11.8, 2.5)
6a	1.50 (m)	1.59 (m)	1.56 (m)	1.44 (m)	1.64 (m)	1.78 (m)
6b	1.41 (m)	1.38 (m)	1.48 (m)	1.40 (m)	1.43 (m)	1.56 (m)
7a	1.61 (m)	1.65 (m)	1.67 (m)	1.52 (m)	1.65 (m)	1.68 (m)
7b	1.49 (m)	1.56 (m)	1.58 (m)	1.45 (m)	1.53 (m)	1.62 (m)
9	1.13 (d, 8.8)	1.29 (d, 8.6)	1.26 (d, 9.0)	1.13 (d, 8.8)	1.21 (d, 8.7)	2.40 (d, 8.5)
11a	1.69 (m)	1.76 (m)	1.76 (m)	1.69 (m)	1.48 (m)	1.92 (m)
11b	1.47 (m)	1.54 (m)	1.52 (m)	1.46 (m)	1.41 (m)	1.72 (m)
12a	1.86 (m)	1.91 (m)	1.92 (m)	1.85 (m)	1.88 (m)	1.95 (m)
12b	1.42 (m)	1.50 (m)	1.50 (m)	1.45 (m)	1.42 (m)	1.59 (m)
13	2.43 (s-like)	2.47 (s-like)	2.50 (s-like)	2.48 (s-like)	2.46 (s-like)	2.51 (s-like)
14a	2.02 (dd, 11.3, 4.1)	2.06 (dd, 11.1, 4.7)	2.10 (m)	1.82 (d, 11.0)	2.04 (dd, 11.3, 4.4)	2.10 (dd, 11.4, 4.4)
14b	1.92 (d, 11.3)	1.97 (d, 11.1)	1.98 (d, 11.4)	1.69 (m)	1.88 (d, 11.3)	2.01 (d, 11.4)
15a	1.82 (d, 14.2)	1.86 (d, 14.3)	1.89 (d, 14.2)	1.74 (d, 14.5)	1.84 (d, 14.1)	1.88 (d, 14.2)
15b	1.72 (d, 14.2)	1.77 (d, 14.3)	1.80 (dd, 14.2, 1.5)	1.48 (d, 14.5)	1.71 (dd, 14.1, 1.6)	1.83 (d, 14.2)
17a	4.11 (dd, 10.8, 4.6)	4.14 (dd, 10.8, 5.0)	4.18 (dd, 10.9, 4.7)	4.08 (dd, 12.3, 4.7)	4.14 (dd, 10.8, 4.0)	4.14 (dd, 10.8, 5.2)
17b	4.05 (dd, 10.8, 4.6)	4.06 (dd, 10.8, 5.0)	4.09 (dd, 10.9, 4.7)	4.00 (dd, 12.3, 4.7)	4.05 (dd, 10.8, 4.0)	4.05 (dd, 10.8, 5.2)
18	5.73 (s)	5.81 (s)	6.00 (d, 1.7)	5.73 (s)	5.91 (s-like)	5.97 (dd, 2.5, 1.6)
20	0.67 (s, 3H)	0.80 (s, 3H)	0.79 (s, 3H)	0.66 (s, 3H)	0.84 (s, 3H)	0.94 (s, 3H)
OMe-3			3.16 (s, 3H)			
OMe-16				3.31 (s, 3H)		
OH-1						6.87 (d, 5.8)
OH-3		9.49 (s)				
OH-16	5.24 (s)	5.22 (s)	5.22 (s)		5.24 (s)	5.21 (s)
OH-17	6.16 (t, 6.2)	6.16 (br s)	6.16 (t, 5.1)	5.72 (br s)	6.16 (br s)	6.12 (t, 5.3)

^a Assignments based on ¹H-¹H COSY, HMBC, and HMQC experiments. Multiplicity and J-values in Hz are given in parentheses.

^b Recorded at 500 MHz.

^c Recorded at 400 MHz.

H-1a/H₃-20, H-1b/H-9, H-3/H-5, H-5/H-9, H-9/H-15b, H-11a/H-15b, H-11a/H-17a, H-11a/H-17b, H-12a/H-17a, H-12a/H-17b, and H-14b/H₃-20 (Fig. 3), its stereostructure was determined to be as shown in Figure 1. Compound **1** corresponds to the aglycone moiety of

tricalysiosides B-E obtained previously from the leaves of the same plant source.¹

The absolute configuration of 1 was determined by the X-ray crystallographic analysis of its *p*-bromobenzoate 7

Table 2. ^{13}C NMR data for tricalysiolides A–F (1–6) in C₅D₅N

Cashan	1 ^a	2 ^a	3 ^a	4 ^a	5 ^b	6 ^a
Carbon	1	2	3	4	5	0
1	35.8	36.1	35.6	35.9	39.6	69.0
2	29.9	35.0	33.6	29.9	107.4	109.3
3	81.4	105.7	107.3	81.4	149.7	151.4
4	175.1	173.7	171.2	175.0	159.9	160.3
5	48.7	47.4	47.5	48.7	45.5	39.2
6	22.0	22.1	21.8	22.0	21.8	21.8
7	40.2	40.3	40.1	40.2	39.9	40.0
8	44.5	44.6	44.5	44.2	44.3	44.3
9	53.5	53.7	53.6	53.5	52.8	44.4
10	42.9	43.8	43.6	42.9	41.9	44.3
11	19.4	19.5	19.5	19.5	18.8	18.4
12	26.3	26.4	26.3	26.0	26.3	26.4
13	45.8	45.9	45.9	42.0	45.9	46.0
14	38.0	38.1	38.0	37.7	37.6	37.6
15	53.8	53.8	53.7	49.8	53.5	53.9
16	81.5	81.5	81.5	87.3	81.5	81.5
17	66.4	66.4	66.3	60.9	66.3	66.4
18	111.4	112.6	115.2	111.4	110.5	111.7
19	173.7	171.5	170.5	173.7	170.6	170.3
20	14.7	14.4	14.5	14.7	15.3	16.1
OMe-3			49.9			
OMe-16				49.3		

^a Recorded at 125 MHz.

^b Recorded at 100 MHz.

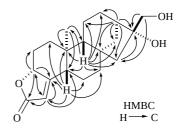


Figure 2. Selected HMBC correlations for 1.

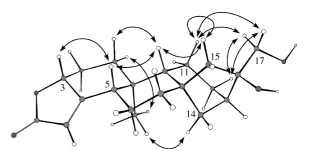
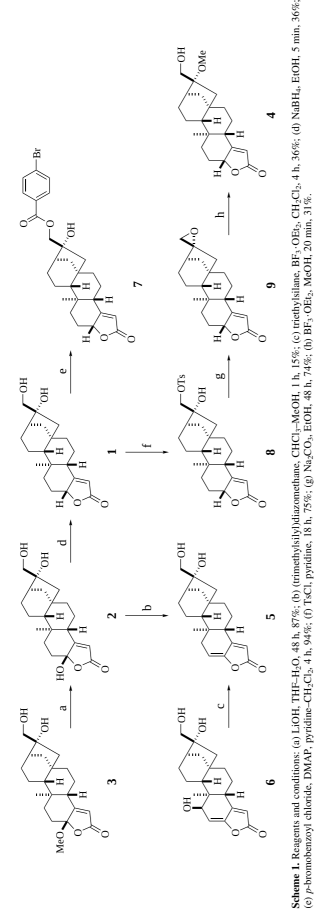


Figure 3. Selected NOESY correlations for 1.



(Scheme 1, Fig. 4). The value of the final refined Flack parameter 0.012(7) confirmed that the absolute configuration of **1** was as depicted in Figure 1.

Tricalysiolide B (2) was isolated as an amorphous powder. Its molecular formula was determined to be C₂₀H₂₈O₅ from the $[M+H]^+$ peak at m/z 349.2033 (calcd for $C_{20}H_{29}O_5$, 349.2015) in the HRESIMS. The ¹H and ¹³C NMR spectra of 2 were similar to those of 1. The differences noted between 1 and 2 were that the C-3 signal, observed as a methine carbon signal (δ 81.4) in **1**, was observed as a quaternary carbon signal (δ 105.7) in **2**. The C-3 signal was correlated with the hydroxyl proton (δ 9.49) in the HMBC spectrum in 2, showing the location of the hydroxyl group was at C-3. The NOESY correlation detected between H-5 and OH-3 indicated that the hydroxyl group at C-3 was of β-orientation. The structure and absolute stereochemistry of 2 were determined to be as shown in Figure 1, by the preparation of 1 from 2 (Scheme 1). Reduction of 2 with sodium borohydride afforded a product, which was shown to be identical to natural 1 by comparison of their spectral data and optical rotations.

Tricalysiolide C (3) was isolated as colorless prisms. Its molecular formula was determined to be $C_{21}H_{30}O_5$ from the $[M+H]^+$ peak at *m/z* 363.2182 (calcd for C₂₁H₃₁O₅, 363.2171) in the HRESIMS. The ¹H and ¹³C NMR spectra of 3 were very similar to those of 2, implying that 2 and 3 had the same basic structure. The only difference noted between 2 and 3 was that the NMR spectra of 3 displayed signals due to a methoxyl group ($\delta_{\rm H}$ 3.16, $\delta_{\rm C}$ 49.9), whose methoxyl protons were correlated with the C-3 signal in the HMBC spectrum, thus demonstrating the location of the methoxyl group being at C-3 in 3. A NOESY correlation detected between H-5 and OCH₃-3 indicated that the methoxyl group was of β -orientation. Thus, compound 3 was determined to be the 3-O-methyl analogue of 2. This structure was confirmed by the X-ray analysis (Fig. 5). Alkaline hydrolysis and subsequent acid treatment of 3 afforded 2 (Scheme 1), establishing that the absolute structure of **3** was as shown in Figure 1.

Tricalysiolide D (4) was isolated as colorless prisms. Its molecular formula, $C_{21}H_{30}O_4$, determined from the [M+ H]⁺ peak at m/z 347.2232 (calcd for C₂₁H₃₁O₄, 347.2222) in the HRESIMS showed that it was higher than that of 1 by one methylene unit. The ¹H and ¹³C NMR spectra of 4 were similar to those of 1, except that a singlet methoxyl proton signal (δ 3.31) and one carbon signal (δ 49.3), which were not observed in the spectra of 1, were observed in the spectra of **4** and that the C-16 signal (δ 87.3) of **4** was in a lower field than the corresponding signal of 1 (δ 81.5). The HMBC correlation between the methoxyl protons and C-16 indicated that the location of the methoxyl group was at C-16 to show that 4 was the 16-O-methyl ether of 1. The NOESY correlation between H-14b and OCH3-16 indicated that the methoxyl group at C-16 was of α -orientation. This structure was confirmed by the X-ray analysis (Fig. 6). Its absolute configuration was established by the preparation of 4 from 1 (Scheme 1). Tosylate 8, produced by treatment of 1 with *p*-toluenesulfonyl chloride, gave, on treatment with sodium carbonate, epoxide 9. Methanolysis of 9 in the presence of boron trifluoride diethyl etherate afforded a

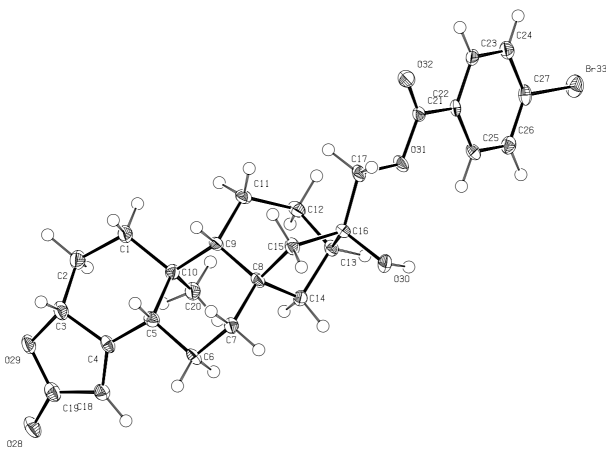


Figure 4. ORTEP representation of 7 as determined by single-crystal X-ray analysis.

product, which was shown to be identical to natural **4** by comparison of their spectral data and optical rotations. Thus, the absolute structure of **4** was determined to be as shown in Figure 1.

Tricalysiolide E (5) was obtained as an amorphous powder. Its molecular formula was determined to be $C_{20}H_{26}O_4$ from the $[M+H]^+$ peak at m/z 331.1901 (calcd for $C_{20}H_{27}O_4$, 331.1909) in the HRESIMS, which

was less than that of **1** by two hydrogen atoms. The ¹H and ¹³C NMR spectra of **5** were generally similar to those of **1**, except for the C-2 and C-3 signals. The C-2 methylene (δ 29.9) and C-3 methine (δ 81.4) signals in **1** were of olefinic methine (δ 107.4) and quaternary (δ 149.7) carbons, respectively, in **5**. When **2** was treated with (trimethylsilyl)diazomethane² in chloroformmethanol (10/1), the product was shown to be identical to natural **5** by their spectral data and optical rotations.

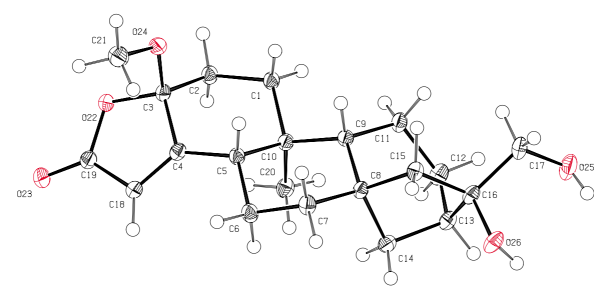


Figure 5. ORTEP representation of tricalysiolide C (3) as determined by single-crystal X-ray analysis.

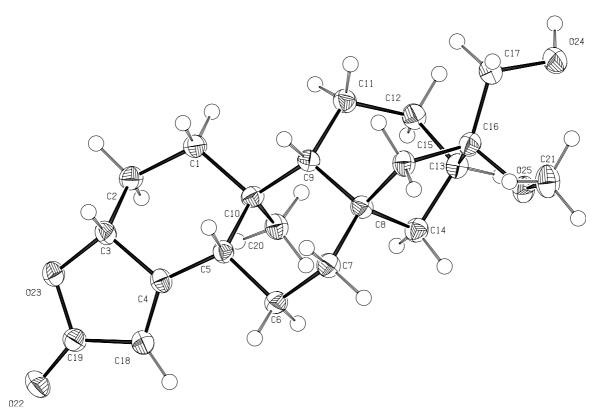


Figure 6. ORTEP representation of tricalysiolide D (4) as determined by single-crystal X-ray analysis.

Thus, the structure of 5 was determined to be as shown in Figure 1.

Tricalysiolide F (**6**) was isolated as an amorphous powder. Its molecular formula was determined to be $C_{20}H_{26}O_5$ from the $[M+H]^+$ peak at m/z 347.1882 (calcd for $C_{20}H_{27}O_5$, 347.1858) in the HRESIMS, which was higher than that of **5** by one oxygen atom. The ¹H and ¹³C NMR spectra of **6** resembled those of **5**, demonstrating that they were of the same basic structure. The differences noted between them were that the C-1 methylene signal (δ 39.6) in **5** was observed as an oxymethine carbon signal (δ 69.0) in **6**. The NOESY correlation between H-1 and H₃-20 indicated that the hydroxyl group at C-1 was of β -orientation. Treatment of **6** with triethylsilane and boron trifluoride diethyl etherate³ afforded a reduction product, which was shown to be identical to natural **5** by comparison of their spectral data and optical rotations. Accordingly, the absolute structure of **6** was determined to be as shown in Figure 1.

Tricalysiolides A–F (1–6) and tricalysiosides A–G¹ are unusual compounds in which one of the geminal methyls is rearranged to the other methyl to form an ethyl unit at C-4 of the *ent*-kaurane skeleton. So far, the only known compounds with this unusual carbon framework are cafestol^{4,5} and several related furokauranes^{6–10} from *Coffea* plants (Rubiaceae). Tricalysiolides A (1), D (4) and tricalysiosides A–G are characterized by an α , β -unsaturated- γ -lactone ring fused to the ring A, and tricalysiolides B (2), C (3), E (5), and F (6) by a further oxidized ring A structure. Tricalysiolides A–F (1–6) showed a weak cytotoxic activity on P-388 murine leukemia cells with IC_{50} values of 40, 40, 50, 40, 17, and 30 µg/mL, respectively.

3. Experimental

3.1. General experimental procedures

Optical rotations were measured on a JASCO P1030 digital polarimeter. IR spectra were recorded on a JASCO FT/IR 620 spectrophotometer. UV spectra were obtained using a JASCO V-530 spectrophotometer. NMR spectra were measured on Bruker DRX-500 and DPX-400 spectrometers at 300 K. The ¹H chemical shifts in C_5D_5N were referenced to the residual C₅D₄HN resonance at 7.21 ppm, and the ¹³C chemical shifts to the solvent resonance at 135.5 ppm. Mass spectra were obtained using a Micromass LCT spectrometer. Preparative HPLC was carried out on a JASCO PU-986 pump unit equipped with a UV-970 UV detector (λ 220 nm) and an Inertsil PREP-ODS column (10 μ m, 20 \times 250 mm), by using a MeOH-H₂O or a MeCN-H₂O solvent system at a flow rate of 10 mL/min. X-ray single crystal analysis was taken on a Mac Science DIP diffractometer with Mo K α radiation $(\lambda = 0.71073 \text{ Å}).$

3.2. Plant material

Wood of *T. dubia* was collected in Iriomote Island, Okinawa, in March 2003, and the plant origin was identified by Dr. T. Kinoshita (Teikyo University, Japan). A voucher specimen has been deposited at the Herbarium of Teikyo University.

1517

3.3. Extraction and isolation

Cut and air-dried wood (1.44 kg) of T. dubia was extracted with hot MeOH $(3 \times 6 L)$. After removal of MeOH under reduced pressure, the residue (104 g) was placed on a column of HP-20 (DIAION, 600 g) and eluted with H_2O , H₂O-MeOH (1/1), H₂O-MeOH (1/4), MeOH, and acetone (each 3 L) sequentially to give five fractions. After removal of the solvent, the residue of the H_2O -MeOH (1/4) fraction (17 g) was subjected to silica gel (Merck Kieselgel 60, 230–400 mesh, 200 g) column chromatography eluting sequentially with EtOAc, CHCl3-MeOH (10/1), CHCl₃-MeOH (3/1), and MeOH (each 500 mL). After evaporation the CHCl₃-MeOH (10/1) fraction (1.2 g) was subjected to aminopropyl-bonded silica gel (Chromatorex, 200-350 mesh, 40 g) column chromatography eluting sequentially with CHCl₃-MeOH (50/1, then 30/1, then 10/1), and MeOH (each 100 mL) to give fractions 1–5. Fraction 2 (16.2 mg) was further applied to ODS HPLC eluting with MeOH-H₂O (43/57) to afford 4 (5.6 mg). Fraction 3 (465.0 mg) was further separated by ODS HPLC eluting with MeOH-H₂O (44/56) to afford 1 (141.5 mg), 3 (50.8 mg), and 5 (6.8 mg). The CHCl₃-MeOH (3/1) fraction (7.15 g) was subjected to aminopropyl-bonded silica gel (35 g) column chromatography eluting sequentially with CHCl₃-MeOH (30/1, then 10/1, then 5/1, then 3/1), and MeOH (each 100 mL) to give fractions A-G. Fraction B (276.7 mg) was further purified by repeated ODS HPLC using MeOH-H₂O (40/60) and then MeCN-H₂O (18/82) to afford 6 (14.8 mg). Fraction E (551.7 mg) was further purified by repeated ODS HPLC using MeOH-H₂O (40/60) and then MeCN– H_2O (17/83) to afford 2 (7.7 mg).

3.4. Characteristics of each terpenoid

3.4.1. Tricalysiolide A (1). Amorphous solid; $[\alpha]_D^{25} - 215$ (*c* 0.11, CHCl₃); UV (MeOH) λ_{max} nm (log ε) 217 (4.29), 276 (1.25); IR (film) ν_{max} 3365, 2932, 2865, 1775, 1742, 1645, 1044, 1021 cm⁻¹; ¹H and ¹³C NMR data, Tables 1 and 2; HRESIMS *m*/*z* 333.2074 [M+H]⁺ (calcd for C₂₀H₂₉O₄, 333.2066).

3.4.2. Tricalysiolide B (2). Amorphous powder; $[\alpha]_{D}^{2D}$ -160 (*c* 0.33, pyridine); UV (MeOH) λ_{max} nm (log ε) 215 (3.93); IR (film) ν_{max} 3394, 2929, 2866, 1754, 1735, 1657, 1451, 1220, 1038 cm⁻¹; ¹H and ¹³C NMR data, Tables 1 and 2; HRESIMS *m/z* 349.2033 [M+H]⁺ (calcd for C₂₀H₂₉O₅, 349.2015).

3.4.3. Tricalysiolide C (3). Colorless prisms (CHCl₃– MeOH); mp 243–246 °C; $[\alpha]_D^{25}$ –243 (*c* 0.11, CHCl₃); UV (MeOH) λ_{max} nm (log ε) 217 (4.46); IR (film) ν_{max} 3419, 2933, 2865, 1762, 1656, 1452, 1194, 1170, 1053 cm⁻¹; ¹H and ¹³C NMR data, Tables 1 and 2; HRESIMS *m/z* 363.2182 [M+H]⁺ (calcd for C₂₁H₃₁O₅, 363.2171).

3.4.4. Tricalysiolide D (4). Colorless prisms (CHCl₃–MeOH); mp 251–253 °C; $[\alpha]_D^{25}$ –198 (*c* 0.25, pyridine); UV (MeOH) λ_{max} nm (log ε) 217 (4.13); IR (film) ν_{max} 3398, 2937, 2864, 1750, 1646, 1448, 1159, 1069, 1047, 1014 cm⁻¹; ¹H and ¹³C NMR data, Tables 1 and 2; HRESIMS *m*/*z* 347.2232 [M+H]⁺ (calcd for C₂₁H₃₁O₄, 347.2222).

3.4.5. Tricalysiolide E (5). Amorphous powder; $[\alpha]_{D}^{25}$ -203 (*c* 0.26, pyridine); UV (MeOH) λ_{max} nm (log ε) 276 (3.99); IR (film) ν_{max} 3389, 2925, 2852, 1746, 1669, 1605, 1464, 1455, 1445, 1259, 1213, 1055, 1021 cm⁻¹; ¹H and ¹³C NMR data, Tables 1 and 2; HRESIMS *m/z* 331.1901 [M+H]⁺ (calcd for C₂₀H₂₇O₄, 331.1909).

3.4.6. Tricalysiolide F (6). Amorphous powder; $[\alpha]_D^{25} - 218$ (*c* 0.14, pyridine); UV (MeOH) λ_{max} nm (log ε) 270 (3.89); IR (film) ν_{max} 3419, 2927, 2867, 1781, 1747, 1672, 1608, 1454, 1395, 1209, 1017 cm⁻¹; ¹H and ¹³C NMR data, Tables 1 and 2; HRESIMS *m/z* 347.1882 [M+H]⁺ (calcd for C₂₀H₂₇O₅, 347.1858).

3.5. Identification of structural relations between the terpenoids by synthetic procedures (Scheme 1)

3.5.1. Preparation of compound 7. To a solution of 1 (20.0 mg, 0.060 mmol) in pyridine-CH₂Cl₂ (1/1, 1 mL)added *p*-bromobenzoyl chloride (66.0 mg, were 0.301 mmol) and 4-(dimethylamino)pyridine (8.0 mg, 0.065 mmol) in one portion. After stirring at room temperature for 4 h, the mixture was diluted with EtOAc. The extract was washed successively with 5% aqueous HCl and brine, dried (MgSO₄), filtered, and concentrated to give a residue, which was then purified by silica gel column chromatography with CHCl₃-MeOH (50/1) to afford p-bromobenzoyl ester 7 (29.0 mg, 94%) as colorless prisms (pyridine); mp 230–232 °C; $[\alpha]_D^{25} = -92$ (c 0.11, CHCl₃); IR (film) ν_{max} 3442, 2924, 2852, 1721, 1679, 1647, 1452, 1396, 1266, 1013 cm⁻¹; ¹H NMR (500 MHz, pyridine- d_5) δ 8.07 (d, 2H, J=8.4 Hz), 7.61 (d, 2H, J= 8.4 Hz), 5.75 (s, 1H), 4.93 (d, 1H, J=11.3 Hz), 4.76 (t, 1H, J=9.3 Hz), 4.71 (d, 1H, J=11.3 Hz), 2.49 (br s, 1H), 2.12 (m, 1H), 2.06 (m, 1H), 1.97–1.92 (m, 2H), 1.89 (d, 1H, J= 14.4 Hz), 1.84 (d, 1H, J = 14.4 Hz), 1.71–1.63 (m, 4H), 1.59–1.43 (m, 6H), 1.16 (d, 1H, J=7.9 Hz), 0.98 (td, 1H, J = 13.8, 3.3 Hz), 0.69 (s, 3H); ¹³C NMR (125 MHz, pyridine-d₅) δ 174.9, 173.7, 166.2, 132.0 (2C), 131.7 (2C), 130.1, 128.1, 111.5, 81.4, 79.3, 70.1, 53.8, 53.4, 48.7, 46.4, 44.7, 42.9, 40.0, 37.9, 35.8, 29.9, 26.2, 22.0, 19.3, 14.7; HRESIMS m/z 515.1410 [M+H]⁺ (calcd for C₂₇H₃₂O₅Br, 515.1433).

3.5.2. Reduction of compound 2. Sodium borohydride (2.5 mg, 0.066 mmol) was added to a solution of **2** (5.0 mg, 0.014 mmol) in EtOH (1.5 mL), and then the mixture was stirred at room temperature for 5 min. After evaporation of the solvent, the residue was purified by ODS HPLC with MeCN-H₂O (23/77) to give a compound [1.7 mg, 36%, $[\alpha]_D^{25} - 186 \ (c \ 0.09, \text{CHCl}_3)]$, which was shown to be identified to the natural product **1**, by comparison of their ¹H NMR and mass spectra, and optical rotations.

3.5.3. Treatment of compound 3 with lithium hydroxide. To a solution of **3** (5.0 mg, 0.014 mmol) in THF–H₂O (10/1, 1.5 mL) was added lithium hydroxide monohydrate (3.0 mg, 0.071 mmol), and the mixture was stirred at room temperature for 48 h. After addition of acetic acid (1 mL) to the mixture, it was evaporated to dryness. The residue was subjected to ODS HPLC with MeCN–H₂O (23/77) to give a compound [4.2 mg, 87%, $[\alpha]_D^{25} - 173$ (*c* 0.11, pyridine)],

which was shown to be identical to natural 2 by comparison of their ¹H NMR and mass spectra, and optical rotations.

3.5.4. Tosylation of compound 1. To a solution of 1 (6.4 mg, 0.019 mmol) in pyridine (1 mL) was added p-toluenesulfonyl chloride (20 mg, 0.10 mmol) at 0 °C. The mixture was kept at room temperature for 18 h, and then was diluted with EtOAc. The solution was washed successively with aqueous Na₂CO₃ and brine, dried (MgSO₄), filtered, and concentrated to give a residue, which, by ODS HPLC with MeCN-H₂O (50/50), gave compound 8 (7.0 mg, 75%) as an amorphous solid: $[\alpha]_{D}^{25} - 152$ (c 0.17, CHCl₃); IR (film) ν_{max} 3445, 2930, 1746, 1644, 1357, 1189, 1175 cm⁻¹; ¹H NMR (500 MHz, pyridine- d_5) δ 8.06 (d, 2H, J=8.0 Hz), 7.27 (d, 2H, J= 8.0 Hz), 5.72 (s, 1H), 4.74 (t, 1H, J=9.0 Hz), 4.58 (d, 1H, J=9.6 Hz), 4.55 (d, 1H, J=9.6 Hz), 2.40 (br s, 1H), 2.22 (s, 3H), 2.19 (m, 1H), 1.98 (m, 1H), 1.86–1.84 (m, 2H), 1.79 (d, 1H, J = 14.3 Hz), 1.71 (d, 1H, J = 14.3 Hz), 1.64–1.55 (m, 3H), 1.51–1.32 (m, 7H), 1.07 (d, 1H, J=8.5 Hz), 0.95 (td, 1H, J = 13.6, 3.6 Hz), 0.62 (s, 3H); ¹³C NMR (125 MHz, pyridine-d₅) δ 174.8, 173.6, 145.1, 133.6, 130.3 (2C), 128.4 (2C), 111.5, 81.3, 78.8, 75.5, 53.3, 53.2, 48.6, 45.9, 44.5, 42.8, 39.9, 37.8, 35.8, 29.8, 25.8, 21.9, 21.3, 19.1, 14.6; HRESIMS m/z 487.2130 [M+H]⁺ (calcd for C₂₇H₃₅O₆S, 487.2154).

3.5.5. Epoxidation of compound 8. A solution of 8 (6.1 mg, 0.013 mmol) in EtOH (3 mL) was stirred with Na₂CO₃ (15 mg, 0.14 mmol) at room temperature for 48 h. The mixture was filtered and the filtrate was evaporated. By ODS HPLC with MeCN-H₂O (50/50), the residue gave compound 9 (2.9 mg, 74%) as an amorphous powder: $[\alpha]_D^{25}$ -223 (c 0.06, CHCl₃); IR (film) ν_{max} 2929, 2861, 1778, 1749, 1645, 1456, 1258, 1180, 1160, 1127, 1048, 1037, 1022 cm⁻¹; ¹H NMR (500 MHz, pyridine- d_5) δ 5.75 (s, 1H), 4.75 (t, 1H, J = 8.7 Hz), 2.89 (d, 1H, J = 5.0 Hz), 2.82 (d, 1H, J=5.0 Hz), 2.21 (m, 1H), 1.92–1.83 (m, 4H), 1.71-1.56 (m, 4H), 1.55-1.30 (m, 8H), 1.11 (d, 1H, J=8.0 Hz), 0.95 (td, 1H, J = 13.8, 3.7 Hz), 0.64 (s, 3H); ¹³C NMR (125 MHz, pyridine-d₅) δ 174.8, 173.6, 111.7, 81.3, 65.7, 52.2, 49.9, 49.0, 48.7, 45.1, 42.7, 42.6, 39.2, 38.9, 35.9, 29.8, 28.7, 21.6, 20.3, 14.9; HRESIMS m/z 315.1949 $[M+H]^+$ (calcd for C₂₀H₂₇O₃, 315.1960).

3.5.6. Methanolysis of compound 9. To a solution of 9 (2.6 mg, 0.0083 mmol) in MeOH (1.5 mL) was added BF₃·OEt₂ (3 µL, 0.024 mmol). The mixture was stirred at room temperature for 20 min. After neutralization with sodium acetate (5 mg), the solvent was evaporated under reduced pressure. By ODS HPLC with MeCN–H₂O (40/60) the residue gave a compound [0.9 mg, 31%, $[\alpha]_D^{25} - 241$ (*c* 0.05, pyridine)], which was shown to be identical to the natural product 4 by comparison of their ¹H NMR spectra and optical rotations.

3.5.7. Treatment of compound 2 with (trimethylsilyl)diazomethane. To a solution of **2** (11.0 mg, 0.032 mmol) in CHCl₃–MeOH (10/1, 3 mL) was added (trimethylsilyl)diazomethane (2.0 M in hexanes, 1 mL, 2.0 mmol). The mixture was stirred at room temperature for 1 h. After removal of the solvent and subsequent ODS HPLC of the residue with MeCN–H₂O (22/78), along with the starting material **2** (1.0 mg, 9%) and **3** (3.8 mg, 33%), another compound [1.6 mg, 15%, $[\alpha]_D^{25} - 136$ (*c* 0.06, pyridine)] was obtained. On the basis of the ¹H NMR and mass spectra, and optical rotations, the third compound was shown to be identical to the natural product **5**.

3.5.8. Reduction of compound 6. Triethylsilane (0.5 mL, 6.1 mmol) and BF₃·OEt₂ (5 μ L, 0.039 mmol) were added to a solution of compound **6** (5.0 mg, 0.014 mmol) in CH₂Cl₂ (2 mL). The mixture was stirred at room temperature for 4 h. After neutralization with sodium acetate (10 mg), the solvent was removed by evaporation and the residue was separated by ODS HPLC with MeCN–H₂O (28/72) to give a compound [1.7 mg, 36%, $[\alpha]_D^{25}$ – 101 (*c* 0.05, pyridine)], which was shown to be identical to the natural product **5**, by comparison of their ¹H NMR and mass spectra, and optical rotations.

3.6. X-ray single crystallographic analysis

The structures of compounds **3**, **4**, and **7** were determined by the direct method using the maXus crystallographic software package¹¹ and the refinement was carried out by the program *SHELXL-97*.¹²

Crystal data for **3**. C₂₁H₃₀O₅; M=362.45; orthorhombic; space group $P2_12_12_1$; a=11.155(3) Å; b=13.087(3) Å; c=12.476(3) Å; V=1821.31(8) Å; Z=4; D_X =1.322 Mg m⁻³; μ (Mo K α)=0.093 mm⁻¹; R=0.0349; R_w =0.0974.

Crystal data for **4**. $C_{21}H_{30}O_4$; M=346.45; orthorhombic; space group $P2_12_12_1$; a=6.899(10) Å; b=11.322(5) Å; c=22.828(10) Å; V=1783.11(11) Å; Z=4; $D_X=1.291$ Mg m⁻³; μ (Mo K α)=0.088 mm⁻¹; R=0.0403; $R_w=0.1064$.

Crystal data for 7. $C_{27}H_{31}BrO_5 \cdot C_5H_5N$; M=594.52; orthorhombic; space group $P2_12_12_1$; a=6.259(10) Å; b=19.99(10) Å; c=22.348(10) Å; V=2796.12(19) Å; Z=4; $D_X=1.412$ Mg m⁻³; μ (Mo K α)=1.512 mm⁻¹; R=0.0501; $R_w=0.067$; Flack parameter=0.012(7).

Crystallographic data for **3**, **4**, and **7** reported in this paper have been deposited at the Cambridge Crystallographic Data Centre, under the reference numbers CCDC 280148, 280149, and 280147, respectively. Copies of the data can be obtained, free of charge, on application to the Director, CCDC, 12 Union Road, Cambridge CB2 1EZ, UK (fax: +44 1223 336033 or e-mail: deposit@ccdc.cam.ac.uk).

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Reactions of (1R,2S)-1,2-di(2-furyl)-1,2-di(3-guaiazulenyl)ethane and (1R,2S)-1,2-di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane with tetracyanoethylene (TCNE) in benzene: comparative studies on the products and their spectroscopic properties

Shin-ichi Takekuma,^{a,*} Kenji Takahashi,^a Akio Sakaguchi,^a Masato Sasaki,^a Toshie Minematsu^b and Hideko Takekuma^a

^aDepartment of Applied Chemistry, Faculty of Science and Engineering, Kinki University, 3-4-1 Kowakae, Higashi-Osaka-shi, Osaka 577-8502, Japan ^bSchool of Pharmaceutical Sciences, Kinki University, 3-4-1 Kowakae, Higashi-Osaka-shi, Osaka 577-8502, Japan

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Abstract—Reactions of the title meso forms, (1R,2S)-1,2-di(2-furyl)-1,2-di(3-guaiazulenyl)ethane (1) and (1R,2S)-1,2-di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane (2), with a two molar amount of TCNE in benzene at 25 °C for 5 h (for 1) and 48 h (for 2) under oxygen give new compounds, 2,2,3,3-tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4-dihydrocyclohepta[c,d]azulene (3) and 2,2,3,3-tetracyano-8-isopropyl-6-methyl-4-(2-thienyl)-1,4-dihydrocyclohepta[c,d]azulene (4), respectively, in 74 and 21% isolated yields. Comparative studies on the above reactions as well as the spectroscopic properties of the unique products 3 and 4, possessing interesting molecular structures, are reported and, further, a plausible reaction pathway for the formation of these products is described. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

In the previous papers,¹⁻⁹ we reported a facile preparation and the crystal structures as well as the spectroscopic, chemical and electrochemical properties of the mono- and dicarbocations stabilized by a 3-guaiazulenyl group. During the course of our investigations, we have recently found (i) that the reactions of naturally occurring guaiazulene (A) with 2-furaldehyde (B) and thiophene-2-carbaldehyde (C) in methanol in the presence of hexafluorophosphoric acid gave (2-furyl)(3guaiazulenyl)methylium and (3-guaiazulenyl)(2-thienyl)methylium hexafluorophosphates (D and E) with the following two representative resonance forms [i.e., 3-guaiazulenylium and 2-furanylium (and 2-thienylium) structures], respectively, in 93 and 98% isolated yields (see Fig. 1),¹⁰ which upon reduction with zinc powder in acetonitrile afforded a mixture of a meso form and two enantiomeric forms of the molecular structures, 1,2-di(2-furyl)-1,2-di(3-guaiazulenyl) ethane (F) and

1,2-di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane (G), respectively, (see Fig. 2),¹⁰ and have quite recently found (ii) that, from a comparative study on the oxidation potentials of the above meso forms, (1R,2S)-1,2-di(2furyl)-1,2-di(3-guaiazulenyl)ethane (1) and (1R,2S)-1,2di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane (2), measured by means of the CV and DPV (Potential/V vs SCE) in CH₂Cl₂ containing 0.1 M [n-Bu₄N]PF₆ as a supporting electrolyte, it could be inferred that 1 and 2 simultaneously underwent two-electron oxidation of their 1,2di(3-guaiazulenyl) groups, respectively, at the potentials of +0.48 (E_{pa} , irreversible) V by CV and +0.48 (E_p) V by DPV for **1**¹¹ and +0.57 (E_{pa} , irreversible) V by CV and +0.56 (E_p) V by DPV for **2**,¹² generating the corresponding di(cation-radical) species **1**^{2(+·)} and **2**^{2(+·)}. Thus, 1 was more susceptible to oxidation than 2 and A [+0.69 (E_{pa} , irreversible) V by CV and +0.65 (E_p) V by DPV],⁸ owing to the difference in ionization potential (corresponding to π -HOMO) based on those π -electron systems. The different oxidation potentials between 1 and 2 are obviously caused by the influence of a different heteroaromatic-ring. Therefore, our interest has been focused on a comparative study on the title chemistry, the reactions of 1 and 2 with TCNE, which serves as an electron acceptor [-0.75 ($E_{1/2}$, quasi-reversible) V by CV and

Keywords: Annulation; C-C bond cleavage; Charge-transfer complex; Furan; Guaiazulene; TCNE; Thiophene; X-ray crystal structure.

^{*} Corresponding author. Tel.: +81 6 6730 5880x4020; fax: +81 6 6727 4301; e-mail: takekuma@apch.kindai.ac.jp

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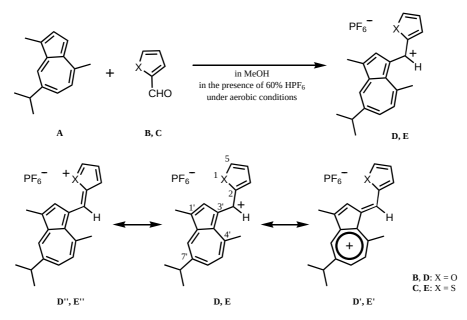


Figure 1. The reactions of A with B and C in CH₃OH with HPF₆ under aerobic conditions, affording the corresponding monocarbocation products D and E with the resonance forms of D', E' and D'', E'', respectively.¹⁰

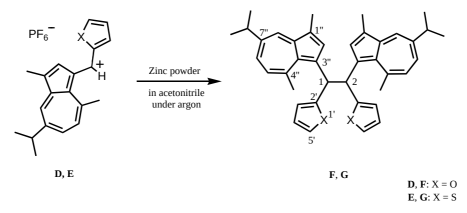


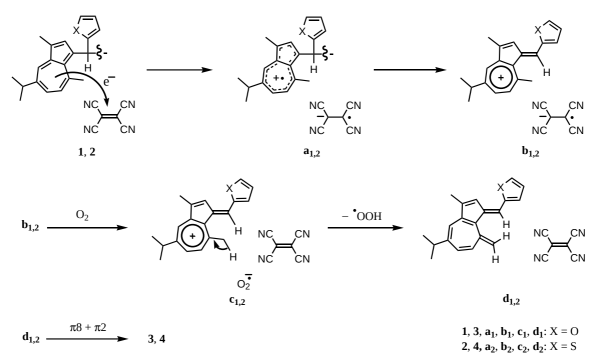
Figure 2. The reductions of **D** and **E** with zinc powder in acetonitrile at 25 $^{\circ}$ C under argon, affording a mixture of a *meso* form and two enantiomeric forms of the molecular structures **F** and **G**, respectively.¹⁰

 $-0.76 (E_p)$ V by DPV].¹³ In relation to our basic studies, in 1961 Hafner and Moritz reported that the reaction of guaiazulene (A) in petroleum ether with TCNE in AcOEt at -20 °C gave a 1:1 π -complex in 98% isolated yield, which was converted into 3-tricyanovinylguaiazulene (68% isolated yield) in DMF at room temperature.¹⁴ Furthermore, the addition and cycloaddition reactions of TCNE in organic chemistry^{15–19} including azulenes^{14,20–22} have been studied to a considerable extent, and a large number of those results and discussion have been well documented. Along with those investigations, we now wish to report the detailed title chemistry, affording the unique products, 2,2,3,3-tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4dihydrocyclohepta[c,d]azulene (3) from 1 and 2,2,3,3tetracyano-8-isopropyl-6-methyl-4-(2-thienyl)-1,4-dihydrocyclohepta[c,d]azulene (4) from 2, possessing interesting molecular structures, presumably via charge-transfer, C-C bond cleavage and, further, $[\pi 8 + \pi 2]$ cycloaddtion reactions (see Scheme 1).

2. Results and discussion

2.1. Reactions of (1R,2S)-1,2-di(2-furyl)-1,2-di(3-guaiazulenyl)ethane (1) and (1R,2S)-1,2-di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane (2) with TCNE; preparation and spectroscopic properties of 2,2,3,3-tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4-dihydrocyclohepta[c,d]azulene (3) and 2,2,3,3-tetracyano-8-isopropyl-6-methyl-4-(2thienyl)-1,4-dihydrocyclohepta[c,d]azulene (4)

Compound **3** was prepared using benzene as a solvent as shown in Figure 3 and Section 4.1.1, whose molecular structure was established on the basis of elemental analysis and spectroscopic data [UV–vis, IR, exact FAB-MS, ¹H and ¹³C NMR including 2D NMR (i.e., H–H COSY, NOESY, HMQC = ¹H detected hetero nuclear multiple quantum coherence and HMBC = ¹H detected hetero nuclear multiple bond connectivity)].



Scheme 1. A plausible reaction pathway for the formation of 3 and 4 yielded by the reactions of the title *meso* forms 1 and 2 with a two molar amount of TCNE in benzene at 25 $^{\circ}$ C for 5 h (for 1) and 48 h (for 2) under oxygen. The partial structures 1, 2 and $a_{1,2}$ are illustrated.

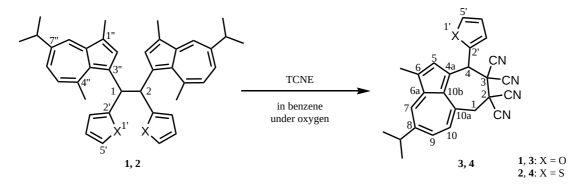


Figure 3. The reactions of the title meso forms 1 and 2 with a two molar amount of TCNE in benzene at 25 °C for 5 h (for 1) and 48 h (for 2) under oxygen.

Compound 3 (74% isolated yield) was blue prisms $[R_f =$ 0.27 on silica-gel TLC (hexane/AcOEt=4:6, vol/vol)], mp 163 °C and decomp. >171 °C [determined by the thermal analysis (TGA and DTA)]. The characteristic UV-vis (CH₂Cl₂) absorption bands based on the azulenyl group were observed and the longest visible absorption wavelength appeared at λ_{max} 622 nm (log $\varepsilon = 2.69$). The IR (KBr) spectrum showed a specific band based on the −C≡N group at 2252 cm⁻¹. The protonated molecular formula $C_{26}H_{21}N_4O~([M+H]^+)$ was determined by the exact FAB-MS (3-nitrobenzyl alcohol matrix) spectrum. The elemental analysis confirmed the molecular formula $C_{26}H_{20}N_4O$. The 500 MHz ¹H NMR (C_6D_6) spectrum showed signals based on the structure of 8-isopropyl-6methyl-1,4-dihydrocyclohepta[c,d]azulene possessing a 2-furyl group at the C-4 position, which signals (δ and J values) were carefully assigned using the H-H COSY technique and the computer simulation analysis (see Section 4.1.1). Moreover, the NOESY spectrum revealed apparent cross peaks between the following signals [i.e., $-C^{1}H_{2}$ - and H-10; δ 3.15 (axial; one of two signals based on the –C¹H₂–) and $C^{4}H$ – (axial)], and suggested that the 2-furyl group

was substituted at the C-4 (equatorial) position. The 125 MHz ¹³C NMR (C₆D₆) spectrum exhibited 25 carbon signals (δ) assigned by the HMQC and HMBC techniques (see Section 4.1.1). Thus, the elemental analysis and these spectroscopic data for **3** led to a new molecular structure, 2,2,3,3-tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4-dihydrocyclohepta[*c*,*d*]azulene (see Fig. 3).

Similarly, as in the case of the preparation of **3**, the target compound **4** was prepared using benzene as a solvent as shown in Figure 3 and Section 4.1.2, whose molecular structure was established on the basis of similar spectroscopic analyses as **3**. Although this reaction afforded numerous products as compared with the reaction of **1** with TCNE (see Section 4.1.1), compound **4** could be isolated in 21% yield. The difference between the yields of the products **3** and **4** is obviously caused by the different oxidation potentials between 1^{11} and 2^{12} (see Section 1), whose oxidation potentials suggest that the reaction of **1** with TCNE readily gives the corresponding 1:2 charge-transfer (CT) complex **a**₁ (see Scheme 1) as compared with the reaction of **2** with TCNE.

Compound 4 was blue blocks $[R_f=0.61 \text{ on silica-gel TLC}]$ (hexane/AcOEt=4:6, vol/vol)], mp 147 °C and decomp. >158 °C [determined by the thermal analysis (TGA and DTA)]. The characteristic UV-vis (CH₂Cl₂) absorption bands based on the azulenyl group, which spectral pattern coincided with that of 3, were observed and the longest visible absorption wavelength appeared at λ_{max} 624 nm (log $\varepsilon = 2.64$). The IR (KBr) spectrum showed a specific band based on the $-C \equiv N$ group at 2252 cm⁻¹, which wavenumber coincided with that of 3. The protonated molecular formula $C_{26}H_{21}N_4S$ ([M+H]⁺) was determined by the exact FAB-MS (3-nitrobenzyl alcohol matrix) spectrum. The 500 MHz ¹H NMR (C₆D₆) spectrum showed signals based on the structure of 8-isopropyl-6-methyl-1,4dihydrocyclohepta[c,d]azulene possessing a 2-thienyl group at the C-4 position, which signals (δ and J values) were carefully assigned using the H-H COSY technique and the computer simulation analysis (see Section 4.1.2). Similarly, as in the case of 3, the NOESY spectrum revealed apparent cross peaks between the following signals [i.e., $-C^{1}H_{2}$ - and H-10; δ 3.18 (axial; one of two signals based on the $-C^{1}H_{2}$ -) and $C^{4}H$ – (axial)], and suggested that the 2-thienyl group was substituted at the C-4 (equatorial) position. The 125 MHz ¹³C NMR (C₆D₆) spectrum exhibited 25 carbon signals (δ) assigned by the HMQC and HMBC techniques (see Section 4.1.2). Thus, these spectroscopic data for 4 led to a new molecular structure, 2,2,3,3-tetracyano-8-isopropyl-6-methyl-4-(2-thienyl)-1,4-dihydrocyclohepta[c,d]azulene (see Fig. 3).

2.2. X-ray crystal structure of 3

Although an X-ray crystallographic analysis of **4** has not yet been achieved because of difficulty in obtaining a single crystal suitable for this purpose, the crystal structure of **3** has been determined by means of the X-ray diffraction (see Section 4.1.3). In the course of refinement, the atoms based on the 2-furyl group at the C-4 (equatorial) position and two methyl groups of the isopropyl group at the C-8 position for **3** were found to be disordered over two sites of equal occupancy. Therefore, those atoms in the occupancy ratio 50:50 were refined. The only ORTEP drawing of **3** noted the above disorder, indicating the molecular structure, 2,2,3,3tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4-dihydrocyclohepta[*c*,*d*]azulene, illustrated in Figure 4a, is shown in Figure 4b. As the result, the crystal structure of **3** clarified the conformation of 2,2,3,3-tetracyano-8-isopropyl-6-

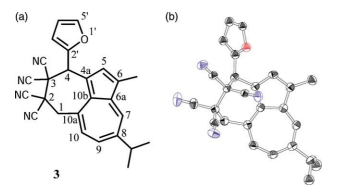


Figure 4. (a) The molecular structure of **3**. (b) The ORTEP drawing of **3** (30% probability thermal ellipsoids).

methyl-1,4-dihydrocyclohepta[c,d]azulene possessing a 2-furyl group at the C-4 (equatorial) position, presumed by means of the NOESY spectrum (see Sections 2.1 and 4.1.1). From a comparative study on the detailed spectroscopic properties of **3** and **4** (see Sections 4.1.1 and 4.1.2), it can be inferred that the crystal structure of **4** assumes similar conformation to that of **3**.

2.3. A plausible reaction pathway for the formation of the products 3 and 4

From the molecular structures of the resulting products **3** and **4**, obtained by the reactions of the *meso* forms **1** and **2** with a two molar amount of TCNE in benzene at 25 °C for 5 h (for **1**) and 48 h (for **2**) under oxygen, a plausible reaction pathway for the formation of compounds **3** and **4** can be inferred as illustrated in Scheme 1; namely, (i) the reactions of **1** and **2** with TCNE gradually give the corresponding 1:2 charge-transfer (CT) complexes **a**_{1,2}, respectively, whose formation was supported by the time-dependent visible spectra of the reaction solutions as shown in Figures 5 and 6. The characteristic CT band based on the formation of **a**₁, generated from the reaction of **1** with TCNE, appeared at λ_{max} 518 nm (the time to reach

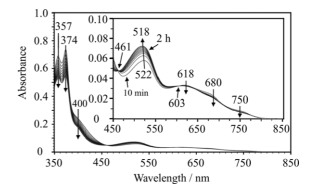


Figure 5. The time-dependent visible spectra for the reaction of **1** (2.0 mg, 3.6 µmol) with TCNE (1.0 mg, 7.8 µmol) in benzene (10 mL) at 25 °C for 2 h under argon. Length of the cell: 0.1 cm. Interval time: 10 min. λ_{max} 357, 374 and 400sh nm: the specific bands based on TCNE; λ_{max} 518 nm: the specific band based on the formation of the CT complex **a**₁ (see Scheme 1) (the time to reach maximum absorption=2 h); λ_{max} 618, 680sh and 750sh nm: the specific bands based on **1**.

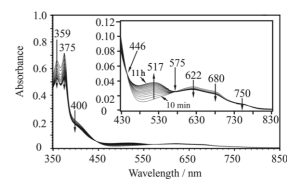


Figure 6. The time-dependent visible spectra for the reaction of **2** (2.0 mg, 3.4 µmol) with TCNE (1.0 mg, 7.8 µmol) in benzene (10 mL) at 25 °C for 11 h under argon. Length of the cell: 0.1 cm. Interval time: 1 h. λ_{max} 359, 375 and 400sh nm: the specific bands based on TCNE; λ_{max} 517 nm: the specific band based on the formation of the CT complex **a**₂ (see Scheme 1) (the time to reach maximum absorption=11 h); λ_{max} 622, 680sh and 750sh nm: the specific bands based on **2**.

maximum absorption =2 h), whose band revealed a bathochromic shift ($\Delta 19$ nm) in comparison with the characteristic visible band of (2-furyl)(3-guaiazulenyl)methylium hexafluorophosphate (**D**) (λ_{max} 499 nm, log $\varepsilon =$ 4.61);¹⁰ and, similarly, the characteristic CT band based on the formation of a_2 , generated from the reaction of 2 with TCNE, appeared at λ_{max} 517 nm (the time to reach maximum absorption = 11 h), whose band showed a bathochromic shift ($\Delta 20$ nm) in comparison with the characteristic visible band of (3-guaiazulenyl)(2-thienyl)methylium hexafluorophosphate ($\mathbf{\tilde{E}}$) (λ_{max} 497 nm, log ε = 4.73).¹⁰ The time to reach maximum absorption indicates that 1 more readily forms the corresponding 1:2 CT complex a_1 than 2, whose result is obviously caused by the different oxidation potentials between 1^{11} and 2^{12} (see Section 1); and (ii) although the CT complexes $a_{1,2}$ generated were stable in benzene at 25 °C for a long time (>72 h) under argon and they did not give 3 and 4, the complexes $a_{1,2}$ were gradually converted into 3 and 4 in benzene at 25 °C under oxygen. Furthermore, the reactions of 1 and 2 with a two molar amount of TCNE in toluene at -20 °C for 24 h under argon gave the corresponding 1:2 CT complexes $a_{1,2}$, respectively, in 18% isolated yield from 1^{23} and 23% isolated yield from 2^{24} , whose complexes were gradually converted into 3 and 4, respectively, in benzene at 25 °C under oxygen. Along with the above results, the reactions of (2-furyl)(3-guaiazulenyl)methylium and (3guaiazulenyl)(2-thienyl)methylium hexafluorophosphates (D and E) with an equimolar amount of TCNE in acetonitrile at 25 °C for 72 h under oxygen did not afford any products. From these results, it can be inferred that the CT complexes $a_{1,2}$ generated are converted into 3 and 4 presumably via the 3-guaiazulenylium-ions $b_{1,2}$ yielded by the C-C bond cleavage of the ethane units of $a_{1,2}$, the azulenequinodimethanes $d_{1,2}$ produced by the deprotonations of the C4-methyl groups of $c_{1,2}$ by the attack of $O_2^$ generated by the electron transfer from the TCNE⁻ to O_2 and, further, the $[\pi 8 + \pi 2]$ cycloaddition reactions of $d_{1,2}$ with TCNE, as illustrated in Scheme 1. On the other hand, in 1988 one of us (S. Takekuma) reported that the autoxidation of guaiazulene (A) under several reaction conditions gave a wide variety of products possessing interesting molecular structures, respectively, through such several types of the reactions as oxidative dimerization, oxidation of side chains, azulenequinone formation, intermolecular onecarbon transfer reactions and rearrangements to naphthalenoid, benzenoid and 1H-inden-1-one derivatives, simultaneously.²⁵ In this case, by the attack of oxygen, A was considered to initially form an electron-transfer complex $(\mathbf{A}^+, \mathbf{O}_2^-)$, which in turn leads to the formation of various products.

3. Conclusion

We have reported the following four points in this paper: (i) the reactions of the title *meso* forms, (1R,2S)-1,2-di(2-furyl)-1,2-di(3-guaiazulenyl)ethane (1) and <math>(1R,2S)-1,2-di(3-guaiazulenyl)-1,2-di(2-thienyl)ethane (2), with a two molar amount of tetracyanoethylene (TCNE) in benzene at 25 °C for 5 h (for 1) and 48 h (for 2) under oxygen gave new compounds 3 and 4, respectively, in 74 and 21% isolated yields; (ii) the detailed spectroscopic analyses of these products led to the molecular structures, 2,2,3,3-tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4-dihydrocyclohepta-[c,d]azulene for **3** and 2,2,3,3-tetracyano-8-isopropyl-6methyl-4-(2-thienyl)-1,4-dihydrocyclohepta[c,d]azulene for **4**; (iii) along with the spectroscopic data for **3** and **4**, we clarified the crystal structure of **3**; and, further, (iv) a plausible reaction pathway for the formation of the unique products **3** and **4**, possessing interesting molecular structures, was described.

4. Experimental

4.1. General

Thermal (TGA/DTA) and elemental analyses were taken on a Shimadzu DTG-50H thermal analyzer and a Yanaco MT-3 CHN corder, respectively. MS spectra were taken on a JEOL The Tandem Mstation JMS-700 TKM data system. UV–vis and IR spectra were taken on a Beckman DU640 spectrophotometer and a Shimadzu FTIR-4200 Grating spectrometer, respectively. NMR spectra were recorded with a JEOL GX-500 (500 MHz for ¹H and 125 MHz for ¹³C) at 25 °C. The ¹H NMR spectra were assigned using the computer-assisted simulation analysis (the software: gNMR developed by Adept Scientific plc) on a DELL Dimension XPS T500 personal-computer with a Pentium III processor. Cyclic and differential pulse voltammograms were measured by an ALS Model 600 electrochemical analyzer.

4.1.1. Reaction of (1R,2S)-1,2-di(2-furyl)-1,2-di(3-guaiazulenyl)ethane (1) with TCNE; preparation and spectroscopic properties of 2,2,3,3-tetracyano-4-(2-furyl)-8isopropyl-6-methyl-1,4-dihydrocyclohepta[c,d]azulene (3). To a solution of TCNE (48 mg, 0.37 mmol) in benzene (10 mL) was slowly added a solution of 1 (97 mg, 0.17 mmol) [$R_f = 0.71$ on silica-gel TLC (hexane/AcOEt = 4:6, vol/vol)] in benzene (30 mL), turning the blue solution into a red solution. The mixture was stirred at 25 °C for 5 h under oxygen, turning the red solution into a green solution. After the reaction, the reaction solution was evaporated in vacuo, giving a green solid. The residue obtained was carefully separated by silica-gel column chromatography with hexane–ethyl acetate (3/2, vol/vol) as an eluant. The crude product 3 thus obtained was recrystallized from benzene-cyclohexane (1/5, vol/vol) (several times) to provide pure 3 as stable crystals (104 mg, 0.26 mmol, 74% yield).

Compound **3**: blue prisms $[R_f=0.27 \text{ on silica-gel TLC}$ (hexane/AcOEt=4:6, vol/vol)], mp 163 °C and decomp. > 171 °C [determined by thermal analysis (TGA and DTA)]; Found: C, 76.89; H, 5.15; N, 13.66%. Calcd for $C_{26}H_{20}N_4O$: C, 77.21; H, 4.98; N, 13.85%; UV–vis λ_{max} (CH₂Cl₂) nm (log ε), 247 (4.46), 296 (4.65), 358 (3.86), 376 (3.86), 622 (2.69), 684sh (2.58) and 754sh (2.12); IR ν_{max} (KBr) cm⁻¹, 2252 (–C=N); FAB-MS (3-nitrobenzyl alcohol matrix), m/z 405 ([M+H]⁺, 100%) and 404 (M⁺, 94); exact FAB-MS (3-nitrobenzyl alcohol matrix), found: m/z 405.1711; calcd for $C_{26}H_{21}N_4O$: [M+H]⁺, m/z 405.1715; 500 MHz ¹H NMR (C₆D₆), signals based on the 1,4-dihydro-8isopropyl-6-methylcyclohepta[c,d]azulene unit at δ 1.05 (6H, d, J=6.9 Hz, (CH₃)₂CH-8), 2.26 (3H, s, Me-6), 2.60 (1H, sept, J=6.9 Hz, (CH₃)₂CH-8), 3.15 (axial), 3.42 (equatorial) (1H each, d, J=15.2 Hz, $-C^{1}H_{2}$ -), 5.11 (1H axial, br d s, $C^{4}H$ –), 6.36 (1H, d, J=10.4 Hz, H-10), 6.98 (1H, dd, J = 10.4, 2.0 Hz, H-9), 7.00 (1H, br d s, H-5) and7.94 (1H, d, J=2.0 Hz, H-7) and signals based on the 2-furyl group at δ 5.99 (1H, dd, J = 3.5, 2.0 Hz, H-4'), 6.05 (1H, ddd, J=3.5, 0.9, 0.7 Hz, H-3') and 7.03 (1H, dd, J=2.0, 0.9 Hz, H-5'); the NOESY spectrum showed cross peaks between the following signals [- C^1H_2 - and H-10; δ 3.15 (axial; one of two signals based on the $-C^{1}H_{2}$ -) and $C^{4}H$ (axial)]; 125 MHz⁻¹³C NMR (C₆D₆), signals based on the 2,2,3,3-tetracyano-8-isopropyl-6-methyl-1,4-dihydrocyclohepta[c,d]azulene unit at δ 12.2 (Me-6), 24.1 $((CH_3)_2CH-8)$, 38.2 $((CH_3)_2CH-8)$, 44.3 $(-C^1H_2-)$, 45.7 $(C^{4}H-)$, 43.8, 49.6 [(NC)₂C-2 or (NC)₂C-3], 111.22, 111.24, 111.4, 112.2 [(NC)₂C-2 or (NC)₂C-3], 118.7 (C-10b), 125.5 (C-6a), 126.9 (C-10), 127.4 (C-4a), 134.9 (C-10a), 135.3 (C-9), 135.5 (C-7), 140.1 (C-6), 140.6 (C-5) and 144.1 (C-8) and signals based on the 2-furyl group at δ 111.1 (C-4'), 112.9 (C-3'), 144.4 (C-5') and 148.8 (C-2').

4.1.2. Reaction of (1*R*,2*S*)-1,2-di(3-guaiazulenyl)-1,2di(2-thienyl)ethane (2) with TCNE; preparation and spectroscopic properties of 2,2,3,3-tetracyano-8-isopropyl-6-methyl-4-(2-thienyl)-1,4-dihydrocyclohepta[c,d]azulene (4). To a solution of TCNE (57 mg, 0.44 mmol) in benzene (10 mL) was slowly added a solution of 2 (130 mg, 0.22 mmol) [$R_f = 0.70$ on silica-gel TLC (hexane/AcOEt = 4:6, vol/vol)] in benzene (20 mL), turning the blue solution into a red solution. The mixture was stirred at 25 °C for 48 h under oxygen, turning the red solution into a green solution. After the reaction, the reaction solution was evaporated in vacuo, giving a green solid. The residue obtained was carefully separated by silica-gel column chromatography with hexane-ethyl acetate (3/2, vol/vol) as an eluant. The crude product 4 thus obtained was recrystallized from dichloromethane-hexane (1/5, vol/vol) (several times) to provide pure 4 as stable crystals (40 mg, 0.10 mmol, 21% vield).

Compound 4: blue blocks $[R_f=0.61$ on silica-gel TLC (hexane/AcOEt=4:6, vol/vol)], mp 147 °C and decomp. >158 °C [determined by thermal analysis (TGA and DTA)]; UV–vis λ_{max} (CH₂Cl₂) nm (log ε), 247 (4.46), 297 (4.59), 360 (3.79), 377 (3.77), 624 (2.64), 683sh (2.55) and 760sh (2.06); IR ν_{max} (KBr) cm⁻¹, 2252 (-C \equiv N); FAB-MS (3-nitrobenzyl alcohol matrix), m/z 421 ([M+H]⁺, 100%) and 420 (M⁺, 80); exact FAB-MS (3-nitrobenzyl alcohol matrix), found: m/z 421.1449; calcd for C₂₆H₂₁N₄S: $[M+H]^+$, m/z 421.1487; 500 MHz ¹H NMR (C₆D₆), signals based on the 1,4-dihydro-8-isopropyl-6-methylcyclohepta[c,d]azulene unit at δ 1.04 (6H, d, J=6.9 Hz, $(CH_3)_2$ CH-8), 2.23 (3H, s, Me-6), 2.59 (1H, sept, J = 6.9 Hz, (CH₃)₂CH-8), 3.18 (axial), 3.24 (equatorial) (1H each, d, J = 15.1 Hz, $-C^{1}H_{2}$ -), 5.30 (1H axial, br d s, $>C^{4}H_{-}$), 6.33 (1H, d, J=10.3 Hz, H-10), 6.96 (1H, dd, J=10.3, 2.0 Hz,H-9), 7.17 (1H, br d s, H-5) and 7.93 (1H, d, J = 2.0 Hz, H-7) and signals based on the 2-thienyl group at δ 6.65 (1H, dd, J=5.2, 3.7 Hz, H-4'), 6.80 (1H, dd, J=5.2, 1.1 Hz, H-5') and 7.11 (1H, dd, J=3.7, 1.1 Hz, H-3'); the NOESY spectrum showed cross peaks between the following signals $[-C^{1}H_{2}-$ and H-10; δ 3.18 (axial; one of two signals based on the $-C^{1}H_{2}$ and $C^{4}H_{-}$ (axial)]; 125 MHz ¹³C NMR (C_6D_6) , signals based on the 2,2,3,3-tetracyano-8-isopropyl6-methyl-1,4-dihydrocyclohepta[c,d]azulene unit at δ 12.3 (Me-6), 24.1 ((CH₃)₂CH-8), 38.2 ((CH₃)₂CH-8), 44.4 (-C¹H₂-), 47.0 (C^{4} H-), 44.3, 51.9 [(NC)₂C-2 or (NC)₂C-3], 111.3, 111.5, 111.7, 112.1 [(NC)₂C-2 or (NC)₂C-3], 120.8 (C-4a), 125.2 (C-6), 126.9 (C-10), 135.0 (C-10a), 135.2 (C-10b), 135.4 (C-9), 135.6 (C-7), 140.2 (C-6a), 141.0 (C-5) and 144.2 (C-8) and signals based on the 2-thienyl group at δ 127.5 (C-5'), 127.8 (C-4'), 129.8 (C-3') and 138.6 (C-2').

4.1.3. X-ray crystal structure of 2,2,3,3-tetracyano-4-(2-furyl)-8-isopropyl-6-methyl-1,4-dihydrocyclohepta-[c,d]azulene (3). A total 5443 reflections with $2\theta_{\text{max}} = 55.0^{\circ}$ were collected on a Rigaku AFC-5R automated four-circle diffractometer with graphite monochromated Mo K α radiation ($\lambda = 0.71069$ Å, rotating anode: 50 kV, 180 mA) at 296 K. The structure was solved by direct methods (SIR92) and expanded using Fourier techniques (DIRDIF99). The atoms based on the 2-furyl group at the C-4 (equatorial) position and two methyl groups of the isopropyl group at the C-8 position for **3** were disordered over two sites of equal occupancy. Therefore, those atoms in the occupancy ratio 50:50 were refined. The non-hydrogen atoms were refined anisotropically. The final cycle of full-matrix least-squares refinement was based on F^2 . All calculations were performed using the Crystal Structure 3.7.0 software package. Crystallographic data have been deposited at the CCDC, 12 Union Road, Cambridge CB2 1EZ, UK and copies can be obtained on request, free of charge, by quoting the publication citation and the deposition number CCDC 248295.

Crystallographic data for **3**: $C_{26}H_{20}N_4O$ (FW=404.47), blue prism (the crystal size, $0.80 \times 0.30 \times 0.30$ mm³), monoclinic, P_{21}/c (#14), a=10.278(2) Å, b=15.3808(19) Å, c=14.2500(19) Å, $\beta=91.831(14)^\circ$, V=2251.5(7) Å³, Z=4, $D_{calcd}=1.193$ g/cm³, μ (Mo K α) =0.750 cm⁻¹, Scan width= $(1.47+0.30 \tan\theta)^\circ$, Scan mode= $\omega - 2\theta$, Scan rate=9.0°/min, measured reflections=5443, observed reflections=5163, No of parameters=270, R1=0.0677, wR2=0.2539 and Goodness of Fit Indicator =1.027.

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- 11. The electrochemical measurement conditions of **1** are as follows: the cyclic and differential pulse voltammograms (Potential/V vs SCE) of **1** (3.0 mg, 5.4 μ mol) in 0.1 M [*n*-Bu₄N]PF₆, CH₂Cl₂ (10 mL) at a glassy carbon (ID: 3 mm) and a platinum wire served as the working and auxiliary electrodes; scan rates 100 mV s⁻¹ at 25 °C under argon, respectively. For comparative purposes, the oxidation potential using ferrocene as a standard material showed +0.42 (E_p) V by DPV and +0.42 ($E_{1/2}$, quasi-reversible) V by CV in 0.1 M [*n*-Bu₄N]PF₆, CH₂Cl₂ under the same electrochemical conditions as **1**.
- The cyclic and differential pulse voltammograms of 2 (3.0 mg, 5.1 μmol) were measured under the same electrochemical conditions as 1.
- 13. The electrochemical measurement conditions of TCNE are as follows: the cyclic and differential pulse voltammograms (Potential/V vs SCE) of TCNE (3.0 mg, 23.4 µmol) in 0.1 M [*n*-Bu₄N]BF₄, CH₃CN (10 mL) at a glassy carbon (ID: 3 mm) and a platinum wire served as the working and auxiliary electrodes; scan rates 100 mV s⁻¹ at 25 °C under argon, respectively. For comparative purposes, the oxidation potential using ferrocene as a standard material showed +0.45 (E_p) V by DPV and +0.42 ($E_{1/2}$, quasi-reversible) V by CV in 0.1 M [*n*-Bu₄N]BF₄, CH₃CN under the same electrochemical conditions as TCNE.
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- 23. Preparation of complex a_1 : To a solution of TCNE (5.7 mg, 45 µmol) in toluene (1 mL) was slowly added a solution of 1 (12.4 mg, 23 μ mol) in toluene (1 mL) at -20 °C under argon, and then was stirred for 2 h. After addition of hexane (8 mL) to the reaction solution, the mixture was allowed to stand at -20 °C for 24 h under argon, giving a precipitation of a dark red solid, which was centrifuged at 2.5 krpm for 1 min. The crude product \mathbf{a}_1 thus obtained was carefully washed with hexane and dried well in vacuum desiccator to provide stable a₁ as a dark red powder (3.2 mg, 3.95 µmol, 18% yield). Complex a_1 : mp > 157 °C [decomp., determined by thermal analysis (TGA and DTA)]; UV-vis λ_{max} (benzene) nm, 295, 340, 357, 372 and 515; IR ν_{max} (KBr) cm⁻¹, 2206 (-C \equiv N); FAB-MS (positive) (3-nitrobenzyl alcohol matrix), m/z 555 $([M-2(TCNE)+H]^+)$; FAB-MS (negative) (3-nitrobenzyl alcohol matrix), *m/z* 128 (TCNE⁻); exact FAB-MS (positive) (3-nitrobenzyl alcohol matrix), found: m/z 555.3244; calcd for $C_{40}H_{43}O_2$: $[M - 2(TCNE) + H]^+$, *m/z* 555.3263.
- 24. Preparation of complex a2: To a solution of TCNE (11.4 mg, 89 µmol) in toluene (2 mL) was slowly added a solution of 2 (26 mg, 44 μ mol) in toluene (2 mL) at -20 °C under argon, and then was stirred for 2 h. After addition of hexane (16 mL) to the reaction solution, the mixture was allowed to stand at -20 °C for 24 h under argon, giving a precipitation of a dark red solid, which was centrifuged at 2.5 krpm for 1 min. The crude product \mathbf{a}_2 thus obtained was carefully washed with hexane and dried well in vacuum desiccator to provide stable \mathbf{a}_2 as a dark red powder (8.3 mg, 9.8 µmol, 23% yield). Complex a_2 : mp > 120 °C [decomp., determined by thermal analysis (TGA and DTA)]; UV-vis λ_{max} (benzene) nm, 297, 342, 358, 375 and 508; IR ν_{max} (KBr) cm⁻¹, 2206 (–C \equiv N); FAB-MS (positive) (3-nitrobenzyl alcohol matrix), m/z 843 $([M+H]^+)$ and 715 $([M-TCNE+H]^+)$; FAB-MS (negative) (3-nitrobenzyl alcohol matrix), m/z 128 (TCNE⁻); exact FAB-MS (positive) (3-nitrobenzyl alcohol matrix), found: m/z 715.2965; calcd for $C_{46}H_{43}N_4S_2$: $[M-TCNE+H]^+$, m/z715.2929.
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Facile synthesis of 3,3-di(heteroaryl)indolin-2-one derivatives catalyzed by ceric ammonium nitrate (CAN) under ultrasound irradiation

Shun-Yi Wang and Shun-Jun Ji*

The Key Lab. of Organic Synthesis of Jiangsu Province, College of Chemistry and Chemical Engineering, Suzhou (Soochow) University, Jiangsu 215123, China

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Abstract—Ceric ammonium nitrate efficiently catalyzes the reaction of isatin with indoles under sonic waves to afford symmetrical 3,3-di(indolyl)indolin-2-ones in excellent yields, as well as the reaction of 3-hydroxy-3-indolylindolin-2-ones with indoles, pyrrole to afford the corresponding adducts in excellent yields, which provides an efficient route to the synthesis of symmetrical and unsymmetrical 3,3-di(indolyl)indolin-2-one derivatives, respectively.

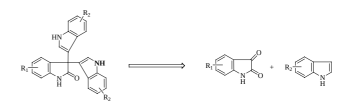
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1. Introduction

Indole fragment is featured widely in a wide variety of pharmacologically and biologically active compounds.¹ Oxindole derivatives are known to possess a variety of biological activity.² The 3,3-diaryloxindoles have been shown to possess mechanism-specific antiproliferative, antibacterial, antiprotozoal, and antiinflammatory activities.³ These compounds have also been used as laxatives⁴ and lead compounds for Ca²⁺-depletion-mediated inhibition of translation initiation.⁵ The 3,3-di(indolyl)indolin-2-ones can be formed by the reaction of isatin and indoles in acid conditions for long reaction times or promoted by KAl(SO₄)₂ under microwave conditions (Scheme 1).⁶ Few methods have been developed for the synthesis of this class of compounds.^{4,7} Especially, 3,3-diheteroaryloxindoles have not been widely explored. And the use of Lewis acid as a catalyst in the synthesis of 3,3-di(indolyl)oxindoles under mild conditions has not been reported.

In recent years, ceric ammonium nitrate (CAN) has been attracted much attention as an inexpensive and easily available catalyst for effecting various organic reactions.⁸ The reaction of indoles with carbonyl catalyzed by CAN afford the symmetrical bisindolymethane derivatives, which has been reported recently.⁹ However, the reaction must be

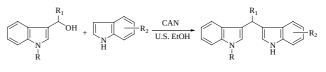
Keywords: 3,3-Di(heteroaryl)indolin-2-one; CAN; Indole; Isatin.



Scheme 1.

performed using the toxic CH_3CN as the solvent under the protection of N_2 atmosphere and was only limited to the synthesis of symmetrical BIAs.

More recently, we described an ultrasound-accelerated reaction of indoles with (1H-indol-3-yl)alkylmethanol using a catalytic amount of CAN, which provided an efficient route to the synthesis of unsymmetrical BIAs (Scheme 2).¹⁰ There was no previous report, which had indicated that two different indole or pyrrole residues could be so incorporated onto an indolin-2-ones derivative.



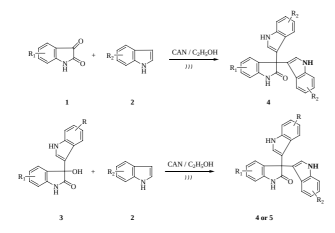
Scheme 2.

As a continue of our work on the synthesis of indole derivatives 10-12 we describe an ultrasound-accelerated reaction of isatin 1 with indoles 2 or 3-hydroxy-3-

^{*} Corresponding author. Tel.: +86 512 65880086; fax: +86 512 65880307; e-mail: shunjun@suda.edu.cn

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indolylindolin-2-ones **3** with indoles **2** using a catalytic amount of CAN, which provide an efficient route to the synthesis of symmetrical and unsymmetrical 3,3-bis(indolyl)oxindole derivatives, respectively (Scheme 3).



Scheme 3.

2. Results and discussion

In our initial research, we carried out the reaction of isatin **1a** with indole **2a** in the presence of CAN at room temperature using different solvents. The results were listed in Table 1. The reaction of **1a** with **2a** in the presence of CAN (10 mol%) and anhydrous C_2H_5OH (2 ml) proceeded smoothly at room temperature, giving 3,3-di(indolyl)indo-

Table 1. The solvent effect of the reaction between isatin 1a and indole 2a^a

Entry	Solvent	Time (h)	Yield (%) ^b
1	Anhydrous C ₂ H ₅ OH	3	95
2		10 ^c	90 ^c
3	Anhydrous CH3OH	3	96
4	CH_2Cl_2	3	80
5	CH ₃ CN	3	75

^a All reactions were carried out using catalytic amount of CAN (10 mol%) at room temperature.

^b Isolated yields.

^c The reaction was carried out under stir condition at room temperature.

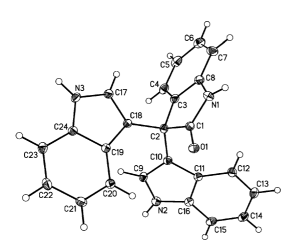
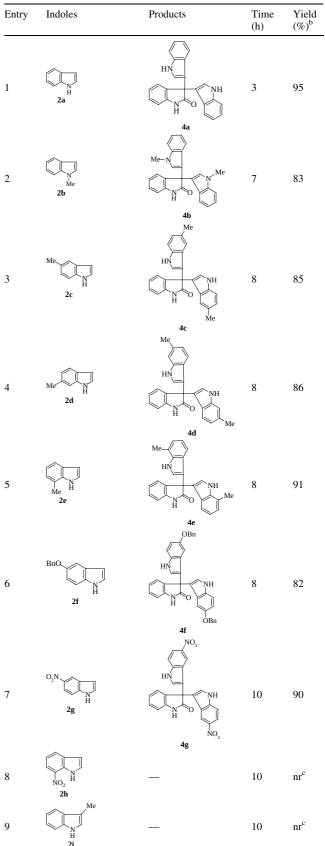


Figure 1. The crystal structure of 4a.

Table 2. The reaction of isatin with indoles catalyzed by CAN under ultrasound irradiation condition $^{\rm a}$



^a All reactions were carried out under sonic conditions.

^b Isolation yields.

^c nr=no reaction was detected.

lin-2-one **4a** in 95% yield (Table 1, entry 1) under sonic waves for 3 h, while 90% yield was found by stiring for 10 h (Table 1, entry 2). Compound **4a** was additionally confirmed by X-ray crystal structure analysis (Fig. 1).¹³ It was found that ultrasound could enhance the reaction rates. This reaction can also work well in anhydrous CH₃OH (Table 1, entry 3). However, the reaction did not progress well in CH₂Cl₂, CH₃CN during 3 h (Table 1, entries 4–5). Considering the toxicity of the methanol, study was continued using CAN/C₂H₅OH system.

CAN was found to be an efficient catalyst in the view of handling, temperature, yields and reaction times as indicated in comparison to the reported methods. This was because of the mild Lewis acidity of CAN and a small quantity of the solvent needed. As shown in Table 2, this method worked with a wide variety of substrates. In most cases, the reaction proceeded smoothly to produce the corresponding 3-(3-oxoalkyl)indole **3** in good yield. The treatment of **1a** with indole **2e** afforded **4e** in 91% yield under identical condition (Table 2, entry 5). The substituents of the indole ring showed some effect on this conversion. When unactivated indoles such as **2h**, **2i** were used, there were no reaction under the same reaction conditions even for 10 h (Table 2, entries 8–9). Indole-2,3-diones **1(b–d)** could also react well with indoles **2a**, **2b**, respectively, which afforded the corresponding symmetrical 3,3-di(indo-lyl)indolin-2-ones **4(i–m)** in good to excellent yields. The results were summarized in Table 3.

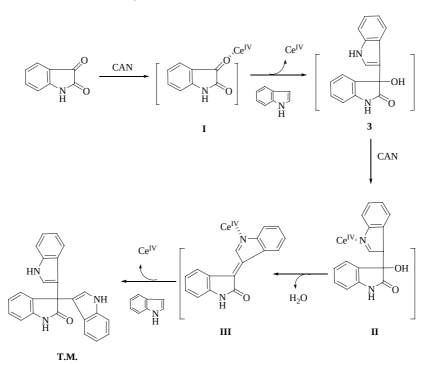
The reaction was probably proceeded through the activation of carbonyl group I as well as the indole moiety by CAN as shown in Scheme 4. 3-Hydroxy-3-(1H-indol-3-yl)indolin-2-one **3** may be formed in situ as a key intermediate, which can not be obtained in this CAN-methanol system. In the following step, the N–H bond of **3** was activated by CAN to

Table 3. The reaction of 1b-1c with indoles catalyzed by CAN under ultrasound irradiation condition^a

Entry	Isatin	Indoles	Products	Time (h)	Yield (%) ^b
1		2a		2	85
2		2ь		2	80
3	$ \begin{array}{c} Br \\ $	2a		3	90
4		2b		3	91
5	$Br \xrightarrow{H} O$	2a		3	88
6		2b	Me-N Br N N Me	3	87

^a All reactions were carried out under sonic conditions.

^b Isolation yields.



Scheme 4.

give intermediate II, which lost of H_2O to afford III. The indole or pyrrole attacked III to give the TM and Ce^{IV} , which could be recycled to catalyze the reaction.

In order to prove the mechanism, intermediates 3-hydroxy-3-(1*H*-indol-3-yl)indolin-2-one **3(a–f)** were synthesized according to the reported methods.¹⁴ We were pleased to find that the reaction of 3-hydroxy-3-(1*H*-indol-3-yl)indolin-2-one **3a** with indole **2a** in the presence of CAN (10 mol%) and anhydrous C₂H₅OH (2 ml) proceeded smoothly giving the 3,3-di(1*H*-indol-3-yl)indolin-2-one **(4a)** in 85% yield (Table 4, entry 1). This reaction can also be performed well by stirring to afford **4a** in 82% yield at room temperature.

Encouraged by this result, a number of other indoles were applied to this reaction (Scheme 5). The results were listed in Table 4. To our delight, **3a** smoothly reacted with substituted indole (**2b–e**) in the presence of CAN under sonic waves to afford the unsymmetrical 3,3-di(indolyl)indolin-2-ones (**5a–d**) in high yields (Table 4, entries 2–5) as expected. The reactions of **3a** with **2b** and **1b** with **2a** afforded the same product **5a** in 95, 84% yields under identical conditions, respectively.

It is well known that indole undergoes electrophilic substitution preferentially at their β -positions, which hold true for indoles but not for 3-substituted indoles such as 3-methyl-1*H*-indole.¹⁵ The reaction of **3(a–f)** with 3-methyl-lindole **2i** proceeded smoothly at 2-position giving the 3-indolyl-3-(3-methyl-1*H*-indol-2-yl)indolin-2-ones **6(a–f)** in good to excellent yields under identical conditions (Table 5, entries 1–6).

To further demonstrate the efficiency and scope of the reaction, pyrrole **7** was investigated. The reaction occurred

at 2-position of pyrrole 7 predominantly while 3-substituted adducts could hardly be detected by analysis of the reaction mixture by ¹H NMR. A variety of (3a-f) and pyrrole 7 were examined to generate the desired products (8a-f) under sonic conditions. The results were summarized in Table 6. It should be noted that pyrrole 7 (3 mmol) and 3 (1 mmol) were used in this reaction.

In addition, the structures of the product 8c was ascertained by spectroscopic methods, and the final proofs for the assigned structures were obtained by single-crystal X-ray analysis, respectively (Fig. 2).¹³

3. Conclusion

In conclusion, we have developed a simple, convenient and efficient protocol for **4**, **5**, **6**, **8** using catalytic amount of CAN under sonic conditions. At the same time, we proposed a plausible mechanism. In addition, using commercially available, easy to handle, inexpensive indole, pyrrole and the very mild conditions make this process a simple and convenient approach to obtain these compounds. It also makes it possible to design and synthesize of appropriately substituted symmetrical 3,3-di(indolyl)indolin-2-one derivatives as well as 3-(1*H*-pyrrol-2-yl)-3(1*H*-indol-3-yl)indolin-2-one derivatives, which are in progress in our laboratories.

4. Experimental

4.1. General

Melting points were recorded on an Electrothermal digital melting point apparatus and uncorrected. ¹H NMR (400 MHz)

Table 4. The reaction of 3a-f with indoles 2 catalyzed by CAN^a

Entry	1	Indole	Product	Time (h)	Yield (%) ^b	
1	3a	2a		4a	1 6 ^c	85 82
2		2b		- 5a	5	95
3	HN HN HN HN H OH H Ja	2c	HN HN HN HN HO HO Me	5b	3	80
4		2d	HN HN HN H	5c Me	5	95
5		2e	HN HN HN HO H	5d ^{Me}	5	84
6		2a		5a	1.5	84
7	HN HN HN OH H G G G	2a		5b	1	90
8		2a		5d	3	90
9	HN HN OH Me 3e	2a		4b	1	92
10	HN HN HN OH H M	2a		4c	1	92

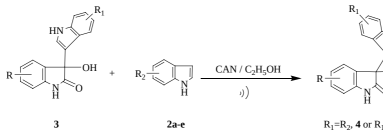
^a All reactions were carried out under sonication conditions.

^b Isolated yields.

^c The reaction was carried out under stir condition at room temperature.

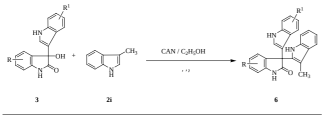
and ¹³C NMR (100 MHz) spectra were recorded on a Varian Mercury MHz spectrometer in CDCl₃ or DMSO- d_6 . IR Spectra were obtained on a Nicolet FT-IR500 spectrophotometer using KBr pellets. Elemental analysis was performed by a Carlo-Erba EA1110 CNNO-S analyzer. High-resolution

mass spectra were obtained using GCT-TOF instrument. Ultrasound irradiation was performed in a KQ-250E ultrasonic cleaner with a frequency of 40 KHz and a normal power of 250 W. The reaction flask was located in the water bath of the ultrasonic cleaner, and the temperature of the water



Scheme 5.

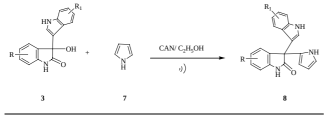
Table 5. The reaction of 3(a-f) with 3-methyl-1H-indole 2i^a



Entry	Sub.	Product	Time (h)	Yield (%) ^b
1	3a	6a	5	82
2	3b	6b	4	86
3	3c	6c	1	77
4	3d	6d	3	84
5	3e	6e	1	85
6	3f	6f	2	60

^a All reactions were carried out under sonication conditions. ^b Isolated yields.

Table 6. The reaction of 3(a-f) with pyrrole 7^a



Entry	Sub.	Product	Time (h)	Yield (%) ^b
1	3a	8a	1	88
2	3b	8b	1	85
3	3c	8c	1	85
4	3d	8d	1	83
5	3e	8e	1	82
6	3f	8f	1	80

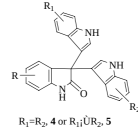
 $^{\rm a}$ Compound 3 (1 mmol) and pyrrole 7 (3 mmol) were used, and all reactions were performed under sonication conditions.

^b Isolated yields.

bath was controlled by 21 circulative water. 3-Hydroxy-3-(1Hindol-3-yl)indolin-2-one derivatives $3(a-f)^{14}$ were prepared according to the literature methods.

4.2. Typical experimental procedure

A mixture of 1d (0.225 g, 1 mmol), indole 2a (0.117 g, 1 mmol), CAN (0.056 g, 0.1 mmol) and anhydrous C₂H₅OH (2 ml) was irradiated by ultrasound in a vessel until the disappearance of the starting isatin (1 h, monitored by TLC). After standing 1 h, the reaction mixture was washed



by cool water $(3 \times 15 \text{ mL})$, warm water $(2 \times 10 \text{ mL})$ and cool ethanol (3×0.5 mL). The crude mixture was purified by flash chromatography to afford the pure product 4m (0.39 g, yield: 88%).

4.2.1. 3,3-Di(1H-indol-3-yl)indolin-2-one, 4a. White solid; mp: > 300 °C; IR (KBr): v 743, 1099, 1467, 1617, 1690, 3056, 3123, 3399 (NH), 3440 (NH) cm^{-1} ; ¹H NMR (400 MHz, CDCl₃): δ 6.93–6.99 (m, 6H), 7.12–7.24 (m, 3H), 7.34–7.40 (m, 5H), 7.74 (br, s, 1H, NH), 8.10 (br, s, 2H, NH); HRMS [Found: m/z 363.1359 (M⁺), calcd for C₂₄H₁₇N₃O: M, 363.1372].

4.2.2. 3,3-Bis(1-methyl-1*H*-indol-3-yl)indolin-2-one, 4b. White solid; mp: >300 °C (lit., 6 330–332 °C); IR (KBr): ν 1209, 1243, 1455, 1616, 1693, 2878, 2939, 3057, 3319 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.71 (s, 6H, CH₃), 6.86 (s, 2H), 6.93–7.42 (m, 12H), 7.81 (br, s, 1H, NH); HRMS [Found: m/z 391.1686 (M⁺), calcd for C₂₆H₂₁N₃O: M, 391.1685].

4.2.3. 3,3-Bis(5-methyl-1*H*-indol-3-yl)indolin-2-one, 4c. White solid; mp: >300 °C; IR (KBr): ν 751, 1107, 1235, 1466, 1614, 1712, 2853, 2914, 3386 (NH), 3413 (NH) cm⁻ ¹: ¹H NMR (400 MHz, CDCl₃): δ 2.40 (s, 6H, CH₃), 6.78 (d, 2H, J=7.6 Hz), 6.93-7.00 (m, 5H), 7.14-7.40 (m, 5H), 7.54 (br, s, 1H, NH), 7.94 (br, s, 2H, NH); HRMS [Found: *m*/*z* 391.1667 (M⁺), calcd for $C_{26}H_{21}N_3O$: M, 391.1685].

4.2.4. 3,3-Bis(6-methyl-1*H*-indol-3-yl)indolin-2-one, 4d. White solid; mp: 297–298 °C; IR (KBr): v 754, 1102, 1235, 1471, 1620, 1698, 2853, 2913, 3172, 3319 (NH), 3433 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.30 (s, 6H, CH₃), 6.94–7.02 (m, 7H), 7.17–7.35 (m, 5H), 7.49 (br, s, 1H, NH), 7.97 (br, s, 2H, NH); HRMS [Found: m/z 391.1681 (M^+) , calcd for C₂₆H₂₁N₃O: M, 391.1685].

4.2.5. 3,3-Bis(7-methyl-1*H*-indol-3-yl)indolin-2-one, 4e. White solid; mp: 196–198 °C; IR (KBr): v 747, 1101, 1343, 1470, 1089, 1455, 1487, 1617, 1699, 3054, 3410 (NH) cm⁻ ¹; ¹H NMR (400 MHz, CDCl₃): δ 2.46 (s, 6H, CH₃), 6.85– 7.00 (m, 8H), 7.18–7.24 (m, 3H), 7.40 (d, 1H, J=7.2 Hz), 7.73 (br, s, 1H, NH), 8.00 (br, s, 2H, NH); HRMS [Found: m/z 391.1671 (M⁺), calcd for C₂₆H₂₁N₃O: M, 391.1685].

4.2.6. 3,3-Bis(5-(benzyloxy)-1H-indol-3-yl)indolin-2-one, **4f.** White solid; mp: 198–200 °C; IR (KBr): v 743, 1101, 1383, 1469, 1481, 1699, 2863, 2914, 3027, 3381 (NH), 3421 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 6.85–7.00 (m, 8H), 7.15–7.49 (m, 19H), 7.91 (br, s, 2H, NH); HRMS

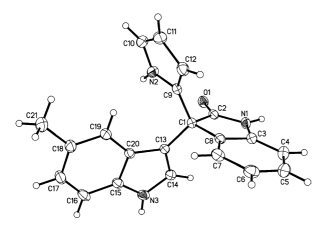


Figure 2. The crystal structure of 8c.

[Found: m/z 575.2178 (M⁺), calcd for C₃₈H₂₉N₃O₃: M, 575.2209].

4.2.7. 3,3-Bis(5-nitro-1*H***-indol-3-yl)indolin-2-one, 4g.** Yellow solid; mp: > 300 °C; IR (KBr): ν 739, 1090, 1250, 1260, 1470, 1620, 1709, 3346 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-*d*₆): δ 7.02–7.09 (m, 2H), 7.20–7.32 (m, 4H), 7.57 (d, 2H, *J*=8.4 Hz), 7.96 (d, 2H, *J*=8.4 Hz), 8.23 (s, 2H), 10.96 (br, s, 1H, NH), 11.81 (br, s, 2H, NH); HRMS [Found: *m/z* 453.1052 (M⁺), calcd for C₂₄H₁₅N₅O₅: M, 453.1073].

4.2.8. 3,3-Di(1*H*-indol-3-yl)-1-methylindolin-2-one, **4**h. White solid; mp: > 300 °C (lit., 310 °C); IR (KBr): ν 742, 1091, 1351, 1610, 1668, 2929, 3051, 3116, 3357 (NH), 3440 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.35 (s, 3H, CH₃), 6.91–7.03 (m, 6H), 7.12 (t, 3H, *J*=7.2 Hz), 7.32 (t, 4H, *J*=6.8 Hz), 7.41 (d, 1H, *J*=7.2 Hz), 8.04 (br, s, 2H, NH); HRMS [Found: *m*/*z* 377.1527 (M⁺), calcd for C₂₅H₁₉N₃O: M, 377.1528].

4.2.9. 1-Methyl-3,3-bis(1-methyl-1*H***-indol-3-yl)indolin-2-one, 4i.** White solid; mp: 233–235 °C (lit.,⁶ 232– 234 °C); IR (KBr): ν 746, 1087, 1469, 1606, 1722, 2924, 3049 cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.35 (s, 3H, CH₃), 3.70 (s, 6H, CH₃), 6.84 (d, 2H, *J*=8.0 Hz), 6.91–7.18 (m, 7H), 7.26–7.46 (m, 5H); HRMS [Found: *m/z* 405.1826 (M⁺), calcd for C₂₇H₂₃N₃O: M, 405.1841].

4.2.10. 4-Bromo-3,3-di(1*H*-indol-3-yl)indolin-2-one, **4**j. White solid; mp: > 300 °C; IR (KBr): ν 733, 1106, 1615, 1702, 3047, 3330 (NH), 3378 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 6.95 (d, 1H, *J*=7.0 Hz), 6.99–7.04 (m, 4H), 7.12–7.19 (m, 4H), 7.38 (d, 2H, *J*=8.0 Hz), 7.52 (d, 2H, *J*=9.2 Hz), 7.98 (br, s, 1H, NH), 8.12 (br, s, 2H, NH); HRMS [Found: *m*/*z* 443.0438 (M⁺), calcd for C₂₄H₁₆⁸BrN₃O: M, 443.0456].

4.2.11. 4-Bromo-3,3-bis(1-methyl-1*H***-indol-3-yl)indolin-2-one, 4k.** White solid; mp: 281–283 °C; IR (KBr): ν 734, 1135, 1429, 1612, 1704, 2934, 3054, 3104, 3351 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.74 (s, 6H, CH₃), 6.92 (s, 1H), 6.96–7.32 (m, 10H), 7.47 (d, 2H, *J*=8.0 Hz), 7.61 (br, s, 1H, NH); HRMS [Found: *m*/*z* 469.0768 (M⁺), calcd for C₂₆H⁷⁹₂₀BrN₃O: M, 469.0790]. **4.2.12. 6-Bromo-3,3-di**(1*H*-indol-3-yl)indolin-2-one, **4**I. White solid; mp: > 300 °C; IR (KBr): ν 743, 1114, 1451, 1607, 1709, 3029, 3110, 3400 (NH), 3442 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO- d_6): δ 6.81 (t, 2H, J=7.6 Hz), 6.85 (d, 2H, J=2.4 Hz), 7.02 (t, 2H, J=7.6 Hz), 7.10–7.16 (m, 3H), 7.21 (d, 2H, J=8.0 Hz), 7.35 (d, 2H, J=8.0 Hz), 10.74 (br, s, 1H, NH), 11.00 (br, s, 2H, NH); ¹³C NMR (100.57 MHz, DMSO- d_6): 52.3, 111.7, 112.4, 113.6, 118.4, 120.3, 120.6, 121.0, 124.1, 124.4, 125.5, 126.7, 133.8, 136.9 143.0, 178.5; HRMS [Found: m/z 443.0450 (M⁺), calcd for C₂₄H⁸₁₆BrN₃O: M, 443.0456].

4.2.13. 6-Bromo-3,3-bis(1-methyl-1*H***-indol-3-yl)indolin-2-one, 4m.** White solid; mp: 248–250 °C; IR (KBr): ν 734, 1337, 1612, 1690, 3049, 3169, 3359 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 3.70 (s, 6H, CH₃), 6.84 (d, 2H, *J*= 8.0 Hz), 6.95–7.34 (m, 11H), 8.39 (br, s, 1H, NH); HRMS [Found: *m/z* 471.0746 (M⁺), calcd for C₂₆H⁸¹₂₀BrN₃O: M, 471.0769].

A mixture of **3a** (0.225 g, 1 mmol), indole **2b** (0.117 g, 1 mmol), CAN (0.056 g, 0.1 mmol) and C_2H_5OH (2 ml) was irradiated by ultrasound in a vessel until the disappearance of the starting isatin (1 h, monitored by TLC). After standing 1 h, the reaction mixture was washed by cool water (3×15 mL), warm water (2×10 mL) and cool ethanol (3×0.5 mL). The crude mixture was purified by flash chromatography to afford the pure product **5a** (0.388 g, yield: 95%).

4.2.14. 3-(1*H***-Indol-3-yl)-3-(1-methyl-1***H***-indol-3-yl)indolin-2-one, 5a. White solid; mp: 298–300 °C; IR (KBr): \nu 743, 1099, 1467, 1617, 1690, 3056, 3123, 3399 (NH), 3440 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-***d***₆): \delta 1.99 (s, 3H), 6.77–6.86 (m, 4H), 6.90–7.03 (m, 3H), 7.06 (s, 1H), 7.17–7.25 (m, 4H), 7.35 (d, 1H,** *J***=8.4 Hz), 10.60 (s, 1H), 10.83 (br, s, 1H, NH), 10.94 (br, s, 1H, NH); HRMS [Found:** *m***/***z* **377.1523 (M⁺), calcd for C₂₅H₁₉N₃O: M, 377.1528].**

4.2.15. 3-(1*H***-Indol-3-yl)-3-(5-methyl-1***H***-indol-3-yl)indolin-2-one, 5b.** White solid; mp: 281–283 °C; IR (KBr): ν 751, 1099, 1468, 1615, 1711, 3047, 2842, 2909, 3123, 3322 (NH), 3392 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.31 (s, 3H), 6.94–7.38 (m, 13H), 7.55 (br, s, 1H, NH), 7.98 (br, s, 1H, NH), 8.07 (br, s, 1H, NH); HRMS [Found: *m*/*z* 377.1412 (M⁺), calcd for C₂₅H₁₉N₃O: M, 377.1528].

4.2.16. 3-(1*H***-Indol-3-yl)-3-(6-methyl-1***H***-indol-3-yl)indolin-2-one, 5c.** White solid; mp: 210–212 °C; IR (KBr): ν 743, 1100, 1470, 1616, 1712, 2858, 2914, 3055, 3405 (NH), 3440 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 2.41 (s, 3H), 6.79 (d, 1H, *J*=7.6 Hz), 6.92–7.00 (m, 5H), 7.12–7.25 (m, 4H), 7.34–7.40 (m, 3H), 7.68 (br, s, 1H, NH), 7.94 (br, s, 1H, NH), 8.07 (br, s, 1H, NH); HRMS [Found: *m*/*z* 377.1523 (M⁺), calcd for C₂₅H₁₉N₃O: M, 377.1528].

4.2.17. 3-(1*H***-Indol-3-yl)-3-(7-methyl-1***H***-indol-3-yl)indolin-2-one, 5d. Solid; mp: 240–242 °C; IR (KBr): \nu 744, 1101, 1470, 1616, 1712, 2858, 2915, 3053, 3123, 3409 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): \delta 2.47 (s, 3H), 6.87 (t, 1H,** *J***=7.6 Hz), 6.94–7.40 (m, 12H), 7.70 (br, s, 1H, NH), 7.99 (br, s, 1H, NH), 8.07 (br, s, 1H, NH); HRMS**

[Found: m/z 377.1518 (M⁺), calcd for C₂₅H₁₉N₃O: M, 377.1528].

4.2.18. 3-(1*H*-Indol-3-yl)-3-(3-methyl-1*H*-indol-2-yl)indolin-2-one, 6a. Colorless needles; mp: 196–198 °C; IR (KBr): ν 738, 1107, 1471, 1615, 1724, 2852, 2970, 3052, 3365 (NH), 3441 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 1.95 (s, 3H), 6.97–7.29 (m, 9H), 7.38 (t, 2H, *J*=7.6 Hz), 7.50 (t, 2H, *J*=8.4 Hz), 7.95 (br, s, 1H, NH), 8.12 (br, s, 1H, NH), 8.18 (br, s, 1H, NH); HRMS [Found: *m/z* 377.1494 (M⁺), calcd for C₂₅H₁₉N₃O: M, 377.1528].

4.2.19. 3-(3-Methyl-1*H***-indol-2-yl)-3-(1-methyl-1***H***-indol-3-yl)indolin-2-one, 6b.** Colorless needles; mp: 153–155 °C; IR (KBr): ν 742, 1470, 1615, 1714, 2852, 2914, 3047, 3350 (NH), 3395 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): δ 1.94 (s, 3H), 2.32 (s, 3H), 6.90 (d, 1H, *J*= 2.4 Hz), 6.94 (d, 1H, *J*=7.6 Hz), 7.02 (t, 2H, *J*=7.6 Hz), 7.07–7.13 (m, 2H), 7.18 (d, 1H, *J*=7.2 Hz), 7.24 (d, 2H, *J*= 8.0 Hz), 7.32 (d, 2H, *J*=7.6 Hz), 7.51 (d, 1H, *J*=7.2 Hz), 8.09 (s, 2H), 8.16 (br, s, 1H, NH); ¹³C NMR (100.57 MHz, CDCl₃):9.2, 22.1, 54.2, 109.0, 110.0, 111.4, 111.6, 113.6, 118.8, 119.5, 121.6, 122.0, 123.3, 124.9, 126.1, 126.6, 128.9, 130.0, 130.4, 131.2, 133.4, 135.1, 135.9, 140.4, 178.4 (C=O); HRMS [Found: *m*/z 391.1673 (M⁺), calcd for C₂₆H₂₁N₃O: M, 391.1685].

4.2.20. 3-(3-Methyl-1*H***-indol-2-yl)-3-(5-methyl-1***H***-indol-3-yl)indolin-2-one, 6c. Colorless needles; mp: 191–193 °C; IR (KBr): \nu 743, 1099, 1470, 1616, 1715, 2852, 3914, 3057, 3407 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): \delta 1.94 (s, 3H), 2.32 (s, 3H), 6.90 (d, 1H,** *J***= 2.4 Hz), 6.94 (d, 1H,** *J***=7.6 Hz), 7.02 (t, 2H,** *J***=7.6 Hz), 7.07–7.13 (m, 2H), 7.18 (d, 1H,** *J***=7.2 Hz), 7.24–7.29 (m, 3H), 7.33 (d, 1H,** *J***=6.4 Hz), 7.51 (d, 1H,** *J***=7.6 Hz), 7.85 (br, s, 1H, NH), 8.10 (br, s, 1H, NH), 8.14 (br, s, 1H, NH); HRMS [Found:** *m/z* **391.1681 (M⁺), calcd for C₂₆H₂₁N₃O: M, 391.1685].**

4.2.21. 3-(3-Methyl-1*H***-indol-2-yl)-3-(7-methyl-1***H***-indol-3-yl)indolin-2-one, 6d. Colorless needles; mp: 232–234 °C; IR (KBr): \nu 737, 1470, 1614, 1716, 2858, 2975, 3057, 3123, 3289 (NH), 3359 (NH), 3439 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): \delta 1.95 (s, 3H), 2.50 (s, 3H), 6.92–7.19 (m, 6H), 7.20 (d, 1H,** *J***=8.4 Hz), 7.39 (d, 1H,** *J***=7.2 Hz), 7.51 d, 1H,** *J***=6.8 Hz), 7.18 (d, 1H,** *J***=7.2 Hz), 7.24 (d, 1H,** *J***=8.0 Hz), 7.32 (d, 1H,** *J***=7.6 Hz), 7.71 (br, s, 1H, NH), 8.10 (br, s, 1H, NH), 8.12 (br, s, 1H, NH); HRMS [Found:** *m***/***z* **391.1696 (M⁺), calcd for C₂₆H₂₁N₃O: M, 391.1685].**

4.2.22. 3-(1*H***-Indol-3-yl)-1-methyl-3-(3-methyl-1***H***-indol-2-yl)indolin-2-one, 6e. Colorless needles; mp: 161–163 °C; IR (KBr): \nu 743, 1090, 1469, 1609, 1703, 2858, 2921, 3054, 3324 (NH), 3407 (NH) cm⁻¹; ¹H NMR (400 MHz, CDCl₃): \delta 1.90 (s, 3H), 3.33 (s, 3H), 6.93 (d, 1H, J=2.0 Hz), 6.96–7.21 (m, 7H), 7.34–7.40 (m, 3H), 7.48 (t, 2H, J=8.0 Hz), 8.10 (br, s, 1H, NH), 8.18 (br, s, 1H, NH); HRMS [Found: m/z 391.1670 (M⁺), calcd for C₂₆H₂₁N₃O: M, 391.1685].**

4.2.23. 6-Bromo-3-(1*H***-indol-3-yl)-3-(3-methyl-1***H***-indol-2-yl)indolin-2-one, 6f.** White solid; mp: 208–

210 °C; IR (KBr): ν 741, 1443, 1607, 1744, 3052, 3123, 3399 (NH), 3440 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSOd₆): δ 1.90 (s, 3H), 6.78–7.39 (m, 13H), 10.33 (br, s, 1H, NH), 10.81 (br, s, 1H, NH); HRMS [Found: *m*/*z* 457.0590 (M⁺), calcd for C₂₅H₁₈⁸BrN₃O: M, 457.0613].

4.2.24. 3-(**1***H*-**Pyrrol-2-yl**)-**3**(1*H*-**indol-3-yl**)**indolin-2-one, 8a.** White solid; mp: 175–177 °C; IR (KBr): ν 737, 759, 1106, 1469, 1614, 1708, 3326 (NH), 3430 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-*d*₆): δ 6.78 (t, 2H, *J*=8.0 Hz), 6.85 (s, 2H), 6.92 (t, 2H, *J*=8.0 Hz), 6.97–7.03 (m, 3H), 7.20 (d, 4H, *J*=8.4 Hz), 7.34 (d, 2H, *J*=8.4 Hz), 10.54 (br, s, 1H, NH), 10.92 (br, s, 2H, NH); HRMS [Found: *m/z* 313.1200 (M⁺), calcd for C₂₀H₁₅N₃O: M, 313.1215].

4.2.25. 3-(1-Methyl-1*H***-indol-3-yl)-3-(1***H***-pyrrol-2-yl)indolin-2-one, 8b.** White solid; mp: 266–268 °C (dec); IR (KBr): ν 743, 799, 1102, 1471, 1621, 1689, 2842, 2919, 3282 (NH), 3368 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-*d*₆): δ 2.17 (s, 3H, CH₃), 5.77 (s, 1H), 5.93 (d, 1H, *J*= 2.8 Hz), 6.62 (s, 1H), 6.69–6.72 (m, 2H), 6.83 (d, 1H, *J*= 8.4 Hz), 6.95 (t, 2H, *J*=7.6 Hz), 7.20–7.24 (m, 2H), 7.31 (d, 1H, *J*=8.0 Hz), 10.59 (br, s, 2H), 10.83 (br, s, 1H, NH); HRMS [Found: *m/z* 327.1363 (M⁺), calcd for C₂₁H₁₇N₃O: M, 327.1372].

4.2.26. 3-(**5**-Methyl-1*H*-Indol-3-yl)-3-(1*H*-pyrrol-2-yl)indolin-2-one, **8c.** White solid; mp: 253–255 °C (dec); IR (KBr): ν 732, 796, 1101, 1619, 1680, 2914, 3279 (NH), 3367 (NH), 3456 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO- d_6): δ 2.17 (s, 3H, CH₃), 5.78 (s, 1H), 5.93 (d, 1H, J= 2.4 Hz), 6.62 (s, 1H), 6.69–6.72 (m, 2H), 6.83 (d, 1H, J= 8.4 Hz), 6.95 (t, 2H, J=7.2 Hz), 7.20–7.22 (m, 2H), 7.31 (d, 1H, J=7.2 Hz), 10.60 (br, s, 2H), 10.84 (br, s, 1H, NH); HRMS [Found: m/z 327.1360 (M⁺), calcd for C₂₁H₁₇N₃O: M, 327.1372].

4.2.27. 3-(7-Methyl-1*H***-indol-3-yl)-3-(1***H***-pyrrol-2-yl)indolin-2-one, 8d. White solid; mp: 154–156 °C; IR (KBr): \nu 742, 1101, 1470, 1619, 1689, 2975, 3055, 3296 (NH), 3388 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-***d***₆): \delta 2.40 (s, 3H, CH₃), 5.78 (s, 1H), 5.93 (dd, 1H,** *J***=2.8, 2.4 Hz), 6.62 (s, 1H), 6.65–6.70 (m, 3H), 6.74 (d, 1H,** *J***=2.4 Hz), 6.80 (d, 1H,** *J***=6.0 Hz), 7.22 (t, 2H,** *J***=7.2 Hz), 7.31 (d, 1H,** *J***=7.2 Hz), 10.60 (br, s, 2H), 10.94 (br, s, 1H, NH); HRMS [Found:** *m/z* **327.1359 (M⁺), calcd for C₂₁H₁₇N₃O: M, 327.1372].**

4.2.28. 3-(1*H***-Indol-3-yl)-1-methyl-3-(1***H***-pyrrol-2-yl)indolin-2-one, 8e.** White solid; mp: 138–140 °C (dec); IR (KBr): ν 742, 1088, 1470, 1610, 1698, 3054, 3410 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-*d*₆): δ 3.25 (s, 3H, CH₃), 5.76 (s, 1H), 5.92 (s, 1H), 6.69 (s, 1H), 6.78–6.83 (m, 3H), 7.00–7.04 (m, 2H), 7.14 (d, 1H, *J*=7.6 Hz), 7.31–7.39 (m, 3H), 10.69 (br, s, 1H), 11.00 (br, s, 1H, NH); HRMS [Found: *m/z* 327.1358 (M⁺), calcd for C₂₁H₁₇N₃O: M, 327.1372].

4.2.29. 6-Bromo-3-(1*H***-indol-3-yl)-3-(1***H***-pyrrol-2-yl)indolin-2-one, 8f. White solid; mp: 184–186 °C (dec); IR (KBr): \nu 742, 1110, 1479, 1610, 1712, 3374 (NH) cm⁻¹; ¹H NMR (400 MHz, DMSO-***d***₆): \delta 5.70 (s, 1H), 5.94 (s, 1H), 6.70 (s, 1H), 6.80–6.89 (m, 3H), 7.01 (t, 2H,** *J***=7.4 Hz),**

7.11–7.15 (m, 2H), 7.24 (d, 1H, J=6.4 Hz), 7.31 (d, 1H, J= 8.4 Hz), 10.63 (br, s, 1H), 10.73 (br, s, 1H), 11.00 (br, s, 1H, NH); HRMS [Found: m/z 391.0320 (M⁺), calcd for C₂₀H₁₄⁷⁹BrN₃O: M, 391.0320].

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Coupling and fast decarboxylation of aryloxyl radicals of 4-hydroxycinnamic acids with formation of stable *p*-quinomethanes

Carmelo Daquino and Mario C. Foti*

Istituto di Chimica Biomolecolare del CNR - Sez. di Catania, Via del Santuario, 110, I-95028 Valverde (CT), Italy

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This paper is dedicated to Keith Usherwood Ingold.

Abstract—The reaction at room temperature of 3,5-di-*tert*-butyl- and 3,5-di-methoxy-4-hydroxycinnamic acids 1 and 2 with the dpph' radical in acetone or other *non-hydroxylic* polar solvents yields interesting dimeric *p*-quinomethanes 10–16 characterized by a broad and strong absorption in the visible region. Although the yields appear to be low to moderate (10–40%), this simple synthesis affords quinones not otherwise obtainable, which contain an unsaturated γ -lactone ring (14–16). The structures have been elucidated by interpretation of ESI-MS, FT-IR and NMR spectral data. In particular, FT-IR spectra in a KBr matrix demonstrate the quinone nature of these compounds because of the presence of strong absorption bands at 1604–1640 cm⁻¹ and allows excluding the presence of carboxylic acid groups in the molecules. Kinetic evidence and molecular structures suggest that the formation of these *p*-quinomethanes is best explained through an 8–8 C–C coupling of the aryloxyl radicals derived from 1 and 2 and a subsequent fast mono- or di-decarboxylation of the initial dimer by an S_E1-type mechanism. Further oxidation of the phenolic intermediates by dpph' yields the final quinones. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Phenols (ArOH) are effective antioxidants because of their ability to react with peroxyl radicals, RO₂, responsible for the autoxidation processes of organic materials.¹ Considerable information as to the antioxidant properties of phenols can be gained from the kinetic parameters and stoichiometry of reactions with the stable and commercially available nitrogen-centered radical, 2,2-diphenyl-1-picrylhydrazyl (dpph⁻), because these parameters are correlated to those of ArOH/RO₂.² This has justified an intense proliferation of studies on ArOH/dpph⁻ reactions.

The mechanism and rate of ArOH/dpph' reactions depend largely upon the nature of phenol (vide infra) and the solvent in which these reactions occur. Recently, Litwinienko and Ingold³ and, independently, Foti et al.⁴ have demonstrated that in *alcohols* these reactions essentially proceed via an electron transfer (ET) step from the phenoxide anion ArO⁻ to dpph' (Reactions 1 and 2). 4-Hydroxycinnamic acids (HCA) are demonstrated to be ideal models to disclose this mechanism since the presence of the carboxylic acid groups strongly influences the ionization of phenolic OH and modulate the contribution of Reaction 2 over the direct H-atom transfer (Reaction 3).⁴ In non-hydroxylic polar solvents, phenols are poorly ionized⁵ and the reaction occurs through the direct and *slow* (since ArOH are hydrogen bonded to the solvent molecules)⁶ transfer of the hydrogen atom from ArOH to dpph[•] (Reaction 3).

$$ArOH \rightleftharpoons ArO^{-} + H^{+} \tag{1}$$

$$ArO^{-} + dpph^{\bullet} \xrightarrow{H^{+}} ArO^{\bullet} + dpph - H$$
 (2)

$$ArOH + dpph^{\bullet} \xrightarrow{slow} ArO^{\bullet} + dpph - H$$
(3)

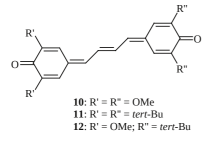
$$2ArO^{\bullet} \xrightarrow{\text{solvents}} \text{various products}$$
 (4)

Afterwards, additional work has revealed that the solvent effect plays another intriguing role in the products of the self-coupling reaction of aryloxyl radicals of HCA's, Reaction 4. For instance, in the case of sinapic acid 1/ dpph', Reaction 4 yielded in *methanol or ethanol* products that were totally transparent to the electromagnetic waves of the spectral region 400–800 nm. However, when this reaction was carried out in *acetone* successive scans of the UV–vis spectrum showed that the course of Reaction 4 was

Keywords: Sinapic acid; Cinnamic acids; dpph' radical; Decarboxylation; Free radical; *p*-Quinomethane; γ -Lactone.

^{*} Corresponding author. Tel.: +39 095 721 2136; fax: +39 095 721 2141; e-mail: mario.foti@icb.cnr.it

different since an intensely coloured product was formed. Chromatographic purification on silica gel of the reaction mixture gave a reasonable amount of this compound (yield 37%), which surprisingly proved to be nearly NMR-silent in many deuterated polar solvents (acetone, methanol, DMSO) (vide infra). In two solvents (CD_2Cl_2 and $CDCl_3$), however, we succeeded in obtaining excellent ¹H and ¹³C NMR spectra, which revealed that this compound was the ethylene bis(*p*-quinomethane) **10**.



The undoubted importance of this compound⁷ and the peculiar mechanism by which it is formed (vide infra) led us to explore the behaviour of other HCA's (**1–8**, see Chart 1) with dpph⁷ (or MnO₂) in various non-hydroxylic polar solvents. We found that HCA **2** gave the corresponding dimeric *p*-quinomethane **11** as well, whereas the others did not except for 3,5-di-bromo*p*-coumaric acid **3** and ferulic acid **6**, which gave traces of their corresponding *p*-quinomethanes. Using HCA's **1** and **2** in a 1:2 ratio (mol/mol) it was also possible to prepare the asymmetrical *p*-quinomethane **12**.

These reactions also afforded other oxidation products, which were isolated and characterized as another interesting group of stable quinones bearing an unsaturated γ -lactone ring (14–16).

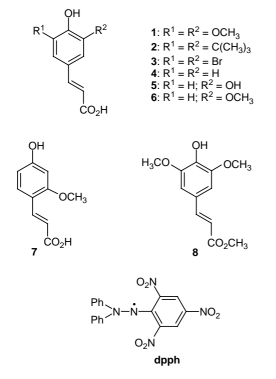
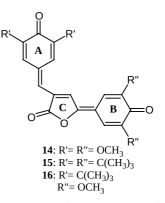


Chart 1. 4-Hydroxy cinnamic acids and relative derivatives employed in the present study.



Formation of C-C coupled dimers is not surprising in view of the fact that aryloxyl radicals undergo many of the reactions typical of oxygen and carbon radicals, such as C-C and/or C-O dimerization, isomerization, disproportionation, hydrogen abstraction and addition.8 In fact, the chemistry of aryloxyl radicals has been investigated intensely during 1960s and early 1970s.^{8,9} Various products of C–C and C–O coupling of the aryloxyl radicals derived from 1 and its methyl ester 8 have already been isolated^{10,11} among which thomasidioic $acid^{12}$ (a phenolic lignan) is one of the major products. However, to the best of our knowledge in no case has formation of decarboxylated dimers been reported in mild oxidative reactions of cinnamic acids. Therefore, the *p*-quinomethanes **10–16** formed under our experimental conditions are even more interesting because they show that a process of mono- or di-decarboxylation has occurred spontaneously at room temperature at some stage of the reaction sequence (vide infra).

We therefore, report herein on the synthesis and spectral characterization of all these new compounds and propose a mechanistic rationale for their formation compatible with our kinetic data.

2. Results and discussion

2.1. Isolation and structure determination

In Figure 1 is reported the time evolution at 25 °C of the UV– vis spectrum of an acetone solution of dpph (0.12 mM) in which sinapic acid 1 was added to a final concentration of 1.28 mM. For comparison, the evolution of this reaction in methanol is also shown in the Inset (Fig. 1). The spectra in acetone show the formation of a compound with broad absorption bands in the visible region (λ_{max} at 508 nm) and which is responsible for the purplish colour of the solution. The UV-vis spectrum of this compound, isolated from the reaction mixture (see Section 4), is shown in Figure 2a. The large values of the molar extinction coefficient and λ_{max} (see Table 1) suggested a highly conjugated structure for 10. The FT-IR spectrum in a KBr matrix also showed the presence of α , β -unsaturated carbonyl groups (1621.9, 1563.4 and 1548.5 cm^{-1} ¹³ and unexpectedly the absence of carboxylic acid groups. HPLC-ESI-MS spectrum of this compound in the positive ion-mode showed three peaks at m/z: 357, 379 (base peak) and 735. Since many compounds form adducts with adventitious alkali metals¹⁴ and some of the above peaks could be due to such adducts, we deliberately used water/ acetonitrile containing 0.1 mM LiCl as eluents in the HPLC-

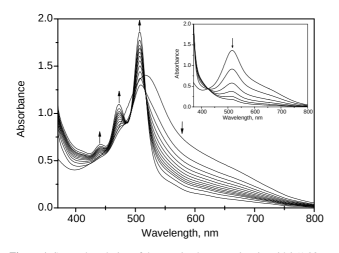


Figure 1. Spectral evolution of the reaction between sinapic acid 1 (1.28×10^{-3} M) and dpph' (1.20×10^{-4} M) in acetone at 25 °C. Spectra were recorded at: 0; 6; 11; 21; 31; 46; and 76 s. Inset: spectral evolution of sinapic acid 1 (1.28×10^{-3} M) + dpph' (1.25×10^{-4} M) in methanol at 25 °C at: 0; 1; 2; 3; 4; and 5 s.

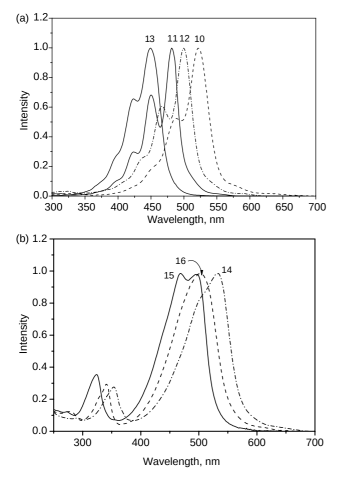


Figure 2. (a) Normalized UV–vis spectra of **10–13** obtained from the HPLC UV–DAD. The eluent composition during the readings was 20:80 water/acetonitrile for **10**; 15:85 water/acetonitrile for **12**; 100% acetonitrile for **11** and **13**. The values of λ_{max} and ε in CH₂Cl₂ are reported in Table 1. (b) Normalized UV–vis spectra of **14–16** obtained from the HPLC UV–DAD. The eluent composition during the readings was 15:85 water/acetonitrile for **14**; 10:90 water/acetonitrile for **16**; 100% acetonitrile for **15**. The values of λ_{max} and ε in CH₂Cl₂ are reported in Table 1.

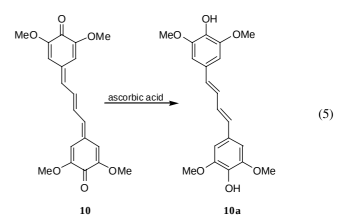
MS analyses. In the presence of LiCl, the spectrum essentially consisted of two peaks at *m*/*z*: 363 (base peak) and 719. Since

Table 1. FT-IR spectra in KBr and UV–vis spectra in CH_2Cl_2 of quinones 10–16

Quinone	$(\text{cm}^{\nu_{\text{CO}}})$	$\frac{\nu_{C=C}}{(cm^{-1})}$	$\lambda_{\rm max} \ ({\rm nm}) \ (\epsilon/10^3 \ {\rm M}^{-1} \ {\rm cm}^{-1})$
10	1621.9	1563.4 1548.5	519 (110); 484 (56); 453 (21)
11	1605.3	1570.9 1557.6	486 (110); 454 (72); 427 (31)
12	1629.9 1607.7	1567.7	502 (98); 468 (57); 440 (26)
13	1602.9		455 (54); 428 (36)
14	1782.6 1639.4 1621.6	1565.1	531 (40); 355 (13)
15	1781.2 1612.0	1572.6	507 (45); 477 (41); 324 (19)
16	1768.7 1640.7 1614.0	1566.2	503 (41); 344 (13)

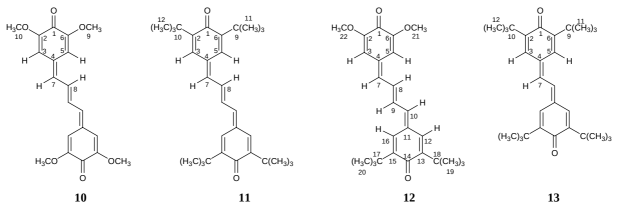
the difference, 379-363=735-719=16 was equal to the difference in the masses of Na⁺ and Li⁺, the above signals were given by the following pseudo-molecular ions: $[M+H]^+$ (*m*/*z* 357); $[M+Na]^+$ (*m*/*z* 379); $[M+Li]^+$ (*m*/*z* 363); $[2M+Na]^+$ (*m*/*z* 735) and $[2M+Li]^+$ (*m*/*z* 719) and hence, we determined that the molecular weight of **10** was 356 (C₂₀H₂₀O₆ 356.38).

When we tried to record the NMR spectra of 10 for the final structure elucidation we unexpectedly observed that the ¹H NMR spectrum showed little information in many solvents (acetone, methanol, DMSO), that is, this quinone was demonstrated to be NMR-silent.¹⁵ Addition of ascorbic acid into the NMR tube caused the bleaching of the solution and the appearance of the ¹H and ¹³C NMR spectra reported in Section 4. These spectra, the pseudo-molecular peak $[M-H]^-$ at m/z357, the appearance in the FT-IR spectrum in CH_2Cl_2 of a fairly sharp band at ca. 3529 cm^{-1} attributable to intra-molecular H-bonded OH's¹⁶ and finally the UV-spectrum, which was similar to that of 1,4-diphenyl-1,3-butadiene^{17–19} prompted us to assign structure 10a to this compound and, consequently, structure 10 to the oxidized form (Reaction 5). Contrary to the behaviour described above, we successively observed that in CD₂Cl₂ and CDCl₃ the *p*-quinomethane 10 gave both the ¹H and ¹³C NMR spectra (see Table 2), which provided a further evidence of its structure.



One interesting aspect of the ¹H NMR spectrum of **10** in CD_2Cl_2 was represented by the splitting of the signals corresponding to the methoxy groups (δ_H 3.80 and 3.86) and

Table 2. ¹H and ¹³C NMR spectra of quinones 10–12 in CD₂Cl₂ and quinone 13 in (CD₃)₂CO at room temperature with respect to TMS



No.	$\delta_{ m C}$	$\delta_{ m H}{}^{ m a}$	$\delta_{ m C}$	$\delta_{ m H}{}^{ m a}$	$\delta_{ m C}$	$\delta_{ m H}{}^{ m a}$	$\delta_{ m C}$	${\delta_{ m H}}^{ m a}$
1	174.4		186.6		174.4		186.6	
2	152.7		149.2		152.6		149.3	
3	111.3	6.38d 1.8	134.3	6.99d 2.3	111.2	6.37d 1.8	135.0	7.29d 2.4
4	132.8		134.6		132.9		136.9	
5	102.5	6.81d 1.8	125.3	7.50d 2.3	102.5	6.79d 1.8	125.5	7.80d 2.4
6	152.8		149.4		152.8		149.8	
7	138.0	6.90dd 2.6, 8.6	140.3	6.93dd 2.8, 8.4	137.8	6.93m	135.4	7.86s
8	133.6	7.33dd 2.6, 8.6	134.8	7.37dd 2.9, 8.4	133.4	7.35m		
9	55.6	3.91	35.4		134.4	7.35m	35.4	
10	55.6	3.85	35.9		139.9	6.93m	35.8	
11			29.6	1.36	133.9		b	1.34s
12			29.7	1.32	133.7	6.99d 2.4	b	1.31s
13					148.6			
14					186.0			
15					148.7			
16					124.7	7.50d 2.4		
17					34.9			
18					35.3			
19					29.0	1.32		
20					29.0	1.36		
21					55.5	3.90		
22					55.5	3.84		

^a The values of $\delta_{\rm H}$ are followed by multiplicity and coupling constants (Hz).

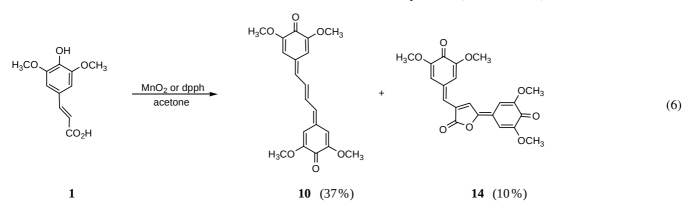
^b The methyl carbon signals of the *tert*-butyl groups fall into the solvent signal (acetone).

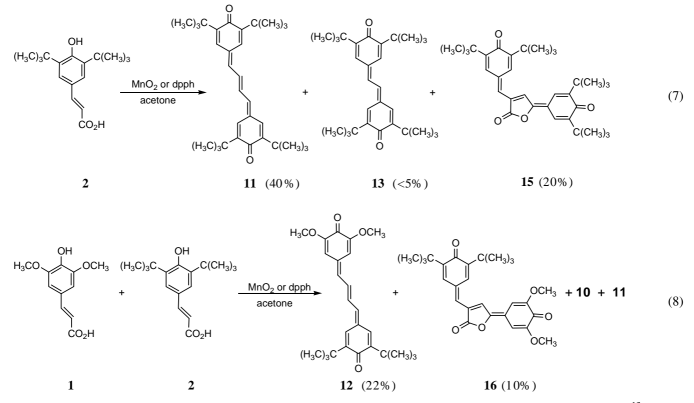
the ring protons ($\delta_{\rm H}$ 6.33 and 6.76). A less pronounced splitting was also observed in the ¹³C NMR spectrum for the pertinent carbon atoms (see Table 2). Similarly, the ¹H and ¹³C NMR spectra of the other three *p*-quinomethanes **11–13** showed an analogous splitting (see Table 2).

This molecular asymmetry has previously been observed²⁰ with compound **13** and rationalized.²¹ It is worth mentioning that in the case of **10a** the ¹H NMR spectrum showed one signal only for the four ring protons at about 6.67 ppm

and only one for the four methoxy groups at about 3.88 ppm down to a temperature of 190 K.

All *p*-quinomethanes **10–12** can also be prepared (yields 30-40%) by oxidation of HCA's at room temperature with activated MnO₂ in various solvents (see Section 4). The oxidation pattern with MnO₂ (evaluated by TLC analysis) was similar to that of dpph' but the slightly higher yields allowed an easier isolation and characterization of the major reaction products (Reactions 6–8).





The resemblance of the UV–vis and NMR spectra of quinone **11** to those of **10** (see Figure 2a and Table 2) allowed a straightforward recognition of its structure. In agreement, the ESI-MS spectrum in the positive ion-mode and in the presence of 1 mM LiCl in the acetonitrile phase showed the following peaks at m/z 483 [M+Na]⁺; 467 [M+Li]⁺ and 461 [M+H]⁺ (C₃₂H₄₄O₂ 460.71).

The ¹H NMR spectrum of the asymmetric p-quinomethane 12 in CD₂Cl₂ solution revealed resonances of two nonequivalent MeO groups at 3.89 and 3.84 ppm and two nonequivalent tert-butyl groups at 1.36 and 1.32 ppm. Additional proton signals were observed at δ 6.93 (2H, m) and at 7.35 (2H, m), which could be assigned to the couples of protons H-7/H-10 and H-8/H-9, respectively, (see Table 2), on the basis of the correspondence with the resonance frequencies of H-7 and H-8 in the symmetrical quinones 10 and 11 (Table 2). The ring protons resonated as doublets at δ 6.37 (1H, J=1.8 Hz), 6.79 (1H, 1.8 Hz), 6.99 (1H, 2.4 Hz) and 7.50 (1H, 2.4 Hz). In this case, the correspondence in terms of coupling constants and chemical shifts with the ring protons of quinones 10 and 11 (see Table 2) allowed to assign the upfield signals, that is, 6.37 and 6.79 ppm, to the protons of the ring bearing the MeO groups and the downfield signals, that is, 6.99 and 7.50 ppm, to the ring with the *tert*-butyl groups.

The two carbonyl groups of **12** at positions 1 and 14 resonated in the ¹³C NMR spectrum in CD_2Cl_2 at 174.4 and 186.0 ppm, respectively, as in the symmetric quinones **10** (174.4 ppm) and **11** (186.6 ppm) (see Table 2). This correspondence was also observed in the FT-IR spectra as the frequencies of CO stretching of the three quinones were 1621.9 cm⁻¹ in **10**, 1606.3 cm⁻¹ in **11**, and 1629.9 and 1607.7 cm⁻¹ in **12** (see Table 1). Aside from the carbon

signals of the *tert*-butyl and methoxyl groups, the ¹³C NMR spectrum of **12** in CD_2Cl_2 showed 16 distinct carbon atoms versus the 8 carbon signals of the symmetric quinones **10** and **11**. HMQC, HMBC and NOESY 2D experiments (see Fig. 3) were done to establish the direct C–H bonds and the C–C connectivity.

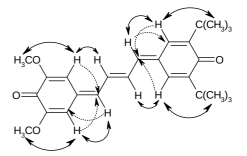


Figure 3. Selected HMBC (dotted line) and NOESY (solid line) correlations observed in the asymmetric quinone 12.

Finally, the ESI-MS spectrum of **12** in the positive ion-mode and in the presence of 0.1 mM LiCl confirmed the structure assignment because of the presence of peaks at m/z 431 $[M+Na]^+$; 415 $[M+Li]^+$ and 409 $[M+H]^+$ (C₂₆H₃₂O₄ 408.54).

Quinones 14–16 were isolated from the reaction mixtures of 1, 2 and 1+2 (1:2 mol/mol) with either dpph⁻ or MnO₂ in acetone at room temperature (yields ca. 10–20%, Reactions 6–8). Their UV–vis spectra are reported in Figure 2b and Table 1. The ¹H and ¹³C NMR spectra reported in Table 3 indicated that compounds 14–16 shared the same gross molecular structure. Indeed, the proton spectrum of 14 in

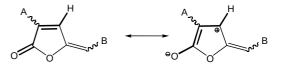
CD₂Cl₂ showed the presence of four distinct signals in the range 3.86–3.94 ppm corresponding to four non-equivalent methoxy groups. In the case of compound **15**, three upfield signals at 1.33, 1.36 and 1.38 ppm (ratio 1:2:1, respectively) were observed and assigned to four *tert*-butyl groups. Two distinct signals of methoxy groups at 3.91 and 3.92 ppm and two distinct signals for two *tert*-butyl groups at 1.33 and 1.38 ppm were present in the spectrum of **16**. These spectral data therefore indicated the presence of two di-substituted benzene rings in each molecule. In addition, all ¹H NMR spectra in CD₂Cl₂ exhibited 4 sharp doublets (4H, $J \sim 1.8-2.4$ Hz), suggestive of two couples of *meta* benzene protons, and 2 sharp singlets (2H) in the spectral region 6.42–8.07 ppm, see Table 3.

The presence of the γ -lactone moiety in **14–16** was deduced by a combination of NMR and IR spectroscopy. The FT-IR spectra in KBr suggested the presence of two different carbonyl groups ($\nu_{CO} \sim 1640-1612$ and $\nu_{CO} \sim 1780-$ 1770 cm⁻¹) one of which could be attributed to a quinone moiety (1640–1612 cm⁻¹) and the other one to a γ -lactone (1780–1770 cm⁻¹).²² Resonance peaks at about 167 ppm in the ¹³C NMR spectra in CD₂Cl₂ of **14–16** confirmed the presence of a γ -lactone moiety (see Table 3). Further, the presence of a considerably deshielded proton at about 8 ppm

14

Table 3. 1 H and 13 C NMR spectra of quinones 14–16 in CD₂Cl₂ with respect to TMS

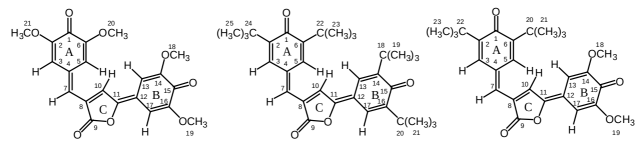
in the ¹H NMR spectra in CD_2Cl_2 and the extended π -electron system of these molecules, which were intensely coloured suggested an unsaturation α to the C=O and an additional exocyclic double bond on the carbon α to the O.



The above spectral data allowed the partial structures, that is, two *p*-quinomethane units and the γ -lactone unit, to be assembled into the structures given for **14–16**. The number of quaternary and C–H carbons observed in the ¹³C NMR and DEPT spectra of **14–16** (i.e., 11 and 6, respectively) was consistent with the proposed structures as well as the HMQC, HMBC and NOESY 2D spectra. Figure 4 shows a few selected correlations observed in the HMBC and NOESY spectra of **15** and **16**.

In the case of **16**, the specific substitution of the rings A and B was deduced from the observed identity of the chemical shifts of the carbons and protons 3 and 5 with those of **15**, and 13 and 17 with those of **14**, see Table 3. This supported the conclusion

16



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No.	$\delta_{ m C}$	$\delta_{ m H}{}^{ m a}$	$\delta_{ m C}$	$\delta_{ m H}{}^{ m a}$	$\delta_{ m C}$	$\delta_{ m H}{}^{ m a}$
1	174.6		186.2		186.2	
2	151.2		149.5		149.3	
3	111.7	6.42d 1.8	134.0	7.06d 2.2	134.1	7.06d 2.3
4	129.4		129.6		128.7	
5	104.4	7.14d 1.8	126.0	7.67d 2.2	126.1	7.68d 2.3
6	152.6		150.8		150.6	
7	129.0	6.80s	126.2	6.86s	126.7	6.87s
8	136.3		138.4		137.8	
9	167.1		167.2		167.1	
10	129.7	7.92s	131.1	8.07s	130.4	8.00s
11	153.6		152.9		151.1	
12	118.7		118.9		118.9	
13	102.4	6.58d 1.8	124.6	7.29d 2.4	102.2	6.56d 1.8
14	153.7		151.5		153.7	
15	174.6		185.9		174.6	
16	153.5		150.5		153.4	
17	102.6	6.97d 1.8	124.6	7.61d 2.4	102.5	6.95d 1.8
18	56.3	3.93s	35.4		56.2	3.91s
19	56.3	3.94s	29.0	1.36s	56.1	3.92s
20	56.1	3.86s	35.6		35.0	
21	55.6	3.92s	29.0	1.36s	29.0	1.38s
22			35.0		35.5	
23			28.9	1.38s	29.1	1.33s
24			35.7			
25			29.1	1.33s		

^a The values of $\delta_{\rm H}$ are followed by multiplicity and coupling constants (Hz).

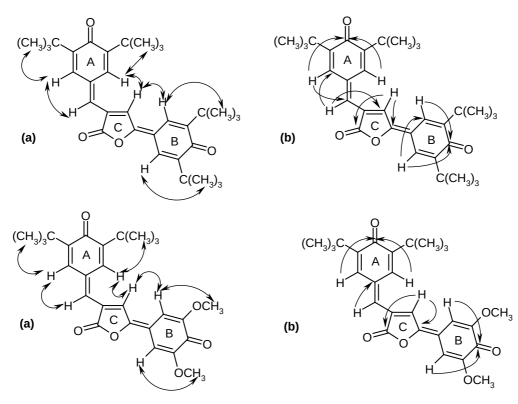


Figure 4. (a) Selected NOESY correlations observed in 15 and 16; (b) selected HMBC correlations observed in 15 and 16.

that the *t*-Bu groups were located on the ring A whereas ring B contained two MeO groups. Consistently, the NOESY spectrum of **16** (see Fig. 4) showed a correlation of H-7 ($\delta_{\rm H}$ =6.87) with the nearest proton of the ring bearing the *t*-Bu groups, that is, H-3, $\delta_{\rm H}$ =7.06. The transoid geometry of the protons 7 and 10 was deduced from the NOESY spectrum, which showed a marked correlation of H-10 with H-5 whereas there was no noticeable correlation between H-7 and H-10.

The ESI-MS spectra of quinones **14–16** were done in the positive ion-mode. For the analyses of **14** and **16** the eluents (water/acetonitrile) were added of 0.1 mM LiCl whereas in the case of **15**, which ionized with difficulty the LiCl concentration in the acetonitrile phase was increased to 1 mM. The MS spectrum of **14** showed peaks at m/z 437 [M+K]⁺; 421 [M+Na]⁺; 405 [M+Li]⁺ and 399 [M+H]⁺, which confirmed the structure assignment (C₂₁H₁₈O₈ 398.37). Analogously, the main peaks in the MS spectra of quinones **15** and **16** were at m/z 525 [M+Na]⁺; 509 [M+Li]⁺, 503 [M+H]⁺ (C₃₃H₄₂O₄ 502.70) and 473 [M+Na]⁺; 457 [M+Li]⁺ and 451 [M+H]⁺ (C₂₇H₃₀O₆ 450.54), respectively.

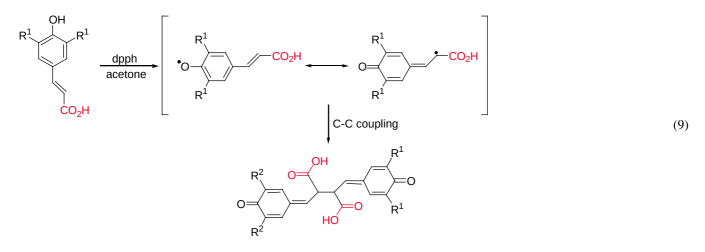
2.2. Kinetic aspects and mechanism

The molecular structures of 10-16 suggest that the formation of these *p*-quinomethanes proceeds through a C-C coupling of ArO' radicals at the positions 8 of the side chains (one exception is represented by 13). The ArO' radicals are produced by H-atom abstraction from the phenolic OH of 1 and 2 by dpph' (or MnO₂), Reaction 3. The absence in the final products of one (14–16) or two (10–12) carboxylic groups is particularly

surprising because of the mild conditions employed in our experiments. Decarboxylation of α , β -unsaturated carboxylic acids is known to take place with difficulty and usually requires vigorous experimental conditions²³ although for cinnamic acids the presence of p-OH seems to accelerate the process, which in any case requires comparatively high temperatures.²⁴ On the contrary, the intermediates involved in our reactions apparently undergo decarboxylation fairly readily. Under our experimental conditions, we observed that the free carboxylic acid group in the reacting HCA's 1 and 2 is of pivotal importance. Indeed, when the methyl ester 8 of sinapic acid 1 was allowed to react with dpph in acetone at room temperature no traces of 10 or 14 were detected in solution.²⁵ The spectral changes observed during the reaction corresponded exclusively to the occurrence of Reaction 3 with a rate constant of $16\pm 2 \text{ M}^{-1} \text{ s}^{-1}$ and a stoichiometric factor²⁶ of ca. 1.0. The latter value demonstrates that once the aryloxyl radicals from 8 were formed in Reaction 3 they exclusively self-quenched (Reaction 4) without reacting further with dpph'. It is interesting to observe that in the case of the reactions 1+dpph' or 2+dpph' carried out in acetone the stoichiometry was 1:2.

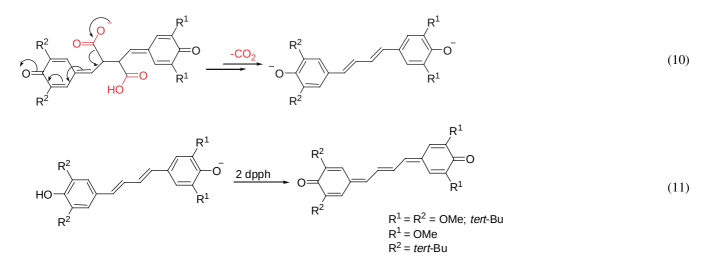
Very recently, Bietti and Capone²⁷ have provided spectroscopic evidence that arylethanoic acids, in the presence of SO_4^- radicals (the oxidizing agent), lose CO_2 in water via an aromatic radical-cation formed after an ET process to SO_4^- (with formation of SO_4^{2-} ions). Subsequently, the aromatic radical-cation undergoes fast intramolecular ET from the carboxylate anion to the ring followed by decarboxylation with formation of a resonance-stabilized benzyl radical either by a concerted or stepwise mechanism.²⁸ Unlike arylethanoic acids, in our case similar reactions cannot be invoked for the process of decarboxylation of the precursors of quinones 10-16 for at least two reasons. Firstly, once the radical-cations 1^{+} or 2^{+} were formed by ET from 1 or 2 to dpph they would rapidly lose a proton from the phenolic OH, affording the aryloxyl radical ArO⁺, long before the process of ionization of the carboxylic acid group since phenol

solution $(k \sim 10^9 \text{ M}^{-1} \times \text{s}^{-1})$.³² If the actual mechanism of formation of **10–16** involved coupling of Ar–CH=CH' in the presence of dioxygen there would not be any formation of C–C dimers.³³ Actually, the experiment showed that the yields of quinones **10** and **11** were unaffected by the presence or absence of oxygen in solution. Thereby, we can conclude that decarboxylation must occur after the 8–8 C–C dimerization of aryloxyls (Reaction 9).³⁵



radical-cations are strong acids³⁰ (in any case, the methyl ether of **1** demonstrated to be inert to dpph' or MnO_2). Secondly, if a fraction of **1**⁺⁺ or **2**⁺⁺ were able to give the acyloxyl radical, that is, Ar–CH=CH–CO₂, by ET from the carboxylate anion to the aromatic ring, this

The initial C–C dimer formed in Reaction 9 may lose CO_2 via an S_E1 mechanism^{36,24} (Reaction 10) followed by a fast oxidation of the phenolic intermediates (Reaction 11).



would likely react with the H-atom donors present in our system, (i.e., 1 or 2, dpph-H and acetone) thus regenerating the parent phenol³¹ since the process of decarboxylation of such a vinylacyloxyl radical is expected to be comparatively slow (lifetime of the order of microseconds).^{31b-e}

In connection with the above arguments, it is important to point out that experiments carried out with dioxygen did not highlight formation of carbon-centered radicals, that is, Ar–CH=CH⁻, in our system. Generally, carbon radicals are known to react very quickly with the dioxygen dissolved in

Reaction 9 is the rate-determining step for the formation of the quinones **10–16** in aprotic solvents and its rate is proportional to $[ArO']^2$ (the rate constant can be close to the diffusion limit³⁷). The steady-state concentration of ArO' is essentially determined by the rates of Reaction 3 and of the overall processes of ArO' quenching. The substituents present on phenols exert strong effects on the rate constant of Reaction 3. Electron-donating (ED) groups in the *ortho* and *para* positions of the phenol ring decrease the bond dissociation enthalpy (BDE) of OH and *increase* the rate of Reaction 3.^{2,38} On the contrary, electron-withdrawing groups increase the OH BDE and *decrease* the rate of ArO' formation.^{2,38} Therefore, the low yields observed in the reactions of HCA's **3** (bromine present in the *ortho* positions), **4** (no ED groups in the *ortho* positions), **6** (one only *ortho* ED group and intramolecular hydrogen-bond)⁶ and **7** (one ED group in *meta* position) with dpph' are readily explained. In the case of caffeic acid **5**, the rate constant of Reaction 3 is comparatively large because of the presence of two *ortho* OHs.² However, the main stabilization pathway of the semiquinone radical of **5** consists of an additional H-atom transfer to the dpph' radical with formation of *ortho*-quinone therefore precluding the dimerization and subsequent reactions.

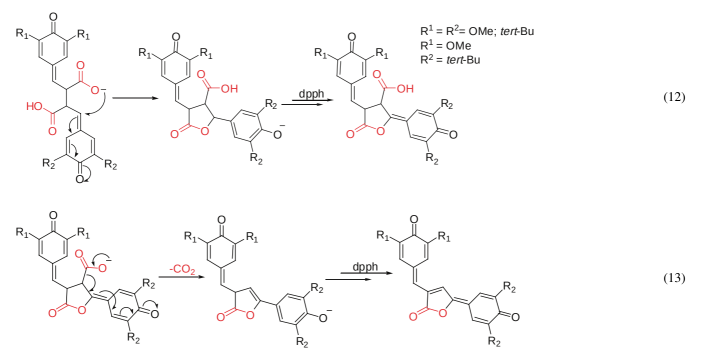
The reaction mechanism suggested above for the formation of quinones 10-12 justifies the presence of a few minor compounds found in solution. Traces of phenol 10a were detected, for instance, during the oxidation of 1 with dpph. or MnO_2 in acetone at room temperature although this phenol reacts quickly with dpph', being $k_3(10a) = 1130 \pm$ $80 \text{ M}^{-1} \text{ s}^{-1}$ and n=2.0. The presence of 10a can be considered a convincing evidence of the occurrence of Reaction 10. Other minor compounds include quinones 14–16 bearing the γ -lactone moiety, which may originate from an intramolecular nucleophilic addition of the carboxylate anion to the opposite terminal methylene of the quinone system (Reaction 12). A subsequent S_{E1} mechanism³⁶ of decarboxylation of the intermediate carboxylated γ -lactone and a stepwise oxidation by dpph radicals may lead to the final quinones bearing the unsaturated γ -lactone ring (Reaction 13).

that hydroxylic solvents support the ionization of Brønsted acids better than non-hydroxylic solvents of similar dielectric constant.^{3–5,40} It seems likely, therefore, that the solvent plays an important role in the process of decarboxylation of the initial dimer (Reaction 10). In fact, similar solvent effects were reported for the unimolecular decarboxylation of substituted benzisoxazole-3-carboxylate anion.⁴¹ Dramatic rate accelerations resulted if protic solvents (water, methanol or ethanol) were replaced by aprotic solvents. For instance, the rate of decarboxylation at 30 °C of 6-nitrobenzisoxazole-3-carboxylate ion in acetone resulted to be ca. 100,000 times larger than in methanol and ca. 3,300,000 times larger than in water!⁴¹

There are a number of potential explanations⁴² for the inhibitory effect of alcohols but we consider it most probable that in these media the strong solvation of negative ions by hydrogen bonding is retarding decarboxylation of the initial dimer (Reaction 10) by stabilizing the carboxylate ion.⁴¹

3. Conclusion

We have described the synthesis and spectral identification of highly conjugated dimeric quinones **10–16** some of which bearing a peculiar unsaturated γ -lactone ring (**14–16**). The synthesis of these quinones consists of oxidizing 4-hydroxy cinnamic acids with dpph[•] (or with MnO₂) in an appropriate solvent at room temperature, the process being most successful when: (i) the solvent is non-



Finally, the polar nature of the intermediates involved in the mechanisms outlined above may also explain the solvent effects observed on the yields and course of Reactions 6–8. We found that non-hydroxylic polar solvents with higher dielectric constants generally promoted the formation of quinones better than solvents with low dielectric constants.³⁹ In alcohols, however, no formation of quinones **10–16** was observed³⁹ at room temperature despite the fact

hydroxylic and of high dielectric constant; and (ii) the *ortho* positions to the phenolic OH of HCA are both occupied by bulky electron-donating groups.

The yields are low to moderate (10–40%) because of the many side reactions, however, the complex structure of these compounds makes these yields acceptable.

Kinetic data along with molecular structures and the presence in solution of a few intermediates suggest that the mechanism of formation of **10–16** with dpph[•] proceeds through four different steps: (1) formation of ArO[•] by H-atom transfer from HCA to dpph[•]; (2) dimerization by 8–8 C–C coupling of two ArO[•] radicals; (3) fast decarboxylation at room temperature of the intermediate dimer by an S_E1-type mechanism; and finally, (4) oxidation of the intermediate phenolate anions by dpph[•].

4. Experimental

4.1. General

Cinnamic acids 1, 4, 6 were purchased from Fluka; ascorbic acid, dpph['], cinnamic acid 2 were obtained from Aldrich and cinnamic acids 5 and 7 from Extrasynthèse. All compounds were used as received. The methyl ester 8 of sinapic acid was available from a previous work, its synthesis is described in Ref. 4. 3,5-Di-bromo-4-hydroxycinnamic acid 3 was donated by Dr. Paolo Bovicelli (ICB-CNR, Università La Sapienza, Roma) and its synthesis and spectral characterization will be reported in a separate paper. All solvents (Carlo Erba and Merck) were of the highest commercially available quality and were used without further purification (except for diethyl ether and THF, which were distilled prior to their use). NMR spectra were recorded at 400.13 MHz (¹H) and 100.62 MHz (¹³C) in CD_2Cl_2 solutions at 298 K on a Bruker AvanceTM 400 spectrometer. Chemical shifts were referenced to the residual signal of CD₂Cl₂. HPLC analyses were done on an instrument (Waters 1525) equipped with ESI-MS (Waters Micromass ZQ) and UV-DAD (Waters 996) detectors (column: Phenomenex[®] Luna, C18, $250 \times 4.6 \text{ mm}$ (5 µm) at 20 °C using as eluent system H₂O/CH₃CN containing 0.1 or 1 mM LiCl). A double-ray Perkin Elmer Lambda 25 spectrophotometer was used for the kinetics and to record the UV-vis spectra whereas the FT-IR spectra were obtained with a Perkin Elmer Spectrum BX FT-IR System spectrophotometer. Analytical and preparative (silica gel, $20 \times$ 20 cm, 0.5-1 mm thick) TLC plates and silica gel (63-200 µm) were purchased from Merck. The syntheses and purification reported in the following paragraphs for 10 and 14 with dpph' and 12 and 16 with MnO₂ are general and apply to all quinones. The purity of quinones 10-16 determined by HPLC analysis was not inferior to 95%.

4.2. Preparation of activated MnO₂

MnSO₄·4H₂O (11 g) were dissolved in 15 ml of distilled water and the solution treated with 11.7 ml of 40% NaOH (solution A); 9.6 g of KMnO₄ were dissolved in 60 ml of hot distilled water (solution B). Then, the two solutions A and B were slowly mixed together in about 1 h under vigorous stirring. The final solution was centrifuged and the precipitate of MnO₂ washed with distilled water until the wash waters were colourless. The solid was then dried at 100–120 °C.

4.2.1. Preparation of 4,4'-(2-butene-1,4-diylidene)bis(2,6-dimethoxy-2,5-cyclohexadien-1-one) 10 and 5-(3,5-dimethoxy-4-oxocyclohexa-2,5-dienylidene)-3-[(3, 5-dimethoxy-4-oxocyclohexa-2,5-dienylidene)methyl]furan-2(5H)-one 14 by using dpph'. Sinapic acid 1 (200 mg, ca. 0.9 mmol) were allowed to react with 1.06 g of dpph' (2.69 mmol) in 300 ml of acetone at 25 °C in the dark for 2 h. After solvent removal, the crude product was purified by column chromatography on silica gel using ethyl acetate–hexane (80/20, v/v), acetone and methanol as eluents to give ca. 60 mg of **10** (final yield 37%) and ca. 20 mg of **14** (final yield 10%) as a dark violet powder. ¹H and ¹³C NMR spectra were performed in dilute CD₂Cl₂ solutions and are reported in Tables 2 and 3. The UV–vis spectra are given in Table 1. The ESI-MS spectrum is discussed in Section 2.

4.2.2. Preparation of 4-[(3,5-di-tert-butyl-4-oxocyclohexa-2,5-dienylidene)but-2-enylidene]-2,6-dimethoxycyclohexa-2,5-dienone 12 and 3-[(3,5-di-tert-butyl-4-oxocyclohexa-2,5-dienylidene)methyl]-5-(3,5-dimethoxy-4-oxocyclohexa-2,5-dienylidene)furan-2(5H)-one 16 by using MnO₂. 3,5-Di-*tert*-butyl-4-hydroxycinnamic acid 2 (180 mg, 0.65 mmol) and sinapic acid 1 (75 mg, 100 mg)0.33 mmol) were solubilized in 10 ml of acetone. The solution was then added with a syringe-pump (200 μ l/min) to an initial suspension of 200 mg of MnO₂ in 1 ml of acetone (in the dark and under stirring) to which aliquots of 100 mg each of MnO₂ per 2 ml of phenol solution pumped were successively added (700 mg in total corresponding to 8 mmol). After 2 h the suspension was filtered and the solvent removed. The crude residue was then purified on a preparative TLC plate (20×20 silica gel, 1 mm thick) using hexane-acetone (90/10) as eluent to give 30 mg of 12 (yield 22%) and 12 mg of 16 (yield 10%). The UV-vis and FT-IR spectra are reported in Figures 2a and b and in Table 1. ¹H and ¹³C NMR spectra performed in dilute CD_2Cl_2 solutions are reported in Tables 2 and 3. The ESI-MS spectra are reported in Section 2.

4.2.3. Purification of 4,4'-(2-butene-1,4-divlidene)bis(2,6-di-*tert*-butyl-2,5-cyclohexadien-1-one) 11, 4,4'-(1,2-ethanediylidene)-bis(2,6-di-tert-butyl-2,5cyclohexadien-1-one) 13 and 5-(3,5-di-tert-butyl-4-oxocyclohexa-2,5-dienylidene)-3-[(3,5-di-tert-butyl-4-oxocyclohexa-2,5-dienylidene)methyl]furan-2(5H)-one 15. These quinones can be obtained with both methods reported above. Their purification can be accomplished by column chromatography on silica gel or preparative TLC (silica gel) using hexane-THF (95/5) (final yields, 40% for 11 and 20% for 15). Quinone 13 was obtained in very low yield (<5%); however, it was possible to characterize this molecule and its UV-vis and FT-IR spectra are reported in Figure 2a and Table 1. The NMR spectra are reported in Table 2; the ESI-MS spectrum obtained in the presence of 1 mM LiCl in the acetonitrile phase showed the following peaks at m/z 441 $[M+Li]^+$ and 435 $[M+H]^+$ (C₃₀H₄₂O₂ 434.67).

4.2.4. Synthesis of 1,4-di(4-hydroxy-3,5-dimethoxyphenyl)-1,3-butadiene 10a. This phenol was obtained by reduction of 10 with ascorbic acid using the following procedure: 4 mg of 10 (0.011 mmol) were dissolved in 2 ml of CH₂Cl₂-CH₃OH (1/1 v/v) then 3.9 mg of ascorbic acid (0.022 mmol) were added. The solution was shaken at 35-45 °C for ca. 1 h in the dark. After solvent removal, compound 10a was extracted from the residue with CH₂Cl₂ (3×1 ml). The evaporation of the solvent yielded 10a as a pale

yellow solid in a quantitative yield. ¹H NMR (CD₂Cl₂) (numbering system: OH on C-1 and C-7 and C-8 on the butadiene chain), δ =3.88 (s, 12H, OCH₃), 5.54 (s, 2H, OH), 6.54 (dd, *J*=11.6, 2.4 Hz, 2H, H₇), 6.67 (s, 4H, H_{3,5}), 6.83 (dd, *J*=11.6, 2.4 Hz, 2H, H₈). ¹³C NMR (CDCl₃) δ =56.08 (OCH₃), 103.14 (C_{3,5}), 127.31 (C₈), 128.88 (C₄), 131.80 (C₇), 134.78 (C₁), 147.13 (C_{2,6}). FT-IR (CH₂Cl₂, cm⁻¹) 3528.97 (m, OH), 1616.10 (m), 1601.0 (m), 1511.79 (s). The UV-vis spectrum in methanol shows a maximum at 354 nm ε =(5.4 ± 0.2)×10⁴ M⁻¹ cm⁻¹. The HPLC-ESI-MS (water/acetonitrile) spectrum of **10a** in the negative ion-mode showed peaks at *m/z* 737 [2M-H]⁻; 357 [M-H]⁻ (base peak); 342 and 327 (C₂₀H₂₂O₆ 358.39).

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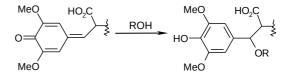
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- 42. For instance, it is possible that in the alcohols the following addition reaction to the dimer takes place:



This reaction is, however, evaluated to be comparatively slow. In the case of a methyl *p*-quinomethane, the rate constant relative to the addition of neutral methanol has been reported to be 0.031 s^{-1} at 23 °C and ca. 500 times lower for the addition to a more stable *p*-quinomethane.²¹ Other evidence in support of the poor importance of this reaction is given by the fact that the yield of formation of **10** is not significantly enhanced in the sterically hindered *tert*-butyl alcohol.³⁹



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Origin of the stereo- and regioselectivities in the Diels–Alder reactions of azaphospholes: a DFT investigation

Raj K. Bansal,* Neelima Gupta and Surendra K. Kumawat

Department of Chemistry, University of Rajasthan, Jaipur 302 004, India

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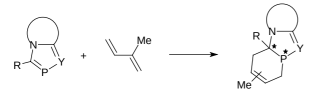
Abstract—Computations of the Diels–Alder (DA) reactions of azaphosphole representative namely, thiazolo[3,2-*d*][1,4,2]diazaphosphole with 1,3-butadiene and isoprene at the density functional theory level reveal concerted mechanisms via asynchronous transition states. The activation energies (B3LYP/6-311++G**// B3LYP/6-311G**), 16–19 kcal mol⁻¹, are much smaller than the value (32.57 kcal mol⁻¹) calculated for the DA reaction of the non-phosphorus analogue, imidazo[2,1-*b*]thiazole with 1,3-butadiene. An electron-withdrawing group at the 3-position of the dienophile enhances both stereo- and regioselectivities, which agree nicely with the experimental values. Inclusion of solvent effect (PCM model) reveals that the stereo- and regioselectivities are not affected appreciably. The relative stabilities of the transition structures corresponding to the *endolexo* stereoisomers and *meta* (P/Me, 1:3)/*para* (P/Me, 1:4) regioisomers have been rationalized on the basis of the secondary molecular orbital interactions.

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1. Introduction

The Diels–Alder (DA) reaction is perhaps the most widely used method for the construction of the six-membered rings.¹ It has also been central in the development of theoretical models of pericyclic reactions.² In recent years the scope of this reaction has been further enlarged by extending it to the organophosphorus compounds having C=P- functionality, namely phosphaalkenes,³ heterophospholes,⁴ anellated azaphospholes⁵ and phosphinines.⁶

The DA reaction of the >C=P- functionality of the azaphosphole is of particular significance, because it generates two stereogenic centres in one step (Scheme 1). The results, however, indicate that these reactions are normally accompanied by complete diastereoselectivity and regioselectivity.^{7,8}



Scheme 1.

0040–4020/\$ - see front matter 2005 Elsevier Ltd. All rights reserved. doi:10.1016/j.tet.2005.11.009

Recently, we have undertaken a systematic study of the DA reactions of some azaphospholes whose syntheses we had reported earlier.^{5,9–11} 1,3-Azaphospholo[5,1-*a*]isoquinoline undergoes DA reactions with 2,3-dimethylbutadiene¹⁰ and with isoprene,¹² all reactions occurring with complete *endo*stereoselectivity and in the latter case with 62-100% regioselectivity dependent on the conditions. The DA reactions of [1,4,2]diazaphospholo[4,5-a]pyridines with 2,3-dimethylbutadiene and with isoprene occur likewise, 80% regioselectivity being observed in the latter case.¹³ The recently reported DA reactions of thiazolo[3,2-d][1,4,2]diazaphospholes and their dihydro and benzo derivatives, and also those of 1,3-azaphospholo[5,1-b]benzothiazole are accompanied by several interesting features: complete endo-stereoselectivity, 70-100% regioselectivity in the reaction with isoprene, and differences in the reactivities of thiazolo[3,2-d][1,4,2]diazaphospholes and 1,3-azaphospholo[5,1-b]benzothiazole, the former reacting at room temperature while the reaction of the latter being completed only at elevated temperature in the presence of an oxidizing agent (O₂, S₈ or Se_n).¹⁴

In more recent publications, results of several DA reactions have been frequently explained on the basis of computational calculations.¹⁵ The reports concerning the theoretical studies of the DA reactions of the phosphorus containing reactants are, however, scarce. Computations of the DA cycloadditions of phospha-1,3-butadiene with ethene,¹⁶ of phosphaethene and of phosphaethyne with butadiene,¹⁷ and those of phosphaethene with 2*H*-phosphole¹⁸ and with 1,3-butadiene,^{19–21}

Keywords: DFT calculations; Azaphospholes; Diels–Alder reactions; Stereoselectivity; Regioselectivity; NICS values.

^{*} Corresponding author. Tel.: +91 1412701868; fax: +91 1412652730; e-mail: rajbns@yahoo.com

revealed low activation energies and exothermicities and also a preference for the endo approach. It was concluded that the presence of phosphorus in a DA reactant lowers the activation energy barrier relative to the analogous hydrocarbon system due to the weaker C=P π bond as compared to the C=C π bond.^{16,17} The DFT calculations of the prototype DA reactions of phosphaethene with 1,3-butadiene and with isoprene at $B3LYP/6-311+G^{**}$ level established a concerted mechanism, but indicated low regioselectivity, and in order to account for the experimentally observed high regioselectivity, an alternative radical cation mechanism was proposed.²² In connection with a study of the aromaticitycontrolled DA reactions, it has been reported that an aromatic stabilization process of the masked diene accelerates the DA reaction through an 'early' TS, while dearomatization of a DA reactant deactivates the cycloaddition via a 'late' TS.²³ In this context, a model DA reaction of the acyclic phosphaethene with 1,3-diene²² is inadequate to explain all the aspects of the DA reactions of aromatic azaphospholes. These results prompted us to compute DA reactions of two representative azaphospholes, namely thiazolo[3,2-d][1,4,2]diazaphosphole and 3-methoxycarbonylthiazolo[3,2-d][1,4,2]diazaphosphole at the DFT (B3LYP/6-311G**) level, with a view to investigate the origin of the stereo- and regioselectivities observed in these reactions. The ratios of the endo/exo stereoisomers and meta (P/Me, 1:3)/para (P/Me, 1:4) regioisomers computed 24 from the Boltzmann distribution are found to be in very good agreement with the experimental values. The aromaticity of the transition states is confirmed by their NICS values.²⁵ Dearomatization accompanying DA reactions of these dienophiles raises the activation energy barriers by 4–5 kcal mol⁻¹ as compared to those for the DA reaction of the acyclic phosphaethene.

Furthermore, as the reactions were carried out in toluene,¹⁴ solvent effect has also been studied using polarizable continuum model, which reveals that the solvent has no effect on stereo- or regioselectivity.

2. Computational methods

All calculations were carried out with the Gaussian 03 suite of programmes.²⁶ Reactants and the products and transition structures resulting from the DA reactions were optimized at B3LYP/6-311G** level. The frequencies were computed at B3LYP/6-311G** level. All minima and transition structures were confirmed to have none or one imaginary frequency, respectively. The normal mode corresponding to the imaginary frequency in the case of a particular transition structure was found to involve vibrations of the new bonds (C3–C12, P2–C9) being formed. Unscaled zero-point energy corrections (ZPE) from B3LYP/6-311G** level were added to the single point energies calculated at B3LYP/6-311++G** level.

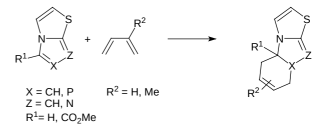
Intrinsic reaction coordinate (IRC) calculations starting at all the transition structures were performed to establish their connection with the respective reactants and products. Natural bond orbital (NBO) analysis²⁷ was used for computing the bond orders (Wiberg bond indices)²⁸ of the transition structures. NICS values were

calculated at GIAO-B3LYP/6-311 + $G^{**}/B3LYP/6-311G^{**}$ level.²⁹

The solvent effect has been studied by calculating single point energy of the B3LYP/6-311G** gas phase optimized stationary points at B3LYP/6-311++G** level using self-consistent reaction field (SCRF) method^{30,31} based on Tomasi's integral equation formalism polarizable continuum model (iefpcm).³² To the energy so obtained was added the unscaled ZPE calculated at B3LYP/6-311G** level for the gas phase.

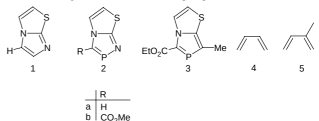
3. Results and discussion

To investigate the origin of the regio- and stereoselectivities in the DA reactions of azaphospholes we have examined ten model reactions, (Scheme 2 and Table 1) including a DA reaction with the non-phosphorus analogue, imidazo-[2,1-b]thiazole (1).



Scheme 2.

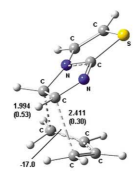
Table 1. Investigated DA reactions of azaphospholes with 1,3-dienes



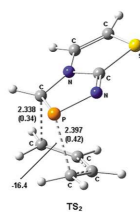
Reaction no.	Reactants (approach)	TS	Product
1	1+4 (endo)	TS ₁	P ₁
2	2a+4 (endo)	TS_2	P_2
3	2a+4 (exo)	TS_3	$\overline{P_3}$
4	2a+5 (endo)	TS_4	P_4 (P/Me, 1:3)
5	2a+5 (endo)	TS ₅	P ₅ (P/Me, 1:4)
6	$2\mathbf{b} + 4$ (endo)	TS ₆	P ₆
7	$2\mathbf{b} + 4$ (exo)	TS ₇	\mathbf{P}_{7}
8	2b + 5 (endo)	TS ₈	P_{8} (P/Me, 1:3)
9	2b+5 (endo)	TS ₉	P_{0} (P/Me, 1:4)
10	3+4 (endo)	TS_{10}	P ₁₀

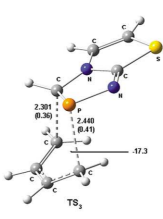
3.1. Geometries

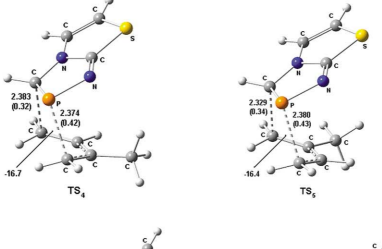
The B3LYP/6-311G^{**} geometries of the transition structures in the DA reactions of imidazo[2,1-*b*]thiazole (1) (TS₁), thiazolo[3,2-*d*][1,4,2]diazaphospholes (2) (TS₂–TS₉), and 1,3-azaphospholo[5,1-*b*]thiazole (3) (TS₁₀) are given in Figure 1 along with the bond distances (in Å), Wiberg bond



TS₁







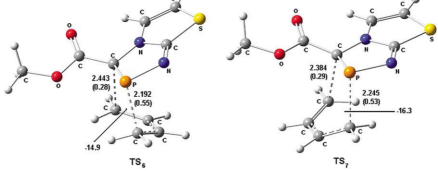


Figure 1. Optimized geometries (B3LYP/6-311G**) of the transition structures (TS₁–TS₁₀), bond distances (in Å), Wiberg bond indices (in parenthesis) and NICS values (in ppm).

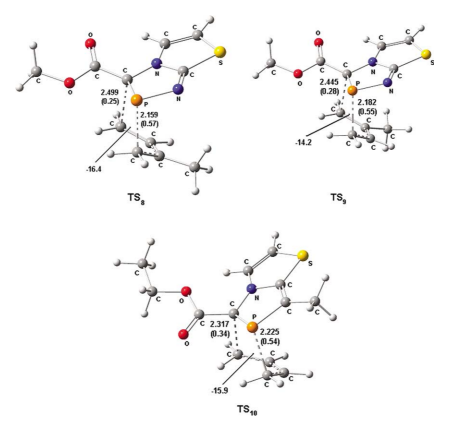


Figure 1. (continued)

indices (in parenthesis) of the two new forming bonds, and also the NICS values (in ppm).

 $\pi_{P2-C3} \rightarrow \pi^*_{C9-O10}$, $LP_{O11} \rightarrow \pi^*_{C9-O10}$ and $\pi^*_{P2-C3} \rightarrow \pi^*_{C9-O10}$ interactions (Table 2).

The pericyclic mechanism of all the DA reactions is established by the aromatic character of the respective transition structures, as revealed by their NICS values (-14 to -17).²⁵

The transition structures are, however, asynchronous, the forming P2–C9 bond being more advanced (WBI=0.41–0.57) than the C3–C12 bond (WBI=0.25–0.36). The asynchronicity of the TS increases further in the DA reactions of 3-methoxycarbonyl substituted thiazolo-[3,2-d][1,4,2]diazaphosphole (TS₆–TS₉) due to steric hindrance caused by the methoxycarbonyl group. There is, however, no appreciable difference in the asynchronicities between *endolexo* transition structures and also between the *meta* (P/Me, 1:3) and *para* (P/Me, 1:4) transition structures in the reactions with isoprene.

On scanning the potential energy surface of 3-methoxycarbonyl-thiazolo[3,2-*d*][1,4,2]diazaphosphole (**2b**), two minima corresponding to the two conformers resulting from rotation around C3–C9 bond are located, which are separated by a saddle point. The optimized geometries (B3LYP/6-311G^{**}) of the two conformers **2bA**, **2bB** and the transition structure ($\Delta E_a = 10.23$ kcal mol⁻¹) are given in Figure 2. The conformer **2bA** (torsion angle PCCO= 180.0) is more stable than **2bB** (torsion angle=0.0) by 2.64 kcal mol⁻¹. The NBO analysis establishes that the antiperiplanar orientation of the C=P and C=O moieties results in greater stabilization of **2bA** due to increased

3.2. Energetics

The total energies $(B3LYP/6-311 + G^{**}//B3LYP/$ 6-311G^{**}), activation (ΔE_a) and reaction (ΔE_{rxn}) energies of the reactions 1-10 are given in Table 3. It has been reported that while the exothermicities of DA reactions of ethene with 1,3-butadiene and with phosphabutadienes are quite similar, ¹⁶ the ΔE_a values for the latter are smaller. Our results are quite parallel: introduction of phosphorus in the dienophilic five-membered ring lowers the activation barrier by 10.0–16.4 kcal mol⁻¹ as compared to ΔE_a for the prototype DA reaction between imidazo[2,1-b]thiazole (1) and 1,3-butadiene (4), but ΔE_{rxn} are comparable. The smaller ΔE_a of the phospha DA reaction as compared to that of the carbocyclic one is due to the weaker $C = P \pi$ bond relative to the C=C π bond.¹⁶ Furthermore, DA reactions of 2 with 1,3-butadiene are accompanied by dearomatization of the azaphosphole ring and it is therefore expected²³ that in contrast to the DA reactions of the acyclic phosphaethene with 1,3-butadiene,²² the TS in the present case will be reached later and the activation energies would be comparatively higher. The present results conform with these predictions, and the activation barriers are higher by 2-5 kcal mol⁻¹ than for the DA reaction of phosphaethene.

3.3. Stereoselectivity

The observed *endolexo* stereoselectivity can be rationalized by an analysis of the molecular orbitals (MOs) of

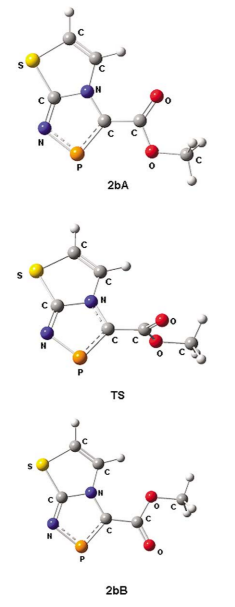


Figure 2. Optimized geometries (B3LYP/6-311G**) of the two conformers of **2b** and the corresponding TS.

Table 2. Stabilization energies E_{ij} from second-order perturbation NBO analysis of the conformers **2bA** and **2bB**

	Significant interactions	Stabilization energies E_{ij} (kcal mol ⁻¹)		
	Donor NBO _i \rightarrow acceptor NBO _j	2bA	2bB	
1	$\pi_{N1-C8} \rightarrow \pi^*_{P2-C3}$	11.03	11.01	
2	$\pi_{P2-C3} \rightarrow \pi^*_{C9-O10}$	27.21	26.56	
3	$\sigma_{N1-P2} \rightarrow \sigma^*_{S7-C8}$	13.21	13.29	
4	$LP_{S7} \rightarrow \pi^*_{C5-C6}$	20.17	20.66	
5	$LP_{S7} \rightarrow \pi^*_{N1-C8}$	28.87	28.58	
6	$LP_{N4} \rightarrow \pi^*_{C5-C6}$	30.32	30.72	
7	$LP_{N4} \rightarrow \pi^*_{P2-C3}$	33.50	32.55	
8	$LP_{N4} \rightarrow \pi^*_{N1-C8}$	51.05	50.09	
9	$LP_{N1} \rightarrow \sigma^*_{N4-C8}$	12.02	12.32	
10	$LP_{O10} \rightarrow \sigma^*_{C3-C9}$	16.46	16.61	
11	$LP_{O10} \rightarrow \sigma^*_{C9-O11}$	31.35	33.57	
12	$LP_{O11} \rightarrow \pi^*_{C9-O10}$	46.08	42.04	
13	$\pi^*_{P2-C3} \rightarrow \pi^*_{N1-C8}$	36.56	37.25	
14	$\pi^*_{P2-C3} \rightarrow \pi^*_{C9-O10}$	44.52	39.01	

Table 3. Total energies and relative energies for the stationary points corresponding to DA reactions 1-10

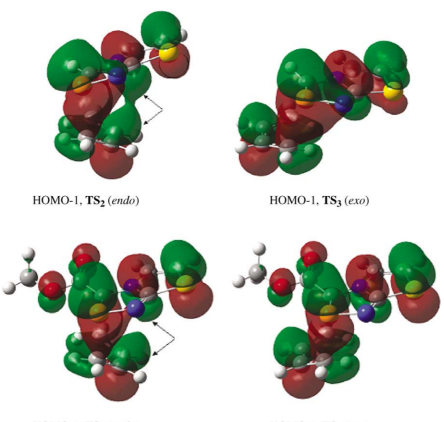
	Total energies (a.u.) ^a	Relative energy (kcal mol ^{-1}	
	Gas phase	Toluene	Gas phase	Toluene
1	-700.621240	-700.627488	0.0	0.0
2a	-1003.305121	-1003.310998	0.0	0.0
2bA	-1231.213654	-1231.219041	0.0	0.0
2bB	-1231.209447	-1231.216064	2.64	1.87
TS ^b	-1231.197345	-1231.204144	10.23	9.35
3	-1293.741951	-1293.747148	0.0	0.0
4	-155.950050	-155.951692	0.0	0.0
5	-195.250518	-195.251835	0.0	0.0
TS_1	- 856.519390	-856.526222	32.57	33.23
P_1	-856.589940	-856.596381	-11.70	-10.79
TS_2	-1159.225971	-1159.232466	18.32	18.96
P_2^{-}	-1159.278582	-1159.285251	-14.69	-14.16
TS ₃	-1159.224753	-1159.231058	19.09	19.85
P_3	-1159.276823	-1159.283354	-13.59	-12.97
TS_4	-1198.526647	-1198.532456	18.19	19.06
P_4	-1198.581192	-1198.587252	-16.03	-15.32
ГŚ ₅	-1198.524968	-1198.531146	19.25	19.88
P ₅	-1198.580282	-1198.586590	-15.46	-14.91
TS ₆	-1387.135906	-1387.142520	17.44	17.70
P ₆	-1387.177023	-1387.184485	-8.36	-8.63
ΓS_7	-1387.133563	-1387.140105	18.91	19.22
P ₇	-1387.173908	-1387.181547	-6.40	-6.78
TS ₈	-1426.438473	-1426.44444	16.13	16.59
P ₈	-1426.479680	-1426.486565	-9.73	-9.84
TS ₉	-1426.435269	-1426.441496	18.14	18.44
P ₉	-1426.478855	-1426.485911	-9.21	-9.43
TS ₁₀	-1449.655999	-1449.661508	22.59	23.43
P ₁₀	-1449.693554	-1449.699441	-0.97	-0.37

^a Energies at B3LYP/6-311++G**+ ZPE at B3LYP/6-311G** level. ^b TS ($2bA \rightarrow 2bB$).

the corresponding transition structures. It is found that in the *endo* transition structures TS_2 and TS_6 , HOMO is localized on the thiazole ring, but in the HOMO-1 molecular orbitals, besides the formation of the new P2–C9 and C3–C12 bonds, secondary molecular orbital (SMO) interactions can be detected between C10–C11 (π) and N1–C8–N4 (π) orbitals (Fig. 3). These SMO interactions are missing in the *exo* transition structures, TS₃ and TS₇, which explains the lower activation energy barrier in the former.

The regio- and stereoselectivities observed in the DA reactions of vinylboranes were evaluated theoretically and the calculated results were found to be in good agreement with the experimental ones.³³

The calculated ratios of the *endolexo* stereoisomers of the DA cycloadducts of **2** are given in Table 4. Experimentally, 3-alkoxycarbonyl substituted thiazolo[3,2-d][1,4,2]diazaphosphole shows complete *endo* stereoselectivity.¹⁴ It can be seen that the computed values of the *endolexo* stereoisomers for the 3-methoxycarbonyl derivative **2b** agree nicely with the experimental values (Table 4). For the unsubstituted reactant **2a**, where experimental stereoselectivity results are not reported so far, computations predict approximately 80% *endo* stereoselectivity. Furthermore, theoretical calculations of the solvent effect indicate that the stereoselectivity in toluene is quite similar to that in the gaseous phase.



HOMO-1, TS_6 (endo)

HOMO-1, TS₇ (exo)

Figure 3. Secondary molecular orbital interactions in *endo* transition structures for DA reactions of thiazolo[3,2-*d*][1,4,2]diazaphospholes (2) with 1,3-butadiene.

3.4. Regioselectivity

We have earlier reported the experimental results of the regioselectivity in the DA reactions of thiazolo-[3,2-d][1,4,2]diazaphospholes with isoprene.¹⁴ 3-Alkoxy-carbonyl substituted dienophiles produced the *meta* regioisomer (P/Me, 1:3) exclusively while an unsubstituted dienophile, [1,4,2]diazaphospholo[5,4-b]benzothiazole gave *meta* (P/Me, 1:3) and *para* (P/Me, 1:4) regioisomers in a 2:1 ratio. Similar regioselectivity was observed in the DA reaction of 2-acetyl-2*H*-1,2,3-diazaphosphole with isoprene where only the *meta* regioproduct was formed.³⁴ To rationalize these results we have computed the DA reactions of thiazolo[3,2-d][1,4,2]diazaphospholes (**2**) with isoprene.

As for *endolexo* stereoselectivity, the SMO interactions appear to determine *metalpara* regioselectivity also.

Table 4. Experimental and calculated²⁴ ratios of the stereo- and regioisomers for the DA reactions between diazaphospholes **2** and dienes **4**, **5**

	Percentages of stereoisomers (endo:exo)			Percentages of regioisomers (<i>meta:para</i>)			
	Calcd		Exp.	Calcd		Exp.	
	Gas phase	Toluene		Gas phase	Toluene		
2a 2b	79:21 92:08	82:18 93:07	100:0	86:14 97:03	80:20 95:05	66:34 ^a 100:0	

^a For DA reaction of [1,4,2]diazaphospholo[5,4-b]benzothiazole.

On close inspection of the HOMO-1 molecular orbital of the transition structure TS₈ resulting from meta approach of **2b** and isoprene, SMO interactions can be detected between the orbital having a lone pair on sulfur and a p orbital on the methyl carbon (Fig. 4). These interactions are missing in the transition structure TS₉ resulting from *para* approach. These SMO interactions account for the exclusive (100%) meta regioselectivity. The unsubstituted dienophile 2a is structurally similar to [1,4,2]diazaphospholo[5,4-b]benzothiazole, which on reaction with isoprene gives both meta and *para* regioisomers in a 2:1 ratio.¹⁴ It is interesting to find that in the HOMO-1 molecular orbitals of the TS_4 and TS_5 (Fig. 4) corresponding to the *meta* and *para* approaches, respectively of 2a and isoprene, SMO interactions are possible in both due to the proximity of lone pair orbital on sulfur and p orbital on the methyl carbon, although the magnitude of this interaction in TS₅ would be much smaller than in TS₄ due to greater distance between the interacting orbitals in the former. This explains the reduced regioselectivity and formation of both regioisomers in 2:1 ratio.

It may be noted that NICS values of the transition structures corresponding to the preferred *meta* approach of isoprene are slightly more negative than for the transition structures resulting from the *para* approach, indicating greater aromatic character of the former (Fig. 1). These differences are, however, very small. Furthermore, it has been pointed out that although transition state aromaticity is important in the DA reaction, it does not determine regioselectivity.³⁵

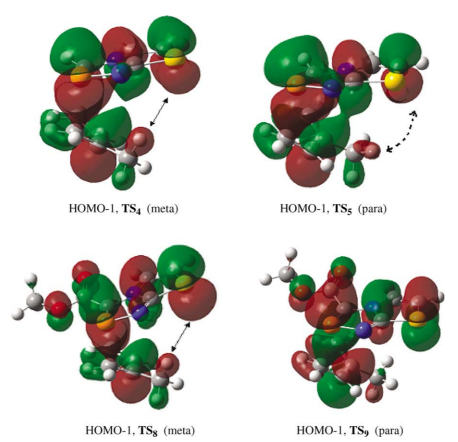


Figure 4. Secondary molecular orbital interactions in the transition structures for DA reactions of thiazolo[3,2-d][1,4,2]diazaphospholes (2) with isoprene.

The calculated ratios of the *meta/para* regioisomers are given in Table 4 and it can be seen that, as for *endo* stereoselectivity, these values for the 3-methoxycarbonyl substituted reactant, **2b** agree nicely with the experimental results. The computations for the 3-unsubstituted dienophile, **2a**, however, slightly overestimate the regioselectivity. The calculations taking into account the solvent effect indicate that toluene does not influence regioselectivity also.

3.5. DA reaction of 3-ethoxycarbonyl-1-methyl-1,3-aza-phospholo[5,1-*b*]benzothiazole

It has been reported that in contrast to thiazolo[3,2-d]-[1,4,2]diazaphospholes, DA reaction of 3-ethoxycarbonyl-1-methyl-1,3-azaphospholo[5,1-b]benzothiazole occurs only at elevated temperature in the presence of an oxidizing agent like O_2 , S_8 or Se_n .¹⁴ To investigate the cause of its lower reactivity and role of the oxidizing agent, we have carried out computational calculations (B3LYP/6-311G**) of a model system of the DA reaction of 3-ethoxycarbonyl-1-methyl-1,3-azaphospholo[5,1-b]thiazole (3) with 1,3-butadiene. The transition structure TS_{10} is shown in Figure 3. As indicated by NICS value, it also occurs via a pericyclic mechanism, but the activation energy barrier ($\Delta E_a =$ 22.59 kcal mol⁻¹) in this case is much higher than for the DA reactions of the thiazolo[3,2-d][1,4,2]diazaphospholes. It therefore appears likely that a reversible DA reaction sets in between 3-ethoxycarbonyl-1-methyl-1,3azaphospholo[5,1-b]benzothiazole and 1,3-butadiene,

which is pushed in the forward direction on raising the temperature. Furthermore, the oxidizing agent helps in pushing the reaction in the forward direction by oxidizing σ^3 -P of the initially formed cycloadduct in low concentration.

4. Conclusion

The DA reactions of azaphospholes with 1,3-dienes occur via a pericyclic mechanism. The aromatic character of the transition structures, which are asynchronous, is confirmed by high negative NICS values. The activation energy barriers ($\Delta E_a = 16-19$ kcal mol⁻¹) are much lower than that for the DA reaction of the non-phosphorus analogue, imidazo[2,1-b]thiazole. The experimentally observed endo stereoselectivity can be rationalized on the basis of secondary molecular orbital interactions in the HOMO-1 molecular orbitals of TS_2 and TS_6 for the DA reactions of 2 with 1,3-butadiene. The experimentally reported regioselectivity can be rationalized on the basis of secondary molecular orbital interactions in the HOMO-1 MOs of TS₄ and TS₈ involved in the DA reactions of 2. In the case of 2a, weak SMO interactions in the HOMO-1 of TS₅ are responsible for the formation of the para regioisomer also as the minor product besides the main meta regioproduct via TS₄.

The calculated ratios of the *endolexo* stereoisomers as well as those of the *metalpara* regioisomers agree well with the experimental values. The activation energy barrier for the DA reaction of 3-ethoxycarbonyl-1-methyl-1,3azaphospholo[5,1-*b*]thiazole is found to be comparatively higher, which suggests a reversible DA reaction and rationalizes its occurrence at elevated temperature in the presence of an oxidizing agent. The computations with solvent effect indicate that toluene does not influence *endol exo* stereoselectivity or *meta/para* regioselectivity vis-a-vis gaseous phase.

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Regioselectivity in arene-catalyzed reductive lithiation of acetals of chlorobenzaldehydes

Ugo Azzena,^{a,*} Giovanna Dettori,^a Giuseppe Sforazzini,^a Miguel Yus^{b,*} and Francisco Foubelo^b

^aDipartimento di Chimica, Università di Sassari, via Vienna 2, I-07100 Sassari, Italy ^bDepartamento de Quimica Orgánica, Facultad de Ciencias, Universidad de Alicante, Apdo. 99, E-03080 Alicante, Spain

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Abstract—The regioselectivity of arene-catalyzed reductive lithiation of acetals of chlorobenzaldehydes strongly depends on the form of lithium metal employed as a reducing agent. According to previous findings, naphthalene catalyzed reductions run in the presence of lithium powder (high Na content) led to competitive metalations of both aromatic carbon–chlorine and benzylic carbon–oxygen bonds. At variance with these results, naphthalene catalyzed reductions run in the presence of lithium wire (either high or low Na content) led to highly regioselective metalation of aromatic carbon–chlorine bonds. These results disclose new possibilities of selective applications of arene-catalyzed reductive lithiation reactions.

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1. Introduction

Arene-catalyzed reductive lithiation of carbon-heteroatom bonds is a powerful approach to the generation of a wide array of functionalized and non-functionalized organolithium compounds.^{1,2} Application of this procedure to arylmethyl alkyl ethers results in highly regioselective cleavage of benzylic carbon–oxygen bonds, thus affording access to a wide array of arylalkyl organolithium derivatives, some of which are not easily accessible by other methodologies.^{3–5} Interestingly, reductive lithiation of aromatic ethylene,⁶ dimethyl⁷ and bis-(2-methoxyethyl)⁸ acetals is a useful approach to the generation of α -alkoxysubstituted benzyllithium derivatives, a procedure successfully extended to arenetricarbonylchromium acetals.⁹

The last finding highlights a potential limitation in applying the reductive metalation procedure to the generation of aromatic organolithium derivatives bearing a masked carbonyl group.

Indeed, Yus et al.¹⁰ reported that reductions of 2-(chlorophenyl)-1,3-dioxolanes, run in the presence of lithium powder and a catalytic amount of naphthalene ($C_{10}H_8$), occurs with regioselective cleavage of the aromatic carbon–chlorine bond.

However, these reactions have to be carried out under Barbier conditions, to avoid decomposition of intermediate organometals, due to competitive cleavage of benzylic carbon–oxygen bonds. Besides affording the desired products in moderate yields, this approach limits the number of electrophiles that can be added to carbanionic intermediates to aldehydes and ketones. Interestingly, an even lower selectivity was observed employing 4,4'-di-*tert*butylbiphenyl (DTBB) as a catalyst.¹⁰

As an alternative approach, Azzena et al. investigated the reductive metalation of 1,3-dimethyl-2-(4-chlorophenyl) imidazolidine,¹¹ taking advantage of the relatively high stability of carbon–nitrogen bonds to reductive metalation.⁴ This procedure allows the generation of stable solutions of a synthetic equivalent of 4-formylphenyllithium, efficiently trapped with a variety of electrophilic reagents; however, resulting 2-(4-substituted)aryl-1,3-dimethyloxazolidines are not completely stable to purification by flash chromatography, thus precluding the recovery of protected benzal-dehydes as reaction products.¹¹

Looking for a more efficient approach to the generation of synthetic analogues of formylphenyllithiums, we investigated the effect of different types of lithium metal (lithium wire vs lithium dispersion,¹² as well as lithium with a high sodium content vs lithium with a low sodium content¹³) on the selectivity of the reductive metalation of several acetals of chlorobenzaldehydes, and wish now to report that the regioselectivity of this reaction strongly depends on the form of lithium metal employed as a reducing agent.

Keywords: Metalation; Organometals; Reduction; Regioselectivity.

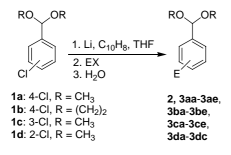
^{*} Corresponding authors. Tel.: + 39 079229549; fax: + 39 079229559 (U.A.); tel.: + 34 965903548; fax: + 34 965903549 (M.Y.); e-mail addresses: ugo@uniss.it; yus@ua.es

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2. Results and discussion

Dimethyl acetals **1a**, **1c** and **1d** were prepared by heating the corresponding aldehydes in MeOH at reflux in the presence of $HC(OCH_3)_3$ and a catalytic amount of NH_4Cl . Ethylene acetal **1b** was prepared by heating 4-chlorobenzaldehyde with 1,2-ethanediol in dry toluene at reflux, in the presence of a catalytic amount of *p*-toluenesulfonic acid.

Reductive metalations were carried out under Ar, with an excess of Li metal, either in the form of Li wire (0.32 mm, low or high Na content) or Li powder (high Na content). All reductions were run in the presence of a catalytic amount of naphthalene ($C_{10}H_8$), at temperature ranging from -40 to -80 °C, followed by addition of an electrophilic reagent and, finally, aqueous work up (Scheme 1).



Scheme 1. Reductive lithiation of acetals 1a–d, and reaction with electrophiles. 2: E=H, $R=CH_3$; 3aa: $E=4-CH_3$, $R=CH_3$; 3ab: E=4-PhCHOH, $R=CH_3$; 3ac: $E=4-(CH_3)_3CCHOH$, $R=CH_3$; 3ad: E=4-PhCOHAr, $R=CH_3$; 3ae: E=4-CHO, $R=CH_3$; 3ba: E=H, $R=(CH_2)_2$; 3bb: $E=4-CH_3$, $R=(CH_2)_2$; 3bc: $E=4-(CH_2)_5COH$, $R=(CH_2)_2$; 3bd: $E=4-Et_2COH$, $R=(CH_2)_2$; 3be: E=4-PhCHOH, $R=(CH_2)_2$; 3bd: E=3-PhCHOH, $R=(CH_2)_2$; 3bd: E=3-PhCHOH, $R=CH_3$; 3ce: $E=3-(CH_3)_2CH]_2$. COH, $R=CH_3$; 3cd: E=3-CHO, $R=CH_3$; 3ce: E=3-ArCHOH, $R=CH_3$; 3da: $E=2-CH_3$, $R=CH_3$; 3db: E=2-PhCHOH, $R=CH_3$; 3dc: $E=2-[(CH_3)_2CH]_2COH$, $R=CH_3$.

Reductive lithiation and reaction with electrophiles of dimethyl acetal **1a**, taken as a model compound, was investigated in detail.

Reduction with 6 equiv of Li wire (low Na content) and 5 mol% of $C_{10}H_8$ led, after 4.5 h stirring at -40 °C and aqueous quenching, to the recovery of benzaldehyde dimethyl acetal, **2**, as the only reaction product, with no evidence of formation of products of cleavage of the benzylic carbon–oxygen bonds (Table 1, entry 1).

Intermediate formation of a protected, stable, 4-formylsubstituted aryllithium was evidenced quenching the reduction mixture with CH_3I : under these conditions, we recovered a reaction mixture containing, besides a relatively small amount of dimethyl acetal **2**, the dimethyl acetal of 4-methylbenzaldehyde, **3aa**, in 91% yield, as determined by ¹H NMR spectroscopy of the crude reaction mixture (Table 1, entry 2).

Similar results were obtained reacting dimethyl acetal **1a** with 6 equiv of Li wire containing a high percentage of Na, in the presence of 5 mol% of $C_{10}H_8$ (Table 1, entries 3 and 4).

Table 1. Reductive lithiation and reaction with electrophiles of dimethyl acetal of 4-chlorobenzaldehyde, $1a^a$

Entry	EX $(T (^{\circ}C))^{b}$	Product, E=	Yield (%) ^c
1	$H_2O(-40)$	2 , H	>95 ^d
2	$CH_{3}I(-40)$	3aa , 4-CH ₃	91 ^d
3	$H_2O(-40)$	2 , H	>95 ^{d,e}
4	$CH_{3}I(-40)$	3aa, 4-CH ₃	84 ^{d,e}
5	PhCHO (-40)	3ab, 4-PhCHOH	63
6	<i>t</i> -BuCHO (-40)	3ac, 4-t-BuCHOH	71
7	PhCOCl (-80)	3ad , 4-PhCOHAr ^f	58
8	$HCOOCH_3 (-80)^g$	3ae, 4-CHO	55

^a All reactions were run at -40 °C, during 4.5 h, in the presence of 6 equiv of Li wire (low Na content, unless otherwise indicated) and a catalytic amount of C₁₀H₈ (5 mol%).

^b Temperature of addition of EX (2 equiv, if not otherwise indicated).

^c Determined on isolated products, if not otherwise indicated.

^d Determined by ¹H NMR analysis of crude reaction mixture.

^e High Na content Li metal was employed.

^f EX (0.45 equiv) were employed; $Ar = 4-(CH_3O)_2CHC_6H_4$.

g Inverse addition.

It is worth noting that using a higher amount of $C_{10}H_8$ (10 mol%) did not significantly affect these results, whatever kind of Li wire was employed as a reducing agent (not reported in Table 1).

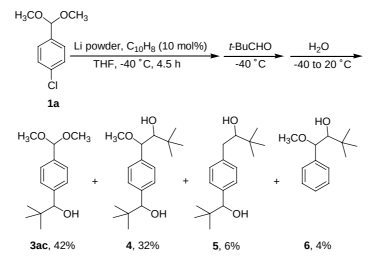
Trapping the intermediate organometal, generated as reported in entries 1 and 2, with PhCHO, *t*-BuCHO, and PhCOCl allowed the synthesis of the corresponding carbinols **3ab**, **3ac** and **3ad**, respectively, in satisfactory yields (Table 1, entries 5–7). Finally, reaction of the same intermediate with HCOOCH₃, under inverse addition reaction conditions, led to the recovery of the monoprotected terephthalaldehyde **3ae**, in 55% yield (Table 1, entry 8).

Different results were obtained performing the reductive lithiation of dimethyl acetal **1a** with Li powder, in the presence of 10 mol% of $C_{10}H_8$, at -40 °C. Indeed, quenching this reduction mixture with *t*-BuCHO, followed by aqueous work up, afforded a complex reaction mixture (Scheme 2).

Chromatographic fractionation, followed by tandem GC/ MS and ¹H NMR analysis of the different fractions, allowed us to identify, besides the expected carbinol **3ac** (42%), significative amounts of products derived from competitive reductive lithiation of the benzylic carbon–oxygen bonds, that is, alcohols **4–6**. It is interesting to observe that this result is in good qualitative agreement with what is already reported for the arene-catalyzed reductive metalation of aromatic ethylene acetals with lithium powder,¹⁰ thus underlining the major role played by the form of the metal on the regioselectivity of these reactions.

To shed more light on the role of the different forms of the metal on regioselectivity, we investigated the reduction with Li wire of the ethylene acetal of 4-chlorobenzaldehyde, **1b**. To get a better comparison with literature data,¹⁰ we carried out these reactions with 13 equiv of Li wire, in the presence of 10 mol% of $C_{10}H_8$, at -80 °C (Table 2).

Contrary to what was observed in the presence of lithium powder,¹⁰ results reported in Table 2 show that employment of Li wire (low Na content) allowed highly regioselective



Scheme 2. Reductive metalation of dimethyl acetal 1a with Li powder and 10 mol% of C₁₀H₈, followed by reaction with *t*-BuCHO and aqueous work up.

Table 2. Reductive lithiation and reaction with electrophiles of ethylene acetal of 4-chlorobenzaldehyde, 1b^a

Ent- ry	EX $(T (^{\circ}C))^{b}$	Product, E=	Yield (%) ^b
1	H ₂ O	3ba , H	>95 ^c
2	CH ₃ I	3bb , 4-CH ₃	93 ^c
3	H_2O	3ba , H	86 ^{d,e}
4	CH ₃ I	3bb , 4-CH ₃	84 ^{c,d}
5	(CH ₂) ₅ CO	3bc , 4-(CH ₂) ₅ COH	61
6	Et ₂ CO	3bd, 4-Et ₂ COH	67
7	PhCHO	3be, 4-PhCHOH	67

^a All reactions were run at -80 °C, during 3 h, in the presence of 13 equiv of Li wire (low Na content, unless otherwise indicated) and a catalytic amount of C10H8 (10 mol%).

^b Determined on isolated products, if not otherwise indicated.

^c Determined by ¹H NMR analysis of crude reaction mixture.

^d High Na content Li metal was employed.

e Ca. 5% of 2-benzyloxyethanol was also recovered.

cleavage of the aromatic carbon-chlorine bond (Table 2, entry 1), via intermediate formation of a stable aromatic organometal, as evidenced by quenching the reduction mixture with CH₃I (Table 2, entry 2).

Similar results were obtained employing Li wire with a high Na content as the reducing agent, although a minor amount (\leq 5%) of 2-benzyloxyethanol, 7, was recovered when quenching the reduction mixture with H₂O (Table 2, entry 3). It is worth noting that alcohol 7 is the product of cleavage of both the aromatic carbonchlorine and a benzylic carbon-oxygen bond. However, no product of cleavage of a benzylic carbon-oxygen bond was evidenced quenching a similar reduction mixture with CH_3I (Table 2, entry 4).

Satisfactory results were obtained trapping the intermediate organometal, generated as reported in entries 1 and 2, with enolizable and non-enolizable carbonyl compounds (Table 2, entries 5–7). Interestingly, besides showing higher selectivity, the two-pot procedure led to an improvement of those results obtained under Barbier-type reaction conditions¹⁰ (Table 2, entries 5 and 6).

Finally, to get a wider picture on the versatility of the proposed methodology, we investigated the reductive lithiation of *meta-* and *ortho-*chloro-substituted dimethyl acetals of benzaldehydes, that is, compounds 1c and 1d. These reactions were carried out in the presence of 6 equiv of Li wire (low Na content), and in the presence of 5 mol% of $C_{10}H_8$ during 4.5 h at -40 °C. Selected results are reported in Table 3.

Table 3. Reductive lithiation and reaction with electrophiles of acetals 1c and 1da

Entry	Sub- strate	EX $(T \circ C)^{b}$	Product, E=	Yield (%) ^c
1 2 3 4 5 6 7 8 9 10	1c 1c 1c 1c 1c 1c 1d 1d 1d	$\begin{array}{l} H_2O\ (-40)\\ CH_3I\ (-40)\\ PhCHO\ (-80)\\ I(CH_3)_2CH]_2CO\ (-80)\\ HCOOCH_3\ (-80)^e\\ HCOOCH_3\ (-80)^f\\ H_2O\ (-40)\\ CH_3I\ (-40)\\ PhCHO\ (-80)\\ I(CH_3)_2CH]_2CO\ (-80) \end{array}$	2, H 3ca, 3-CH ₃ 3cb, 3-PhCHOH 3cc, 3-[(CH ₃) ₂ CH] ₂ COH 3cd, 3-CHO 3ce, 3-ArCHOH ^g 2, H 3da, 2-CH ₃ 3db, 2-PhCHOH 3dc, 2-[(CH ₃) ₂ CH] ₂ COH	$> 95^{d}$ 82^{d} 55 45 56 62 $> 95^{d}$ 83^{d} 56 80^{d} $(52)^{h}$

All reactions were run at -40 °C, during 4.5 h, in the presence of 6 equiv of Li (low Na content) and a catalytic amount of C₁₀H₈ (5 mol%). Temperature of addition of EX (2 equiv, if not otherwise indicated).

Determined on isolated products, if not otherwise indicated. Determined by ¹H NMR analysis of crude reaction mixture.

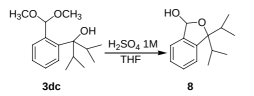
Inverse addition

EX (0.45 equiv) were employed. $Ar = 3 - (CH_3O)_2 CHC_6H_4.$

Determined on the corresponding cyclic hemiacetal, 8, obtained by acidic hydrolysis of alcohol 3dc.

Under the above mentioned reaction conditions, cleavage of meta-substituted acetal 1c afforded, after aqueous work up, the dimethyl acetal of benzaldehyde, 2, as the only reaction product (Table 3, entry 1). Furthermore, intermediate formation of 3-dimethoxymethylphenyllithium was evidenced quenching the reduction mixture with CH₃I (Table 3, entry 2), with carbonyl compounds (Table 3, entries 3 and 4), and with HCOOCH₃, either under inverse or conventional addition of the electrophile to the reduction mixture (Table 3, entries 5 and 6, respectively).

Comparable results were obtained in the reductive lithiation of the *ortho*-derivative **1d**. Indeed, under identical reaction conditions, its reductive lithiation allowed the generation of stable solutions of 2-dimethoxymethylphenyllithium; reaction of this intermediate with H₂O, CH₃I and PhCHO, afforded the expected derivatives **2**, **3da–3db** in good to satisfactory yields (Table 3, entries 7–9). Finally, reaction of the same organometals with $[(CH_3)_2CH]_2CO$ led to the formation of alcohol **3dc** (Table 3, entry 10); although it was not possible to purify the latter by flash chromatography, its structure was established by acidic hydrolysis to hemiacetal **8** (Scheme 3).



Scheme 3. Synthesis of hemiacetal 8, by acidic hydrolysis of alcohol 3dc.

3. Conclusions

Our results clearly show that regioselective reductive lithiation of acetals of chlorobenzaldeydes strongly depends on the form of the lithium metal. Indeed, the actual form of a metal is known to affect, to various degree, its reactivity, and the higher reactivity of powders towards massive metals is well documented.¹⁴

Reductions carried out with Li wire (either with a high or low Na content) led to the generation of stable solutions of *ortho-*, *meta-* or *para-*substituted protected formylphenyllithium, independent from the nature of the protective group (dimethyl or ethylene acetal), via a highly regioselective cleavage of the aromatic carbon–chlorine bond. Accordingly, it is possible to trap these organometals with a range of electrophilic reagents, as well as to enhance the versatility of this methodology by applying an inverse addition technique.

A relatively lower regioselectivity was observed in a single experiment, that is, in the reaction of ethylene acetal **1b** with Li wire containing a high Na content, followed by aqueous work up (Table 2, entry 3). Under these conditions, competitive cleavage of the benzylic carbon–oxygen bond, although limited to a very low percentage, was also observed.

Completely different results were obtained employing lithium powder (high Na content) as a reducing agent. Indeed, under these conditions, dimethyl acetal **1a** underwent competitive benzylic carbon–oxygen bond(s) cleavage leading, as a consequence, to the formation of significant amounts of disubstituted reaction products, as already reported for several aromatic ethylene acetals.¹⁰

In conclusion, we have shown that aromatic acetals are effective protective groups in the reductive lithiation of chlorine-substituted aromatic carbonyls, provided that lithium wire (in the presence of $C_{10}H_8$) is employed as a reducing agent.

In a broader sense, however, our results underline the very powerful reducing power of lithium powder (either in the presence of $C_{10}H_8$ or DTBB).

4. Experimental

4.1. General

Boiling and melting points are uncorrected; the air bath temperature on bulb-to-bulb distillation are given as boiling points. Commercially available Li wire (Ø 3.2 mm) was 99.9% purity (low Na content) or 99% purity (high Na content). Li powder (high Na content) was prepared as described in Ref. 12. Other starting materials were of the highest commercial quality and were purified by distillation or recrystallization immediately before use. THF was distilled from Na/K alloy under N₂ immediately before use. ¹H NMR spectra were recorded at 300 MHz and ¹³C NMR spectra were recorded at 75 MHz with a Bruker AC-300, or with a Varian VXR 300, in CDCl₃ with SiMe₄ as internal standard. CDCl3 for recording spectra of acetals was stored over K₂CO₃ in the refrigerator. IR spectra were measured with a Nicolet Impact 400 D-FT Spectrometer, or with a Jasco FT/IR-480 Plus. LRMS and HRMS were measured with a Shimadzu GC/HS QP-5000 and with a Finingan MAT95 S spectrometers, respectively. Elemental analysis were performed by the Microanalytical Laboratory of the Dipartimento di Chimica, Università di Sassari. Flash chromatography was performed on silica gel 60 (40-63 µm), and TLC analyses on silica gel precoated plastic sheets (0.20 mm).

4.2. Starting materials

Dimethyl acetals **1a** (88% yield),¹⁵ **1c** (81% yield),¹⁶ and **1d** (80% yield),¹⁶ were prepared by heating the corresponding chlorobenzaldehyde (0.07 mol, 10 g) in a mixture of 50 mL of CH₃OH and 20 mL of HC(OCH₃)₃ at reflux overnight, in the presence of a catalytic amount of NH₄Cl (0.1 g, 1.9 mmol). Reaction mixtures were chilled to 0 °C, made basic by slow dropwise addition of Et₃N (2 mL), evaporated, then diluted with Et₂O (20 mL) and NaHCO₃ (20 mL) and worked up as usual.

Ethylene acetal **1b** $(72\% \text{ yield})^{10}$ was prepared distilling with fractionation a solution of 2-chlorobenzaldehyde (0.036 mol, 5 g) and 1,2-ethanediol (2.46 g, 0.04 mol) in 50 mL of dry toluene in the presence of a catalytic amount of *p*-toluenesulfonic acid (0.53 mmol, 100 mg), until the azeotrope toluene–H₂O was distilled away, followed by work up as described above.

All acetals were purified by vacuum distillation and characterized by comparison with literature data.

4.3. Reductive cleavage of acetals, and reactions with electrophiles. General procedure

Li (6–13 equiv) was placed under Ar in a 50 mL twonecked flask equipped with reflux condenser and magnetic stirrer, and suspended in dry THF (5 mL). A catalytic amount of $C_{10}H_8$ (5–10 mol%) was added to the suspended metal. When using Li wire, each metal piece was cut into 2-3 smaller pieces with a spatula. The resulting mixture was stirred at rt until a dark green colour appeared. The mixture was chilled to the reported temperature (Tables 1-3) and a solution of the appropriate acetal (2.7 mmol) dissolved in THF (2.5 mL) was added dropwise. Reaction mixtures were stirred for the reported time (Tables 1-3), and a solution of the appropriate electrophile (2 equiv), dissolved in THF (2 mL), was added dropwise. After stirring at the reported temperature until an evident chromatic variation occurred, the mixture was quenched by slow dropwise addition of H_2O (5 mL) dissolved in THF (5 mL) (caution!), the cold bath removed, and the resulting mixture extracted with Et₂O (3×10 mL). Organic phases were collected, washed with saturated NaHCO₃ (20 mL), dried (K_2CO_3) and the solvent evaporated.

Compounds 2, 3aa,¹⁵ 3ba, 3bb,¹⁷ 3bc,¹⁰ 3bd,¹⁰ 3ca,¹⁵ 3cd,¹⁸ 3da,¹² 3db,¹⁹ 5^{20} and 7 were purified by flash chromatography (petroleum ether/AcOEt/Et₃N) and characterized by comparison with commercially available samples (compounds 2, 3ba and 7) or literature data.

Other products were purified and characterized as follows.

4.3.1. (4-Dimethoxymethylphenyl)phenylmethanol (3ab). Purified by flash chromatography (petroleum ether/AcOEt/Et₃N=8:2:1), colourless oil; [Found: C, 74.2; H, 7.2. C₁₆H₁₈O₃ requires C, 74.39; H, 7.02]; $R_{\rm f}$ 0.64 (petroleum ether/AcOEt/Et₃N=8:2:1); $\nu_{\rm max}$ (liquid film) 3467 cm⁻¹; $\delta_{\rm H}$ 7.45–7.27 (9H, m, 9×ArH), 5.86 (1H, s, OCHO), 5.36 (1H, s, OCHAr), 3.12 (6H, s, 2×CH₃); $\delta_{\rm C}$ 144.0, 143.7, 137.3, 128.5, 127.6, 126.8, 126.5, 126.4, 103.0, 76.0, 52.7.

4.3.2. 1-(4-Dimethoxymethylphenyl)-2,2-dimethylpropan-1-ol (3ac). Purified by flash chromatography (petroleum ether/AcOEt/Et₃N=8:2.5:1), colourless oil; [Found: C, 70.4; H, 9.5. $C_{14}H_{22}O_3$ requires C, 70.56; H, 9.30]; R_f 0.44 (petroleum ether/AcOEt/Et₃N=8:2.5:1); ν_{max} (liquid film) 3465 cm⁻¹; δ_H 7.43–7.37 (2H, m, 2×ArH), 7.35–7.29 (2H, m, 2×ArH), 5.38 (1H, s, CHAr), 4.41 (1H, d, J=3.0 Hz, OCHO), 3.33 (6H, s, 2×CH₃O), 1.87 (1H, d, J=3.0 Hz, OH), 0.92 (9H, s, 3×CH₃); δ_C 143.4, 137.2, 127.4, 126.8, 103.2, 82.6, 52.6, 35.8, 25.8.

4.3.3. Bis-(4-dimethoxymethylphenyl)phenylmethanol (3ad). Purified by flash chromatography (petroleum ether/AcOEt/Et₃N=8:2:1), colourless oil; [Found: C, 73.4; H, 6.7. $C_{25}H_{28}O_5$ requires C, 73.51; H, 6.91]; R_f 0.42 (petroleum ether/AcOEt/Et₃N=8:2:1); ν_{max} (liquid film) 3426 cm⁻¹; δ_H 7.42–7.36 (4H, m, 4×ArH), 7.32–7.23 (9H, m, 9×ArH), 5.38 (2H, s, 2×OCHO), 3.34 (12H, s, 4× CH₃); δ_C 147.0, 146.7, 137.0, 127.9, 127.9, 127.8, 127.3, 126.2, 103.1, 81.7, 52.9.

4.3.4. 4-Dimethoxymethylbenzaldehyde (3ae). Purified by flash chromatography (petroleum ether/Et₃N = 10:1), colourless oil; [Found: C, 66.5; H, 6.9. $C_{10}H_{12}O_3$ requires C, 66.65; H, 6.71]; R_f 0.48 (petroleum ether/Et₃N = 10:1); ν_{max} (liquid film) 1703 cm⁻¹; δ_H 10.04 (1H, s, CHO), 7.93–7.87 (1H, m, ArH), 7.67–7.61 (2H, m, 2×ArH), 5.46 (1H, s, CHAr), 3.34 (6H, s, 2×CH₃O); δ_C 192.0, 144.5, 136.3, 129.6, 127.4, 102.1, 52.7.

4.3.5. [4-(1,3)Dioxolan-2-ylphenyl]phenylmethanol (3be). Purified by flash chromatography (petroleum ether/AcOEt/Et₃N=5:5:1), colourless oil, which solidifies upon standing; [Found: C, 75.1; H, 6.5. C₁₆H₁₆O₃ requires C, 74.98; H, 6.29]; $R_{\rm f}$ 0.58 (petroleum ether/AcOEt/Et₃N=5:5:1); $\nu_{\rm max}$ (Nujol) 3401 cm⁻¹; $\delta_{\rm H}$ 7.47–7.25 (m, 9H, 9×ArH), 5.85 (br s, 1H, CH), 5.79 (s, 1H, CH), 4.15–3.99 (m, 4H, 2×CH₂), 2.23 (1H, br s, OH); $\delta_{\rm C}$ 144.8, 143.6, 137.0, 128.4, 127.6, 126.6, 126.5, 126.5, 103.5, 75.9, 65.2.

4.3.6. (3-Dimethoxymethylphenyl)phenylmethanol (3cb). Purified by flash chromatography (petroleum ether/ AcOEt/Et₃N=8:2.5:1.5), colourless oil; [Found: C, 74.2; H, 7.3. C₁₆H₁₈O₃ requires C, 74.39; H, 7.02]; $R_{\rm f}$ 0.46 (petroleum ether/AcOEt/Et₃N=8:2.5:1.5); $\nu_{\rm max}$ (liquid film) 3422 cm⁻¹; $\delta_{\rm H}$ 7.52–7.48 (1H, m, ArH), 7.40–7.26 (8H, m, 8×ArH), 5.85 (1H, s, CHOH), 5.37 (1H, s, CHAr), 3.30 (6H, s, 2×CH₃); $\delta_{\rm C}$ 143.8, 143.7, 138.3, 128.5, 128.4, 127.6, 126.6, 126.5, 125.9, 124.8, 103.0, 76.2, 52.7.

4.3.7. 3-(3-Dimethoxymethylphenyl)-2,4-dimethylpentan-3-ol (**3cc**). Purified by flash chromatography (petroleum ether/AcOEt/Et₃N = 10:1), colourless oil; [Found: C, 71.9; H, 10.1. $C_{16}H_{26}O_3$ requires C, 72.14; H, 9.84]; R_f 0.32 (petroleum ether/Et₃N = 10:1); ν_{max} (liquid film) 3502 cm⁻¹; δ_H 7.48–7.43 (1H, s, ArH), 7.37–7.29 (3H, m, 3×ArH), 5.41 (1H, s, CHAr), 3.32 (6H, s, 2×CH₃O), 2.32 (2H, hept, J=6.9 Hz, 2×CH), 1.55 (1H, s, OH), 0.85–0.80 (6H, d, J=6 Hz, 2×CH₃), 0.65–0.80 (6H, d, J=6 Hz, 2×CH₃), 81.0, 52.6, 33.8, 17.5, 16.5.

4.3.8. Bis-(3-dimethoxymethylphenyl)methanol (3ce). Purified by flash chromatography (petroleum ether/ $AcOEt/Et_3N=9:1:1$), colourless oil; [Found: C, 68.5; H, 7.4. $C_{19}H_{24}O_5$ requires C, 68.66; H, 7.28]; R_f 0.18 (petroleum ether/AcOEt/Et_3N=9:1:1); ν_{max} (liquid film) 3425 cm⁻¹; δ_H 7.51–7.47 (2H, m, 2×ArH), 7.38–7.32 (6H, m, 6×ArH), 5.88 (1H, s, CH), 5.37 (2H, s, 2× CHAr), 3.31 (12H, s, 4×CH₃O); δ_C 143.8, 138.3, 128.4, 126.7, 126.0, 125.0, 103.0, 76.2, 52.7.

4.3.9. 1-[4-(1-Hydroxy-2,2-dimethylpropyl)phenyl]-1methoxy-3,3-dimethylbutan-2-ol (4). Diastereomeric mixture, purified by column chromatography (hexane/ AcOEt=20:1), colourless oil; first diastereomer, $R_{\rm f}$ 0.20 (hexane/AcOEt=5:1); second diastereomer, $R_{\rm f}$ 0.19 (hexane/AcOEt=5:1); $\nu_{\rm max}$ (liquid film) 3474, 3048, 2953, 2908, 2868, 1396, 1361, 1270, 1105, 1065, 1010, 745 cm⁻¹; first diastereomer, $\delta_{\rm H}$ 7.32–7.26 (4H, m, ArH), 4.41 (1H, s, ArCHOH), 4.17 (1H, d, J=3.9 Hz, CHOCH₃), 3.35 (1H, m, CHOH), 3.22 (3H, s, OCH₃), 2.91 (1H, d, J=5.7 Hz, OH), 0.92 (9H, s, 3×CH₃), 0.89 (9H, s, 3×CH₃); second diastereomer, $\delta_{\rm H}$ 7.32–7.24 (4H, m, ArH), 4.41 (1H, s, ArCHOH), 4.13 (1H, d, *J*=6.0 Hz, CHOCH₃), 3.52 (1H, d, *J*=6.0 Hz, CHOH), 3.14 (3H, s, OCH₃), 2.85 (1H, br s, OH), 0.92 (9H, s, 3×CH₃), 0.91 (9H, s, 3×CH₃); diastereomeric mixture, $\delta_{\rm C}$ 141.2, 141.8, 139.7, 138.1, 127.8, 127.6, 126.5, 82.2, 82.1, 80.7, 56.2, 56.0, 35.6, 34.8, 34.5, 26.5, 25.9; diastereomeric mixture, MS (E.I., 70 eV) *m*/*z* 237 (M⁺ – *t*-Bu, 2), 208 (31), 207 (100), 190 (56), 175 (20), 151 (89), 122 (37), 91 (16), 57 (31%); HRMS: found 237.1488. C₁₄H₂₁O₃ requires 238.1491.

4.3.10. 1-Methoxy-3,3-dimethyl-1-phenylbutan-2-ol (6). Diastereomeric mixture, purified by column chromatography (hexane/AcOEt=40:1), colourless oil; first diastereomer, R_f 0.39 (hexane/AcOEt = 10:1); second diastereomer, $R_{\rm f}$ 0.38 (hexane/AcOEt = 10:1); $\nu_{\rm max}$ (liquid film) 3514, 3088, 3063, 3033, 2958, 2903, 1481, 1451, 1401, 1371, 1075, 1015 cm⁻¹; first diastereomer, $\delta_{\rm H}$ 7.37–7.24 (5H, m, ArH), 4.17 (1H, d, J=3.9 Hz, CHOCH₃), 3.30 (1H, dd, J=5.8, 3.9 Hz, CHOH), 3.22 $(3H, s, OCH_3), 2.92 (1H, d, J=5.8 Hz, OH), 0.90 (9H, s, OH), 0.90 (9H, s, OH), 0.90 (9H, s, OH)$ $3 \times CH_3$); second diastereomer, δ_H 7.37–7.24 (5H, m, ArH), 4.11 (1H, d, J=6.2 Hz, CHOCH₃), 3.57-3.51 (1H, m, CHOH) 3.14 (3H, s, OCH₃), 2.84 (1H, br s, OH), 0.93 (9H, s, $3 \times CH_3$); diastereometric mixture, δ_C 140.7, 139.9, 128.6, 128.5, 128.4, 127.8, 127.3, 82.3, 82.0, 80.6, 80.5, 56.2, 56.0, 34.8, 34.5, 26.5; first diastereomer, MS (E.I., 70 eV) m/z 176 (M⁺ – MeOH, 0.5%), 122 (74), 121 (100), 91 (27), 77 (13%); second diastereomer, MS (E.I., 70 eV) m/z 176 (M⁺ – MeOH, 0.5%), 122 (75), 121 (100), 91 (29), 77 (13%); first diastereomer, HRMS: found 176.1183. C12H16O requires 176.1201; second diastereomer, HRMS: found 176.1201. C12H16O requires 176.1201.

4.3.11. 3,3-Diisopropyl-1,3-dihydro-isobenzofuran-1-ol (8). Acetal 1d (500 mg, 2.7 mmol) was lithiated as described in the Section 4.3. To this mixture, chilled to -80 °C, a solution of 2,4-dimethylpentan-3-one (620 mg, 5.4 mmol) dissolved in THF (2 mL) was added dropwise. After stirring at the same temperature until almost complete decolouration, the reaction mixture was quenched and elaborated as described in the Section 4.3. The crude reaction product was dissolved in a mixture of THF/1 M H₂SO₄=1:1 (5 mL) and stirred at rt for 8 h. The resulting mixture was extracted with Et₂O $(3 \times 10 \text{ mL})$, the organic phases were collected, washed with saturated NaHCO₃ (20 mL), dried (K₂CO₃) and the solvent evaporated. Purification by flash chromatography (petroleum ether/AcOEt/Et₃N=7:3:1), afforded a colourless oil (310 mg, 1.4 mmol, 52%), which solidifies upon standing, and was characterized as following: [Found: C, 76.1; H, 9.5. C₁₄H₂₀O₂ requires C, 76.33; H, 9.15]; R_f 0.66 (petroleum ether/AcOEt/Et₃N=7:3:1); ν_{max} (liquid film) 3342 cm^{-1} ; δ_{H} 7.30–7.12 (3H, m, 3×ArH), 7.17-7.09 (1H, m, ArH), 6.44 (1H, s, OCHO), 3.10 (1H, br s, OH), 2.38-2.16 (2H, m, 2×CH), 0.95 (3H, d, J=6.9 Hz, CH₃), 0.82 (3H, d, J=6.9 Hz, CH₃), 0.82 $(3H, d, J=6.9 \text{ Hz}, CH_3), 0.73 (3H, d, J=6.9 \text{ Hz}, CH_3);$ $\delta_{\rm C}$ 142.6, 140.5, 129.0, 127.9, 122.8, 121.7, 100.5, 96.7, 34.2, 32.6, 18.1, 17.2.

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Stereoselective synthesis and functionalization of 4-heterosubstituted β-lactams

Luigino Troisi,* Ludovico Ronzini, Catia Granito, Luisella De Vitis and Emanuela Pindinelli

Dipartimento di Scienze e Tecnologie Biologiche ed Ambientali, University of Lecce, Via Prov.le Lecce-Monteroni, 73100 Lecce, Italy

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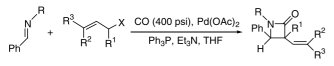
Abstract—Polyfunctionalized β -lactams were prepared with high stereoselectivity in an efficient manner. A palladium-catalyzed [2+2] carbonylative cycloaddition of allyl bromide with heteroaryliden-anilines afforded 2-azetidinones *N*-phenyl substituted, with an heteroaryl moiety linked at the C-4 carbon, and an alkenyl group at the C-3 carbon. The C-3 and the C-4 positions could be further functionalized inserting alkyl and hydroxyl groups in the azetidinone ring, through the generation of a stable azetidinyl anion then captured by various electrophiles.

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1. Introduction

After the discovery of the penicillins and the cephalosporins, the past few decades have witnessed a remarkable growth in the field of β -lactam chemistry, as this heterocycle is a strategic component of various antibacterial agents.^{1,2} The need for potent and effective β -lactam antibiotics, as well as more effective enzyme inhibitors, has motivated synthetic organic chemists to design new functionalized 2-azetidinones.^{3,4} Applications of β -lactams in medicinal chemistry include their use as therapeutic agents for lowering the cholesterol level in plasma,^{5,6} as *anti*-cancer agents,^{4,7–9} and as enzyme inhibitors (for examples inhibitors of HLE¹⁰ and cysteine proteases).¹¹

Among the numerous synthetic protocols reported, a versatile and effective approach to the β -lactams preparation is the transition metal-catalyzed carbonylation of imines with allyl phosphate.^{12,13} Recently, we reported the syntheses of alkenyl β -lactams in good yields and high selectivity by Pd-catalyzed [2+2] cycloaddition of allyl halides and simple imines under CO pressure¹⁴ (Scheme 1).



Scheme 1.

* Corresponding author. Tel./fax: +39 0832 298 701;

e-mail: luigino.troisi@unile.it

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In our opinion, the presence of an heterocycle as an additional substituent of the azetidinone ring should increase the solubility of these structures in polar medium. Moreover, the resulting greater susceptibility to synthetic elaborations should favour an increased and various biological activity. To our knowledge, only few examples of 4-heterosubstituted β -lactams are reported in the literature, such as the preparation of *N*-unsubstituted β -lactams bearing 2-furyl substituent at the C-4 carbon.¹⁵ Therefore, in this paper we report the synthesis of novel alkenyl *N*-phenyl-4-heterosubstituted β -lactams, following the synthetic protocol described above, and especially the further and various functionalization of the β -lactam ring.

2. Results and discussion

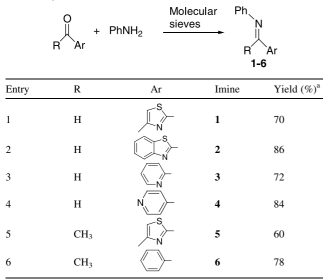
The heteroaryl imines used in these reactions were prepared, in good yields, by coupling reactions of aniline with the appropriate aldehydes, according to Taguchi's method.¹⁶ The results are collected in Table 1. The compound with a phenyl substituent was prepared and is reported for comparison with the heteroaryl groups (entry 6).

The imines **1–6** (1 mmol) were reacted with allyl bromide (1.5 mmol) by [2+2] cycloaddition, under CO pressure (400 psi), in the presence of Et₃N (2 mmol) and 2 mol% of Pd(OAc)₂ complexed by 8 mol% of Ph₃P, for 30–35 h. The catalytic species involved in the process is Pd(0), according to the mechanism previously reported.¹⁷ The cycloaddition results are collected in Table 2.

The alkenyl 4-heterosubstituted β -lactams were isolated with high stereoselectivity: the obtained trans/cis ratios

Keywords: Alkenyl β -lactams; Electrophiles; Cabonylative cycloaddition; Stereoselectivity.

 Table 1. Synthesis of imines 1–6



^a Isolated yields.

were always high, except for entries 5 and 6 (Table 2). In these cases, the presence of two groups (methyl and aryl) on the starting imines reduced the stereoselectivity of the cycloaddition reaction. Moreover, when the heterocycle was benzothiazole (entry 2), two new products **2c** and **2d** were observed, together with the expected compounds **2a** and **2b**. The **2c** and **2d** amounts were small, but they increased for longer reaction times. Their formation should be due to the isomerization of **2a** and **2b** to the more stable α - β -unsaturated carbonyl structures. For instance, when **2a** and **2b** were warmed up with Et₃N in THF, an analogous transformation was observed.

The trans and cis configurations of the β -lactam ring, have been assigned on the basis of the ${}^{3}J_{\text{H-H}}$ coupling constants between the two protons at the C-3 and the C-4 carbon atoms, $(J_{cis} > J_{trans})$.^{18,19} Moreover, the spectroscopic data have been compared to those obtained for similar β -lactams previously characterized by X-ray crystallography.¹⁷

Table 2. Synthesis of 4-heterosubstituted β-lactams (1a-1d)-(6a-6d)

For compounds showing a methyl group linked at C-4 or C-3, the relative configuration was assigned from the coupled ¹³C NMR spectra. A very small or negligible ³ J_{CH_3-H} coupling constant corresponded to a trans configuration, while a larger ${}^{3}J_{CH_3-H}$ (~0.5 ÷ 1.7 Hz) corresponded to a cis configuration.²⁰ These latter configurations were confirmed also by the 400 MHz-NOESY spectra. The differentiation between the two isomers **2c** and **2d** was made from the ¹H NMR spectra: the *Z* isomer displayed its vinylic proton with an upfield chemical shift, whereas the *E* compound showed a downfield chemical shift as this proton is in the deshielding region of the neighbouring carbonyl group.^{21–23}

The 2-azetidinones **1a–4a** and **1b–4b** show two types of acidic protons, linked to the C-3 and the C-4 carbon atoms. The deprotonation of either of them would lead to the formation of an azetidinyl anion stabilized by a large conjugation: by structures A, B, and C in the deprotonation of the C-3 carbon, by structures D, E and by an additional inductive effect of the β -lactam nitrogen, in the deprotonation of the C-4 (Scheme 2).

However, when the *trans*-(**1a–3a**) β -lactams (1 mmol) were treated with LDA (1.2 mmol) in THF, at -78 °C, we noticed that the deprotonation occurred exclusively at the C-3 allylic carbon. Then, adding an electrophile (E⁺, 1 mmol), the carbanion was trapped affording four different quenching products, resulting from A and/or C anions, according to the suggested mechanism of Scheme 2.

The results of the functionalization of **1a–3a** with various electrophiles are collected in Table 3.

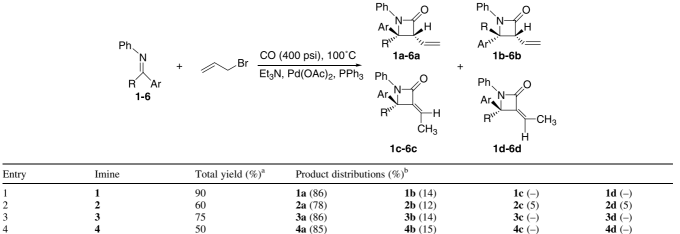
All the reactions performed with 1a showed high yields. Using small electrophiles such as H⁺ or D⁺, equimolecular mixtures of cis and trans diastereomers were observed (entries 1 and 2). When alkyl halides were used, the reaction became highly diastereoselective, the cis isomer being the major reaction product (entries 3 and 5). With benzyl chloride (entry 4), in addition to the expected cis product **9b**,

5c (-)

6c (-)

5d (-)

6d (-)



5b (53)

6b (35)

5a (47)

6a (65)

^a Isolated yields.

5

6

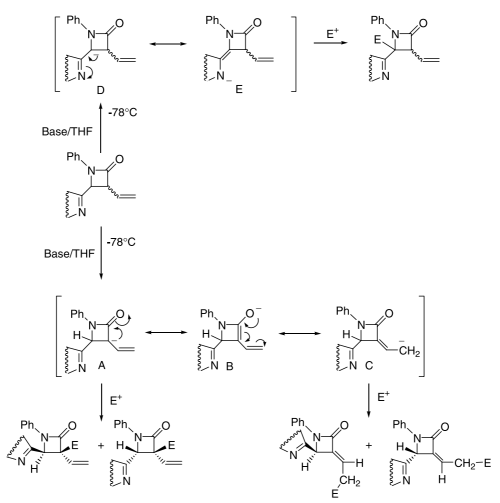
^b Diasteromeric ratios evaluated by GC and ¹H NMR spectroscopy.

40

80

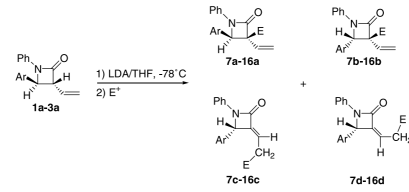
5

6



Scheme 2.

Table 3. Functionalization of 1a–3a with various electrophiles (H^+, D^+, R^+)



Entry	β-Lactam	Е	Total yield (%) ^a 90	Product distributions (%) ^b							
1	1a	H ₂ O		1a (50)	1b (50)	1c (-)	1d (-)	_	_	_	_
2	1a	$\overline{D_2O}$	90	1a (-)	1b (–)	1c (-)	1d (-)	7a (42)	7b (58)	7c (–)	7d (-)
3	1a	CH ₃ I	99	1a (16)	1b (8)	1c (-)	1d (-)	8a (–)	8b (76)	8c (-)	8d (-)
4	1a	PhCH ₂ Cl	85	1a (40)	1b (24)	1c (5)	1d (5)	9a (-)	9b (26)	9c (-)	9d (-)
5	1a	CH ₂ CHCH ₂ Br	95	1a (13)	1b (6)	1c (-)	1d (-)	10a (-)	10b (81)	10c (-)	10d (-)
6	1a	$(CH_3)_2CO$	87	1a (50)	1b (26)	1c (-)	1d (-)	11a (-)	11b (-)	11c (12)	11d (12)
7	1a	PhCHO	86	1a (50)	1b (25)	1c (-)	1d (-)	12a (-)	12b (-)	12c (3)	12d (22)
8	2a	D_2O	85	2a (-)	2b (-)	2c (-)	2d (-)	13a (50)	13b (50)	13c (-)	13d (-)
9	2a	CH ₃ I	90	2a (15)	2b (5)	2c (-)	2d (-)	14a (-)	14b (80)	14c (-)	14d (-)
10	3a	D_2O	85	3a (-)	3b (–)	3c (-)	3d (-)	15a (50)	15b (50)	15c (-)	15d (-)
11	3a	CH ₃ I	90	3a (-)	3b (–)	3c (-)	3d (-)	16a (10)	16b (90)	16c (-)	16d (-)

^a Isolated yields. ^b Diasteromeric ratios evaluated by GC and ¹H NMR spectroscopy.

two more isomers were also generated, **1c** and **1d**, showing an unsaturation at the C-3 carbon. These latter α - β unsaturated β -lactams should result from water quenching the resonance structure C (Scheme 2). With carbonyl compounds as electrophiles (acetone or benzaldehyde), the quenching occurred at the terminal carbon atom of the vinylic chain, on the γ position (structure C, Scheme 2) affording products **11c**, **12c** and **11d**, **12d**, respectively (entries 6 and 7, Table 3).

The diastereomeric mixtures of 1a,1b and 2a,2b isolated in almost every reaction performed with 1a (entries 1-7) and 2a (entry 9), could be formed from quenching of any carbanion not captured by the electrophile. The results confirm a planar structure for the carbanion generated by the deprotonation at C-3 (Scheme 2). While a small electrophile such as H⁺ or D⁺, can bind indifferently from both sides of the molecule, bulkier electrophiles, such as alkyl halides prefer an anti type attack with respect to the heterocycle bonded at the C-4, leading stereoselectively to the cis 2-azetidinones. Moreover, the tridentate nature of the reacting anion could influence the regioselectivity of the electrophilic attack, which may be directed to the α or the γ position of the allylic moiety.^{24–26} Thus α attack predominates in the irreversible reaction with alkyl halides, while γ -adducts are obtained with carbonyl compounds. The lower conversion yields observed for ketone and aldehyde with respect to alkyl halides could be due to the reversible nature of these latter addition reactions.^{25,27–31} Furthemore, in order to verify if the regioselectivity was influenced by electronic effects, the ¹³C NMR spectra of the azetidinyl anion in THF, generated deprotonating 1a with *n*-BuLi, were recorded and data are summarized in Table 4. As recently reported for dienediolates, ³² ¹³C NMR chemical shifts can be related with the π -electron density. The chemical shift displacements to higher fields, listed in Table 4, reveal higher π -electron density at the C α with respect to the C γ carbon atom, which may account for a preferential electrophilic attack to the α position of the allylic moiety, even if sterically more hindered than the γ position.

Table 4.	¹³ C NMR	data of	the azetidinyl	anion
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H ₃ C N H	NHH <u>nBuLi</u> 1a	/THF //	$ \begin{array}{c} Ph \\ S \\ H \\ H \end{array} $ $ \begin{array}{c} 0 \\ - Li^+ \\ \gamma \\ \beta \\ \end{array} $
Carbon atom	δ (ppm)	m)	
	2-Azetidinone 1a	Azetidinyl anion	
Са	62.13	56.47	5.66
Cβ	136.51	136.45	0.06
Сү	115.39	115.10	0.29

Bulky electrophiles such as ketone and aldehyde, however, seem to prefer the less hindered γ position, affording products functionalized at the C γ carbon atom (entries 6 and 7, Table 3). ¹³C NMR investigations of the azetidinyl anion generated with *n*-BuLi from **1b** in THF, afforded similar results observed for **1a**. For instance, for **1b** we noticed

chemical shift displacements towards the same values of the azetidinyl anion reported in Table 4. This behaviour strongly supports the generation of a unique planar anion either starting from the 2-azetidinone **1a** or **1b**. Similarly, **2a** and **3a** deprotonated with LDA, at -78 °C in THF, produced the azetidinyl anion after losing the proton at the C-3. Then, quenching with D₂O led in both cases to an equimolecular mixture of trans and cis isomers (entries 8 and 10). A large stereoselectivity was instead found when the carbanion was quenched with CH₃I, having isolated only the *cis*-**14b** isomer and a cis/trans mixture in the 9:1 ratio, respectively (entries 9 and 11).

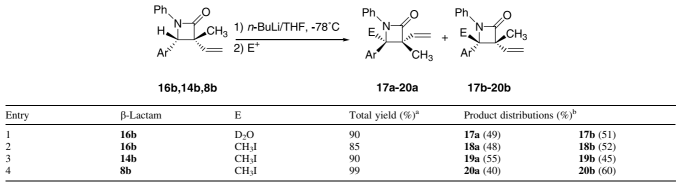
None of the reactions carried out with these substrates showed formation of products derived from deprotonation of the C-4 carbon atom, even using stronger bases like sec-BuLi or n-BuLi. The deprotonation, therefore, seems to depend on the strong difference of acidity between the hydrogens linked to the C-3 and the C-4 carbons. Deprotonation of the C-4 does not occur after treating the substrates 2c and 2d with LDA, which do not have protons at the C-3. For instance, quenching with D₂O produced compounds 13a and 13b arising from the same planar carbanion in the form A, B, and C (Scheme 2). Finally, it is worth noting that the same isomeric ratios and transformation yields were obtained deprotonating β -lactams **1b**, **2b**, and **3b** of cis configuration and trapping them with D₂O or CH₃I. This behaviour provides further support to the hypothesis of a planar structure of the carbanion stabilized by the resonance structures A, B, and C of Scheme 2.

The functionalization of the C-4 carbon atom was achievable only with β -lactams doubly functionalized at C-3, as those isolated from the above reported reactions. In particular, when **16b**, **14b**, and **8b** were treated with *n*-BuLi, in THF at -78 °C, the carbanion was formed at the C-4 with planar or configurationally unstable tetrahedral structure (structures D and E, Scheme 2), since subsequent quenching with electrophiles, such as D₂O or CH₃I, led to products functionalized exclusively at C-4 (entries 1–4, Table 5). No relevant diastereoselectivity was noticed, probably due to the two groups linked at the nearby C-3, which did not allow the electrophile to distinguish between the two sides of the carbanion.

3. Conclusion

In summary, we have synthesised novel β -lactams functionalized with several heterocycles. We exploited the possibility of inserting more functions and groups at the C-3 and C-4 carbon atoms, without performing new cyclizations, but through the generation of stable carbanions and subsequent trapping with electrophiles. The presence on the β -lactam ring of various functionalities susceptible to further synthetic elaborations, such as heterocycles, unsaturated fragments, alkyl and hydroxyl groups make this class of compounds particularly interesting for the study of their potential biological and pharmacological activities.

Table 5. Functionalization of 16b, 14b, and 8b with D_2O and CH_3I



^a Isolated yields.

^b Diasteromeric ratios evaluated by GC and ¹H NMR spectroscopy.

4. Experimental

4.1. General

n-BuLi was a commercial solution in hexanes (Aldrich) and was titrated with *N*-pivaloyl-*o*-toluidine prior to use.³³ THF, triethylamine, palladium(II) acetate, triphenylphosphine, allyl bromide, 2-pyridinecarboxaldehyde, 4-pyridinecarboxaldehyde, acetophenone, 4-methyl-thiazole, 2-aminothiophenol, glycolic acid, lithium diisopropylamide (LDA), deuterium oxide and all other chemicals were of commercial grade (Aldrich) and were used without further purification. Acetaldehyde, benzaldehyde, methyl iodide, allyl bromide, benzyl chloride, and acetone of commercial grade (Aldrich), were purified by distillation prior to use. Petroleum ether refers to the 40–60 °C boiling fraction. The ¹H and the ¹³C NMR spectra were recorded on a Bruker Avance 400 apparatus (400.13 and 100.62 MHz, for ¹H and ¹³C, respectively) with CDCl₃ as solvent and TMS as internal standard (δ =7.24 for ¹H spectra; δ =77.0 for ¹³C spectra). The IR spectra were recorded on a Perkin Elmer spectrometer Model 283. GC-MS analyses were performed with Hewlett-Packard HP-5890 series II gas chromatograph (5% diphenyl/95% dimethylpolysiloxane capillary column, 30 m, 0.25 mm i.d.), equipped with an HP-5971 massselective detector operating at 70 eV (EI). The electrospray ionisation (HR-ESI-MS) experiments were carried out in a hybrid OqTOF mass spectrometer (PE SCIEX-OSTAR) equipped with an ion spray ionisation source. MS (+) spectra were acquired by direct infusion (5 µL/min) of a solution containing the appropriate sample (10 pmol/ μ L), dissolved in a solution 0.1% acetic acid, methanol/water 50:50 at the optimum ion voltage of 4800 V. The nitrogen gas flow was set at 30 psi (pounds per square inch) and the potentials of the orifice, the focusing ring and the skimmer were kept at 30, 50, and 25 V relative to ground, respectively. Elemental analyses were performed on a Carlo Erba C, H, N analyzer. Melting points were determined using an Electrothermal melting point apparatus and are uncorrected. TLC were performed on Merck silica gel plates with F-254 indicator; viewing was by UV light (254 nm). Column chromatographies were performed on silica gel (63-200 µm) using petroleum ether/diethyl ether (Et₂O) mixtures as eluents. All reactions involving airsensitive reagents were performed under nitrogen, in ovendried glassware using syringe/septum cap techniques.

4.2. General procedure for the preparation of heteroaryliden-anilines 1–6

The heteroaryliden-anilines were prepared by coupling reactions of 1 mmol of aniline with the appropriate aldehydes (1 mmol) according to Taguchi's method.¹⁶

4.2.1. (4-Methyl-thiazol-2-yl-methylene)-phenyl-amine **1.** Yield 141 mg (70%), oil. ¹H NMR (400.13 MHz): δ 2.53 (s, 3H), 7.07 (s, 1H), 7.26–7.43 (m, 5H), 8.65 (s, 1H). ¹³C NMR (100.62 MHz): δ 17.0, 117.0, 121.2, 127.2, 129.3, 150.0, 152.8, 154.7, 166.3. GC–MS (70 eV) *m*/*z* (rel int.): 202 (90, M⁺), 201 (94), 174 (84), 125 (14), 104 (78), 77 (100). IR (CHCl₃): 3060, 2960, 1620, 1590, 1500, 1440, 1200 cm⁻¹. HR-ESI-MS: *m*/*z* calcd for C₁₁H₁₁N₂S: 203.0644, [M+H]⁺; found: 203.0644.

4.2.2. Benzothiazol-2-ylmethylene-phenyl-amine 2. Yield 205 mg (86%), yellow solid, mp 99.0–101.0 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 7.33–7.56 (m, 7H), 7.96 (d, J= 7.7 Hz, 1H), 8.12 (d, J=8.2 Hz, 1H), 8.80 (s, 1H). ¹³C NMR (100.62 MHz): δ 121.4, 122.1, 124.3, 126.6, 126.8, 127.8, 129.4, 135.4, 149.6, 153.4, 153.8, 167.4. GC–MS (70 eV) m/z (rel int.): 238 (69, M⁺), 237 (94), 210 (43), 135 (23), 104 (30), 77 (100). IR (CHCl₃): 3050, 2960, 1620, 1590, 1430, 1310, 1200 cm⁻¹. Anal. Calcd for C₁₄H₁₀N₂S: C, 70.56; H, 4.23; N, 11.75. Found: C, 70.36; H, 4.18; N, 11.80.

4.2.3. Phenyl-pyridin-2-ylmethylene-amine 3. Yield 127 mg (72%), oil. ¹H NMR (400.13 MHz): δ 7.23–7.41 (m, 6H), 7.74 (dd, J=7.8, 1.2 Hz, 1H), 8.17 (d, J=7.8 Hz, 1H), 8.60 (s, 1H), 8.68 (d, J=4.7 Hz, 1H). ¹³C NMR (100.62 MHz): δ 120.9, 121.6, 124.9, 126.5, 129.0, 136.4, 149.4, 150.8, 154.4, 160.4. GC–MS (70 eV) m/z (rel int.): 182 (79, M⁺), 181 (100), 155 (67), 154 (77), 105 (53), 77 (86). IR (film): 3050, 2900, 1630, 1590, 1430, 1200, 780, 690 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₂H₁₁N₂: 183.0923, [M+H]⁺; found: 183.0924.

4.2.4. Phenyl-pyridin-4-ylmethylene-amine **4.** Yield 153 mg (84%), yellow solid, mp 71.8–72.3 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 7.24–7.45 (m, 5H), 7.76 (d, J= 5.8 Hz, 2H), 8.46 (s, 1H), 8.76 (d, J=5.8 Hz, 2H). ¹³C NMR (100.62 MHz): δ 120.9, 122.3, 126.9, 129.3, 142.8, 150.6, 151.0, 157.9. GC–MS (70 eV) *m*/*z* (rel int.): 182 (94, M⁺), 181 (78), 104 (73), 79 (61), 77 (100). IR (CHCl₃): 3060,

2960, 1630, 1600, 1480, 1410 cm⁻¹. Anal. Calcd for $C_{12}H_{10}N_2$: C, 79.10; H, 5.53; N, 15.37. Found: C, 78.98; H, 5.48; N, 15.35.

4.2.5. [1-(4-Methyl-thiazol-2-yl)-ethylidene)]-phenylamine **5.** Yield 130 mg (60%), oil. ¹H NMR (400.13 MHz): δ 2.36 (s, 3H), 2.51 (s, 3H), 7.02 (s, 1H), 7.13–7.19 (m, 3H), 7.36 (t, J=8.2 Hz, 2H). ¹³C NMR (100.62 MHz): δ 16.8, 17.3, 115.0, 119.6, 124.1, 128.9, 149.7, 153.8, 155.0, 169.4. GC–MS (70 eV) *m*/*z* (rel int.): 216 (35, M⁺), 201 (19), 174 (23), 118 (30), 77 (100), 51 (48). IR (CHCl₃): 3060, 2960, 1620, 1590, 1500, 1440, 1200 cm⁻¹. HR-ESI-MS: *m*/*z* calcd for C₁₂H₁₃N₂S: 217.0801, [M+H]⁺; found: 217.0802.

4.2.6. Phenyl-(1-phenyl-ethylidene)-amine 6. Known compound previously reported.¹⁴

4.3. General procedure for the preparation of alkenyl β-lactams 4-heterosubstituted (1a–1d)–(6a–6d)

A mixture of 1.0 mmol of **1–6**, 1.5 mmol of allyl bromide, 0.08 mmol of PPh₃, 0.02 mmol of Pd(AcO)₂, and 2 mmol of Et₃N were dissolved in 10 mL of solvent (THF) and placed in a 45 mL autoclave. The autoclave was purged, pressurized (400 psi CO), and then heated to 100 °C for 30–35 h. The reaction was then cooled to room temperature, and worked up by addition of water (15 mL) and extraction with Et₂O (3×5 mL). The combined organic layers were dried (Na₂SO₄) and concentrated in vacuo. The crude products were purified by column chromatography (silica gel, petroleum ether/Et₂O=7:3) to afford the pure β -lactams (**1a–1d**)–(**6a–6d**); yields: 40–90%.

4-(4-Methyl-thiazol-2yl)-1-phenyl-3-vinyl-4.3.1. azetidin-2-one 1a,1b. Overall yield 243 mg (90%). *Compound* **1a**. Yield 208 mg (77%), yellow solid, mp 62.0–63.6 °C (petroleum ether). ¹H NMR (400.13 MHz): δ 2.46 (s, 3H), 4.01 (dd, J=7.4, 2.3 Hz, 1H), 5.20 (d, J=2.3 Hz, 1H), 5.35 (d, J=10.3 Hz, 1H), 5.45 (d, J=17.1 Hz, 1H), 6.00–6.08 (m, 1H), 6.89 (s, 1H), 7.07 (t, J=7.3 Hz, 1H), 7.25–7.36 (m, 4H). 13 C NMR (100.62 MHz): δ 16.9, 58.4, 63.3, 114.6, 116.9, 120.4, 124.3, 129.1, 129.5, 137.0, 153.3, 164.5, 166.9. GC-MS (70 eV) m/z (rel int.): 270 (59, M⁺), 202 (55), 201 (60), 174 (34), 150 (100), 77 (72). IR (CHCl₃): 3040, 2920, 1750, 1595, 1495, 1370 cm⁻¹. Anal. Calcd for C₁₅H₁₄N₂OS: C, 66.64; H, 5.22; N, 10.36. Found: C, 66.72; H, 5.24; N, 10.34. Compound 1b. Yield 35 mg (13%), yellow solid, mp 90.5–91.5 °C (petroleum ether). ¹H NMR (400.13 MHz): δ 2.47 (s, 3H), 4.41 (dd, J=6.2, 6.1 Hz, 1H), 5.23 (dd, J=8.6, 1.8 Hz, 1H), 5.43–5.57 (m, 2H), 5.62 (d, J=6.1 Hz, 1H), 6.87 (s, 1H), 7.10 (t, J= 7.3 Hz, 1H), 7.29–7.39 (m, 4H). ¹³C NMR (100.62 MHz): δ 17.0, 56.9, 58.5, 114.6, 117.1, 121.9, 124.4, 127.4, 129.1, 136.9, 153.5, 164.8, 165.4. GC-MS (70 eV) m/z (rel int.): 270 (60, M⁺), 202 (57), 201 (62), 174 (33), 150 (100), 77 (79). IR (CHCl₃): 3040, 2920, 1750, 1595, 1495, 1370 cm⁻¹. Anal. Calcd for C₁₅H₁₄N₂OS: C, 66.64; H, 5.22; N, 10.36. Found: C, 66.40; H, 5.21; N, 10.33.

4.3.2. 4-Benzothiazol-2-yl-1-phenyl-3-vinyl-azetidin-2-one 2a–2d. Overall yield 184 mg (60%). *Compound* **2a**. Yield 144 mg (47%), yellow solid, mp 100.0–102.0 °C

(*n*-hexane). ¹H NMR (400.13 MHz): δ 4.10 (dd, J=6.5, 2.2 Hz, 1H), 5.33 (d, J=2.2 Hz, 1H), 5.40 (d, J=10.4 Hz, 1H), 5.49 (d, J = 17.1 Hz, 1H), 6.04–6.12 (m, 1H), 7.08 (t, J = 7.4 Hz, 1H), 7.25–7.43 (m, 5H), 7.85 (t, J = 8.0 Hz, 1H), 7.85 (d, J = 8.0 Hz, 1H), 8.05 (d, J = 8.2 Hz, 1H). ¹³C NMR (100.62 MHz): δ 59.0, 63.2, 116.9, 120.9, 122.0, 123.4, 124.6, 125.8, 126.5, 129.2, 129.3, 134.9, 137.0, 153.0, 164.2, 168.9. GC-MS (70 eV) *m/z* (rel int.): 306 (30, M⁺), 237 (25), 186 (100), 77 (29). IR (CHCl₃): 3050, 2970, 1760, 1600, 1490, 1370, 1310 cm^{-1} . Anal. Calcd for C₁₈H₁₄N₂OS: C, 70.57; H, 4.61; N, 9.14. Found: C, 70.55; H, 4.59; N, 9.17. Compound **2b**. Yield 22 mg (7%), oil. ¹H NMR (400.13 MHz): δ 4.52 (dd, J=6.9, 6.2 Hz, 1H), 5.19 (dd, J=9.4, 1.0 Hz, 1H), 5.48–5.64 (m, 2H), 5.73 (d, J=6.2 Hz, 1H), 7.10 (t, J = 6.5 Hz, 1H), 7.27–7.43 (m, 5H), 7.52 (t, J=7.4 Hz, 1H), 7.83 (d, J=8.0 Hz, 1H), 8.05 (d, J = 8.2 Hz, 1H). ¹³C NMR (100.62 MHz): δ 57.3, 58.7, 117.1, 121.9, 122.4, 123.3, 124.6, 125.6, 126.4, 127.0, 129.2, 134.9, 137.0, 153.3, 164.5, 167.5. GC-MS (70 eV) m/z (rel int.): 306 (20, M⁺), 237 (26), 186 (100), 77 (31). IR $(CHCl_3)$: 3050, 2970, 1760, 1600, 1490, 1370, 1310 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₈H₁₅N₂OS: 307.0906, [M+H]⁺; found: 307.0907. 4-Benzothiazol-2-yl-3-ethylidene-1-phenyl-azetidin-2-one 2c. Yield 9 mg (3%), yellow solid, mp 103.0-105.0 °C (petroleum ether). ¹H NMR (400.13 MHz): δ 1.77 (d, J=7.1 Hz, 3H), 5.97 (s, 1H), 6.49 (q, J=7.1 Hz, 1H), 7.06 (t, J=7.5 Hz, 1H), 7.26-7.45 (m, 5H), 7.53 (t, J=8.0 Hz, 1H), 7.84 (d, J=8.0 Hz, 1H), 8.08 (d, J=8.1 Hz, 1H). ¹³C NMR (100.62 MHz): δ 29.7, 60.5, 116.7, 122.1, 123.4, 124.3, 125.8, 126.3, 126.7, 129.3, 135.3, 137.4, 140.3, 152.9, 160.5, 169.2. GC-MS (70 eV) m/z (rel int.): 306 (100, M⁺), 277 (70), 263 (15), 186 (94), 77 (85). IR (CHCl₃): 3080, 2009, 1740, 1600, 1450, 1360, 1090 cm^{-1} . Anal. Calcd for C₁₈H₁₄N₂OS: C, 70.57; H, 4.61; N, 9.14. Found: C, 70.50; H, 4.60; N, 9.11. Compound 2d. Yield 9 mg (3%), yellow solid, mp 113.0–115.0 °C (petroleum ether). ¹H NMR (400.13 MHz): δ 2.13 (d, J =7.2 Hz, 3H), 5.81 (s, 1H), 5.95 (q, J = 7.2 Hz, 1H), 7.07 (t, J = 7.5 Hz, 1H), 7.26–7.46 (m, 5H), 7.52 (t, J = 7.3 Hz, 1H), 7.83 (d, J = 8.0 Hz, 1H), 8.05 (d, J = 8.1 Hz, 1H). ¹³C NMR (100.62 MHz): δ 29.7, 60.6, 116.2, 122.0, 123.3, 124.3, 125.7, 126.3, 129.3, 129.8, 135.2, 137.4, 139.4, 153.0, 160.9, 169.5. GC–MS (70 eV) m/z (rel int.): 306 (100, M⁺), 277 (65), 263 (15), 186 (86), 77 (75). IR (CHCl₃): 3080, 2009, 1740, 1600, 1450, 1360, 1090 cm⁻¹. Anal. Calcd for C₁₈H₁₄N₂OS: C, 70.57; H, 4.61; N, 9.14. Found: C, 70.55; H, 4.62; N, 9.18.

4.3.3. 1-Phenyl-4-pyridin-2-yl-3-vinyl-azetidin-2-one 3a,3b. Overall yield 187 mg (75%). *Compound* **3a**. Yield 161 mg (64%), white solid, mp 107.0–109.0 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 3.88 (dd, J=7.5, 2.4 Hz, 1H), 4.98 (d, J=2.4 Hz, 1H), 5.35 (d, J=10.4 Hz, 1H), 5.44 (d, J=17.1 Hz, 1H), 6.06–6.13 (m, 1H), 7.05 (t, J=7.0 Hz, 1H), 7.23–7.34 (m, 6H), 7.70 (t, J=7.6 Hz, 1H), 8.64 (d, J=4.3 Hz, 1H). ¹³C NMR (100.62 MHz): δ 61.9, 62.4, 116.9, 120.0, 120.4, 123.3, 124.0, 129.1, 130.3, 137.2, 137.5, 150.1, 157.1, 165.1. GC–MS (70 eV) *m/z* (rel int.): 250 (13, M⁺), 181 (22), 155 (6), 130 (100), 77 (30). IR (CHCl₃): 3025, 2930, 1750, 1590, 1490, 1370 cm⁻¹. Anal. Calcd for C₁₆H₁₄N₂O: C, 76.78; H, 5.64; N, 11.19. Found: C, 76.55; H, 5.65; N, 11.22. *Compound* **3b**. Yield 26 mg (10%), white solid, mp 101.0–103.0 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 4.41 (dd, J=6.6, 6.4 Hz, 1H), 5.09 (d, J=9.5 Hz, 1H), 5.26–5.45 (m, 3H), 7.08 (t, J=7.2 Hz, 1H), 7.20–7.45 (m, 6H), 7.65 (t, J=7.5 Hz, 1H), 8.63 (d, J=4.6 Hz, 1H). ¹³C NMR (100.62 MHz): δ 58.0, 59.8, 117.1, 121.1, 121.7, 123.0, 124.1, 128.1, 129.2, 136.7, 140.9, 149.8, 155.3, 165.2. GC–MS (70 eV) m/z (rel int.): 250 (18, M⁺), 181 (37), 155 (10), 130 (100), 77 (46). IR (CHCl₃): 3025, 2930, 1750, 1590, 1490, 1370 cm⁻¹. Anal. Calcd for C₁₆H₁₄N₂O: C, 76.78; H, 5.64; N, 11.19. Found: C, 76.48; H, 5.62; N, 11.15.

4.3.4. 1-Phenyl-4-pyridin-4-yl-3-vinyl-azetidin-2-one 4a,4b. Overall yield 125 mg (50%). Compound 4a. Yield 106 mg (42%), oil. ¹H NMR (400.13 MHz): δ 3.72 (dd, J =7.9, 2.5 Hz, 1H), 4.82 (d, J=2.5 Hz, 1H), 5.37 (d, J=10.2 Hz, 1H), 5.41 (d, J = 16.6 Hz, 1H), 6.00–6.10 (m, 1H), 7.03–7.12 (m, 1H), 7.24–7.34 (m, 6H), 8.64 (d, J = 5.5 Hz, 2H). ¹³C NMR (100.62 MHz): δ 59.8, 63.7, 116.9, 120.7, 124.5, 129.3, 129.8, 137.1, 146.3, 146.8, 150.4, 164.4. GC-MS (70 eV) *m/z* (rel int.): 250 (5, M⁺), 181 (10), 130 (100), 104 (25), 77 (70). IR (CHCl₃): 3030, 2920, 1750, 1600, 1495, 1375 cm⁻¹. HR-ESI-MS: m/z calcd for $C_{16}H_{15}N_2O: 251.1185, [M+H]^+$; found: 251.1184. Compound **4b**. Yield 19 mg (7%, measured by GC), oil. 1 H NMR (400.13 MHz): δ 4.44 (dd, J = 6.8, 6.4 Hz, 1H), 5.21– 5.51 (m, 4H), 7.03–7.40 (m, 5H), 7.52 (d, J = 5.5 Hz, 2H), 8.65 (d, J=5.5 Hz, 2H). GC-MS (70 eV) m/z (rel int.): 250 (5, M⁺), 181 (12), 130 (100), 104 (25), 77 (70). IR (CHCl₃): 3030, 2920, 1750, 1600, 1495, 1375 cm⁻¹. HR-ESI-MS: *m*/ z calcd for $C_{16}H_{15}N_2O$: 251.1185, $[M+H]^+$; found: 251.1185.

4.3.5. 4-Methyl-4-(4-methyl-thiazol-2yl)-1-phenyl-3-vinyl-azetidin-2-one 5a,5b. Overall yield 114 mg (40%). Compound 5a. Yield 54 mg (19%), oil. ¹H NMR (400.13 MHz): δ 1.99 (s, 3H), 2.46 (s, 3H), 4.09 (d, J= 7.7 Hz, 1H), 5.43 (d, J = 10.5 Hz, 1H), 5.47 (d, J = 17.4 Hz, 1H), 5.89–5.98 (m, 1H), 6.89 (s, 1H), 7.06 (t, J=7.3 Hz, 1H), 7.24–7.44 (m, 4H). ¹³C NMR (100.62 MHz): δ 17.2, 19.0, 63.9, 67.1, 114.6, 117.7, 122.4, 124.4, 127.6, 129.0, 136.6, 153.4, 166.0, 171.8. GC-MS (70 eV) m/z (rel int.): 284 (7, M⁺), 216 (11), 164 (100), 118 (13), 77 (60). IR (CHCl₃): 3060, 2925, 1740, 1600, 1490, 1365 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₆H₁₇N₂OS: 285.1063, $[M+H]^+$; found: 285.1063. *Compound* **5b**. Yield 60 mg (21%), oil. ¹H NMR (400.13 MHz): δ 2.18 (s, 3H), 2.46 (s, 3H), 4.00 (d, J=6.9 Hz, 1H), 5.11–5.14 (m, 1H), 5.34–5.47 (m, 2H), 6.86 (s, 1H), 7.08 (t, J=7.4 Hz, 1H), 7.29 (t, J=7.4 Hz, 2H), 7.39 (t, J=7.6 Hz, 2H). ¹³C NMR (100.62 MHz): δ 17.2, 23.0, 65.6, 67.1, 114.6, 117.9, 121.5, 124.1, 127.5, 128.9, 136.5, 153.6, 164.7, 168.7. GC-MS (70 eV) m/z (rel int.): 284 (14, M⁺), 216 (18), 164 (100), 118 (18), 77 (62). IR (CHCl₃): 3060, 2925, 1740, 1600, 1490, 1365 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₆H₁₇N₂OS: 285.1063, [M+H]⁺; found: 285.1064.

4.3.6. 4-Methyl-1,4-diphenyl-3-vinyl-azetidin-2-one 6a,6b. Overall yield 211 mg (80%). *Compound* **6a.** Yield 137 mg (52%), oil. ¹H NMR (400.13 MHz): δ 1.91 (s, 3H), 3.80 (d, J=8.2 Hz, 1H), 5.36–5.43 (m, 2H), 5.90–5.99 (m, 1H), 7.04 (t, J=7.4 Hz, 1H), 7.22–7.39 (m, 9H). ¹³C NMR (100.62 MHz): δ 19.5, 64.2, 67.8, 117.7, 122.0, 123.7, 124.8, 127.8, 128.2, 129.0, 129.1, 137.0, 141.0, 165.7.

GC–MS (70 eV) m/z (rel int.): 263 (8, M⁺), 196 (5), 195 (23), 180 (48), 144 (60), 129 (100), 77 (85). IR (CHCl₃): 3030, 2990, 2920, 1735, 1600, 1590, 1370 cm⁻¹. HR-ESI-MS: m/z calcd for $C_{18}H_{18}NO$: 264.1389, $[M+H]^+$; found: 264.1390. *Compound* **6b**. Yield 74 mg (28%), oil. ¹H NMR (400.13 MHz): δ 2.09 (s, 3H), 3.87 (d, J=7.9 Hz, 1H), 5.01 (d, J=10.4 Hz, 1H), 5.09–5.17 (m, 1H), 5.25 (d, J= 16.0 Hz, 1H), 7.06 (t, J=7.5 Hz, 1H), 7.24–7.45 (m, 9H). ¹³C NMR (100.62 MHz): δ 23.6, 65.7, 66.9, 117.8, 120.4, 123.7, 126.5, 127.7, 128.6, 129.0, 129.2, 137.1, 138.7, 165.2. GC–MS (70 eV) m/z (rel int.): 263 (8, M⁺), 196 (7), 195 (33), 180 (63), 144 (63), 129 (100), 77 (90). IR (CHCl₃): 3030, 2990, 2920, 1735, 1600, 1590, 1370 cm⁻¹. HR-ESI-MS: m/z calcd for $C_{18}H_{18}NO$: 264.1389, $[M+H]^+$; found: 264.1390.

4.4. General procedure for the functionalization of alkenyl β -lactams 4-heterosubstituted 1a–3a

To a stirred solution of 1 mmol of **1a–3a** in THF (30 mL) at -78 °C, LDA (2.0 M in hexanes, 0.6 mL, 1.2 mmol) was added dropwise under nitrogen. The resulting mixture was stirred at -78 °C for 5 min, and then the electrophile was added (1.5 mmol). The reaction was warmed up to room temperature and quenched with saturated aq NH₄Cl. The aqueous layer was extracted with Et₂O (3×20 mL) and the combined organic layers were dried over anhydrous Na₂SO₄ and concentrated in vacuo. The crude products were purified by column chromatography (silica gel, petroleum ether/Et₂O, 1:1) to afford the pure functionalized β -lactams (**7a–7d**)–(**16a–16d**); yields: 85–99%.

4.4.1. 3-Deutero-4-(4-methyl-thiazol-2-yl)-1-phenyl-3vinyl-azetidin-2-one 7a. Yield 103 mg (38%), (>80%D), yellow solid, mp 60.0–62.0 °C (n-hexane). The IR, ¹H and ¹³C NMR data are the same of those reported for **1a**. In the ¹H NMR spectrum the double doublet at 4.01 ppm almost disappears, while the doublet at 5.20 becomes a singlet. GC–MS (70 eV) *m/z* (rel int.): 271 (72, M⁺), 202 (70), 201 (75), 174 (51), 151 (100) 77 (85). IR (CHCl₃): 3020, 2910, 2850, 2220, 1750, 1595, 1500, 1370 cm⁻¹. HR-ESI-MS: m/ z calcd for $C_{15}H_{14}DN_2OS$: 272.0969, $[M+H]^+$; found: 272.0970.Compound 7b. Yield 141 mg (52%), (>80%D), yellow solid, mp 90.0–92.0 °C (*n*-hexane). The IR, ¹H and ¹³C NMR data are the same of those reported for **1b**. In the ¹H NMR spectrum the double doublet at 4.41 ppm almost disappears, while the doublet at 5.23 becomes a singlet. GC–MS (70 eV) *m/z* (rel int.): 271 (57, M⁺), 202 (65), 201 (76), 174 (40), 151 (100) 77 (90). IR (CHCl₃): 3020, 2910, 2850, 2220, 1750, 1595, 1500, 1370 cm⁻¹. HR-ESI-MS: m/ z calcd for $C_{15}H_{14}DN_2OS$: 272.0969, $[M+H]^+$; found: 272.0970.

4.4.2. 3-Methyl-4-(4-methyl-thiazol-2yl)-1-phenyl-3-vinyl-azetidin-2-one 8b. Yield 213 mg (75%), yellow solid, mp 99.4–100.9 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 1.69 (s, 3H), 2.47 (s, 3H), 5.12 (dd, J= 9.3, 2.4 Hz, 1H), 5.24 (s, 1H), 5.42–5.49 (m, 2H), 6.86 (s, 1H), 7.09 (t, J=7.2 Hz, 1H), 7.26–7.35 (m, 4H). ¹³C NMR (100.62 MHz): δ 17.0, 21.0, 62.9, 64.8, 114.6, 117.3, 118.4, 124.3, 129.1, 133.1, 137.1, 153.3, 165.8, 168.2. GC–MS (70 eV) *m*/*z* (rel int.): 284 (58, M⁺), 269 (13), 202 (66), 201 (76), 174 (50), 165 (73), 164 (100) 77 (82). IR (CHCl₃): 3020, 2970, 2920, 1750, 1600, 1590, 1360, 1310 cm⁻¹. Anal. Calcd for C₁₆H₁₆N₂OS: C, 67.58; H, 5.67; N, 9.85. Found: C, 67.24; H, 5.71; N, 9.84.

4.4.3. 3-Benzyl-4-(4-methyl-thiazol-2yl)-1-phenyl-3-vinyl-azetidin-2-one 9b. Yield 79 mg (22%), yellow solid, mp 109.8–110.7 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 2.46 (s, 3H), 3.21 (d, *J*=14.1 Hz, 1H), 3.37 (d, *J*=14.1 Hz, 1H), 5.15 (d, *J*=10.4 Hz, 1H), 5.30 (s, 1H), 5.39–5.47 (m, 1H), 5.12 (d, *J*=17.3 Hz, 1H), 6.82 (s, 1H), 7.02 (t, *J*=7.1 Hz, 1H), 7.15–7.38 (m, 9H). ¹³C NMR (100.62 MHz): δ 17.1, 41.1, 61.4, 67.3, 114.7, 117.4, 119.1, 124.3, 127.1, 128.4, 128.9, 130.3, 132.5, 135.2, 136.5, 153.2, 166.0, 167.2. GC–MS (70 eV) *m/z* (rel int.): 360 (17, M⁺), 269 (27), 241 (78), 240 (100), 202 (26), 201 (36) 91 (40), 77 (62). IR (CHCl₃): 3020, 2950, 1750, 1600, 1490, 1370 cm⁻¹. Anal. Calcd for C₂₂H₂₀N₂OS: C, 73.30; H, 5.59; N, 7.77. Found: C, 73.35; H, 5.58; N, 7.79.

4.4.4. 3-Ethylidene-4-(4-methyl-thiazol-2yl)-1-phenyl-**3-vinyl-azetidin-2-one 1c.** Yield 11 mg (4%), oil. ¹H NMR (400.13 MHz): δ 1.74 (d, J=7.2 Hz, 3H), 2.48 (s, 3H), 5.85 (s, 1H), 6.42 (q, J = 7.2 Hz, 1H), 6.90 (s, 1H), 7.06 (t, J=7.2 Hz, 1H), 7.26–7.31 (m, 2H), 7.43 (d, J=7.2 Hz, 2H). ¹³C NMR (100.62 MHz): δ 17.0, 29.7, 59.9, 115.2, 116.8, 124.1, 125.9, 129.2, 133.9, 140.0, 153.0, 160.7, 176.5. GC-MS (70 eV) m/z (rel int.): 270 (93, M⁺), 241 (45), 227 (14), 178 (47), 150 (100), 77 (56). IR (CHCl₃): 3020, 2930, 1740, 1595, 1500, 1360 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₅H₁₅N₂OS: 271.0906, [M+H]⁺; found: 271.0907. Compound 1d. Yield 11 mg (4%), oil. ¹H NMR (400.13 MHz): δ 2.13 (d, J=7.2 Hz, 3H), 2.48 (s, 3H), 5.69 (s, 1H), 5.90 (q, J=7.2 Hz, 1H), 6.89 (s, 1H), 7.07 (t, J=7.4 Hz, 1H), 7.30 (t, J=7.4 Hz, 2H), 7.41 (d, J=7.4 Hz, 2H). ¹³C NMR (100.62 MHz): δ 16.9, 29.7, 59.9, 115.0, 116.8, 124.2, 125.7, 129.2, 129.3, 137.4, 152.9, 160.7, 177.0. GC-MS (70 eV) m/z (rel int.): 270 (31, M⁺), 241 (26), 227 (7), 178 (38), 150 (100), 77 (98). IR (CHCl₃): $3020, 2930, 1740, 1595, 1500, 1360 \text{ cm}^{-1}$. HR-ESI-MS: *m*/ z calcd for $C_{15}H_{15}N_2OS$: 271.0906, $[M+H]^+$; found: 271.0907.

4.4.5. 3-Allyl-4-(4-methyl-thiazol-2yl)-1-phenyl-3-vinylazetidin-2-one 10b. Yield 239 mg (77%), oil. ¹H NMR (400.13 MHz): δ 2.47 (s, 3H), 2.74–2.77 (m, 2H), 5.15–5.54 (m, 6H), 5.87–.93 (m, 1H), 6.86 (s, 1H), 7.09 (t, *J*=7.0 Hz, 1H), 7.26–7.34 (m, 4H). ¹³C NMR (100.62 MHz): δ 17.0, 39.4, 61.4, 66.2, 114.7, 117.3, 119.0, 120.1, 124.3, 129.1, 131.7, 132.1, 136.7, 153.2, 165.9, 167.0. GC–MS (70 eV) *m*/*z* (rel int.): 310 (15, M⁺), 269 (36), 202 (40), 201 (59), 191 (69), 190 (100) 77 (85). IR (CHCl₃): 3060, 2920, 1750, 1695, 1490, 1360 cm⁻¹. HR-ESI-MS: *m*/*z* calcd for C₁₈H₁₉N₂OS: 311.1220, [M+H]⁺; found: 311.1220.

4.4.6. 3-(3-Hydroxy-3-methyl-butylidene)-4-(4-methyl-thiazol-2yl)-1-phenyl-3-vinyl-azetidin-2-one 11c+11d. Overall yield 69 mg (21%), oil. Inseparable mixture of two trans and cis-configurated diasteromers (dr=1/1 by ¹H NMR). ¹H NMR, ¹³C NMR, GC–MS, HR-ESI-MS and IR data were measured on the mixture. ¹H NMR (400.13 MHz): δ 1.09 (s, 3H), 1.17 (s, 3H), 1.27 (s, 3H), 1.30 (s, 3H), 1.85 (s, 1H+1H, broad), 2.28–2.33 (m, 2H), 2.48 (s, 3H+3H), 2.74–2.77 (m, 2H), 5.77 (s, 1H), 5.87

(s, 1H), 6.03 (t, J=7.4 Hz, 1H), 6.51 (t, J=7.0 Hz, 1H), 6.89 (s, 1H+1H), 7.07 (t, J=7.4 Hz, 1H+1H), 7.27–7.43 (m, 4H+4H). ¹³C NMR (100.62 MHz): δ 15.3, 16.9, 29.0, 29.1, 29.5, 29.7, 41.9, 42.5, 60.1, 60.2, 70.7, 71.3, 115.1, 115.2, 116.8, 116.9, 124.3, 126.7, 129.1, 129.2, 130.0, 137.2, 137.3, 141.6, 141.9, 152.9, 153.0, 160.5, 161.2, 167.29, 167.3. IR (CHCl₃): 3400 (broad), 3020, 2960, 2920, 1735, 1600, 1490, 1370, 1100 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₈H₂₁N₂O₂S: 329.1325, [M+H]⁺; found: 329.1326. Isomer I: GC–MS (70 eV) m/z (rel int.): 328 (22, M⁺), 310 (3), 269 (83), 236 (6), 178 (100), 150 (90), 77 (75), 59 (90). Isomer II: GC–MS (70 eV) m/z (rel int.): 328 (27, M⁺), 310 (3), 269 (100), 236 (75), 178 (95), 150 (85), 77 (73), 59 (90).

4.4.7. 3-(3-Hydroxy-3-phenyl-propylidene)-4-(4-methylthiazol-2yl)-1-phenyl-3-vinyl-azetidin-2-one 12c+12d. Overall yield 79 mg (21%), oil. Inseparable mixture of two trans and cis-configurated diasteromers (dr = 1/7 by ¹H NMR). Compound **12c**. ¹H NMR (400.13 MHz): δ 2.45 (s, 3H), 2.50?2.51 (m, 2H), 2.80 (s, 1H, broad), 4.62–4.68 (m, 1H), 5.77 (s, 1H), 6.32–6.38 (m, 1H), 6.88 (s, 1H), 7.06 (t, J=7.3 Hz, 1H), 7.23–7.39 (m, 9H). Compound **12d**. ¹H NMR (400.13 MHz): δ 2.43 (s, 3H), 2.80 (s, 1H, broad), 2.93-3.08 (m, 2H), 4.85-4.93 (m, 1H), 5.68 (s, 1H), 5.90-5.93 (m, 1H), 6.86 (s, 1H), 7.06 (t, J=7.3 Hz, 1H), 7.23-7.39 (m, 9H). Compound 12c+12d. ¹³C NMR (100.62 MHz): δ 15.2, 16.9, 37.7, 38.2, 60.1, 62.1, 73.1, 73.3, 115.1, 115.3, 116.8, 120.2, 120.7, 124.2, 124.3, 125.5, 125.7, 125.8, 126.6, 127.6, 127.7, 128.4, 128.5, 129.1, 129.3, 129.7, 137.2, 141.1, 141.2, 141.8, 143.2, 143.3, 152.9, 153.0, 160.5, 161.0, 167.1, 169.4. GC-MS (70 eV) *m*/*z* (rel int.): 376 (3, M⁺), 358 (3), 270 (41), 269 (45), 178 (100), 150 (60), 77 (70). IR (CHCl₃): 3370 (broad), 2950, 1730, 1600, 1490, 1360, 1100 cm⁻¹. HR-ESI-MS: m/zcalcd for $C_{22}H_{21}N_2O_2S$: 377.1325, $[M+H]^+$; found: 377.1326.

4.4.8. 4-Benzothiazol-2-yl-3-deutero-1-phenyl-3-vinylazetidin-2-one 13a. Yield 129 mg (42%), (>90%D), yellow solid, mp 100.6–102.1 °C (*n*-hexane). The IR, ¹H and ¹³C NMR data are the same of those reported for 2a. In the ¹H NMR spectrum the double doublet at 4.10 ppm almost disappears, while the doublet at 5.33 becomes a singlet. GC-MS (70 eV) m/z (rel int.): 307 (25, M⁺), 237 (30), 187 (100), 77 (85). IR (CHCl₃): 3050, 2970, 2220, 1760, 1600, 1490, 1370, 1310 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₈H₁₄DN₂OS: 308.0969, [M+H]⁺; found: 308.0970. Compound 13b. Yield 129 mg (42%), (>90%D), oil. The IR, ¹H and ¹³C NMR data are the same of those reported for **2b**. In the ¹H NMR spectrum the double doublet at 4.52 ppm almost disappears, while the doublet at 5.73 becomes a singlet. GC-MS (70 eV) m/z (rel int.): 307 (25, M⁺), 237 (30), 187 (100), 77 (85). IR (CHCl₃): 3050, 2970, 2220, 1760, 1600, 1490, 1370, 1310 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₈H₁₄DN₂OS: 308.0969, [M+H]⁺; found: 308.0969.

4.4.9. 4-Benzothiazol-2-yl-3-methyl-1-phenyl-3-vinyl-azetidin-2-one 14b. Yield 230 mg (72%), yellow solid, mp 142.4–143.6 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 1.75 (s, 3H), 5.10 (dd, *J*=10.0, 1.7 Hz, 1H), 5.35 (s, 1H), 5.45–5.60 (m, 2H), 7.10 (t, *J*=7.4 Hz, 1H), 7.24–7.52

(m, 6H), 7.81 (d, J=8.0 Hz, 1H), 8.05 (d, J=8.0 Hz, 1H). ¹³C NMR (100.62 MHz): δ 21.1, 63.2, 65.0, 117.2, 118.8, 121.9, 123.2, 124.5, 125.5, 126.3, 129.2, 130.0, 132.6, 134.9, 137.1, 153.0, 167.9. GC–MS (70 eV) m/z (rel int.): 320 (12, M⁺), 237 (22), 200 (100), 77 (60). IR (CHCl₃): 3050, 2970, 1760, 1600, 1490, 1370, 1310 cm⁻¹. Anal. Calcd for C₁₉H₁₆N₂OS: C, 71.22; H, 5.03; N, 8.74. Found: C, 71.08; H, 5.05; N, 8.71.

3-Deutero-1-phenyl-4-pyridin-2-yl-3-vinyl-4.4.10. azetidin-2-one 15a. Yield 105 mg (42%), (>90%D), white solid, mp 107.0-109.0 °C (n-hexane). The IR, ¹H and ¹³C NMR data are the same of those reported for **3a**. In the ¹H NMR spectrum the double doublet at 3.88 ppm almost disappears, while the doublet at 4.98 becomes a singlet. GC-MS (70 eV) m/z (rel int.): 251 (19, M⁺), 182 (15), 181 (37), 131 (100), 77 (54). IR (CHCl₃): 3025, 2930, 2220, 1750, 1590, 1490, 1370 cm⁻¹. HR-ESI-MS: m/zcalcd for $C_{16}H_{14}DN_2O$: 252.1248, $[M+H]^+$; found: 252.1248. Compound 15b. Yield 105 mg (42%), (>90%D), white solid, mp 101.0–103.0 °C (*n*-hexane). The IR, ¹H and ¹³C NMR data are the same those reported for **3b**. In the ¹H NMR spectrum the double doublet at 4.41 ppm almost disappears, while the doublet at 5.40 becomes a singlet. GC-MS (70 eV) m/z (rel int.): 251 (25, M⁺), 182 (12), 181 (43), 131 (100), 77 (51). IR (CHCl₃): 3025, 2930, 2220, 1750, 1590, 1490, 1370 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₆H₁₄DN₂O: 252.1248, [M+H]⁺; found: 252.1249.

4.4.11. 3-Methyl-1-phenyl-4-pyridin-2-yl-3-vinyl-azetidin- 2-one 16a. Yield 24 mg (9%), white solid, mp 105.0-106.0 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 1.00 (s, 3H), 5.15 (s, 1H), 5.31 (d, J = 10.6 Hz, 1H) 5.47 (d, J =17.3 Hz, 1H), 6.20 (dd, J=17.3, 10.6 Hz, 1H), 7.05 (t, J=7.1 Hz, 1H), 7.25–7.35 (m, 6H), 7.66 (t, J=7.6 Hz, 1H), 8.64 (d, J=4.3 Hz, 1H). ¹³C NMR (100.62 MHz): δ 15.6, 61.7, 67.0, 116.3, 117.2, 121.4, 122.8, 124.0, 129.1, 136.6, 137.3, 137.5, 149.7, 155.5, 168.7. GC-MS (70 eV) m/z (rel int.): 264 (15, M⁺), 181 (36), 154 (9), 144 (100), 77 (45). IR (CHCl₃): 3020, 2940, 2870, 1740, 1590, 1480, 1440, 1370 cm⁻¹. Anal. Calcd for $C_{17}H_{16}N_2O$: C, 82.22; H, 6.49; N, 11.28. Found: C, 82.17; H, 6.46; N, 11.30. Compound 16b. Yield 214 mg (81%), white solid, mp 100.0–101.1 °C (*n*-hexane). ¹H NMR (400.13 MHz): δ 1.71 (s, 3H), 4.97–4.99 (m, 1H), 5.04 (s, 1H), 5.25–5.32 (m, 2H), 7.07 (t, J=7.0 Hz, 1H), 7.20–7.35 (m, 6H), 7.63 (t, J=7.1 Hz, 1H), 8.63 (d, J=4.5 Hz, 1H). ¹³C NMR $(100.62 \text{ MHz}): \delta 21.1, 62.1, 67.8, 117.2, 117.5, 121.6,$ 122.9, 129.09, 130.5, 133.8, 136.5, 137.2, 149.7, 155.6, 168.6. GC-MS (70 eV) m/z (rel int.): 264 (15, M⁺), 181 (33), 154 (10), 144 (100), 77 (48). IR (CHCl₃): 3020, 2940, 2870, 1740, 1590, 1480, 1440, 1370 cm⁻¹. Anal. Calcd for C₁₇H₁₆N₂O: C, 82.22; H, 6.49; N, 11.28. Found: C, 82.20; H, 6.48; N, 11.29.

4.5. General procedure for the preparation of the azetidinyl anion

For the NMR measurements compound **1a** or **1b** (0.1 mmol) was dissolved in 0.5 ml of a mixture of THF/CDCl₃ in a ratio of 8:2. A ¹³C NMR experiment was performed on these solvent mixture. The NMR tube containing the mixture was

cooled to -78 °C and then *n*-BuLi (2.5 M in hexane, 40 mL, 0.1 mmol) was added to the tube. Then mixture was vigorously stirred and placed into the instrument probe wich is at the constant temperature of 25 °C. The ¹³C NMR spectra was then acquired.

4.6. General procedure for the functionalization of 16b,14b, and 8b

To a stirred solution of 1 mmol of the 2-azetidinone in THF (30 mL) at -78 °C, *n*-BuLi (2.5 M in hexanes, 0.5 mL, 1.2 mmol) was added dropwise under nitrogen. The resulting mixture was stirred at -78 °C for 30 min, and then the electrophile was added (1.5 mmol). The reaction was warmed up to room temperature and quenched with saturated aq NH₄Cl. The mixture was worked up and purified as reported in the Section 4.4. The pure functionalized β -lactams **17a–20a** and **17b–20b** were isolated with yields of 85–99%.

4.6.1. 4-Deutero-3-methyl-1-phenyl-4-pyridin-2-yl-3vinyl-azetidin-2-one 17a,17b. Overall yield 238 mg (90%). Compound 17a. Yield 117 mg (44%), (>90%D), white solid, mp 105.0–106.5 °C (*n*-hexane). The IR, ¹H and ¹³C NMR data are the same of those reported for **16a**. In the ¹H NMR spectrum the singlet at 5.15 ppm almost disappears. \hat{GC} -MS (70 eV) m/z (rel int.): $2\hat{65}$ (21, M⁺), 183 (15), 182 (34), 145 (100), 77 (32). IR (CHCl₃): 3020, 2940, 2870, 2220, 1740, 1590, 1480, 1440, 1370 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₇H₁₆DN₂O: 266.1405, [M+H]⁺; found: 266.1405. Compound 17b. Yield 122 mg (46%), (>90%D), white solid, mp 100.1–101.4 °C (*n*-hexane). The IR, ¹H and ¹³C NMR data are the same of those reported for **16b.** In the ¹H NMR spectrum the singlet at 5.04 ppm almost disappears. GC-MS (70 eV) m/z (rel int.): 265 (15, M⁺), 183 (11), 182 (31), 145 (100), 77 (43). IR (CHCl₃): 3020, 2940, 2870, 2220, 1740, 1590, 1480, 1440, 1370 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₇H₁₆DN₂O: 266.1405, [M+H]⁺; found: 266.1405.

4.6.2. 3,4-Dimethyl-1-phenyl-4-pyridin-2-yl-3-vinylazetidin-2-one 18a+18b. Overall yield 236 mg (85%), oil. Inseparable mixture of two trans and cis-configurated diasteromers (dr = 1/1 by ¹H NMR and GC–MS). ¹H NMR, ¹³C NMR, GC-MS, HR-ESI-MS and IR data were measured on the mixture. ¹H NMR (400.13 MHz): δ 0.87 (s, 3H), 1.55 (s, 3H), 1.96 (s, 3H), 2.04 (s, 3H), 4.80 (dd, J =10.3, 1.5 Hz, 1H), 5.11–5.29 (m, 2H), 5.34 (d, J=10.8 Hz, 1H), 5.50 (d, J=17.4 Hz, 1H), 6.07 (dd, J=17.4, 10.8 Hz, 1H), 7.10–7.65 (m, 16H), 8.64 (d, J = 4.7 Hz, 1H), 8.67 (d, J = 4.7 Hz, 1H). ¹³C NMR (100.62 MHz): δ 17.2, 18.0, 19.1, 20.5, 64.6, 64.8, 70.1, 70.2, 115.9, 117.5, 118.0, 118.1, 121.6, 121.9, 122.1, 122.2, 123.7, 123.8, 129.0, 135.1, 135.4, 136.0, 136.1, 137.1, 137.2, 149.3, 149.5, 159.8, 159.9, 169.1, 169.2. GC-MS (70 eV) m/z (rel int.): 278 (22, M⁺), 196 (60), 195 (70), 181 (20), 158 (80), 118 (44), 77 (100). IR (CHCl₃): 3040, 2920, 1750, 1590, 1500, 1365 cm⁻¹. HR-ESI-MS: m/z calcd for C₁₈H₁₉N₂O: 279.1499, [M+H]+; found: 279.1499.

4.6.3. 4-Benzothiazol-2-yl-3,4-dimethyl-1-phenyl-3-vinyl-azetidin-2-one 19a,19b. Overall yield 301 mg(90%). Compound 19a. Yield 166 mg (49%), oil. ¹H

1573

NMR (400.13 MHz): δ 1.22 (s, 3H), 2.08 (s, 3H), 5.39 (d, J=10.7 Hz, 1H), 5.54 (d, J=17.4 Hz, 1H), 6.02 (dd, J=10.7, 17.4 Hz, 1H), 7.11 (t, J=7.4 Hz, 1H), 7.26–7.51 (m, 6H), 7.84 (d, J=8.1 Hz, 1H), 8.05 (d, J=8.1 Hz, 1H). ¹³C NMR (100.62 MHz): δ 17.4, 21.5, 65.9, 68.8, 117.5, 118.9, 122.1, 123.4, 124.5, 125.7, 126.5, 129.7, 130.2, 132.8, 137.3, 153.2, 167.8, 168.1. GC-MS (70 eV) m/z (rel int.): 334 (51, M⁺), 252 (75), 251 (89), 237 (24), 214 (100), 118 (49), 77 (57). IR (CHCl₃): 3050, 2970, 1760, 1600, 1490, 1370, 1310 cm⁻¹. HR-ESI-MS: m/z calcd for C₂₀H₁₉N₂OS: 335.1220, [M+H]⁺; found: 335.1220. Compound **19b**. Yield 134 mg (40%), oil. ¹H NMR (400.13 MHz): δ 1.60 (s, 3H), 2.17 (s, 3H), 4.91 (dd, J=2.9, 9.0 Hz, 1H), 5.34–5.40 (m, 2H), 7.15 (t, J=7.4 Hz, 1H), 7.25–7.50 (m, 6H), 7.82 (d, J=8.0 Hz, 1H), 8.06 (d, J=8.0 Hz, 1H). ¹³C NMR (100.62 MHz): δ 18.5, 21.6, 66.0, 69.1, 117.3, 119.5, 122.3, 123.3, 124.5, 125.7, 126.7, 129.8, 130.8, 132.9, 137.5, 153.5, 168.0, 168.5. GC-MS (70 eV) m/z (rel int.): 334 (30, M⁺), 252 (52), 251 (63), 237 (16), 214 (100), 118 (43), 77 (49). IR (CHCl₃): 3050, 2970, 1760, 1600, 1490, 1370, 1310 cm⁻¹. HR-ESI-MS: m/z calcd for C₂₀H₁₉N₂OS: 335.1220, [M+H]⁺; found: 335.1219.

4.6.4. 3,4-Dimethyl-4-(4-methylthiazolyl)-1-phenyl-3-vinyl-azetidin-2-one 20a,20b. Overall yield 295 mg (99%). Compound **20a**. Yield 119 mg (40%), oil. ¹H NMR (400.13 MHz): δ 1.08 (s, 3H), 1.99 (s, 3H), 2.46 (s, 3H), 5.33 (d, J = 10.7 Hz, 1H), 5.50 (d, J = 17.5 Hz, 1H), 5.98 (dd, J = 10.7, 17.5 Hz, 1H), 6.85 (s, 1H), 7.09 (t, J =7.4 Hz, 1H), 7.27–7.31 (m, 2H), 7.47–7.49 (m, 2H). ¹³C NMR (100.62 MHz): δ 17.2, 17.3, 21.2, 65.8, 68.9, 113.9, 117.9, 118.5, 124.1, 128.2, 134.8, 136.7, 153.5, 168.6, 169.9. GC-MS (70 eV) m/z (rel int.): 298 (33, M⁺), 216 (100), 215 (55), 201 (29), 178 (91), 174 (45), 118 (31), 77 (43). IR (CHCl₃): 3066, 2980, 2930, 1750, 1600, 1500, ¹. HR-ESI-MS: m/z calcd for C₁₇H₁₉N₂OS: 1362 cm⁻ 299.1220, [M+H]⁺; found: 299.1219. Compound **20b**. Yield 176 mg (59%), oil. ¹H NMR (400.13 MHz): δ 1.55 (s, 3H), 2.08 (s, 3H), 2.45 (s, 3H), 4.95 (dd, J=3.7, 8.1 Hz, 1H), 5.35–5.38 (m, 2H), 6.81 (s, 1H), 7.09 (t, J=7.4 Hz, 1H), 7.27–7.31 (m, 2H), 7.46–7.48 (m, 2H). ¹³C NMR (100.62 MHz): δ 17.2, 17.3, 19.5, 65.7, 68.8, 114.17, 119.9, 118.4, 124.1, 128.9, 134.1, 136.6, 153.2, 168.8, 170.6. GC-MS (70 eV) m/z (rel int.): GC-MS (70 eV) m/z (rel int.): 298 (30, M⁺), 216 (76), 215 (44), 201 (35), 178 (100), 174 (37), 118 (25), 77 (34). IR (CHCl₃): 3066, 2980, 2930, 1750, 1600, 1500, 1362 cm⁻¹. HR-ESI-MS: m/z calcd for $C_{17}H_{19}N_2OS$: 299.1220, $[M+H]^+$; found: 299.1220.

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3-Aza-8,10-dioxa-bicyclo[5.2.1]decane (9-*exo* BTKa) carboxylic acid as a new reverse turn inducer: synthesis and conformational analysis of a model peptide

Dina Scarpi, Daniela Stranges, Andrea Trabocchi and Antonio Guarna*

Dipartimento di Chimica Organica 'U. Schiff', Università di Firenze, Via della Lastruccia 13, 50019 Sesto Fiorentino, Italy

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Abstract—Dipeptide isostere 5, belonging to the class of 9-*exo* BTKa, was synthesised starting from *R*,*R*-tartaric acid and 4-nitro-1-(3-nitrophenyl)butan-1-one. The nine-membered lactam showed interesting structural features and was inserted in a 5-residue model peptide. The conformational properties of this modified peptide have been studied by NMR and molecular modelling, indicating that compound 5 acted as a reverse turn inducer.

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1. Introduction

Reverse turns are structural motifs commonly found in proteins and bioactive peptides that play a central role as molecular recognition elements for many biological processes, by virtue of presenting up to four side chains in a well defined spatial arrangement.¹ In particular, β -turns consist of a tetrapeptide sequence (defined as i-i+1-i+2i+3) in a non-helical region in which the peptide chain direction is reversed. These turns are often stabilised by an intramolecular hydrogen bond between the carbonyl oxygen of the first residue (i) and the amide proton of the fourth one $(i+3)^2$ thus being a key template for the design of the socalled 'turn-mimetics' in the drug discovery area.³ During the last decade many efforts have been dedicated to synthesise new reverse turn inducers and to study the conformational preferences of turn-analogues within protein secondary structure models.⁴ In recent years, we have been developing a new class of aza-dioxa[3.2.1]bicyclic compounds named BTAa⁵ or BTKa,⁶ the synthesis of which is based on the combination of a tartaric acid derivative and either α -amino aldehydes or α -amino ketones.⁷ We have previously described the applications of these scaffolds as dipeptide isosteres when inserted in both cyclic⁸ and linear⁹ peptide sequences, acting as mimetics of i+1-i+2 central dipeptidic sequence of a typical β -turn motif. Moreover,

bicyclic proline mimetics have been explored as reverse turn inducers in model peptides.¹⁰

In a recent paper,¹¹ we described the synthesis of two classes of enantiopure molecular scaffolds, whose lactam structure formally derives from the coupling between tartaric acid and β - or γ -ketoamines. By analogy with the previously reported 7-*exo* BTKa,^{7c} we named these compounds as 8-*exo* and 9-*exo* BTKa, indicating the lactam size (eight- and nine-membered ring, respectively). The general structure is reported in Figure 1.

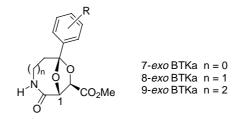


Figure 1. General structure of BTKa.

The ring enlargement of the rigid 7-*exo* BTKa scaffolds afforded, as expected, more flexible compounds that represent a new class of dipeptide isosteres, prone to take different conformations and potentially useful as turn inducers. Preliminary molecular modelling calculations¹¹ explained the experimental upfield shift of the carbomethoxy group observed in the ¹H NMR when passing from

Keywords: Conformational analysis; Peptidomimetic; Turn inducer; Tartaric acid.

^{*} Corresponding author. Tel.: +39 055 4573481; fax: +39 055 4573569; e-mail: antonio.guarna@unifi.it

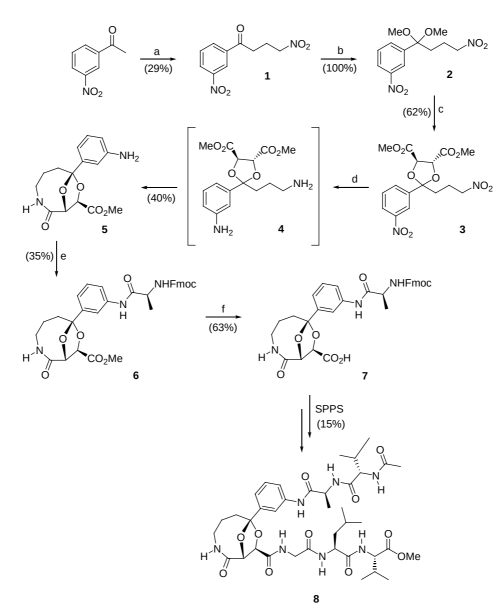
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2. Results and discussion

the 7-exo BTKa to the 8- and 9-exo ones¹² on the basis of the change of the average distance between this group and the aromatic ring on the bridgehead carbon, that decreases from 3.8 to 3.0 Å as the ring enlarges. This interesting structural characteristic prompted us to further investigate these molecules and their application as peptide turn inducers. In particular, we concentrated our attention on the 9-exo BTKa class (Fig. 1, n=2), that, as a consequence of the above-mentioned shortest distance of 3.0 Å between two easily functionalisable groups, could force two peptidic fragments to face each other, allowing the formation of a hydrogen bond network between them. We envisaged the ideal candidate for peptide modification in the compound bearing a *m*-NH₂ substituent on the aromatic ring. The 9-exo BTKa was inserted in the model sequence Ac-VA-BTKa-GLV-OMe and its effect on the peptide conformation is reported in detail in the next section.

2.1. Synthesis

The synthesis of the target 9-*exo* BTKa (Scheme 1) started from γ -nitroketone **1**, that was obtained from commercially available 3'-nitroacetophenone in 29% yield over three steps. Compounds **2** and **3** were straightforwardly obtained in quantitative and 62% yield, respectively, following the previously reported experimental procedures.¹¹ Reduction of the nitro groups was first performed by the reported method, that is, hydrogenation on Raney-Ni in methanol at room temperature. By analogy to the synthesis of other 9-*exo* BTKa, a high yield of the reaction was not expected; however, reduction took place but neither lactam **5** nor diamine **4** were recovered. After a few more experiments, lactam **5** was obtained by hydrogenation with ammonium



Scheme 1. (a) (i) (HCHO)_n, Me₂NH·HCl, EtOH, H⁺, reflux, 2 h; (ii) NaOH aq; (iii) CH₃NO₂, TRITON B, reflux, 2 h; (b) HC(OMe)₃, *p*-TsOH cat., MeOH, reflux, 16 h; (c) BF₃·Et₂O, EtOAc, 0 °C, 4 h; (d) NH₄HCO₂, Pd/C 10%, MeOH, reflux, 16 h; (e) Fmoc-Ala-OH (1 equiv), PyBROP (1 equiv), DIEA (1 equiv), CHCl₃, rt, 24 h; (f) LiOH aq (1.0 equiv), 1,4-dioxane/H₂O 1:1, 0 °C, 30'.

NH	CDCl ₃			CD ₃ CN	DMSO-d ₆	
	δ	$\Delta \delta / \Delta T$	δ	$\Delta \delta / \Delta T$	δ	$\Delta \delta / \Delta T$
Val-1	7.27	-6.83	6.82	-5.54	7.95	-4.61
Ala	6.92	-1.48	7.08	-6.07	8.25	-6.38
NH-3'	9.54	-5.64	8.81	-3.44	9.99	-5.17
Gly	7.10	-2.90	6.99	-2.27	7.78	-4.45
Leu	7.12	-5.64	6.69	-3.26	8.01	-4.64
Val-2	7.95	-5.64	7.08	-4.00	8.19	-7.21

Table 1. Temperature dependence of amide proton chemical shifts for 8

 δ are expressed in ppm and $\Delta \delta / \Delta T$ values in ppb/K.

formate over 10% Pd/C in refluxing methanol for 16 h (40% yield after purification). Compound **5** was then coupled with Fmoc-Ala-OH using PyBrop[†] as the activating agent and the resultant ester **6** was hydrolysed by LiOH in 1,4-dioxane/ water at 0 °C to afford the Fmoc-amino acid **7**.

Peptide Ac-Val-Ala-BTKa-Gly-Leu-Val-OMe (8) was prepared by means of solid-phase techniques using Fmoc protocol and a HMBA-AM polystyrene resin, that afforded the title peptide with the C-terminus protected as methyl ester by a nucleophilic cleavage. Fmoc-Ala-BTKa 7 was incorporated into the growing peptide in the third coupling step. All amide couplings were monitored with bromophenol blue as internal colorimetric indicator. Nucleophilic cleavage from the resin was achieved by trans-esterification, heating a suspension of the resin at 50 °C overnight in a 9:1 MeOH/triethylamine mixture. The crude peptide was purified by semi-preparative HPLC, giving pure 8 in 15% yield.

2.2. NMR studies in CDCl₃

Conformational studies on peptide 8 were performed by NMR, using firstly a relatively non-polar solvent (i.e., CDCl₃), and successively more competitive solvents such as CD_3CN and $DMSO-d_6$, to investigate the solvent effect on the conformational preference of 8. Solutions (4.2 mM) of 8 were used to achieve sufficient dilution to prevent aggregation. TOCSY and ROESY spectra were recorded to assign proton resonances and investigate both sequential and long-range NOE's that provide evidences of preferred conformations and give insight into stable reverse turn and sheet conformations.¹³ Temperature dependence experiments were carried out, since the amide proton chemical shifts are sensitive to temperature and dilution variations, thus giving further insight into the conformational preferences of peptides.¹⁴ The combination of chemical shift and $\Delta\delta/\Delta T$ coefficient of the amide protons provides information on the extent of hydrogen bonding.²

¹H NMR analysis of compound **8** in CDCl₃ solution was complicated by the flexibility and size of the peptidomimetic, and the attempted structural determination of **8** turned out to be complex, as spectral data of sufficient quality for structure determination were not obtained in CDCl₃. Moreover, signals in the ROESY spectrum indicated slow to intermediate exchange rates between several conformers. All amide values except NH-3' were found between 7 and 8 ppm, with Val-2 NH being the most deshielded (Table 1). A temperature dependent NMR experiment indicated the presence of an equilibrium between hydrogen bonded and non-hydrogen bonded states, as many amide protons showed large $\Delta \delta / \Delta T$ values. The low $\Delta \delta / \Delta T$ coefficient of Ala NH, compared with the other amide protons, indicated the presence of a hydrogen bonded environment for this amide proton in CDCl₃ solutions. The glycine amide coefficient also showed a relatively temperature stable chemical shift. In this case, the low coefficient may be due to the anti O-8 orientation of carbonyl group at C-9 of the scaffold rather than to the existence of a hydrogen bond. The high chemical shift of Val-2 NH in conjunction with the corresponding $\Delta \delta / \Delta T$ value suggested an equilibrium of conformers in which this amide proton experienced both non-hydrogen bonded states and hydrogen bonds with different carbonyl groups.

2.3. NMR studies in CD₃CN

Experiments carried out in CD₃CN showed marked differences, suggesting that a more competitive solvent induces peptide 8 to become more organised. As a consequence of the greater solvating effect of CD₃CN, though still remaining non-competitive relatively to hydrogen bonding, the chemical shift values of the amide protons experienced significant changes, except for glycine and alanine, that appeared slightly upfield and downfield relative to signals recorded in CDCl₃, respectively. Moreover, the same trend of temperature coefficients as observed for CDCl₃ solutions was followed, with the glycine amide proton showing the lowest $\Delta \delta / \Delta T$. In contrast the alanine NH signal increased significantly, confirming the poor hydrogen bonding character of this proton in CD₃CN, which is itself a moderate hydrogen bond acceptor. The amide NH-3' coefficient also suggested the presence of an equilibrium between hydrogen bonded and non-hydrogen bonded structures. NOE experiments suggested the existence of β -strand organisation of the main chain, as all the amino acids showed strong α , N(i, i+1) sequential peaks. Moreover, ROESY spectra of 8 showed some cross-peaks between protons on non-adjacent residues. These NOE's were indicative of an equilibrium between more equivalent conformations (Fig. 2). A strong cross-strand ROESY peak between glycine NH and H-2' clearly demonstrated the reverse turn inducing role of the scaffold. Moreover, other cross-strand ROESY peaks between Ala NH and Leu α -H, and between Val-2 NH and both Val-1 β - and γ -H confirmed the existence of a well-ordered reverse turn structure (Fig. 2, left). Interestingly, the presence of crossstrand ROESY peaks experienced by Ala α -H with Leu β and γ -H indicated the existence of a second conformer

 $^{^\}dagger$ Bromotripyrrolidinophosphonium hexafluorophosphate.

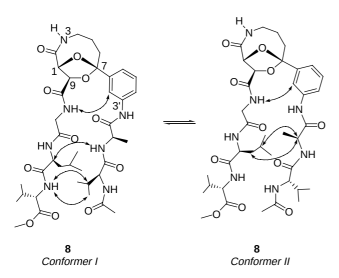


Figure 2. Peptide 8: the arrows indicate significant cross-strand ROESY correlations in CD₃CN.

(Fig. 2, right), in which the N-terminal half was found to be flipped, thus orienting Ala NH or Ala α -H inside the turn structure, respectively. These NOE interactions demonstrated that the scaffold **5** acts as a mimetic of the central dipeptidic unit of a β -turn, thus generating an unusual reverse turn. The existence of these two highly ordered structures was in agreement with $\Delta\delta/\Delta T$ data, indicating the presence of equilibrating weak hydrogen bonds. The absence of stable hydrogen bonds characteristic of reverse turn structures indicated that the principal driving force for reverse turn formation is a consequence of the structure of the scaffold. Further stabilisation is provided by intramolecular hydrogen bonding and hydrophobic interactions that contribute to the overall organisation of the peptide into β -strands conformations.

2.4. NMR studies in DMSO- d_6

Experiments carried out in DMSO- d_6 showed all the amide protons deshielded with respect to CDCl₃ solutions, as expected. Nevertheless, Val-2 NH only showed a small chemical shift variation, changing from 7.96 to 8.19. Thus, the high chemical shift observed in CDCl₃ solution, in conjunction with low influence of solvent composition on chemical shift, indicated the Val-2 amide bond to be engaged in different hydrogen bonds, although its high temperature coefficients in both solvents were indicative of the presence of non-hydrogen bonded states, and of the absence of a specific hydrogen bond of significant strength. Gly NH showed a small temperature coefficient compared to the other protons, confirming the hypothesis that the role of the scaffold was to control the position of the adjacent species. Moreover, Val-1 NH proved to lower its coefficient on moving to a more competitive solvent, probably due to the presence of more structured conformations in the more highly solvating system. The existence of a large temperature coefficient did not prevent the hypothesis of multiple hydrogen bonds, and was in agreement with the presence of two or more equilibrating structures (due in particular to flipping of the N-terminal chain), which in turn generated different patterns of hydrogen bonds. This conformational equilibrium was particularly evident when

CD₃CN and DMSO- d_6 were used as solvents. ROESY experiments in DMSO- d_6 confirmed the β -strand organisation of the reverse turn peptide, as suggested by strong α ,N(i, i+1) sequential peaks experienced by all the amino acids. Moreover, cross-strand ROESY peaks between Val-2 NH and both Val-1 β - and γ -H were also maintained in this solvent, confirming the turn structure.

2.5. Molecular modelling

Molecular modelling using AMBER* as a force field¹⁵ was carried out to gain further insight into the conformational preferences of peptide **8**. Full unconstrained Monte Carlo conformational search¹⁶ using CHCl₃ as explicit solvent resulted in all the conformers having a marked tendency of adopting a reverse turn conformation, in which the scaffold occupies the central turn position.

The distance d and the virtual torsion angle β were computed to investigate the turn propensity The distance dbetween the C- α of the first and fourth residue of a β -turn is diagnostic of the presence of a reverse turn when its value is lower than 7 Å, ^{1a} and the virtual torsion angle β is indicative of a reverse turn when it assumes a value within the range of $0 \pm 30^{\circ}$.¹⁷ For peptide **8**, the chain reversing property of this unusual turn structure was assessed computing parameter das the distance between Gly C- α and C-3^{*i*}, and considering β as the dihedral angle formed by C-3'-C-5-C-9-Gly C- α . In all the conformers, d and β values fell within the diagnostic range for a turn structure, confirming the hypothesis of the scaffold acting as a nucleator of tight reverse turns. All the conformers produced by the Monte Carlo calculation showed that the turn structure of 8 was stabilised by two or more hydrogen bonds in a different fashion, producing two main groups of conformers, in agreement with the NMR data (Fig. 3). The first group of conformers included structures A, C and D, all having in common the same orientation of the two peptidic halves. The second group, represented by structure B, showed the N-terminal main chain flipped with respect to the former one. Specifically, the first group of conformers encompassing the global minimum conformer (E = -545.2 kJ/mol) were found in 8% abundance, and showed two hydrogen bonds, between Val-2 NH and Ala CO, and between Ala NH and Val-2 CO, thus giving a bent turn structure stabilised by a distorted β -sheet structure (Fig. 3, structure A). An additional set of structures belonging to this group were found in 36% abundance (Fig. 3, structure C), and displayed a twisted turn conformation stabilised by Ala NH and Gly CO, and Val-2 NH and Ala CO hydrogen bonds. A higher energy conformer was present in 15% abundance (Fig. 3, structure D), and showed a distorted β -strand structure of peptide halves stabilised by three hydrogen bonds formed by Gly NH and Ala CO, Ala NH and Gly CO, and between Val-2 NH and acetyl CO.

The second group of structures (18%) was represented by the second conformer in order of increasing energy (E = -542.5 kJ/mol), which showed a well-organised sheet structure stabilised by three hydrogen bonds: NH-3' and Gly CO, Val-2 NH and Val-1 CO, Val-1 NH and Val-2 CO (Fig. 3, structure B). In this structure, NH-3' was found to be engaged in a hydrogen bond with Gly CO, in agreement

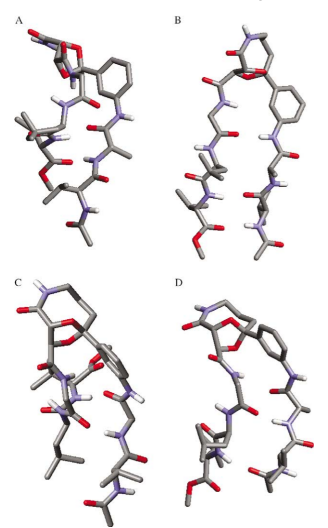


Figure 3. Low energy conformers obtained by Monte Carlo calculation.

with the lower $\Delta\delta/\Delta T$ coefficient in CD₃CN, confirming the hypothesis that such a structure was particularly relevant in this solvent (see also ROESY correlations as in Fig. 2, right). All the low energy conformers displayed a β -sheet structure with the scaffold acting as external reverse turn inducer. The presence of different hydrogen bonds experienced by Ala and Val-2 amide protons were in agreement with NMR data. In particular, the low $\Delta\delta/\Delta T$ value found in CDCl₃ for Ala NH was compatible with its hydrogen bonding character in a non-competitive solvent like chloroform. In the case of Val-2 NH, however, the high chemical shift and $\Delta\delta/\Delta T$ value of Val-2 amide proton in CDCl₃ and the slight chemical shift variation from CDCl₃ to DMSO-d₆ suggested an equilibrium between hydrogen bonded and non-hydrogen bonded states in all the solvents considered.

Molecular dynamic calculations over 1 ns were carried out on all of the four structures found by Monte Carlo calculations to gain information on the vicinity of local minima of found structures, and to verify the flexibility of the structures found in the conformational search. The global minimum conformer showed poor stability under MD calculations, as the hydrogen bond percentages dropped to 4.5%, giving an open turn structure, in agreement with the hypothesis of equilibrating structures, as suggested by NMR data. The population of hydrogen bonded structures was found at a lower percentage with respect to Monte Carlo calculation, confirming the high flexibility of peptide **8** and agreeing with the presence of more structures, though *d* and β values lowered only to about 60%, demonstrating the strong reverse turn propensity of model peptide **8**.

3. Conclusions

In this work, we realised the synthesis of a new dipeptide isostere belonging to the class of 9-*exo* BTKa, starting from a suitable aromatic γ -nitroketone. The title compound (5) was inserted in a linear peptide chain and the structural effects on the conformation of the model were studied.

We verified that compound 5 generated a tight reverse turn stabilised by its rigid structure and by the preferred anti orientation of carbonyl group at C-9 position. Moreover, the ring size allowed the aromatic ring to bend towards the carbonyl at C-9, thus producing an unusual reverse turn inducer. Conformational analysis by NMR in different solvent systems showed the existence of equilibrating structures, stabilised by different patterns of hydrogen bonds or simply by hydrophobic interactions. Sequential and cross-strand ROESY peaks revealed well-ordered turn structures, especially in more competitive solvents such as CD_3CN and $DMSO-d_6$. Moreover, when moving to a more competitive solvent, although the labile hydrogen bonds were disrupted, the β -sheet organisation of the reverse turn peptide was further stabilised, as was particularly evident in ROESY spectra in CD₃CN. The absence of stable hydrogen bonds, characteristic of reverse turn structures, indicated that the principal driving force for reverse turn formation is due to the particular structural form of the scaffold. Further stabilisation is provided by weak intramolecular hydrogen bonding and hydrophobic interactions that contribute to the overall organisation of peptide into β -strand conformations. Molecular modelling confirmed the existence of equilibrating structures, specifically due to a flip of N-terminal chain, which allowed the arrangement of the amide proton NH-3' to the inside or outside of the turn. Finally, molecular dynamic calculations confirmed the low stability of hydrogen bond networks present in the low energy conformers, in accordance with NMR data. Introduction of chemical functionalities on amide at position 3 of the scaffold and on the aromatic ring might allow side chains isosteres to be appended at the i+1 and i+2 positions of the turn.

4. Experimental

4.1. General

Melting points are uncorrected. Chromatographic separations were performed under pressure on silica gel by flashcolumn techniques; R_f values refer to TLC carried out on 25-mm silica gel plates (Merck F254), with the same eluent as indicated for the column chromatography. IR spectra were recorded with a Perkin-Elmer 881 spectrophotometer in CDCl₃ solution. Apart from peptide **8**, ¹H NMR (200 MHz) and ¹³C NMR (50.33 MHz) spectra were recorded with a Varian XL 200 instrument in CDCl₃ solution. NMR spectra of peptide **8** were performed on a Varian MercuryPlus 400 spectrometer operating at 400 MHz for ¹H. The spectra were obtained in 4.2 mM CDCl₃ or CD₃CN solution where aggregation was not significant. One-dimensional ¹H NMR spectra for determining temperature coefficients were obtained at 298–328 K with increments of 5 K. Sample temperatures were controlled with the variable-temperature unit of the instrument. Complete proton resonance assignments were made with the aid of gCOSY, TOCSY, HSQC and ROESY experiments.

Mass spectra were carried out by EI at 70 eV, unless otherwise stated, on 5790A-5970A Hewlett-Packard and QMD 1000 Carlo Erba instruments. Microanalyses were carried out with a Perkin-Elmer 2400/2 elemental analyser. Optical rotations were determined with a JASCO DIP-370 instrument.

All the solid-phase reactions were carried out on a shaker, using solvents of HPLC quality. HPLC purification was performed with an HPLC system equipped with semipreparative C-18 10 μ m, 250×10 mm, reverse-phase column using H₂O–CH₃CN eluent buffered with 0.1% TFA. Peptide **8** was characterised by ESI-MS, 2D-NMR and HPLC system equipped with an analytical C-18 10 μ m, 250×4.6 mm, reverse-phase column.

4-Nitro-1-(3-nitrophenyl)butan-1-one (1) was synthesised as already reported.¹⁸

4.1.1. 1-(1,1-Dimethoxy-4-nitrobutyl)-3-nitrobenzene (2). Synthesised using the procedure previously reported for closely related compounds,¹¹ starting from **1** (1.56 g, 6.55 mmol) an refluxing for 16 h. After filtration and evaporation of the solvent, crude **2** was obtained in quantitative yield and used in the next step without further purification.

Compound **2**. Yellow oil. ¹H NMR δ (ppm): 8.36 (s, 1H), 8.20 (d, J=8.0 Hz, 1H), 7.79 (d, J=7.8 Hz, 1H), 7.57 (t, J=7.8 Hz, 1H), 4.25 (t, J=6.6 Hz, 2H), 3.17 (s, 6H), 2.05–1.97 (m, 2H), 1.74–1.57 (m, 2H). ¹³C NMR δ (ppm): 148.4 (s), 142.5 (s), 132.9 (d), 129.4 (d), 123.2 (d), 122.2 (d), 102.1 (s), 74.8 (t), 48.9 (q, 2C), 33.6 (t), 21.5 (t). MS *m/z* (%): 284 (M⁺, 1).

4.1.2. (4*R*,5*R*)2-(3-Nitrophenyl)-2-(3-nitropropyl)-[1,3]dioxolane-4,5-dicarboxylic acid dimethyl ester (3). Synthesised using the procedure previously reported for closely related compounds,¹¹ starting from 2 (1.8 g, 6.33 mmol). After purification by chromatography (eluent: EtOAc/petroleum ether, 1:4, R_f =0.25), pure 3 was obtained as pale yellow oil (1.56 g, 62%).

Compound 3. $[\alpha]_D^{25}$ +81.9 (c 1.0, CHCl₃). ¹H NMR δ (ppm): 8.36 (s, 1H), 8.20 (d, J=8.0 Hz, 1H), 7.83 (d, J=8.0 Hz, 1H), 7.55 (t, J=8.0 Hz, 1H), 4.81 (AB system, J_{AB} =5.6 Hz, 2H) 4.48 (t, J=6.6 Hz, 2H), 3.86 (s, 3H), 3.58 (s, 3H), 2.24–2.04 (m, 4H). ¹³C NMR δ (ppm): 168.6 (s), 168.5 (s), 148.1 (s), 142.6 (d), 132.0 (d), 129.4 (d), 123.8

(d), 120.9 (d), 113.0 (s), 77.7 (d), 76.5 (d), 74.9 (t), 53.0 (q), 52.7 (q), 36.9 (t), 21.2 (t). MS m/z (%): 399 (M⁺ + 1, 1), 310 (M⁺ - (CH₂)₃NO₂, 100). Anal. Calcd for C₁₆H₁₈N₂O₁₀: C, 48.25; H, 4.55; N, 7.03. Found: C, 48.36; H, 4.52; N, 6.96.

4.1.3. (1*R*,7*R*,9*R*)7-(3-Aminophenyl)-2-oxo-8,10-dioxa-3azabicyclo[5.2.1]decane-9-carboxylic acid methyl ester (5). Ammonium formate (1.58 g, 25.1 mmol) and Pd/C 10% (150 mg) were added to a solution of 3 (500 mg, 1.26 mmol) in MeOH (300 mL). The mixture was heated under reflux for 16 h and, after cooling, filtered through a Celite layer and finally evaporated to give crude 5, that was purified by chromatography (eluent: EtOAc/petroleum ether 0.1% Et₃N, 3:1, R_f =0.17) affording pure 5 (154 mg, 40%) as a white solid.

Compound **5**. $[\alpha]_{D}^{25} - 12.3$ (*c* 1.0, CHCl₃). ¹H NMR δ (ppm): 7.11 (t, *J*=8.0 Hz, 1H), 6.87 (s, 1H), 6.82 (d, *J*=2.2 Hz, 1H), 6.61 (dd, *J*=8.0, 1.4 Hz, 1H), 6.52 (br s, 1H), 5.30 (d, *J*=1.8 Hz, 1H), 4.94 (d, *J*=1.8 Hz, 1H), 4.15–4.05 (m, 1H), 3.46 (s, 3H), 3.40–3.20 (m, 1H), 2.21–2.13 (m, 1H), 2.07–1.84 (m, 3H). ¹³C NMR δ (ppm): 174.0 (s), 169.4 (s), 146.0 (s), 142.5 (s), 128.9 (d), 115.5 (d), 115.0 (d), 114.9 (s), 112.0 (d), 79.5 (d), 78.7 (d), 52.4 (q), 42.1 (t), 36.2 (t), 25.6 (t). MS *m*/*z* (%): 306 (M⁺, 30), 247 (M⁺ – CO₂CH₃, 9), 137 (82), 120 (100). Anal. Calcd for C₁₅H₁₈N₂O₅·2H₂O: C, 52.63; H, 6.48; N, 8.18. Found: C, 52.87; H, 6.50; N, 8.29.

4.1.4. (1R,2S,7R,9R)7-{3-[2-(9H-Fluoren-9-ylmethoxycarbonylamino)propionylamino]phenyl}-2-oxo-8,10dioxa-3-azabicyclo[5.2.1]decane-9-carboxylic acid methyl ester (6). Fmoc-Ala-OH (312 mg, 1.08 mmol), PyBrop (468 mg, 1.08 mmol) and DIEA (344 µL, 2.16 mmol) were added to a solution of 5 (300 mg, 1.08 mmol) in anhydrous CHCl₃ (5 mL). The mixture was left at room temperature, under stirring and nitrogen atmosphere. After 24 h AcOEt (20 mL) was added and the organic phase was washed with water $(2 \times 10 \text{ mL})$, satd NaHCO₃ (2×10 mL), and dried over Na₂SO₄. After filtration and evaporation of the solvent, the obtained crude 6 was purified by flash chromatography (eluent: AcOEt/petroleum ether, 3:1; $R_f = 0.22$), affording pure 6 (230 mg, 36%) as a yellowish solid.

Compound **6**. Mp 154–155 °C. $[\alpha]_{D}^{25}$ –44.3 (*c* 0.25, CH₃OH). ¹H NMR δ (ppm): 8.46 (s, 1H), 7.78–7.74 (m, 12H), 6.58 (br s, 1H), 5.47 (br s, 1H), 5.39 (s, 1H), 4.96 (s, 1H), 4.46–4.40 (m, 3H), 4.22 (t, *J*=7.0 Hz, 1H), 4.19–4.00 (m, 1H), 3.39 (s, 3H), 3.20–3.09 (m, 1H), 2.17–2.06 (m, 1H), 1.95–1.83 (m, 3H), 1.48 (d, *J*=7.0 Hz, 3H). ¹³C NMR δ (ppm): 173.9 (s), 170.7 (s), 169.2 (s), 156.4 (s), 143.6 (s, 2C), 142.5 (s), 141.3 (s, 2C), 137.5 (d), 128.7 (d), 127.8 (d, 2C), 127.1 (d, 2C), 125.0 (d, 2C), 121.2 (d), 120.0 (d, 2C), 119.9 (d), 116.8 (d), 114.5 (s), 79.6 (d), 78.8 (d), 67.3 (t), 52.3 (q), 51.2 (d), 47.0 (t), 42.1 (d), 36.9 (t), 29.7 (t), 25.7 (q). MS *m*/*z* (%): 599 (M⁺, 20). Anal. Calcd for C₃₃H₃₃N₃O₈·2H₂O: C, 62.35; H, 5.87; N, 6.61. Found: C, 62.62; H, 5.86; N, 6.69.

4.1.5. (1*R*,2*S*,7*R*,9*R*)7-{3-[2-(9*H*-Fluoren-9-ylmethoxycarbonylamino)propionylamino]phenyl}-2-oxo-8,10dioxa-3-azabicyclo[5.2.1]decane-9-carboxylic acid (7). A 0.40 M solution of LiOH in H₂O (0.25 mL) was added dropwise to a solution of **6** (62 mg, 0.10 mmol) in 1,4dioxane (0.5 mL) and water (0.5 mL) cooled at 0 °C. The resulting mixture was stirred at 0 °C for 30 min and then 5% KHSO₄ was added reducing the pH to 5. After concentration to a small volume, the product was extracted with CHCl₃ (4×10 mL), adjusting the pH to 5 after every extraction. The combined organic phases were dried over Na₂SO₄. After filtration and evaporation of the solvent, pure **7** (38 mg, 63%) was obtained as a yellowish solid.

Compound 7. $[\alpha]_D^{25}$ –48.8 (c 0.3, CHCl₃). ¹H NMR δ (ppm): 9.13 (s, 1H), 7.76–7.67 (m, 2H), 7.57–7.40 (m, 3H), 7.40–7.14 (m, 7H), 6.25 (m, 1H), 5.23 (s, 1H), 4.89 (s, 1H), 4.42 (m, 1H), 4.29–4.11 (m, 4H), 3.17–2.74 (m, 1H), 2.03–1.45 (m, 4H), 1.37 (m, 3H). ¹³C NMR δ (ppm): 175.1 (s), 172.0 (s), 171.7 (s), 156.4 (s), 143.8 (s), 143.5 (s, 2C), 141.2 (s, 2C), 137.7 (s), 128.7 (d), 127.7 (d, 2C), 127.0 (d, 2C), 125.1 (d, 2C), 121.2 (d), 120.1 (d), 119.9 (d, 2C), 117.0 (d), 114.5 (s), 79.2 (d), 78.7 (d), 67.0 (t), 51.3 (d), 46.9 (t), 41.4 (d), 35.7 (t), 29.6 (t), 24.8 (q). MS *m*/*z* (%): 586 (M⁺, 10). Anal. Calcd for C₃₂H₃₁N₃O₈·2H₂O: C, 61.83; H, 5.67; N, 6.76. Found: C, 61.98; H, 5.72; N, 6.58.

4.2. Peptide synthesis

Peptide **8** was prepared by means of solid-phase techniques using a HMBA-AM polystyrene resin (100 mg, 0.08 mmol). A five equivalent excess of Fmoc amino acids and DIPC/ HOBt carboxylic-activating mixture were used throughout the synthesis, and DMF was used as solvent. Compound **7** was used in 2 equiv excess. Final acetylation was performed with Ac₂O in DMF using catalytic 4-dimethylaminopyridine. All amide couplings were monitored with bromophenol blue as an internal colorimetric indicator.¹⁹ Nucleophilic cleavage from the resin was achieved by transesterification, heating at 50 °C overnight a suspension of the resin in a 9:1 MeOH/triethylamine mixture. Crude peptide was purified by semi-preparative HPLC using 10–90% ACN/55 min as gradient, giving pure **8** as a white solid (9.5 mg, 15%).

Compound 8. $t_{\rm R}$ =20.8 min (91% HPLC purity) using 0% ACN/5 min, 0–10% ACN/5 min, then 10–90% ACN/20 min as gradient. ESI-MS m/z (%): 788.27 (M⁺+1, 35), 810.46 (M⁺+Na, 100), 826.40 (M⁺+K, 45). ¹H and ¹³C NMR data are shown in Table 2.

4.3. Computational methods

Molecular mechanics calculations were carried out on a SGI IRIX 6.5 workstation, using MacroModel (v6.5) molecular modelling software,²⁰ with AMBER* as a force field¹⁵ and the implicit chloroform GB/SA solvating system.²¹ Monte Carlo conformational search¹⁶ was carried out without imposing any constraint and including amide bonds among all rotatable bonds. Two thousand structures were generated and minimised until the gradient was less than 0.05 kJ/Å/ mol using the TNCG gradient implemented in Macro-Model.²² All the conformers having an energy of 6 kcal/mol above the global minimum conformer were discarded. Molecular dynamic (MD) hybrid simulation algorithm was used to assess stability of low energy conformers. AMBER*

Table 2. Proton and carbon chemical shifts of peptide 8 (δ values are expressed in ppm, and in parentheses are reported J values in Hz)

	¹ H (DMSO- d_6)	1 H (CD ₃ CN)	¹ H (CDCl ₃)	¹³ C (CDCl ₃)
Ac	1.92	1.99	2.08	22.9
/al-1 NH	7.95 (d, 8.2)	6.82 (d, 6.8)	7.27	_
/al-1H-α	4.20 (t, 6.8)	3.99 (dd, 6.1; 6.1)	4.33	58.9
′al-1H-β	2.00	2.02	2.04	31.4
al-1H-γ	0.90	0.87 (d, 6.9)	0.95	19.4
la NH	8.25 (d, 7.0)	7.08 (d, 6.2)	6.79	—
la α-H	4.42	4.39 (dq, 7.1; 6.2)	4.82	49.8
la β-H	1.34 (d, 7.0)	1.31 (d, 7.1)	1.46 (d, 6.6)	19.3
5TK H-1	4.99 (d, 2.7)	4.99 (d, 2.7)	5.31	80.0
3TK H-3	7.91	6.37 (t, 7.1)	6.20	—
STK H-4	4.02, 3.09	4.00 and 3.07	4.20 and 3.19	40.9
5TK H-5	1.60, 2.07	1.65 and 2.03	2.13 and 1.88	25.3
3TK H-6	2.0, 2.07	2.05	2.46 and 2.13	22.2
3TK H-9	4.88 (d, 2.8)	4.80 (d, 2.7)	5.01	79.8
5TK H-2'	7.74	7.71	7.62	118.5
TK NH-3'	9.99	8.81	9.49	_
3TK H-4′	7.70 (d, 8.0)	7.19 (d, 8.0)	7.26	121.0
TK H-5'	7.33 (t, 7.8)	7.26 (t, 7.8)	7.36 (dd, 8.0; 7.6)	129.0
TK H-6′	7.24 (d, 7.9)	7.73 (d, 8.0)	8.22	104.3
ly NH	7.78 (t, 5.6)	6.99 (t, 5.9)	7.10	_
ily α-H	3.73 (dd, 16.6, 6.0)	3.54 (t, 5.9)	4.33 and 3.73	42.4
	3.53 (dd, 16.8, 4.9)			
.eu NH	8.01 (d, 8.2)	6.69 (d, 8.0)	7.10	_
eu α-H	4.43	4.35	4.84	52.1
.eu β-H	1.36, 1.60	1.45	1.50 and 1.35	42.1
.eu γ-H	1.60	1.65	1.58	24.7
.eu δ-H	0.90	0.79 (d, 7.1)	0.89 (d, 5.9)	22.8 and 19.1
al-2 NH	8.19 (d, 8.0)	7.08 (d, 6.2)	8.02	_
/al-2H-α	4.14 (t, 7.8)	4.19 (dd, 8.3; 6.2)	4.55 (dd, 8.7; 5.8)	57.4
∕al-2H-β	2.07	2.01	2.10	31.7
al-2H-γ	0.91	0.79 (d, 7.1)	0.89 (d, 5.9)	18.9
OMe	3.64	3.58	3.75	52.2

Experiments in CDCl₃ solution have been carried out at 303 K, and at 298 K for DMSO-d₆ and CD₃CN solutions.

was used as force field, as implemented in Macromodel (v6.5). A time step of 0.75 fs was used and the total simulation was 2000 ps; samples were taken at 1 ps intervals, yielding 2000 conformations for analysis.

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Formation of polysubstituted chlorocyclopropanes from electrophilic olefins and activated trichloromethyl compounds

Sylvain Oudeyer,^a Eric Léonel,^a Jean Paul Paugam,^{a,*} Christine Sulpice-Gaillet^b and Jean-Yves Nédélec^a

^aLaboratoire d'Electrochimie, Catalyse et Synthèse Organique, UMR 7582, Institut des Sciences Chimiques Seine-Amont, IFR 1780,

CNRS-Université Paris XII, 2, rue Henri Dunant, B.P. 28, F-94320 Thiais, France

^bLaboratoire de Recherche sur les Polymères, UMR 7581, Institut des Sciences Chimiques Seine-Amont, IFR 1780,

CNRS-Université Paris XII, 2, rue Henri Dunant, B.P. 28, F-94320 Thiais, France

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Abstract—Chlorocyclopropanes and bicyclic chlorocyclopropanes are prepared in non basic conditions by electroreductive or Mgpromoted Barbier activation of PhCCl₃ or Cl₃CCO₂Me in the presence of acyclic or cyclic α,β -unsaturated carbonyl compounds. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Cyclopropane containing molecules usually display interesting specific structural and physico-chemical properties. The presence of substituents on the C3 ring enables further transformations such as functional group interconversions or couplings with other molecules. Thus, 1-chlorocyclopropanecarboxylic acids are precursors of various aminocyclopropanecarboxylic acids^{1a,b} known for their biological activity² whereas 2-chlorocyclopropanecarboxylic acids are precursors of agrochemicals,³ and have also been used recently in the synthesis of Callipeltoside A, a novel antitumor agent, with the aim of elucidating its structure and notably the C-20 and C-21 configurations.⁴

The formation of polysubstituted chlorocyclopropanes from the coupling of acyclic α , β -unsaturated esters or cyclic α , β -unsaturated ketones with α , α -dichlorocarbanions, or equivalent nucleophilic organometallic species stabilized by an electron withdrawing group such as CO₂R or Ph, has already been reported in the literature. These nucleophilic intermediates are generated either by basic treatments (i.e., sodium hydride,⁵ LDA,⁶ electrogenerated bases,⁷ two-phase-solid–liquid system⁸ or LiHMDS-DBU⁹) of alkyl dichloroacetates and α , α -dichlorotoluene, or by an oxidative addition of a carbon–chlorine bond of the

* Corresponding author. Fax: +33 1 49781148;

e-mail: paugam@glvt-cnrs.fr

corresponding trichloromethyl compounds (Cl₃C–Y: Y = CO_2R , Ph) onto a soluble Cu(0)–isonitrile complex.¹⁰ These preparations of chlorocylopropanes involve either a conjugate nucleophilic addition followed by subsequent ring closure (MIRC reaction^{11a,b}) or carbenoid intermediates. Cyclocondensation to olefins is also mentioned with the ambiphilic chloroaryl carbenes photolytically generated from 3-chloro-3-aryldiazirines.¹² Moreover it must be noted that substituted 1-chlorocyclopropanecarboxaldehydes, precursors of methyl 1-chlorocyclopropanecarboxylates are synthesized via a semi-benzilic Favorski rearrangement of substituted 2,2-dichlorocyclobutanols obtained by reduction of the corresponding cyclobutanones.¹³

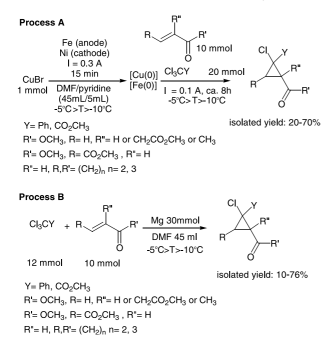
We have already investigated the synthesis of methyl 2,2diphenylcyclopropanecarboxylates and of 2-acyl-1,1diphenylcyclopropanes.^{14a-c} We have notably reported two methods: one is an indirect electroreductive coupling between dichlorodiphenylmethane and cyclic or acyclic α,β -unsaturated carbonyl compounds (referred to below as process A),^{14a,b} whereas the other one is a Mg-mediated Barbier type reaction in DMF (referred to below as process B).^{14c} This last route uses the same couples of reagents as those involved in process A, but it does not apply to α,β -unsaturated methyl ketones.

2. Results and discussion

In this paper, we report the preparation of polysubstituted chlorocyclopropanes from α , β -unsaturated acyclic esters or

Keywords: Chlorocyclopropanes; Mg-Barbier activation; Electro-organic synthesis; Carbenoids; Bicyclic compounds.

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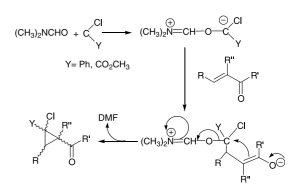
Scheme 1.

from cyclic α , β -unsaturated ketones and methyl trichloroacetate or α , α , α -trichlorotoluene (Scheme 1). It offers the opportunity to use and study both methods (processes A and B) and to compare their respective advantages and limitations, which proved to be rather complementary. The results are listed in Table 1.

These results first show that both methods generate nucleophilic intermediates, which add more or less efficiently to the olefin depending on its nature. More interestingly, these two methods are complementary. Thus, methacrylic acid esters show low reactivity in the electrochemical process (A) while yields obtained from the chemical method (B) are high (Table 1, entries 5 and 6). Such behaviour has already been observed with crotonic and methacrylic acids esters in other electrochemical reactions.¹⁶ On the contrary, yields are higher from the electrochemical method than from the chemical one when maleic or fumaric acid esters are involved (Table 1, entries 7–10). This may indicate the occurrence, in process B, of side reactions at the olefins due to their reducibility, whereas in the electrochemical process, the cathode potential is selfcontrolled according to the most easily reduced species, in this case the copper salts. All the other cases studied gave similar results from both methods.

The mechanisms involved in either process have not been fully elucidated so far. The occurrence of a non complexed carbene species can, however, be ruled out in both cases, due notably to the absence of stereocontrol in the ring formation (Table 1, entries 7, 9 and 8, 10). In addition, would the carben be formed (chlorophenylcarbene and chloromethoxycarbonylcarbene) it would be rather electrophilic, as described in the literature,^{12a,b,17,18} and should therefore react with electron-rich olefins like tetramethyl-ethylene, or cyclohexene, which has never been observed.

In the Mg-Barbier type process (B), a route via α , α -dichloromagnesium compounds, which are known to lose



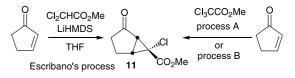
Scheme 2.

rapidly MgX₂ to form carbene intermediates,¹⁹ is not likely since no reaction was observed in the presence of nucleophilic olefins. So, we think that a first formed carben species reacts with DMF to form a nucleophilic intermediate in a process similar to the formation of the DMF–SOCl₂ complex described by Newman²⁰ (Scheme 2). The role of DMF is even crucial in this process. Indeed, very surprisingly, no reaction occurred in diethylether or in THF instead of DMF as solvent. On the contrary, addition of an equal amount of DMF to an ether solution of PhCCl₃ and methyl acrylate induced the cyclopropanation to start.

With reference to the complementarity of both processes (A and B), it is clear that they do not involve the same type of nucleophilic species derived from the trichloromethyl compounds. In the electrochemical process (A), the reactive intermediate could be a copper–iron bi-metallic nucleophilic complex, which is not yet identified.

In the presence of acyclic α , β -unsaturated esters, chlorocyclopropanes are prepared, according to both methods, with a low to moderate diastereoselectivity (Table 1, entries 1–6) but, when cyclic enones are used as electrophilic olefins, the diastereoselectivity of the cyclopropanation becomes very high (Table 1, entries 11–14): only one of the two possible structures (*endo*-chlorine or *exo*-chlorine adduct) is obtained.

We have assigned to the compound **11** an *endo*-chlorine structure by comparison with the results obtained by Escribano et al.⁹ Actually, whatever the route used (process A or B, or Escribano's process⁹) (Scheme 3), the same bicyclic compound is formed, as determined by GC-analysis, and from the ¹H and ¹³C NMR spectra.



Scheme 3.

The *endo*-chlorine structure was established by Escribano⁹ from X-ray diffraction experiments. Our 1D ¹H NOE-Difference NMR experiments, using selective excitation with a shaped pulse (gradient version) on the methoxy group, are consistent with the assignment given by Escribano. Indeed, the NOE effect (Fig. 1) is mainly seen at the H-1 and H-5 bridge-head protons. However, our measurement of the ³J (¹H–¹³C) coupling constant between

Table 1. Formation of polysubstituted chlorocyclopropanes by electroreductive or Mg-promoted coupling of α , β -unsaturated carbonyl compounds and α , α , α -trichloromethyl derivatives (Cl₃C–Y)

Entry	α,β-Unsaturated carbonyl compound ^a E_{red} (V/sce) ^b	Cl ₃ CY	Polysubstituted chlorocyclopropane ^a	n	Process A electrochemical process isolated yield (%)	Process B chemical process isolated yield (%)
1	E (-2.15)	Cl ₃ CCO ₂ CH ₃	CI	1	70 R*S*/R*R* 17/83	76 R*S*/R*R* 7/93
2	E (-2.15)	PhCCl ₃	CI Ph	2	35 R*S*/R*R* 60/40	68 <i>R*S*/R*R*</i> 57/43
3	E (-2.05)	Cl ₃ CCO ₂ CH ₃		3	57 R*S*/R*R* 30/70	65 R*S*/R*R* 28/72
4	E (-2.05)	PhCCl ₃	CI Ph E E	4	41 <i>R*S*/R*R*</i> 35/65	57 <i>R*S*/R*R*</i> 36/64
5	E (-2.30)	Cl ₃ CCO ₂ CH ₃	CI	5	<10 ^c	70 R*S*/R*R* 45/55
6	E (-2.30)	PhCCl ₃	CI Ph	6	<10 ^c	73 R*S*/R*R* 25/75
7	EE (-1.60)	Cl ₃ CCO ₂ CH ₃	E	7	67 <i>R*R*</i>	33 <i>R*R*</i>
8	EE (-1.60)	PhCCl ₃	E ^w E	8	52 <i>R*R*</i>	23 <i>R*R</i> *
9	E (-1.45)	Cl ₃ CCO ₂ CH ₃	E	7	40 <i>R*R*</i>	24 <i>R*R*</i>
10	E (-1.45)	PhCCl ₃	E ^{wi}	8	46 <i>R*R*</i>	10 R*R *
11	O (-2.15)	Cl ₃ CCO ₂ CH ₃	O E	9	58 1 <i>RS</i> ,6 <i>RS</i> ,7 <i>RS</i>	53 1 <i>RS</i> ,6 <i>RS</i> ,7 <i>RS</i>
12	(-2.15)	PhCCl ₃	O Ph	10	30 1 <i>RS</i> ,6 <i>RS</i> ,7 <i>RS</i>	50 ^d 1 <i>RS</i> ,6 <i>RS</i> ,7 <i>RS</i>

Entry	α,β-Unsaturated carbonyl compound ^a E_{red} (V/sce) ^b	Cl ₃ CY	Polysubstituted chlorocyclopropane ^a	п	Process A electrochemical process isolated yield (%)	Process B chemical process isolated yield (%)
13	O (-2.20)	Cl ₃ CCO ₂ CH ₃	CI E	11	30 1 <i>RS</i> ,5 <i>RS</i> ,6 <i>RS</i>	40 ^d 1 <i>RS</i> ,5 <i>RS</i> ,6 <i>RS</i>
14	O (-2.20)	PhCCl ₃	O Ph	12	20 1 <i>RS</i> ,5 <i>RS</i> ,6 <i>RS</i>	40 ^d 1 <i>RS</i> ,5 <i>RS</i> ,6 <i>RS</i>

^a $E = CO_2CH_3$.

^b See Ref. 15.

^c Determined by GC without internal standart.

^d Reagents ratio: activated olefin/ α , α , α -trichlorotoluene, 20 mmol/10 mmol.

the bridge-head protons and the carbon of the carbonyl of the C-6 methyl ester substituent gives a value of 3.7 Hz, and not 7.2 Hz as reported by Escribano.⁹ This result was obtained by using a simple pulse sequence, which selectively decouples protons from the CH_3 of the methyl

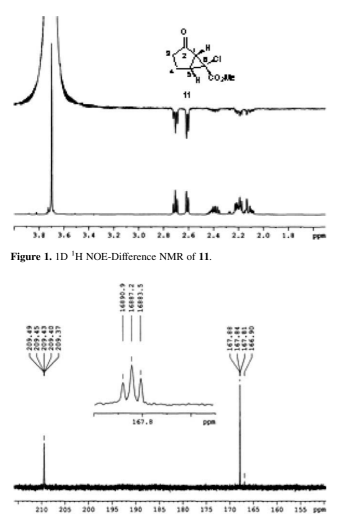


Figure 2. 13 C NMR decoupled –OCH₃ of **11**, ${}^{3}J {}^{1}H - {}^{13}C$: H-1 and H-5/CO₂R = 3.7 Hz.

ester (Fig. 2), and was confirmed by 2D ¹³C/JCH NMR experiment (Fig. 3). Our idea on the discrepancy between Escribano's work and our NMR measurements is that the Karplus relationship used by Escribano is convenient for a ³*J* (¹H–Csp³–Csp³–¹³Csp³) like in the propane¹⁹ but not for a ³*J* (¹H–Csp³–Csp³–¹³Csp²) like in the compound **11**. So, we agree with the structure proposed by Escribano, but not with the NMR data. Now, regarding the other bicyclic compounds **9**, **10**, and **12** (see Table 1) we prepared, they all have ³*J* (¹H–Csp³–Csp³–¹³Csp²) values close to 4 Hz, as for the compound **11** and by using the same NMR methods. So we think that we can reasonably assign an *endo*-Cl structure to these four bicyclic compounds.

The cyclopropanations described here are regiospecific. Indeed, no addition onto the carbonyls of the activated olefins

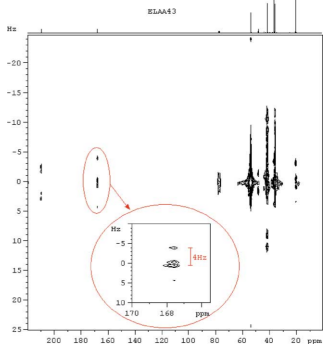
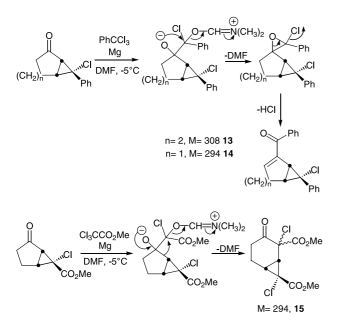


Figure 3. 2D ¹³C/JCH NMR of 11: ³J ¹H–¹³C: H-1 and H-5/CO₂R ~ 4 Hz.

was observed during or at the near end of the reaction, though the trichloromethyl compound is used in excess. Side products coming from the halocompounds are their reduced forms and traces of the dimers (YCCl=CClY). However, with process B, and in the case of cyclic enones and Cl_3C-Y (Table 1, entries 12, 13 and 14), we could observe, at the near end of the reaction, the formation of three by-products showing parent ions at m/e = 308, 294, 294, respectively, in their mass spectra. We thus made the assumption that the nucleophilic species generated in situ could react on the carbonyl of the bicyclic products, according to reactions described by Larson⁶ and by Schäfer.^{21,22} The structures 13, 14, 15 have been postulated for these by-products (Scheme 4). To prevent this side reaction in the preparation of the compounds 10, 11, 12 (see Table 1), we modified process B in a way to keep the electrophilic olefins in excess vs the gem-polyhalocompound all over the reaction. However, surprisingly, in the preparation of the bicyclic compound 9 (Table 1, entry 11), no 1,2-addition was observed. Up to now, we have no explanation for this result.



Scheme 4.

3. Conclusions

We have described in this paper two simple, efficient and complementary methods (processes A and B) for the preparation of polysubstituted chlorocyclopropanes using electrophilic olefins and activated trichloromethyl compounds as starting materials. These new routes do not make use of strong bases or very expensive copper carbenoid tertbutyl isocyanides. Also, we have noticed that, in DMF, the nucleophilic species resulting from the Mg reduction of α, α, α -trichlorotoluene were able to react with ketones leading to benzoylated olefins. So far, the only other reductive route reported involves an electrochemical reduction of α, α, α -trichlorotoluene in a double-walled glass cell with a mercury pool cathode.^{22a,c} We are now extending this Mg-Barbier reaction in DMF to the preparation of cycloalk-1-en-1-yl and alk-1-en-1-ylphenyl ketones.

4. Experimental

Melting points were determined with an Electrothermal IA 9100 digital melting point apparatus. ¹H, ¹³C NMR spectra were recorded on a Bruker AC-200 (200, 50 MHz, respectively) or Bruker Avance 300 (300, 75 MHz, respectively) or Bruker DRX-400 (400, 100 MHz, respectively) spectrometers. Mass spectra (electron impact) were obtained on a GCQ Thermoquest spectrometer equipped with a DB 5MS capillary column. Infrared spectra were recorded on a Perkin Elmer Spectrum BX FT-IR spectrometer. High-resolution mass spectral analyses and elemental analyses were carried out at 'Service Central d'Analyse du CNRS', Vernaison, France. Gas chromatography was performed on a Varian 3300 chromatograph fitted with a SIL-5 CP capillary column. Solvents and chemicals were used as received. The XC10 Fe rod (iron with 0.1% of carbon) and Mg grits (50-150 mesh) were purchased, respectively, from Weber Métaux and Fluka.

4.1. General procedure

Process A, indirect electrochemical process with Fe anode in the presence of CuBr. The reactions are conducted in an undivided cell fitted with an Fe rod as the anode and a nickel foam as the cathode (area: ca. 40 cm²). A solution of CuBr (144 mg, 1 mmol) and Bu_4NBr (300 mg) in DMF (45 mL) and pyridine (5 mL) is electrolysed at constant current intensity (0.3 A) during 15 min at $-5 \degree C > T > -10 \degree C$. Then, the activated olefin (10 mmol) and the α, α, α trichloromethyl compound (20 mmol) are added and electrolysed (0.1 A) until the complete consumption of the olefin (about 8 h). The DMF is evaporated under reduced pressure. The reaction mixture is poured into a cold mixture of 1 M HCl (50 mL) and diethyl ether (50 mL). The layers are separated and extracted with diethyl ether (three portions of 25 mL). The combined ethereal extracts are washed with a saturated solution of ammonium chloride and brine, dried over MgSO₄. Products are isolated either by column chromatography on silica gel (230-400 mesh) or aluminium oxide (70-230 mesh) using pentane-ether as eluent.

Process B, Mg-promoted Barbier type reaction in the presence of DMF. Magnesium grits (50–100 mesh) (30 mmol) are suspended in DMF (40 mL) in a three-neck flask fitted with a thermometer and a dropping funnel, and cooled at -5 °C. Half of the solution containing olefin (10 mmol), α, α, α -trichloromethyl compound (12 mmol) and DMF (5 mL) is rapidly introduced in the flask. The beginning of the reaction is clearly indicated by the temperature rising up to +5 °C, and the mixture turning yellow. The remaining of the reactants was then added within 5 min, and the reaction is allowed to proceed up to complete consumption of the olefin. After the usual work-up, the product is isolated by column chromatography on silica gel (230–400 mesh) using pentane–ether as eluent.

4.2. Isolated products

4.2.1. Dimethyl 1-chlorocyclopropane-1,2-dicarboxylate (1).^{14a} CAS RN: 39822-02-1 (*R**,*S**), 39822-01-0 (*R**,*R**).

4.2.2. Methyl 2-chloro-2-phenylcyclopropane-1carboxylate (2).^{14a} CAS RN: 39822-09-8 (R^* , S^*), 39822-10-1 (R^* , R^*).

4.2.3. Dimethyl 2-chloro-1-methoxycarbonylmethylcyclopropane-1,2-dicarboxylate (3).^{14a} CAS RN: 424790-89-6 (*R**,*S**), 424790-88-5 (*R**,*R**).

4.2.4. Methyl 2-chloro-1-methoxycarbonylmethyl-2phenylcyclopropane-1-carboxylate (4) (new compound). (C₁₄H₁₅ClO₄); MW: 282.723. Anal. Calcd for C₁₄H₁₅ClO₄: C, 59.48; H, 5.35; O, 22.63; Cl, 12.54. Found: C, 59.28; H, 5.33; O, 22.63; Cl, 12.66. Pentane-ether (95/5) to (90/10); obtained: 1.16 g (yield: 41%, (R*,S*)/(R*,R*): 35:65, process A), 1.61 g (yield: 57%, (R*,S*)/(R*,R*): 36:64, process B); (R^*, S^*) : oil, (R^*, R^*) : mp = 76–78 °C. ¹H NMR (200 MHz, CDCl₃) δ (R*,S*): 7.3-7.2 (Ph, 5H, m); 3.75 (OCH₃, 3H, s); 3.5 (OCH₃, 3H, s); 3.1 (CH₂, 1H, d, J = 17.6 Hz); 2.35 (H-3 or H-3', 1H, d, J = 7.4 Hz); 1.65 (H-3 or H-3', 1H, d, J = 7.4 Hz); 1.3 (CH₂, 1H, d, J = 17.6 Hz); for the couple H-3/H-3' ($\Delta \nu/J = 19.0$ AX system); for the methylene group ($\Delta \nu/J = 21.0$ AX system). (R^*, R^*): 7.4–7.2 (Ph, 5H, m); 3.7 (OCH₃, 3H, s); 3.4 (CH₂, 1H, d, J=17.6 Hz); 3.2 (OCH₃, 3H, s); 2.9 (CH₂, 1H, d, J=17.6 Hz); 2.5 (H-3 or H-3', 1H, d, J=6.9 Hz); 1.5 (H-3 or H-3', 1H, d, J=6.9 Hz); for the couple H-3/H-3' $(\Delta \nu/J = 29.0 \text{ AX system})$; for the methylene group $(\Delta \nu/J =$ 5.5 AB system). ¹³C NMR (50 MHz, CDCl₃) δ (R^{*} , S^{*}): CO: 170.9, 169.9; C(Ph): 138.5, 128.5; C-2: 53.3; OCH₃: 52.0, 51.9; CH₂: 37.3; C-1: 33.3; C-3: 25.6. (*R**,*R**): CO: 171.7, 169.7; C(Ph): 137.8, 128.7; C-2: 53.4; OCH₃: 51.7, 49.7; CH2: 37.5; C-1: 34.3; C-3: 23.7. EI-MS m/z (R*,S*): 282 (M, 1), 220 (32), 219 (13), 218 (base peak), 192 (14), 191 (20), 190 (46), 187 (13), 165 (26), 164 (16), 163 (78), 162 (17), 159 (20), 155 (11), 149 (24), 145 (20), 129 (17), 128 (56), 127 (30), 115 (11). (R*,R*): 282 (M, 1), 220 (30), 219 (12), 218 (base peak), 192 (16), 191 (19), 190 (41), 187 (14), 165 (26), 164 (13), 163 (71), 162 (15), 159 (21), 155 (10), 149 (22), 145 (20), 129 (20), 128 (64), 127 (30), 115 (11). IR ν (cm⁻¹) (CDCl₃) 3080, 3030, 2990, 2970, 2900, 1735, 1600, 1570, 1470.

4.2.5. Dimethyl 1-chloro-2-methylcyclopropane-1,2dicarboxylate (5). (C₈H₁₁ClO₄); MW: 206.625; CAS RN: 42392-04-1 (R^*,S^*), 132785-43-4 (R^*,R^*). Pentane (100) to pentane-ether (95/5); obtained: 1.45 g (yield: 70%, $(R^*, S^*)/$ (R^*, R^*) : 45:55, process B; (R^*, S^*) and (R^*, R^*) : oil. ¹H NMR (200 MHz, CDCl₃) δ (*R**,*S**): 3.5 (OCH₃, 3H, s); 3.4 $(OCH_3, 3H, s)$; 2.0 (H-3 or H-3', 1H, d, J=6.5 Hz); 1.3 $(CH_3, 3H, s)$; 1.0 (H-3 or H-3', 1H, d, $J_{gem} = 6.5$ Hz); for the couple H-3/H-3' ($\Delta \nu/J = 31.4$ AX system). (R^*, R^*): 3.65 (OCH₃, 3H, s); 3.6 (OCH₃, 3H, s); 1.85 (H-3 or H-3['], 1H, d, J=6.6 Hz); 1.7 (H-3 or H-3', 1H, d, J=6.6 Hz); 1.2 (CH₃, 3H, s); for the couple H-3/H-3' ($\Delta \nu/J = 5.6$ AB system). ¹³C NMR (50 MHz, CDCl₃) δ (*R**,*S**): CO: 170.8, 167.8; OCH₃: 53.1, 52.4; C-1: 48.5; C-2: 33.7; C-3: 27.9; CH₃: 17.3. (R*,R*): CO: 169.0, 166.9; OCH₃: 53.0, 52.1; C-1: 45.2; C-2: 35.1; C-3: 25.5; CH₃: 14.8. EI-MS *m*/*z* (*R**,*S**): 206 (M, <1), 177 (13), 176 (15), 175 (37), 174 (32), 171 (22), 170 (51), 148 (35), 147 (17), 146 (base peak), 139 (16), 133 (12), 131 (31), 127 (11), 119 (18), 115 (20), 111 (12), 87 (15), 83 (15). (R*,R*): 206 (M, 1), 176 (11), 175 (22), 174 (26), 171 (13), 170 (36), 148 (34), 147 (17), 146 (base peak),

139 (18), 131 (26), 119 (18), 115 (15), 111 (15), 83 (13), 55 (10). IR ν (cm⁻¹) (film) 3100, 2990, 2970, 1750, 1730, 1440.

4.2.6. Methyl 2-chloro-1-methyl-2-phenylcyclopropane-1-carboxylate (6). (C12H13ClO2); MW: 224.687; CAS RN: 91433-96-4 (R*,S*), 91434-02-5 (R*,R*). Pentane (100) to pentane-ether (95/5); obtained: 1.64 g (yield: 73%, (R^*, S^*) / (R^*, R^*) : 25:75, process B; (R^*, S^*) and (R^*, R^*) : oil. ¹H NMR (200 MHz, CDCl₃) δ (*R**,*S**): 7.4–7.6 (Ph, 5H, m); 4.0 (OCH₃, 3H, s); 2.4 (H-3 or H-3['], 1H, d, J=6.8 Hz); 1.7 (H-3 or H-3['], 1H, d, J=6.8 Hz); 1.2 (CH₃, 3H, s); for the couple H-3/H-3' ($\Delta \nu/J = 20.0$ AX system). (R^*, R^*): 7.5-7.45 (Ph, 5H, m); 3.5 (OCH₃, 3H, s); 2.6 (H-3 ou H-3', 1H, d, J = 6.5 Hz); 1.95 (CH₃, 3H, s); 1.6 (H-3 or H-3', 1H, d, J = 6.5 Hz); for the couple H-3/H-3' ($\Delta \nu / J = 32.3$ AX system). ¹³C NMR (50 MHz, CDCl₃) δ (*R**,*S**): CO: 171.0; C(Ph): 137.9, 128.8, C-2: 52.1; OCH₃: 49.9; C-1: 33.1; C-3: 24.5; CH₃: 17.6. (*R**,*R**): C-4: 171.4; C-7: 139.6; other aromatic C: 128.4; C-2: 53.9; C-5: 51.8; C-1: 32.4; C-3: 26.0; C-6: 18.0. EI-MS m/z (R*,S*): 225 (M, 6), 189 (35), 167 (12), 165 (34), 161 (8), 157 (8), 131 (15), 130 (12), 129 (base peak), 128 (28), 105 (10). (R*,R*): 225 (M, 9), 189 (35), 167 (10), 165 (35), 161 (10), 157 (10), 131 (16), 130 (15), 129 (base peak), 128 (27), 105 (10). IR ν (cm⁻¹) (film) 3030, 2920, 1720, 1580, 1500, 1450.

4.2.7. *trans*-Trimethyl 1-chlorocyclopropane-1,2,3-tricarboxylate (7).^{14d} CAS RN: 205320-46-3.

4.2.8. *trans*-Dimethyl 3-chloro-3-phenylcyclopropane-**1,2-dicarboxylate (8).**^{14d} CAS RN: 205320-44-1.

4.2.9. (*1RS*,*6RS*,*7RS*)-Methyl 7-chloro-2-oxobicyclo [**4.1.0**]heptane-7-carboxylate (9). ($C_9H_{11}ClO_3$); MW: 202.637; CAS RN: 406217-16-1. Pentane–ether (90/10) to (80/20); obtained: 1.17 g (yield: 58%, process A), 1.07 g (yield: 53%, process B); oil. ¹H NMR (200 MHz, CDCl₃) 3.5 (OCH₃, 3H, s); 2.3–2.1 (2H, m); 2.1–1.8 (3H, m); 1.7–1.5 (3H, m). ¹³C NMR (50 MHz, CDCl₃) δ COR: 202.5; COOR: 168.7; OCH₃: 53.5; C-7: 48.7; C-3: 38.9; C-1: 34.1; C-6: 30.1; C-4 and C-5: 23.9, 17.6. EI-MS *m*/*z* 202 (M, 10), 176 (23), 174 (73), 172 (28), 171 (13), 170 (88), 148 (10), 147 (31), 146 (14), 145 (34), 144 (36), 143 (51), 142 (base peak), 139 (13), 135 (32), 117 (12), 116 (10), 115 (21), 111 (11), 107 (41), 106 (10), 87 (13), 81 (14), 80 (11), 79 (99), 78 (15), 77 (43), 53 (11), 51 (40). IR ν (cm⁻¹) (CDCl₃) 1750, 1720.

4.2.10. (*1RS*,*6RS*,*7RS*)-7-Chloro-7-phenylbicyclo[4.1.0]-heptane-2-one (10). ($C_{13}H_{13}ClO$); MW: 220.699; CAS RN: 126252-39-9. Pentane (100) to pentane–ether (95/5); obtained: 0.662 g (yield: 30%, process A), 1.10 g (yield: 50%, process B); mp=69–70 °C. ¹H NMR (200 MHz, CDCl₃) 7.6–7.2 (Ph, 5H, m); 2.3–1.6 (H-1 to H-6, 8H, m). ¹³C NMR (50 MHz, CDCl₃) δ CO: 204.9; C(Ph): 141.9, 128.3, 127.5; C-7: 54.9; C-3: 39.1; C-1: 33.6; C-6: 29.0; C-4 and C-5: 24.9, 18.6. EI-MS *m*/*z* 220 (M, 8), 192 (15), 185 (10), 157 (28), 141 (8), 130 (12), 129 (base peak), 128 (27), 127 (9), 115 (15). IR ν (cm⁻¹) (CDCl₃) 3080, 3020, 2980, 1700, 1600, 1580, 1500.

4.2.11. (1*RS*,5*RS*,6*RS*)-Methyl 6-chloro-2-oxobicyclo [3.1.0]hexane-6-carboxylate (11). ($C_8H_9ClO_3$); MW:

188.610; CAS RN: 2158-08-1. Pentane (100) to pentaneether (85/15); obtained: 0.566 g (yield: 30%, process A), 0.754 g (yield: 40%, process B); mp=41-42 °C. ¹H NMR (300 MHz, CDCl₃) 3.7 (OCH₃, 3H, s); 2.7 (H-5, 1H, t, ³*J*=6.3 Hz); 2.6 (H-1, 1H, d, ³*J*=6.3 Hz); 2.5–2.05 (H-3 and H-4, 4H, m). ¹³C NMR (75 MHz, CDCl₃) δ COR: 209.4; CO₂R: 167.8; OCH₃: 53.8; C-6: 48.3; C-1: 41.6; C-3: 36.6; C-5: 36.0; C-4: 20.4. EI-MS *m*/*z* 162 (15), 160 (38), 156 (37), 149 (11), 148 (23), 147 (48), 146 (65), 145 (44), 134 (35), 133 (20), 132 (base peak), 131 (49), 129 (13), 128 (11), 125 (31), 124 (12), 118 (14), 117 (25), 116 (26), 115 (19), 111 (23), 109 (11), 101 (15), 100 (13), 93 (30), 87 (14), 80 (14), 79 (17), 73 (11), 69 (14), 65 (61), 51 (16). IR ν (cm⁻¹) (CDCl₃) 3068, 3050, 3010, 2956, 2873, 1730, 1703, 1440.

4.2.12. (1*RS*,5*RS*,6*RS*)-6-Chloro-6-phenylbicyclo[3.1.0] hexane-2-one (12) (new compound). ($C_{12}H_{11}ClO$); MW: 206.672. ES-HR-MS calcd for $C_{12}H_{11}ONaCl m/z$ 229.0396, found 229.0399. Pentane (100) to pentane–ether (95/5); obtained: 0.413 g (yield: 20%, process A), 0.811 g (yield: 40%, process B); mp=86–87 °C. ¹H NMR (200 MHz, CDCl₃) 7.4–7.15 (Ph, 5H, m); 2.6–2.15 (H-1 to H-5, 6H, m). ¹³C NMR (50 MHz, CDCl₃) δ O: 211.1; C(Ph): 140.8, 128.9, 128.6, 127.6; C-6: 54.5; C-1: 41.4; C-3: 37.4; C-5: 34.7; C-4: 21.1. EI-MS *m*/*z* 164 (20), 143 (10), 130 (11), 129 (base peak), 128 (31), 127 (8), 115 (15). IR ν (cm⁻¹) (CDCl₃) 3150, 3040, 2980, 2940, 1730, 1600, 1580, 1500.

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Enantiocontrolled synthesis of the epoxycyclohexenone moieties of scyphostatin, a potent and specific inhibitor of neutral sphingomyelinase

Tadashi Katoh,^{a,*} Takashi Izuhara,^b Wakako Yokota,^c Munenori Inoue,^c Kazuhiro Watanabe,^a Ayaka Nobeyama^a and Takeyuki Suzuki^a

> ^aDepartment of Chemical Pharmaceutical Science, Tohoku Pharmaceutical University, 4-4-1 Komatsushima, Aoba-ku, Sendai 981-8558, Japan

> ^bDepartment of Electronic Chemistry, Tokyo Institute of Technology, Nagatsuta, Yokohama 226-8502, Japan ^cSagami Chemical Research Center, 2743-1 Hayakawa, Ayase, Kanagawa 252-1193, Japan

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Abstract—The epoxycyclohexenone moieties 2 and 3b of scyphostatin (1), a potent and specific inhibitor of neutral sphingomyelinase, were synthesized in enantiomerically pure forms starting from (-)-quinic acid (11). The synthetic method features (i) the preparation of the olefin masked enones 25 and 29, the precursors for the key aldol-type coupling reaction, (ii) the efficient and stereocontrolled aldol-type coupling reactions between 25 (or 29) and benzaldehyde (8) and Garner's aldehyde analogue 9 to deliver alcohols 23 and 24, respectively, both of which possess the requisite asymmetric quaternary carbon center at the C6 position, and (iii) the stereospecific S_N2-type epoxide ring formation of the mesylates 35 and 47 under mild basic conditions to produce the targeted compounds 2 and 3b, respectively. © 2005 Elsevier Ltd. All rights reserved.

1. Introduction

Recently, sphingomyelinase (SMase) inhibitors have received considerable attention from the biological and pharmacuetical standpoints.¹ SMase is the enzyme that specifically hydrolyzes the phosphoester linkage of sphingomyelin (SM), one of the most abundant sphingolipid species, to generate ceramide and phoshocholine.^{2,3} The SM-derived ceramide is believed to be an intracellular lipid second messenger in cell membranes and to play important roles in the regulation of cell proliferation, differentiation, and apoptosis.^{2,3} SMase inhibitors, therefore, are considered as valuable tools for the investigation of the biological function of the enzyme and the catabolite ceramide in signal transduction.³ In addition, selective SMase inhibitors are highly anticipated to be promising candidates for the treatment of ceramide-mediated pathogenic states such as AIDS,⁴ inflammation,⁵ and immunological and neurological disorders.⁶

In 1997, Ogita et al. at the Sankyo research group reported the isolation and structure elucidation of a novel natural product, scyphostatin (1, Fig. 1), from the mycelial extract

of *Trichopeziza mollissima* SANK 13892.^{7,8} This natural product was found to be a powerful and specific inhibitor of membrane-bound neutral sphingomyelinase (N-SMase).⁸ It has been reported that **1** inhibits N-SMase and acidic SMase (A-SMase) with IC₅₀ values of 1.0 and 49.3 μ M, respectively.^{7,8} Remarkably, scyphostatin is the most potent and specific one among the many low molecular weight N-SMase inhibitors of natural sources⁹ or of synthetic substances¹⁰ known to date.

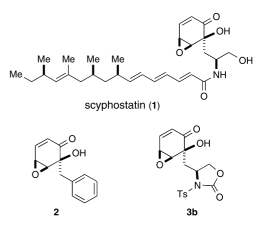


Figure 1. Structures of scyphostatin $\left(1\right)$ and the epoxycyclohexeone moieties 2 and 3b.

Keywords: Sphingomyelinase; Aldol-type coupling; Diels–Alder reaction. * Corresponding author. Tel.: +81 22 234 4183; fax: 81 22 275 2013; e-mail: katoh@tohoku-pharm.ac.jp

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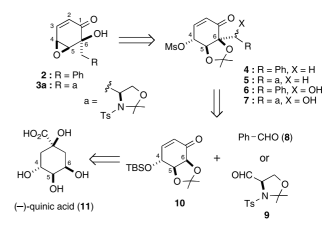
The gross structure of scyphostatin (1) was revealed by extensive and incisive spectroscopic studies.⁷ It consists of a novel, highly oxygenated cyclohexenone ring incorporated with a C-20 unsaturated fatty acid-substituted aminopropanol side chain. This initial structure elucidation only established the relative and absolute stereochemistry of the cyclohexenone moiety of 1.⁷ In 2001, Kogen et al. at the Sankyo research group determined the relative and absolute configurations of the three stereogenic centers present in the fatty acid side chain.¹¹ At the almost same time, Hoye et al. disclosed an enantioselective synthesis of the C-20 unsaturated fatty acid moiety and provided alternative proof of its stereostructure including the absolute configuration.¹²

The remarkable biological properties and unique structural features make **1** an exceptionally intriguing and timely target for total synthesis. So far, a number of synthetic approaches toward scyphostatin (**1**) have been reported by Gurujar's group,¹³ Taylor's group,¹⁴ Ohkata's group,¹⁵ Kita's group,¹⁶ Maier's group,¹⁷ Negishi's group,¹⁸ and Pitsino's group.¹⁹ We have already reported our own preliminary results concerning the enantioselective synthesis of the epoxycylohexenone substructures **2** and **3b**²⁰ (Fig. 1). Additionally, we have also disclosed an efficient method for the introduction of a fatty acid side chain at the amino propanol moiety.²¹ In 2004, our assiduous endeavors culminated in the completion of the first total synthesis of (+)-**1**.²² In this paper, we wish to disclose the full details of our first-generation synthesis of the epoxycylohexenone moieties **2** and **3b** of scyphostatin (**1**).

2. Results and discussion

2.1. Primary synthetic plan for the epoxycyclohexenone moieties 2 and 3a

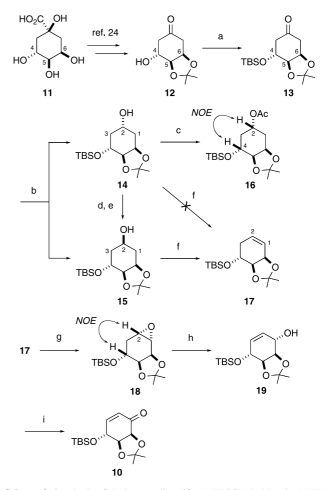
Our primary synthetic plan for the epoxycyclohexenone moieties 2 and 3a is outlined in Scheme 1. The key feature of this plan is aldol-type coupling reactions between the cyclohexenone 10 and the aldehydes 8 and 9 to form the coupling products 6 and 7, respectively $(10+8\rightarrow 6 \text{ and } 10+$ $9 \rightarrow 7$). In these reactions, we envisioned that electrophiles 8 and 9 would approach exclusively from the less hindered α -face of the enolate, generated in situ from 10, under the influence of the β -oriented *O*-isopropylidenedioxy moiety, thus leading to establishment of the requisite asymmetric quaternary carbon center at the C6 position (cyclohexeneone numbering)²³ in 6 and 7. This type of coupling reaction is considerably challenging at the synthetic chemistry level, because the substrate 10 possesses unusual trihydroxy functionalities at the C4, C5, and C6 positions, and in addition, an electrophilic enone system. The coupling products 6 and 7 would be converted to the target molecules 2 and 3a through the advanced key intermediates 4 and 5, respectively, by sequential functional group manipulation and deprotection, or vice versa; the sequence involves deoxygenation of the secondary hydroxy group in the side chain and stereospecific S_N 2-type epoxide ring formation as the crucial steps. The cyclohexenone 10 having three contiguous oxygen functionalities at the C4, C5, and C6 positions with correct stereochemistries would be derived from commercially available (-)-quinic acid (11).



Scheme 1. Primary synthetic plan for the epoxycyclohexeone moieties 2 and 3a.

2.2. Synthesis of the intermediate 10

At first, as shown in Scheme 2, we pursued the synthesis of the intermediate 10, a substrate for the key aldol-type coupling reaction, starting from commercially available (-)- quinic acid (11). The known cyclohexanone 12^{24} was



Scheme 2. Synthesis of the intermediate 10. (a) TBSCL, imidazole, DMF, rt, 98%; (b) NaBH₄, THF–H₂O, $-5 \degree C \rightarrow rt$, 53% for 14, 44% for 15; (c) Ac₂O, pyridine, DMAP, $0\degree C \rightarrow rt$, 98%; (d) DEAD, Ph₃P, benzonic acid, THF, $0\degree C \rightarrow 98\%$; (e) 2 M KOH–MeOH, rt, quant.; (f) DEAD, Ph₃P, THF, rt, 67% for 15 \rightarrow 17, 0% for 14 \rightarrow 17; (g) mCPBA, NaHCO₃, CH₂Cl₂, $0\degree C \rightarrow rt$, 92%; (h) Se₂Ph₂, NaBH₄, EtOH, $0\degree C \rightarrow reflux$; H₂O₂, THF, $0\degree C \rightarrow reflux$, 78%; (i) Dess–Martin periodinane, CH₂Cl₂, rt, 95%.

readily and sufficiently prepared from **11** in three steps [(1) dimethoxypropane/*p*-TsOH/acetone, reflux, 80%; (2) LiAlH₄/THF, reflux, quant.; (3) NaIO₄/*t*-BuOH–THF–AcOH, room temperature, quant.] according to the reported procedure.²⁴ After protection of the hydroxy group in **12** as its *t*-butyldimethylsilyl (TBS) ether, the carbonyl function of the resulting TBS ether **13** was subjected to reduction with sodium borohydride to furnish an epimeric mixture of the alcohols **14** (53%) and **15** (44%) that were separated by silica gel column chromatography. The newly formed C2 stereochemistry of the two isomers was assigned on the basis of spectroscopic studies. The NOESY experiment of the acetate **16** derived from **14** showed a clear interaction between C2–H and C4–H.

We next examined installation of an olefinic double bond by dehydration of 14 and 15. Thus, reaction of 15 with diethyl azodicarboxylate (DEAD) and triphenylphosphine provided the desired olefin 17 in 67% yield with complete regioselectivity at the C1-C2 position. On the contrary, treatment of the C2 epimeric alcohol 14 under the same dehydration conditions afforded none of the desired olefinic product 17, and the unreacted starting material 14 was recovered unchanged. Therefore, the alcohol 14 was converted to 15 by employing the Mitsunobu inversion procedure²⁵ (98% overall yield). The difference of the reactivity between 14 and 15 under the dehydration conditions can be rationalized by conformational analyses of both 14 and 15 (Fig. 2). Thus, the NOESY experiment of 15 indicated that the cyclohexane ring takes a boat-form, which places the C2 hydroxy group in an axial position; this conformation may facilitate E2 elimination to afford the desired $\Delta^{1,2}$ olefin 17. On the other hand, the NOESY experiment of 14 indicated that the C2 hydroxy group is in equatorial orientation within the boat-formed cyclohexane ring; this conformation would preclude any possibility of E2 elimination.

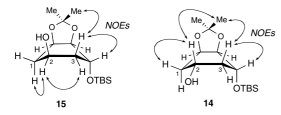


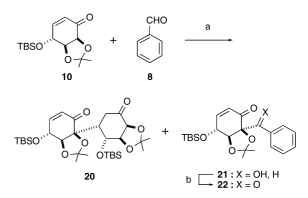
Figure 2. Conformational analyses of the alchohols 14 and 15.

To continue the synthesis, the olefin **17** was oxidized with *m*-chloroperbenzoic acid (*m*CPBA) to give the epoxide **18** as a single diastereomer in 92% yield, whose stereochemistry was assigned based on the NOE experiment. The stereoselectivity can be explained by the consideration that the oxidizing reagent (*m*CPBA) accessed exclusively from the less hindered α -face of the molecule under the influence of the β -oriented *O*-isopropylidenedioxy moiety. Conversion of the epoxide **18** to the allyl alcohol **19** was successfully achieved by employing a reliable Sharpless protocol.²⁶ Thus, treatment of **18** with the phenylselenyl anion, generated in situ from diphenyl diselenide and sodium borohydride, caused the regioselective epoxide ring opening at the sterically and electrostatically favored C2 position to form the corresponding

phenylselenide, which was then oxidized by excess 30% aqueous hydrogen peroxide to provide the allyl alcohol **19** in 95% overall yield via elimination of the intermediary phenylselenoxide. Finally, Dess–Martin oxidation²⁷ of **19** furnished the requisite intermediate **10** in 95% yield.

2.3. Initial attempts to achieve the coupling reaction of the cyclohexenone 10 with benzaldehyde (8)

Having obtained the intermediate 10, we next investigated the crucial aldol-type coupling reaction between 10 and benzaldehyde (8) as shown in Scheme 3. Initial attempts to achieve this coupling reaction, unfortunately, turned out to be fruitless. Thus, reaction of the lithium enolate of 10, generated in situ by reaction with $LiN(SiMe_3)_2$, with 8 in THF at -78 °C resulted in the predominant formation of the unexpected dimerized product 20 (38%) as a single stereo isomer and the desired coupling product 21 (12%) as an epimeric mixture with respect to the benzilic hydroxy group. Since the coupling product 21 was very unstable during isolation and purification by silica gel column chromatography, assignment of the structure and stereochemistry of 21 was performed by spectroscopic analyses (COSY, HMBC, and NOESY experiments) of the corresponding carbonyl compound 22, readily prepared by Dess-Martin oxidation (78%).

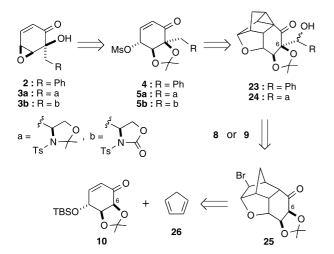


Scheme 3. Aldol-type coupling reaction of the cyclohexenone 10 and benzaldehyde (8). (a) LiN(SiMe₃)₂, THF, -78 °C, 38% for 20, 12% for 21; (b) Dess–Martin periodinane, CH₂Cl₂, rt, 78%.

These preliminary studies demonstrated that the enone olefin function present in **10** was extremely susceptible to nucleophilic attack of the enolate generated from **10** itself. In order to circumvent this problem, we decided to mask the highly reactive enone system of **10** in the form of the bromo ether **25** (cf. Scheme 4) during the aldol-type coupling reaction. We anticipated that **25** would behave as a promising substrate for the designed coupling reaction. Further investigations concerning the synthesis of **25** and subsequent coupling reaction with the aldehydes **8** and **9** are the subject of the following sections.

2.4. Modified synthetic plan for the epoxycyclohexenone moieties 2 and 3a

Our initial attempts to achieve the direct coupling between the cyclohexenone **10** and benzaldehyde (**8**) met with failure; therefore, we settled on modifying our original synthetic plan. Thus, as shown in Scheme 4, the bromo ether

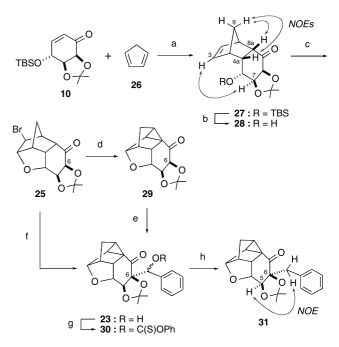


Scheme 4. Modified synthetic plan for the epoxycyclohexenone moieties 2 and 3.

25, a synthetically equivalent of the cyclohexenone **10**, was envisaged to be prepared by Diels–Alder reaction of **10** with cyclopentadiene (**26**) followed by desilylation and bromo etherification. The crucial aldol-type reaction of **25** with the aldehydes **8** and **9** would produce the coupling products **23** and **24**, respectively, with correct stereochemistry at the C6 position. The intermediates **23** and **24** would be converted to the cyclohexenones **4** and **5**, the potential key intermediates of the target molecules **2** and **3**, via sequential functional group manipulation. As will be mentioned later (cf. Sections 2.7 and 2.8), the *N*,*O*-isopropylidene group at the C6 side chain in **24** turned out to be labile during the regeneration of the enone olefin moiety (cf. **24**→**5a**); therefore, the *N*,*O*-isopropylidene group was replaced with a sturdy cyclic carbamate group (cf. **5b**).

2.5. Synthesis of the intermediate 31 for the epoxycyclohexenone moiety 2: preparation of the masked enone 25 and subsequent aldol-type coupling reaction with benzaldehyde (8)

As shown in Scheme 5, we next carried out the synthesis of the intermediate 31 for the first target compound 2; the sequence involved the preparation of the olefin masked cyclohexenone 25 and subsequent coupling reaction with benzaldehyde (8) as the crucial steps. Diels-Alder reaction of 10 with cyclopentadiene (26) in the presence of diethylaluminium chloride proceeded smoothly and cleanly in a completely diastereofacial- and endo-selective manner to provide the corresponding cycloadduct 27 as a single isomer in almost quantitative yield (97%). The structure and stereochemistry of the Diels-Alder adduct 27 was assigned based on the NMR spectral analysis including NOESY experiments; thus, clear NOE interactions between C9-H and C8a-H, C4a-H and between C3-H and C7-H were observed, respectively. After deprotection of the TBS group of 27 with tetrabutylammonium fluoride (TBAF) (75%), the resulting alcohol 28 was subjected to bromo etherification using N-bromosuccinimide $(NBS)^{28}$ to provide the desired tetracyclic bromo ether 25 in 86% yield.



Scheme 5. Aldol-type coupling reaction of the masked enone 25 with benzaldehyde (8) and the synthesis of the intermediate 31. (a) Et₂AlCl, CH₂Cl₂, $-78 \rightarrow 0^{\circ}$ C, 97%; (b) TBAF, THF, 0° C $\rightarrow rt$, 75%; (c) NBS, CH₂Cl₂, 0° C $\rightarrow rt$, 86%; (d) LiN(SiMe₃)₂, THF, 98%; (e) LiN(SiMe₃)₂, THF, -78° C; at -78° C add. benzaldehyde (8), 98%; (f) LiN(SiMe₃)₂, THF, -78° C; at -78° C add. benzaldehyde (8), 98%; (g) phenyl chlorothionoformate, DMAP, MeCN, rt, 92%; (h) *n*-Bu₃SnH, AIBN, toluene, 110 °C, 79%.

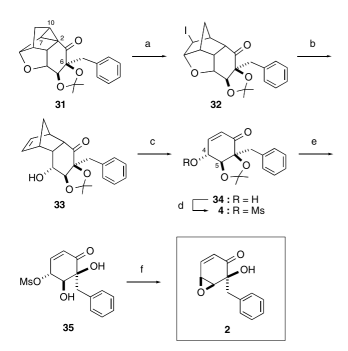
The crucial aldol-type coupling reaction between 25^{29} and benzaldehyde (8) was next conducted to establish the requisite C6 asymmetric quaternary carbon center. During the optimization of the reaction conditions, we found that the bromo ether 25 exhibited an interesting and unprecedented reactivity. Thus, treatment of 25 with 1.1 equiv of $LiN(SiMe_3)_2$ in THF at -78 °C for 30 min resulted in the formation of the unexpected cyclopropane derivative **29** in almost quantitative yield (98%), whose structure was confirmed by extensive spectroscopic studies including COSY, HMBC, and NOESY experiments in the 500 MHz NMR spectra. Subsequent treatment of 29 with 1.1 equiv of $LiN(SiMe_3)_2$ in THF at -78 °C followed by addition of benzaldehyde (8) (2.2 equiv) afforded the desired coupling product 30 in excellent yield (98%) as a hardly separable mixture of the epimeric alcohols (6:1 by 500 MHz⁻¹H NMR). The C6 stereochemistry of the product 23 turned out to be completely controlled as we expected; the assignment was later confirmed by NOE study of the transformed compound **31** (vide infra).

Encouraged by these successful results, we next examined a more efficient one-pot procedure for the direct coupling of **25** and **8**. Thus, treatment of **25** with 2.2 equiv of LiN(SiMe₃)₂ followed by reaction with 2.2 equiv of **8** furnished the requisite coupling product **23** in 98% yield. The secondary hydroxy group in **23** was deleted by using Robin's modification³⁰ of the Barton method.³¹ Thus, treatment of **23** with phenyl thionochloroformate in acetonitrile in the presence of 4-dimethylaminopyridine (DMAP) at ambient temperature afforded the corresponding phenoxythionocarbonate **30** in 92% yield. Compound **30**

was then allowed to react with tri-*n*-butyltin hydride in toluene in the presence of a catalytic amount of 2,2'azobisisobutyronitrile (AIBN) at 110 °C, giving rise to the desired deoxygenated product **31** in 79% yield. At this stage, the C6 stereochemistry could be unambiguously confirmed by NOESY experiments in the 500 MHz ¹H NMR spectrum of **31**, in which a clear NOE interaction between C5–H and the benzylic proton was observed.

2.6. Synthesis of the epoxycyclohexenone moiety 2

Having succeeded in introduction of the benzyl substituent at the C6 position with the correct stereochemistry, we next executed conversion of 31 into the epoxycyclohexenone moiety 2 (Scheme 6); the sequence involved regeneration of the enone system and subsequent epoxide ring formation as the pivotal steps. Regioselective cleavage of the cyclopropane ring in 31 was successfully achieved by treatment with trimethylsilyl iodide (TMSI)³² in carbon tetrachloride at $-20 \rightarrow -10$ °C to give the desired γ -iodo ketone 32 in 89% yield as the sole product. The regioselectivitiy observed for this ring opening reaction can be explained by the so-called stereoelectronic effect. Thus, the $\sigma_{\rm C2-C7}$ orbital efficiently overlaps with the $\pi_{C=O}$ orbital, while the overlap between the σ_{C2-C10} orbital and the $\pi_{C=O}$ orbital is insufficient due to the geometrical factor. An attack of the iodo anion, therefore, occurred predominantly at the C7 position in **31**. Conversion of the γ -iodo ketone 32 to the requisite cyclohexenone 34 was effectively achieved by applying the Ogasawara procedure.²⁸ Thus, treatment of **32** with zinc powder in methanol containing a small amount of acetic acid gave the tricyclic compound 33 in 91% yield, which was then subjected to retro-Diels-Alder reaction by heating at 230 °C in diphenyl ether to produce 34 in 81% yield.



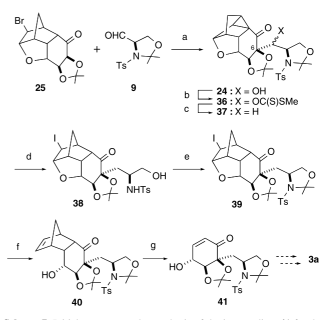
Scheme 6. Synthesis of the epoxycyclohexenone moiety 2. (a) TMSI, CCl₄, $-20 \rightarrow -10$ °C, 89%; (b) Zn, AcOH, MeOH, 60 °C, 91%; (c) Ph₂O, 230 °C, 81%; (d) MsCl, Et₃N, DMAP, CH₂Cl₂, 0 °C → rt, 85%; (e) TFA, H₂O, 0 °C, 85%; (f) 0.2 M NaOH, Et₂O, 0 °C, 90%.

The remaining task to complete the synthesis of the first target compound 2 involved the critical epoxide ring formation utilizing the two oxygen functionalities at the C4 and C5 positions in 34. Toward this end, mesylation of the hydroxy group in 34 under the standard conditions (MsCl, Et₃N, DMAP, CH₂Cl₂, 0 °C \rightarrow room temperature) (85%) followed by hydrolysis of the acetonide moiety of the resulting mesylate 4 by treatment with aqueous trifluoro-acetic acid (TFA), furnished the desired diol 35 in 85% yield. Finally, the expected epoxide ring formation was successfully achieved by brief exposure of 35 to 0.2 M sodium hydroxide in ether at 0 °C for 10 min, providing the epoxycyclohexenone moiety 2 in 90% yield.

2.7. Initial attempts on the synthesis of the fully functionalized epoxycyclohexenone moiety 3a

Having established the synthetic route to the epoxycyclohexenone moiety **2**, we next undertook the synthesis of the fully functionalized epoxycyclohexenone moiety **3a** (cf. Scheme 4), which possesses the *N*,*O*-protected amino propanol side chain and the requisite asymmetric carbon centers. We envisaged that the targeted compound **3a** would be elaborated starting from the bromo ether **25** and D-serinal derivative **9**³³ [(*R*)-*N*-(*p*-toluenesulfonyl)-*N*,*O*-isopropylidene serinal], readily accessible from D-serine, based on the explored synthetic route to the epoxycyclohexenone moiety **2**.

As shown in Scheme 7, the synthesis started with the crucial aldol coupling reaction between **25** and **9**. The enolate anion, generated in situ by treatment of **25** with $\text{LiN}(\text{SiMe}_3)_2$ (2.2 equiv) in THF at -78 °C, was allowed to react with **9** (2.5 equiv) to furnish an excellent yield



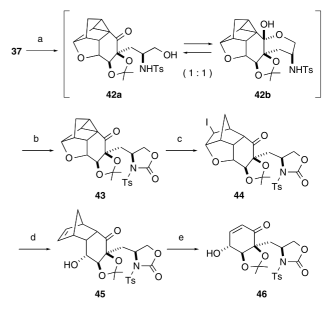
Scheme 7. Initial attempts on the synthesis of the intermediate 41 for the epoxycyclohexenone moiety 3a. (a) LiN(SiMe₃)₂, THF, -78 °C; at -78 °C, add. (*R*)-*N*-(*p*-toluenesulfonyl)-*N*,0-isopropylidene serinal (9), 98%; (b) NaN(SiMe₃)₂, THF, -78 °C; CS₂, $-78 \rightarrow -50$ °C; MeI, $-78 \rightarrow -50$ °C, 88%; (c) *n*-Bu₃SnH, Et₃B, toluene, rt, 95%; (d) TMSI, CCl₄, -10 °C, 91%; (e) 2,2-dimethoxypropane, *p*-TsOH, benzene, 60 °C, 83%; (f) Zn, AcOH, MeOH, 60 °C, 98%; (g) Ph₂O, 230 °C, 25%.

(98%) of the desired coupling product 24 as an inseparable mixture of the epimeric alcohols (9:1 by 500 MHz⁻¹H NMR). Removal of the hydroxy group in 24 was initially attempted by employing the same reaction conditions [ClC(S)OPh, DMAP, MeCN] described for the preparation of 30 from 23 (cf. Scheme 5), which, unfortunately, ended in failure and the starting material 24 was recovered unchanged even under heating conditions. This is presumably due to the steric hindrance around the hydroxy group in 24. Therefore, we looked at the Barton procedure to achieve the requisite deoxygenation of the sterically hindered hydroxy group. Employing the original Barton conditions (NaH, CS₂, THF; MeI, $0 \degree C \rightarrow room$ temperature), the reaction gave a poor yield ($\sim 30\%$) of the desired methyl xanthate 36. In order to improve the yield, some modifications were made of the reaction conditions. After several trials, to our delight, we found that treatment of 24 with NaN(SiMe₃)₂ (1.2 equiv) in THF at -78 °C followed by addition of carbon disulfide (10 equiv) and iodomethane (10 equiv) at the same temperature furnished the methyl xanthate 36 in 88% yield. The resulting methyl xanthate 36 was further treated with tri-n-butyltinhydride and triethylborane³⁴ in toluene at ambient temperature to afford the requisite deoxygenated product 37 in 95% yield.

With the intermediate 37 possessing the requisite N,O-protected amino propanol side chain and the correct stereochemistry in hand, our next efforts were devoted to regeneration of the cyclohexenone olefin moiety. Toward this end, regioselective cleavage of the cyclopropane ring in 37 was conducted by treatment with TMSI to give the iodo ether 38 in 91% yield. In this reaction, the N,Oisopropylidene group was simultaneously hydrolyzed; therefore, regeneration of the N,O-isopropylidene moiety of the resulting aminopropanol 38 was carried out under conventional conditions (2,2-dimethoxypropane, p-TsOH, benzene, 60 °C) to furnish the acetonide **39** in 83% yield. Further treatment of 39 with zinc powder in methanol containing acetic acid at 60 °C furnished the tricyclic compound 40 in 98% yield. Retro-Diels-Alder reaction of 40 by the thermolysis at 230 °C in diphenyl ether, to our disappointment, provided a poor yield (25%) of the cyclohexenone derivative 41. This is presumably due to the instability of the N,O-isopropylidene group under the harsh reaction conditions. Fortunately, this problem was solved by replacement of the N,O-isopropylidene group with a robust cyclic carbamate group prior to subjection to the retro-Diels-Alder reaction (cf. Scheme 8). This is the subject of the following section.

2.8. Successful synthesis of the fully functionalized epoxycyclohexenone moiety 3b

The synthesis of the fully functionalized epoxycyclohexenone moiety **3b** was successfully achieved by exchanging the *N*,*O*-isopropylidene moiety in **37** with the corresponding cyclic carbamate functionality. Thus, as shown in Scheme 8, the *N*,*O*-isopropylidene moiety in **37** was selectively deprotected by exposure to aqueous hydrogen chloride in THF at 55 °C, which furnished an equilibrium mixture of the *N*-Ts- β -amino alcohol **42a** and the cyclic hemiacetal **42b** (ca. 1:1 by ¹H NMR). This equilibrium mixture was then treated with phosgene dimer

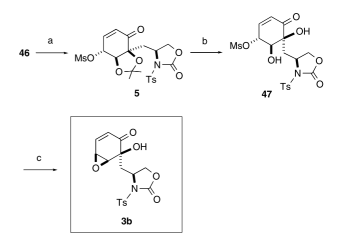


Scheme 8. Synthesis of the intermediate 46 for the epoxycyclohexenone 3. (a) 1.0 M HCl, THF, 55 °C; (b) phosgen dimer, pyridine, THF, rt, 67% (two steps); (c) TMSI, CCl₄, -20 °C, 74%; (d) Zn, AcOH, MeOH, 60 °C, 95%; (e) Ph₂O, 230 °C, 59%.

(trichloromethyl chloroformate) in the presence of pyridine in THF, providing the desired cyclic carbamate **43** in 67% yield for the two steps.

To forward the synthesis, regeneration of the cyclohexenone olefin moiety was next investigated. Thus, regioselective cleavage of the cyclopropane ring in 43 by reaction with TMSI afforded the expected iodide 44 in 74% yield. Further treatment of 44 with zinc powder in methanol containing acetic acid furnished the alcohol 45 in 95% yield. Retro-Diels-Alder reaction of 45 proceeded effectively by thermolysis at 230 °C in diphenyl ether. The desired cyclohexenone 46 was obtained in an acceptable 59% yield.

The final route that led to completion of the synthesis of the targeted molecule **3** is summarized in Scheme 9. The hydroxy group in **46** was mesylated under standard conditions (MsCl, Et₃N, DMAP, CH₂Cl₂, 0 °C \rightarrow room



Scheme 9. Synthesis of the intermediate 3b. (a) MsCl, Et₃N, DMAP, CH₂Cl₂, $0^{\circ}C \rightarrow rt$, 83%; (b) TFA, H₂O, $0^{\circ}C$, quant.; (c) 0.2 M NaOH, Et₂O, $0^{\circ}C$, 75%.

temperature) to give the corresponding mesylate **5** in 83% yield. The *O*-isopropylidene moiety of **5** was then hydrolyzed by reaction with aqueous trifluoroacetic acid at 0 °C to provide the requisite diol **47** in quantitative yield. Finally, brief exposure of **47** to aqueous sodium hydroxide in ether at 0 °C, led to the formation of **3b** in 75% yield. The structure and stereochemistry of **3b** were unambiguously confirmed by single X-ray crystallographic analysis as depicted in Figure 3.³⁵

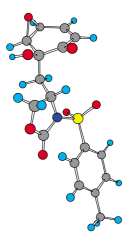


Figure 3. X-ray structure of the epoxycyclohexenone moiety 3b. Red, O; navy, N; yellow, S; blue, H.

3. Conclusion

In conclusion, we have succeeded in developing an efficient and enantioselective synthetic pathway to the epoxycyclohexenone moieties 2 and 3b of scyphostatin (1). The explored method features (i) the preparation of the key intermediate cyclohexene 10 and its olefin masked enone 25 starting from commercially available (-)-quinic acid (11), (ii) the aldol-type coupling reaction of the ketone 25 with benzaldehyde (8) or Garner's aldehyde analogue 9 to install the requisite asymmetric quaternary carbon center at the C6 position with complete stereoselectivity $(25+8\rightarrow 23 \text{ and }$ $25+9\rightarrow 24$), and (iii) the facile epoxide ring formation of the β -hydroxymesylates 35 and 47 under mild basic conditions $(35 \rightarrow 2 \text{ and } 47 \rightarrow 3b)$. Further investigation toward the synthesis of scyphostatin analogues based on the present study is now in progress and will be reported in due course.

4. Experimental

4.1. General methods

All reactions involving air- and moisture-sensitive reagent were carried out using oven-dried glassware and standard syringe-septum cap techniques. Routine monitorings of reaction were carried out using glass-supported Merck silica gel 60 F_{254} TLC plates. Flash column chromatography was performed on Kanto Chemical Silica Gel 60N (spherical, neutral 40–50 µm) with the solvents indicated.

All solvents and reagents were used as supplied with the following exceptions. Tetrahydrofuran (THF) and ether

were freshly distilled from sodium/benzophenone under argon. Dichloromethane, acetonitrile, and *N*,*N*-dimethyl-formamide (DMF) were distilled from calcium hydride under argon.

Melting points were taken on a Yanaco MP-3 micro melting point apparatus and are uncorrected. Measurements of optical rotations were performed with a JASCO P-1020 automatic digital polarimeter. ¹H and ¹³C NMR spectra were measured with a Brucker DRX-500 (500 MHz) spectrometer or a Brucker DRX-250 (250 MHz). Chemical shifts were expressed in ppm using tetramethylsilane ($\delta = 0$) as an internal standard. The following abbreviations are used: singlet (s), doublet (d), triplet (t), multiplet (m), and broad (br). Infrared (IR) spectral measurements were carried out with a JASCO FT/IR-5300 spectrometer. Low-resolution mass (MS) spectra was measured on Shimadzu GCMS-QP2010. High-resolution mass (HRMS) spectra was measured on JEOL MStation JMS-700 mass spectrometer. Elemental analyses were performed with a Perkin Elmer 2400II apparatus.

4.1.1. (1R,2R,3R)-3-tert-Butyldimethylsiloxy-1,2-(O-isopropylidenedioxy)cyclohexan-5-one (13). tert-Butyldimethylsilyl chloride (24.4 g, 0.16 mol) was added to a stirred solution of 12^{24} (10.0 g, 54 mmol) in dry DMF (120 ml) containing imidazole (14.7 g, 0.22 mol) at room temperature. After 15 h, the mixture was diluted with ethyl acetate (800 ml). The organic layer was successively washed with 3% aqueous hydrochloric acid $(2 \times 250 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2×250 ml), and brine $(2 \times 250 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 14:1) to give **13** (16.2 g, 98%) as a colorless oil. $[\alpha]_D^{20}$ +105.3 (*c* 1.00, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.05 (3H, s, Si-Me), 0.09 (3H, s, Si-Me), 0.83 (9H, s, Si-t-Bu), 1.35 (3H, s, C-Me), 1.42 (3H, s, C-Me), 2.37 (1H, ddd, J=1.9, 3.5, 17.5 Hz, C4–H), 2.64 (1H, dd, J=2.2, 17.4 Hz, C4–H), 2.65 (1H, dd, J=2.5, 17.5 Hz, C6–H), 2.75 (1H, dd, J=3.5, 17.5 Hz, C6–H), 4.16 (1H, m, C3–H), 4.22 (1H, dt, J=2.2, 7.2 Hz, C2–H), 4.69 (1H, m, C1–H); ¹³C NMR $(125 \text{ MHz}, \text{CDCl}_3) \delta -5.06, -5.04, 17.9, 23.9, 25.6 (3)$ carbons), 26.3, 40.1, 41.9, 68.7, 72.4, 75.1, 108.7, 207.7; IR (neat) 440, 520, 690, 780, 810, 840, 870, 910, 980, 1010, 1060, 1090, 1140, 1180, 1210, 1250, 1380, 1470, 1720, 2860, 2930, 2960 cm⁻¹; HREIMS (m/z) calcd for $C_{14}H_{25}O_4Si [(M-Me)^+]: 285.1522$, found 285.1253.

4.1.2. (1*R*,2*R*,3*R*,5*R*)- and (1*R*,2*R*,3*R*,5*S*)-3-tert-Butyldimethylsiloxy-5-hydroxy-1,2-(*O*-isopropylidenedioxy)cyclohexane (14) and (15). Sodium borohydride (1.30 g, 34 mmol) in water (15 ml) was added dropwise to a stirred solution of 13 (9.40 g, 31 mmol) in THF (400 ml) at -5 °C, and stirring was continued for 1 h at room temperature. The reaction was quenched with saturated aqueous ammonium chloride (30 ml) at 0 °C, and then the mixture was diluted with ethyl acetate (1000 ml). The organic layer was washed with saturated aqueous ammonium chloride (2×300 ml) and brine (2×300 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo gave a residue, which was separated by column chromatography (hexane/ethyl acetate, $5:1 \rightarrow 3:1$) to give **14** (5.02 g, 53%) as a more polar product and **15** (4.16 g, 44%) as a less polar product.

Compound **14**. Colorless prism, mp 47–48 °C; $[\alpha]_{D}^{20} - 41.7$ (*c* 1.01, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.10 (3H, s, Si-Me), 0.11 (3H, s, Si-Me), 0.89 (9H, s, *t*-Bu), 1.34 (3H, s, C-Me), 1.46 (3H, s, C-Me), 1.60 (1H, m, C4–H), 1.83 (1H, m, C6–H), 2.00 (1H, m, C4–H), 2.20 (1H, m, C6–H), 2.26 (1H, d, *J*=6.8 Hz, OH), 3.89–3.97 (2H, m, C2–H, C3–H), 4.02 (1H, br, C5–H), 4.40 (1H, dd, *J*=4.8, 9.8 Hz, C1–H); ¹³C NMR (125 MHz, CDCl₃) δ –4.81, –4.72, 18.0, 25.8 (3 carbons), 25.9, 28.1, 35.5, 37.7, 65.1, 71.1, 72.7, 78.87, 108.5; IR (KBr) 510, 550, 630, 660, 690, 780, 840, 870, 920, 940, 960, 1020, 1040, 1060, 1120, 1190, 1220, 1240, 1260, 1370, 1380, 1460, 2860, 2890, 2930, 2990, 3420 cm⁻¹; CIMS (*m*/*z*) 303 [(M+H)⁺]; HREIMS (*m*/*z*) calcd for C₁₄H₂₇O₄Si [(M−Me)⁺]: 287.1679, found 287.1682.

Compound **15**. Colorless oil. $[\alpha]_{D}^{20} - 33.7$ (*c* 1.02, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.08 (3H, s, Si-Me), 0.10 (3H, s, Si-Me), 0.88 (9H, s, *t*-Bu), 1.35 (3H, s, C-Me), 1.52 (3H, s, C-Me), 1.73 (1H, m, C4–H), 1.90 (1H, ddd, *J*=3.9, 6.3, 13.7 Hz, C4–H), 2.04 (2H, t, *J*=4.4 Hz, C6–H), 2.27 (1H, d, *J*=8.2 Hz, OH), 3.90 (1H, t, *J*=5.2 Hz, C2–H), 4.04 (1H, dd, br, *J*=7.7, 10.7 Hz, C5–H), 4.09 (1H, m, C3–H), 4.41 (1H, dd, *J*=4.6, 9.7 Hz, C1–H); ¹³C NMR (125 MHz, CDCl₃) δ –4.81, –4.72, 18.0, 25.7, 25.8 (3 carbons), 28.2, 33.8, 37.9, 65.3, 68.7, 74.1, 78.9, 108.6; IR (neat) 520, 660, 780, 840, 910, 940, 960, 1000, 1050, 1070, 1120, 1150, 1220, 1250, 1380, 1460, 2860, 2890, 2930, 2960, 2990, 3440 cm⁻¹; HREIMS (*m*/*z*) calcd for C₁₄H₂₇O₄Si [(M–Me)⁺]: 287.1679, found 287.1680.

4.1.3. (1R,2R,3R,5R)-5-Acetoxy-3-tert-butyldimethylsiloxy-1,2-(O-isopropylidenedioxy)cyclohexane (16). Acetic anhydride (0.1 ml, 1.1 mmol) was added to a stirred solution of 14 (27 mg, 89 µmol) in pyridine (1.0 ml) containing 4-dimethylaminopyridine (1.0 mg, 8 µmol) at 0 °C, and stirring was continued for 3 h at room temperature. The reaction was quenched with saturated aqueous sodium hydrogen carbonate (1 ml) at 0 °C, and the mixture was diluted with ether (30 ml). The organic layer was successively washed with 3% aqueous hydrochloric acid $(2 \times 15 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 15 \text{ ml})$, and brine $(2 \times 10 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 7:1) to give **16** (30 mg, 98%) as a colorless oil. $[\alpha]_D^{20} - 29.6$ (c 1.01, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.07 (3H, s, Si-Me), 0.09 (3H, s, Si-Me), 0.89 (9H, s, Si-t-Bu), 1.34 (3H, s, C-Me), 1.45 (1H, m, C4–H), 1.46 (3H, s, C-Me), 1.82 (1H, m, C6-H), 2.03 (3H, s, Ac), 2.12 (1H, m, C4-H), 2.35 (1H, m, C6–H), 3.80 (1H, m, C3–H), 3.87 (1H, t, J = 5.9 Hz, C2–H), 4.39 (1H, m, C1–H), 5.04 (1H, m, C5–H); ¹³C NMR $(125 \text{ MHz}, \text{CDCl}_3) \delta - 4.85, -4.86, 18.0, 21.3, 25.4, 25.7$ (3 carbons), 27.8, 31.2, 36.1, 66.8, 70.6, 73.0, 79.4, 108.5, 170.4; IR (neat) 410, 510, 610, 660, 700, 780, 840, 870, 900, 920, 940, 990, 1060, 1120, 1150, 1220, 1240, 1370, 1460, 1740, 2860, 2890, 2930, 2960, 2990 cm⁻¹; HREIMS (*m/z*) calcd for $C_{16}H_{29}O_5Si$ [(M-Me)⁺]: 329.1784, found 329.1796.

4.1.4. Conversion of 14 to 15. Diethyl azodicarboxylate in toluene (40% solution, 14.5 ml, 34 mmol) was added dropwise to a stirred solution of 14 (5.00 g, 17 mmol) in dry THF (150 ml) containing triphenylphosphine (8.68 g, 34 mmol) and benzoic acid (4.15 g, 34 mmol) at 0 °C under argon. The mixture was stirred for 3 h at room temperature. Concentration of the mixture in vacuo afforded a residue, which was purified by column chromatography (hexane/ ethyl acetate, 13:1) to give the corresponding benzoate (6.61 g, 98%) as a colorless oil. $[\alpha]_{D}^{20}$ +15.9 (c 1.06, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.09 (3H, s, Si-Me), 0.10 (3H, s, Si-Me), 0.90 (9H, s, Si-t-Bu), 1.37 (3H, s, C-Me), 1.55 (3H, s, C-Me), 1.92-2.10 (3H, m, C4-H, C4-H, C6–H), 2.24 (1H, dt, J=5.1, 14.5 Hz, C6–H), 3.96 (1H, t, J = 4.9 Hz, C2–H), 4.20 (1H, m, C3–H), 4.43 (1H, q, J =5.5 Hz, C1–H), 5.33 (1H, m, C5–H), 7.43 (2H, t, J=7.8 Hz, Ar-H), 7.55 (1H, t, J=7.4 Hz, Ar-H), 8.05 (2H, d, J= 7.1 Hz, Ar-H); ¹³C NMR (125 MHz, CDCl₃) δ -5.02, -4.84, 17.9, 25.7 (4 carbons), 30.9, 33.95, 33.99, 67.5, 68.8, 69.6, 72.8, 128.3 (2 carbons), 129.6 (2 carbons), 130.5, 132.9, 165.8, 207.1; IR (neat) 520, 710, 780, 840, 920, 940, 970, 1010, 1030, 1070, 1110, 1220, 1280, 1320, 1370, 1380, 1450, 1540, 1600, 1720, 1780, 2860, 2890, 2930 cm⁻¹; HREIMS (m/z) calcd for $C_{21}H_{31}O_5Si$ $[(M-Me)^+]$: 391.1941, found 391.1910.

2.0 M Potassium hydroxide solution (22.4 ml, 45 mmol) was added dropwise to a stirred solution of the above benzoate (6.50 g, 16 mmol) in methanol (280 ml) at 0 °C, and stirring was continued for 3 h at room temperature. The mixture was concentrated in vacuo to give a residue, which was diluted with ethyl acetate (800 ml). The organic layer was washed with brine (2×300 ml) and then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 3:1) to give **15** (4.82 g, quant.) as a colorless oil. The IR, ¹H NMR, and mass spectra of this material were identical with those recorded for preparation of **15** (see, Section 4.1.2).

4.1.5. (3R,4R,5R)-5-tert-Butyldimethylsiloxy-3,4-O-isopropylidenedioxy-1-cyclohexene (17). Diethyl azodicarboxylate in toluene (40% solution, 21.6 ml, 50 mmol) was added dropwise to a stirred solution of 15 (5.00 g, 17 mmol) in dry THF (150 ml) containing triphenylphosphine (13.1 g, 51 mmol) at room temperature. After 3 h, the mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, 13:1) to give 17 (3.15 g, 67%) as a colorless oil. $[\alpha]_{D}^{20} = 87.1$ (c 1.05, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.07 (3H, s, Si-Me), 0.10 (3H, s, Si-Me), 0.89 (9H, s, Si-t-Bu), 1.38 (3H, s, C-Me), 1.46 (3H, s, C-Me), 2.01 (1H, m, C6-H), 2.28 (1H, m, C6-H), 3.83 (1H, m, C5–H), 3.98 (1H, t, J=6.8 Hz, C4–H), 4.60 (1H, d, J=6.2 Hz, C3-H), 5.80 (2H, m, C1-H, C2-H);¹³C NMR (125 MHz, CDCl₃) δ -4.8, -4.5, 18.1, 25.8 (3 carbons), 26.1, 28.3, 31.9, 69.6, 72.8, 78.7, 108.6, 124.6, 128.5; IR (neat) 670, 780, 840, 910, 1010, 1060, 1120, 1220, 1250, 1380, 1460, 1690, 1730, 2860, 2930, 2960 cm⁻¹; HREIMS (m/z) calcd for C₁₄H₂₅O₃Si $[(M-Me)^+]$: 269.1573, found 269.1570.

4.1.6. (1S,2S,3R,4R,5R)-5-tert-Butyldimethylsiloxy-1,2-epoxy-3,4-(O-isopropylidenedioxy)cyclohexane (18). 3-Chloroperoxybenzoic acid (mCPBA) (7.53 g, 45 mmol) was added in small portions to a stirred solution of 17 (4.95 g, 17 mmol) in dry dichloromethane (180 ml) containing sodium hydrogen carbonate (7.53 g, 45 mmol) at 0 °C, and stirring was continued for 24 h at room temperature. The reaction was diluted with ethyl acetate (400 ml). The organic layer was washed with saturated aqueous sodium hydrogen carbonate $(2 \times 200 \text{ ml})$ and brine $(2 \times 200 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo affoded a residue, which was purified by column chromatography (hexane/ethyl acetate, 13:1) to give **18** (4.81 g, 92%) as a colorless oil. $[\alpha]_D^{20}$ -29.1 (c 1.04, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.06 (3H, s, Si-Me), 0.07 (3H, s, Si-Me), 0.88 (9H, s, Si-t-Bu), 1.38 (3H, s, C-Me), 1.46 (3H, s, C-Me), 1.90 (1H, ddd, J=1.6, 6.1, 15.6 Hz, C6–H), 2.18 (1H, ddd, J=4.0, 5.1, 15.4 Hz, C6–H), 3.14 (1H, d, J = 3.6 Hz, C2–H), 3.23 (1H, s, C1–H), 3.87 (1H, dd, J=5.8, 11.2 Hz, C5-H), 3.94 (1H, m, C4-H), 4.53 (1H, d, J=5.6 Hz, C3–H); ¹³C NMR (125 MHz, $CDCl_3$) $\delta - 4.77, -4.75, 18.0, 25.7$ (3 carbons), 26.0, 28.0, 29.4, 51.3, 51.7, 66.3, 71.8, 76.8, 109.2; IR (neat) 510, 710, 780, 840, 870, 910, 940, 1000, 1110, 1220, 1250, 1380, 1470, 2860, 2890, 2930, 2990 cm⁻¹; HREIMS (*m/z*) calcd for $C_{14}H_{25}O_4Si [(M-Me)^+]$: 285.1522, found 285.1508.

(1S,4R,5R,6R)-4-tert-Butyldimethylsiloxy-1-4.1.7. hydroxy-5,6-O-isopropylidenedioxy-2-cyclohexene (19). Sodium borohydride (663 mg, 18 mmol) was added in small portions to a stirred suspension of diphenyl diselenide (2.74 g, 8.8 mmol) in dry ethanol (30 ml) at 0 °C under argon. After 30 min, a solution of 18 (4.80 g, 16 mmol) in dry ethanol (30 ml) was added dropwise to the mixture at room temperature. The mixture was heated at reflux for 1 h. After cooling, the mixture was diluted with dry THF (24 ml). Hydrogen peroxide in water (30% solution, 19.5 ml, 0.17 mol) was added dropwise to the mixture at 0 °C. The resulting mixture was further stirred for 5 min at 0 °C and slowly heated at reflux for 1 h. After cooling, the mixture was diluted with ether (300 ml). The organic layer was washed with saturated aqueous sodium hydrogen carbonate $(2 \times 150 \text{ ml})$ and brine $(2 \times 100 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo gave a residue, which was purified by column chromatography (hexane/ethyl acetate, 3:1) to give **19** (3.74 g, 78%) as a colorless oil. $[\alpha]_D^{20}$ -42.4 (*c* 1.03, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.11 (3H, s, Si-Me), 0.13 (3H, s, Si-Me), 0.90 (9H, s, Si-t-Bu), 1.34 (3H, s, C-Me), 1.40 (3H, s, C-Me), 2.90 (1H, d, J=8.3 Hz, OH), 4.12 (1H, m, C1-H), 4.21 (1H, t, J=3.7 Hz, C4–H), 4.29 (1H, dd, J=4.1, 7.5 Hz, C5-H), 4.33 (1H, dd, J=4.1, 7.5 Hz, C6-H), 5.95 (1H, dd, J=4.2, 9.8 Hz, C3-H), 6.07 (1H, dd, J=4.2, 9.8 Hz, C2–H); ¹³C NMR (125 MHz, CDCl₃) δ –4.81, –4.75, 18.0, 24.6, 25.8 (3 carbons), 26.6, 67.9, 68.7, 78.8, 79.0, 108.7, 132.3, 132.3; IR (neat) 410, 480, 520, 640, 660, 690, 780, 840, 890, 940, 960, 990, 1010, 1060, 1120, 1160, 1210, 1250, 1380, 1460, 1640, 2860, 2900, 2930, 2960, 2990, 3050, 3450 cm⁻¹; HREIMS (m/z) calcd for $C_{14}H_{25}O_4Si [(M-Me)^+]: 285.1522$, found 285.1534.

4.1.8. (4R,5R,6S)-4-tert-Butyldimethylsiloxy-5,6-O-isopropylidenedioxy-2-cyclohexen-1-one (10). Dess-Martin periodinane (14.5 g, 34 mmol) was added in small portions to a stirred solution of **19** (5.15 g, 17 mmol) in dry dichloromethane (200 ml) at room temperature. After 2 h, the mixture was diluted with ethyl acetate (500 ml). The organic layer was washed successively with 20% aqueous sodium thiosulfate (2×200 ml), saturated aqueous sodium hydrogen carbonate $(2 \times 200 \text{ ml})$, and brine $(2 \times 200 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 7:1) to give 10 (4.86 g, 95%) as a white solid. Recrystallization from hexane/dichloromethane (5:1) afforded colorless prisms, mp 55–56 °C; $[\alpha]_{\rm D}^{20}$ – 84.7 (c 1.02, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.14 (3H, s, Si-Me), 0.17 (3H, s, Si-Me), 0.92 (9H, s, Si-t-Bu), 1.40 (3H, s, C-Me), 1.42 (3H, s, C-Me), 4.41 (1H, m, C5–H), 4.44 (1H, d, J = 5.9 Hz, C6–H), 4.54 (1H, m, C4–H), 6.08 (1H, d, J=10.3 Hz, C2–H), 6.76 (1H, ddd, J=1.0, 3.8, 10.3 Hz, C3-H); ¹³C NMR (125 MHz, CDCl₃) δ -4.74 (Si-Me), -4.73 (Si-Me), 18.1 (C-Me₃), 25.7 (3 carbons, C-Me₃), 25.9 (Me of O-isopropylidene), 27.4 (Me of O-isopropylidene), 67.1 (C4), 74.4 (C5), 79.6 (C6), 110.2 (C-Me₂ of O-isopropylidene), 127.9 (C2), 148.5 (C3), 194.5 (C1); IR (KBr) 470, 520, 630, 670, 730, 780, 840, 890, 940, 980, 1010, 1080, 1170, 1250, 1330, 1380, 1460, 1630, 1700, 2710, 2740, 2860, 2900, 2930, 2990, 3040, 3370, 3550 cm^{-1} ; EIMS (*m*/*z*) 298 (M⁺), 283 [(M-Me)⁺], 241 $[(M-t-Bu)^+]$. Anal. Calcd for C₁₅H₂₆O₄Si: C, 60.37; H, 8.78. Found C, 60.03; H, 8.56.

4.1.9. (1*S*,5*R*,6*S*,1'*R*,2'*R*,3'*R*,4'*S*)-5,2'-Bis(tert-butyldimethylsiloxy)-1,6:3',4'-bis(O-isopropylidenedioxy)bicyclohexyl-3-ene-2,5'-dione (20) and (4R,5S,6S)-6benzoyl-4-tert-butyldimethylsiloxy-5,6-O-isopropylidenedioxy-2-cyclohexen-1-one (22). Lithium bis(trimethylsilyl)amide in THF (1.0 M solution, 1.7 ml, 1.7 mmol) was added dropwise to a stirred solution of 10 (50 mg, 0.17 mmol) and benzaldehyde (8) (82 μ l, 0.77 mmol) in dry THF (4 ml) at -78 °C under argon, and the stirring was continued for 30 min at the same temperature. The reaction was guenched with saturated aqueous ammonium chloride (1 ml) at 0 °C, and the mixture was diluted with ether (50 ml). The organic layer was washed successively with saturated aqueous ammonium chloride $(2 \times 20 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2×20 ml), and brine $(2 \times 20 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethylacetate, 7:1) to give 20 (19 mg, 38%) as a white amorphous solid and 21 (7.4 mg, 12%) as colorless oil. Since compound 21 was unstable, this was immediately subjected to the following oxidation reaction.

Compound **21** (7.4 mg, 18 µmol) was treated with Dess-Martin periodinane (23.0 mg, 54 µmol) in dichloromethane (0.5 ml) at room temperature for 1 h. The reaction mixture was diluted with ethyl acetate (30 ml). The organic layer was washed successively with 20% aqueous sodium thiosulfate (2×10 ml), saturated aqueous sodium hydrogen carbonate (2×10 ml), and brine (2×10 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded

a residue, which was purified by column chromatography (hexane/ethyl acetate, 7:1) to give **22** (5.7 mg, 78%) as a colorless viscous oil.

Compound **20**. $[\alpha]_D^{20} - 30.0$ (*c* 1.10, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.10 (3H, s, Si-Me of C6'–OTBS), 0.15 (3H, s, Si-Me of C5-OTBS), 0.17 (3H, s, Si-Me of C5–OTBS), 0.22 (3H, s, Si-Me of C6'–OTBS), 0.84 (9H, s, *t*-Bu of C6'–OTBS), 0.92 (9H, s, *t*-Bu of C5–OTBS), 1.27 (3H, s, Me of O-isopropylidene), 1.30 (3H, s, Me of O-isopropylidene), 1.35 (3H, s, Me of O-isopropylidene), 1.45 (3H, s, Me of *O*-isopropylidene), 2.58 (1H, dd, J=7.0, 16.8 Hz, C2'–H), 2.70 (1H, dd, J=10.5, 17.8 Hz, C2'–H), 2.98 (1H, dd, J=7.1, 10.3 Hz, C1'-H), 4.05 (1H, br s, C6'-H), 4.10 (1H, t, J=1.7 Hz, C6-H), 4.26 (2H, br, C4'-H, C5'-H), 4.61 (1H, dd, J=1.0, 4.3 Hz, C5-H), 6.01 (1H, d, J = 10.2 Hz, C3–H) 6.58 (1H, br, C4–H); ¹³C NMR $(125 \text{ MHz}, \text{CD}_2\text{Cl}_2) \delta - 4.89, -4.80, -4.44, -4.30, 18.3,$ 18.4, 24.5, 25.9 (6 carbons), 26.5, 26.6, 27.1 (2 carbons), 30.1, 65.9, 78.2 (2 carbons), 79.3, 80.2, 83.5, 109.2, 111.9, 127.7, 143.5, 198.8, 204.5; IR (neat) 520, 670, 780, 810, 840, 940, 960, 980, 1010, 1040, 1070, 1100, 1130, 1180, 1210, 1230, 1260, 1380, 1470, 1700, 1730, 2860, 2930, 2950, 2990 cm⁻¹; HREIMS (m/z) calcd for C₂₉H₄₉O₈Si₂ $[(M-Me)^+]$, 581.2966, found 581.2949.

Compound **22**. $[\alpha]_{D}^{20} - 3.27$ (*c* 1.00, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.14 (3H, s, Si-Me), 0.15 (3H, s, Si-Me), 0.92 (9H, s, *t*-Bu), 1.27 (3H, s, C-Me), 1.46 (3H, s, C-Me), 4.58 (1H, dt, *J*=1.6, 2.2 Hz, C4–H), 4.79 (1H, dd, *J*= 1.2, 3.0 Hz, C5–H), 6.13 (1H, dd, *J*=1.6, 10.3 Hz, C2–H), 6.79 (1H, ddd, *J*=1.2, 3.4, 10.3 Hz, C3–H), 7.42 (2H, t, *J*= 7.8 Hz, Ph-H), 7.55 (1H, d, *J*=7.8 Hz, Ph-H), 8.22 (2H, d, *J*=7.3 Hz, Ph-H); ¹³C NMR (125 MHz, CDCl₃) δ -4.82, -4.72, 18.1, 25.7 (3 carbons), 26.3, 27.4, 68.1, 82.8, 89.7, 111.3, 127.1, 128.1 (2 carbons), 130.8 (2 carbons), 133.3, 134.9, 148.8, 192.8, 196.9; IR (neat) 690, 780, 840, 870, 900, 1050, 1100, 1180, 1260, 1380, 1450, 1460, 1580, 1600, 1680, 1700, 2860, 2890, 2930, 2960, 2990 cm⁻¹; HREIMS (*m*/*z*) calcd for C₂₂H₃₀O₅Si (M⁺): 402.1863, found 402.1835.

4.1.10. (1R,4S,4aR,5R,6S,7S,8aS)-5-tert-Butyldimethylsiloxv-6.7-O-isopropylidenedioxy-1.4.4a.5.6.7.8.8a-octahydro-endo-1,4-methanonaphthalen-8-one (27). Diethylaluminum chloride in hexane (1.0 M solution, 2.68 ml, 0.27 mmol) was added dropwise to a stirred solution of 10 (4.00 g, 13 mmol) and cyclopentadiene (11.1 ml, 0.13 mol) in dry dichloromethane (140 ml) at -78 °C under argon. The mixture was gradually warmed up to 0 °C over 1 h, and stirring was continued for 1 h at 0 °C. The mixture was diluted with ether (400 ml). The organic layer was washed with saturated aqueous sodium hydrogen carbonate $(2 \times 200 \text{ ml})$ and brine $(2 \times 200 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 6:1) to give 27 (4.74 g, 97%) as a white solid. Recrystallization from hexane/dichloromethane (10:1) afforded colorless prisms, mp 92–93 °C; $[\alpha]_D^{20}$ +46.7 (*c* 1.01, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.08 (3H, s, Si-Me), 0.09 (3H, s, Si-Me), 0.89 (9H, s, Si-t-Bu), 1.30 (3H, s, C-Me), 1.37 (1H, d, J=8.4 Hz, C9–H), 1.47 (3H, s, C-Me), 1.54 (1H, d, *J*=8.4 Hz, C9–H), 2.90 (1H, ddd, *J*=3.3, 5.6, 10.2 Hz, C4a–H), 3.08 (1H, s, C1–H), 3.11 (1H, s, C4–H), 3.18 (1H, dd, *J*=3.8, 10.2 Hz, C8a–H), 3.99 (1H, t, *J*=6.2 Hz, C5–H), 4.12 (1H, d, *J*=8.4 Hz, C7–H), 4.22 (1H, dd, *J*= 7.0, 8.3 Hz, C6–H), 6.12 (1H, dd, *J*=3.0, 5.6 Hz, C2–H), 6.20 (1H, dd, *J*=3.0, 5.4 Hz, C3–H); ¹³C NMR (125 MHz, CDCl₃) δ –4.93, –4.53, 18.0, 24.0, 25.8 (3 carbons), 26.5, 45.2, 45.7, 46.6, 49.6, 51.6, 71.6, 78.1, 79.7, 109.9, 133.1, 137.0, 208.7; IR (KBr) 560, 680, 730, 780, 840, 850, 900, 940, 970, 1010, 1040, 1080, 1110, 1160, 1210, 1260, 1380, 1460, 1720, 2860, 2900, 2930, 2960 cm⁻¹; EIMS (*m*/*z*) 349 [(M–Me)⁺], 307 [(M–*t*-Bu)⁺], 249 [(M–TBS)⁺]; CIMS (*m*/*z*) 365 [(M+H)⁺]; HREIMS (*m*/*z*) calcd for C₁₉H₂₉O₄Si [(M–Me)⁺]: 349.1835, found 349.1825.

4.1.11. (1R,4S,4aR,5R,6S,7S,8aS)-5-Trihydroxy-6,7-Oisopropylidenedioxy-1,4,4a,5,6,7,8,8a-octahydro-endo-1,4-methanonaphthalen-8-one (28). Tetrabutylammonium fluoride in THF (1.0 M solution, 15.0 ml, 15 mmol) was added to a stirred solution of 27 (3.51 g, 9.6 mmol) in dry THF (100 ml) at 0 °C, and stirring was continued for 2 h at room temperature. The mixture was diluted with ether (400 ml). The organic layer was successively washed with saturated aqueous ammonium chloride $(2 \times 150 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2×150 ml), and brine $(2 \times 150 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 5:3) to give 24 (1.81 g, 75%) as a white solid. Recrystallization from hexane/dichloromethane (10:1) afforded colorless needles, mp 137–138 °C; $[\alpha]_D^{20}$ +112.9 (*c* 1.00, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.34 (3H, s, C-Me), 1.42 (1H, d, J = 8.4 Hz, C9–H), 1.53 (3H, s, C-Me), 1.58 (1H, d, J =8.4 Hz, C9-H), 2.16 (1H, s, OH), 3.08-3.16 (3H, m, C1-H, C4–H, C5–H), 3.38 (1H, dd, J=3.3, 10.5 Hz, C8a–H), 4.15 (1H, t, J=4.1 Hz, C4a–H), 4.20 (1H, d, J=8.0 Hz, C7–H), 4.43 (1H, dd, J=5.9, 8.0 Hz, C6–H), 6.25 (1H, dd, J=2.6, 5.5 Hz, C3–H), 6.37 (1H, dd, J=3.0, 5.5 Hz, C2–H); ¹³C NMR (125 MHz, CDCl₃) δ 23.9, 26.4, 44.6 (2 carbons), 45.3, 48.4, 51.7, 70.2, 78.1, 79.7, 111.0, 134.9, 137.4, 208.6; IR (KBr) 550, 740, 860, 890, 1040, 1060, 1160, 1210, 1260, 1380, 1630, 1710, 2940, 2980, 3440 cm⁻¹; EIMS (*m/z*) 250 (M^+) , 235 $[(M-Me)^+]$. Anal. Calcd for $C_{14}H_{18}O_4$: C, 67.18; H, 7.25. Found: C, 67.35, H, 7.24.

(1S,2S,3S,4R,4aR,5R,6S,7S,8aR)-2-Bromo-4.1.12. 3,5-epoxy-6,7-(O-isopropylidenedioxy)perhydro-endo-1,4-methanonaphthalen-8-one (25). N-Bromosuccinimide (2.78 g, 16 mmol) was added in small portions to a stirred solution of 28 (3.02 g, 12 mmol) in dry dichloromethane (180 ml) at 0 °C, and stirring was continued for 1 h at room temperature. The mixture was diluted with ether (400 ml). The organic layer was washed successively with 20% aqueous sodium thiosulfate $(2 \times 150 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2 \times 150 ml), and brine (2 \times 150 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 4:1) to give 25 (3.41 g, 86%) as a white solid. Recrystallization from hexane/ether (5:1) afforded pale yellow prisms, mp 121- $122 \text{ °C}; [\alpha]_D^{20} + 65.9 (c \ 1.00, \text{ CHCl}_3); ^1\text{H NMR} (500 \text{ MHz},$ CDCl₃) & 1.33 (3H, s, C-Me), 1.52 (3H, s, C-Me), 1.67 (1H, d, J=11.2 Hz, C9-H), 1.22 (1H, d, J=11.2 Hz, J=11.2

C9–H), 2.82 (1H, br, C1–H), 2.97 (1H, br, C4–H), 3.00– 3.10 (2H, m, C4a–H, C8a–H), 3.82 (1H, d, J=1.2 Hz, C2–H), 3.88 (1H, t, J=1.9 Hz, C5–H), 4.28 (1H, d, J=6.3 Hz, C7–H), 4.44 (1H, dd, J=0.8, 5.3 Hz, C3–H), 4.54 (1H, dd, J=1.7, 6.3 Hz, C6–H); ¹³C NMR (125 MHz, CDCl₃) δ 24.7, 26.6, 33.7, 41.4, 42.7, 46.1, 47.5, 55.0, 77.5, 77.7, 78.1, 87.3, 111.2, 207.0; IR (KBr) 540, 710, 750, 770, 810, 840, 860, 890, 940, 970, 1060, 1090, 1160, 1210, 1270, 1310, 1380, 1460, 1720, 1790, 2890, 2940, 2990 cm⁻¹; EIMS (m/z) 330 [(M+2)⁺, ⁸¹Br], 328 (M⁺, ⁷⁹Br), 315 [(M–Me+2)⁺, ⁸¹Br], 313 [(M–Me)⁺, ⁷⁹Br]. Anal. calcd for C₁₄H₁₇BrO₄: C, 51.08; H, 5.21; Br, 24.27. Found: C, 51.33; H, 5.23; Br, 24.50.

4.1.13. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4,5-(*O*-isopropylidenedioxy)tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one (29). Lithium bis(trimethylsilyl)amide in THF (1.0 M solution, 67 µl, 67 µmol) was added dropwise to a stirred solution of 25 (20 mg, 61 µmol) in dry THF (1.5 ml) at -78 °C under argon, and the stirring was continued for 30 min at the same temperature. The reaction was quenched with saturated aqueous ammonium chloride (1 ml) at -78 °C, and the mixture was diluted with ether (40 ml). The organic layer was washed successively with saturated aqueous ammonium chloride $(2 \times 15 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2 \times 15 ml), and brine (2 \times 15 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 2:1) to give 29 (15 mg, 98%) as a white solid. Recrystallization from hexane/dichloromethane (10:1) afforded colorless prisms, mp 77–78 °C; $[\alpha]_D^{20}$ –61.6 (*c* 1.05, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.39 (3H, s, C-Me); 1.45 (3H, s, C-Me); 1.67 (1H, dd, J=1.3, 5.0 Hz, C10–H); 1.80 (1H, d, J= 11.4 Hz, C11–H); 1.86 (1H, d, J=11.4 Hz, C11–H); 2.42 (1H, d, *J*=4.3 Hz, C1–H); 2.46 (1H, s, C8–H); 2.87 (1H, t, *J*=2.4 Hz, C7–H); 4.40–4.45 (3H, m, C4–H, C6–H, C9–H); 4.66 (1H, dd, J=1.3, 5.7 Hz, C5–H); ¹³C NMR (125 MHz, CDCl₃) δ 21.5, 25.0, 27.2, 30.4, 32.5, 34.3, 41.7, 44.7, 77.6, 77.9, 79.4, 82.5, 110.1, 201.7; IR (KBr) 520, 580, 630, 830, 860, 880, 920, 940, 960, 990, 1020, 1040, 1070, 1160, 1210, 1270, 1300, 1330, 1380, 1700, 2880, 2900, 2940, 2990 cm⁻¹; EIMS (m/z) 248 (M⁺), 233 [(M-Me)⁺], 190 $[(M - Me_2CO)^+]$. Anal. Calcd for C₁₄H₁₆O₄: C, 67.73; H, 6.50. Found: C, 67.51; H, 6.57.

4.1.14. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4-[hydroxy(phenyl)methy]-4,5-(O-isopropylidenedioxy)tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one (23). Lithium bis(trimethylsilyl)amide in THF (1.0 M solution, 94 µl, 94 µmol) was added dropwise to a stirred solution of 29 (21.0 mg, 85 μ mol) in dry THF (1.0 ml) at -78 °C under argon. After 30 min, a solution of benzaldehyde (8) (18 µl, 0.17 mmol) in dry THF (0.5 ml) was added slowly at -78 °C, and the stirring was continued for 1 h at the same temperature. The reaction was quenched with saturated aqueous ammonium chloride (1 ml) at 0 °C, and the mixture was diluted with ether (15 ml). The organic layer was washed successively with saturated aqueous ammonium chloride $(2 \times 7 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 7 \text{ ml})$, and brine $(2 \times 7 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography

(hexane/ethyl acetate, $2:1 \rightarrow 1:1$) to give **23** (29.3 mg, 98%) as a hardly separable epimeric mixture (6:1 by 500 MHz ¹H NMR). In order to obtain analytical samples, a small amount of the epimeric mixture **23** was further subjected to column chromatography (hexane/ethyl acetate, $2:1 \rightarrow 1:1$) to provide pure samples of **23a** (major, more polar) and **23b** (minor, less polar).

Compound **23a**. Colorless prisms; mp 181–182 °C; $[\alpha]_D^{20}$ -68.9 (c 1.17, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.98 (3H, s, C-Me), 1.38 (3H, s, C-Me), 1.61 (1H, d, J=5.2 Hz, OH), 1.78 (1H, d, J=11.5 Hz, C11-H), 1.84 (1H, d, J= 11.5 Hz, C11–H), 2.30 (1H, d, J = 5.2 Hz), 2.45 (1H, s), 2.90 (1H, t, J=2.3 Hz), 3.13 (1H, d, J=4.1 Hz), 4.44 (1H, d, J=3.0 Hz), 4.51 (1H, t, J=2.3 Hz), 4.52 (1H, t, J=2.8 Hz), 5.05 (1H, d, J=8.5 Hz), 7.28-7.36 (3H, m, Ph), 7.42-7.46 (2H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 20.9, 26.2, 28.0, 30.3, 30.4, 32.4, 41.1, 46.6, 76.4, 77.7, 79.6, 83.2, 88.1, 110.1, 127.8 (2 carbons), 128.4, 129.2 (2 carbons), 138.0, 205.6; IR (KBr) 510, 600, 700, 730, 830, 880, 920, 1020, 1060, 1110, 1170, 1250, 1290, 1380, 1450, 1720, 2940, 2990, 3380 cm⁻¹; EIMS (*m/z*) 337 [(M-OH)⁺]; CIMS (m/z) 355 $[(M+H)^+]$. Anal. Calcd for $C_{21}H_{22}O_6$: C, 71.17; H, 6.26. Found: C, 71.24; H 6.35.

Compound **23b**. Colorless viscous oil. $\left[\alpha\right]_{D}^{20} - 85.2$ (c 1.20, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.91 (3H, s, C-Me), 1.38 (3H, s, C-Me), 1.78 (1H, d, J=11.5 Hz, C11-H), 1.84 (1H, d, J=11.5 Hz, C11–H), 1.92 (1H, dd, J=1.5, 5.1 Hz, OH), 2.41 (1H, d, J = 5.1 Hz), 2.46 (1H, s), 2.89 (1H, t, J =2.3 Hz), 3.23 (1H, d, J = 8.6 Hz), 4.51 (2H, m), 4.69 (1H, d, J=2.7 Hz), 5.06 (1H, d, J=8.5 Hz), 7.27–7.31 (1H, m, Ph), 7.34 (2H, t, J=7.4 Hz, Ph), 7.44 (2H, d, J=7.2 Hz, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 22.4, 26.5, 28.5, 30.3, 33.3, 33.3, 41.0, 45.2, 75.3, 77.0, 78.6, 83.3, 86.9, 110.5, 127.8 (2 carbons), 128.1, 128.5 (2 carbons), 139.2, 204.2; IR (neat) 510, 590, 710, 730, 830, 880, 920, 1060, 1170, 1240, 1290, 1380, 1450, 1700, 2940, 2990, 3470 cm⁻¹; EIMS (*m/z*) 337 $[(M-OH)^+]$, 248 $[(M-PhCHO)^+]$ CIMS (m/z) 355 $[(M+H)^+]$; HREIMS (*m/z*) calcd for C₁₄H₁₆O₄ [(M-PhCHO)⁺]: 248.1049, found 248.1057.

4.1.15. One-pot procedure for the preparation of 23 from 25. Lithium bis(trimethylsilyl)amide in THF (1.0 M solution, 1.43 ml, 1.4 mmol) was added dropwise to a stirred solution of 25 (213 mg, 0.65 mmol) in dry THF (8 ml) at -78 °C under argon. After 30 min, a solution of benzaldehyde (8) (0.20 ml, 2.0 mmol) in dry THF (1 ml) was added slowly to the mixture at -78 °C, and the stirring was continued for 1 h at the same temperature. The reaction was quenched with saturated aqueous ammonium chloride (2 ml) at 0 °C, and the mixture was diluted with ether (50 ml). The organic layer was washed successively with saturated aqueous ammonium chloride $(2 \times 20 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2×20 ml), and brine $(2 \times 20 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, $2:1 \rightarrow$ 1:1) to give 23 (225 mg, 98%) as an epimeric mixture (6:1 by 500 MHz ¹H NMR). The IR, ¹H NMR, and mass spectra of this material were identical with those recorded for the preparation of 23 (see, Section 4.1.14).

[(1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4.1.16. 4,5-(O-isopropylidenedioxy)tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one-4-yl](phenyl)methyl O-phenyl carbonothioate (30). Phenyl thionochloroformate (0.18 ml, 1.3 mmol) was added to a stirred solution of 23 (6:1 epimeric mixture) (230 mg, 0.65 mmol) in dry acetonitrile (10 ml) containing 4-dimethylaminopyridine (DMAP) (316 mg, 2.6 mmol) at room temperature. After 12 h, the mixture was diluted with diethyl ether (100 ml). The organic layer was successively washed with 3% aqueous hydrochloric acid $(2 \times 50 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 50 \text{ ml})$, and brine $(2 \times 50 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 5:2) to give 30 (292 mg, 92%) as a hardly separable epimeric mixture (6:1 by 500 MHz ¹H NMR), as a colorless oil. In order to obtain analytical samples, a small amount of the epimeric mixture 30 was further subjected to column chromatography (hexane/ethyl acetate, $4:1 \rightarrow 3:1$) to provide pure samples of 30a (major, more polar) and 30b (minor, less polar).

Compound **30a**. Colorless viscous oil. $[\alpha]_{20}^{20} - 33.4$ (*c* 1.05, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.09 (3H, s, C-Me), 1.44 (3H, s, C-Me), 1.68 (1H, br), 1.75 (1H, d, *J*=11.5 Hz), 1.83 (1H, d, *J*=11.5 Hz), 2.23 (1H, d, *J*=5.2 Hz), 2.44 (1H, s), 2.90 (1H, t, *J*=2.4 Hz), 4.50 (1H, br), 4.53 (1H, t, *J*=2.7 Hz), 4.60 (1H, br d, *J*=2.2 Hz), 6.70 (1H, s), 6.99–7.03 (2H, m), 7.22–7.27 (1H, m), 7.32–7.41 (5H, m), 7.51–7.55 (2H, m); ¹³C NMR (125 MHz, CDCl₃) δ 20.5, 26.2, 28.1, 30.1, 30.3, 32.2, 41.2, 46.9, 76.5, 80.0, 83.2, 86.2, 87.1, 110.8, 121.9 (2 carbons), 126.5, 128.0 (2 carbons), 129.2, 129.4 (2 carbons), 130.2 (2 carbons), 133.8, 153.5, 194.1, 202.2; IR (neat) 510, 690, 750, 850, 880, 940, 1030, 1070, 1120, 1200, 1270, 1380, 1460, 1490, 1590, 1710, 2940, 2990 cm⁻¹; HREIMS (*m*/*z*) calcd for C₂₈H₂₆O₆S (M⁺): 490.1450, found 490.1428.

Compound **30b**. Colorless viscous oil. $[\alpha]_{20}^{20} - 64.7$ (*c* 1.14, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.90 (3H, s, C-Me), 1.39 (3H, s, C-Me), 1.82 (1H, d, J=11.4 Hz), 1.88 (1H, d, J=11.4 Hz), 2.15 (1H, dd, J=1.8, 5.1 Hz), 2.46 (1H, d, J= 5.0 Hz), 2.50 (1H, s), 2.94 (1H, t, J=2.4 Hz), 4.55 (1H, t, J=2.5 Hz), 4.58 (1H, t, J=2.3 Hz), 4.90 (1H, d, J= 2.5 Hz), 6.55 (1H, s), 6.98–7.03 (2H, m, Ph), 7.22–7.27 (1H, m, Ph), 7.32–7.46 (5H, m, Ph), 7.49–7.55 (2H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 22.2, 26.5, 28.7, 30.3, 33.5, 33.6, 41.3, 45.2, 77.2, 79.1, 83.1, 83.9, 85.8, 110.5, 121.9 (2 carbons), 126.6, 128.1 (2 carbons), 128.9, 129.3 (2 carbons), 129.5 (2 carbons), 134.2, 153.4, 192.9, 202.7; IR (neat) 690, 750, 880, 1020, 1070, 1200, 1270, 1380, 1460, 1490, 1590, 1700, 2940, 2990 cm⁻¹; HREIMS (m/z) calcd for C₂₈H₂₆O₆S (M⁺): 490.1450, found 490.1476.

4.1.17. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-4-Benzyl-6, 9-epoxy-4, 5-(*O*-isopropylidenedioxy)tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one (31). Tri-*n*-butyltin hydride (0.35 ml, 1.3 mmol) and azobisisobutyronitrile (AIBN) (21 mg, 0.13 mmol) were added to a solution of 30 (6:1 epimeric mixture) (213 mg, 0.43 mmol) in dry toluene (7.5 ml). For the deaeration of the reaction mixture, it was frozen using liquid nitrogen, and the reaction vessel was evacuated in vacuo for 30 min and then filled with dry argon. The mixture was heated at reflux for 2 h. After cooling, the mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, $1:0 \rightarrow 3:1$) to give **31** (116 mg, 79%) as a white solid. Recrystallization from hexane/ether (5:1) afforded colorless prisms, mp 107–108 °C; $\left[\alpha\right]_{\rm D}^{20}$ – 69.6 (c 0.97, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.91 (3H, s, C-Me), 1.36 (3H, s, C-Me), 1.67 (1H, dd, J=1.6, 5.1 Hz, C10-H), 1.80 (1H, d, J=11.4 Hz, C11-H), 1.85 (1H, d, J= 11.4 Hz, C11–H), 2.33 (1H, d, J=4.9 Hz, C1–H), 2.46 (1H, s, C8–H), 2.90 (1H, t, J=2.3 Hz, C7–H), 2.99 (1H, d, J= 13.7 Hz, CH_aH_bPh), 3.22 (1H, d, J=13.7 Hz, CH_aH_bPh), 4.39 (1H, d, J=3.0 Hz, C5-H), 4.54 (1H, t, J=2.8 Hz, C6-H), 4.57 (1H, t, J=2.3 Hz, C9-H), 7.22-7.34 (5H, m, Ph); 13 C NMR (125 MHz, CDCl₃) δ 20.6, 26.3, 28.1, 30.3, 31.9, 31.9, 41.2, 43.3, 46.1, 76.8, 78.5, 83.2, 85.7, 109.2, 127.0, 127.9 (2 carbons), 131.9 (2 carbons), 135.2, 207.5; IR (KBr) 510, 620, 700, 770, 830, 880, 920, 940, 990, 1030, 1060, 1080, 1100, 1120, 1140, 1170, 1240, 1300, 1330, 1380, 1450, 1490, 1710, 2870, 2930, 2990 cm⁻¹; EIMS (*m*/ z) 338 (M⁺), 280 [(M-Me₂CO)⁺], 247 [(M-PhCH₂)⁺]. Anal. Calcd for C21H22O4: C, 74.54; H 6.55. Found: C, 74.65; H, 6.61.

4.1.18. (1S,2S,3S,4R,4aR,5R,6S,7S,8aR)-7-Benzyl-3,5epoxy-2-iodo-6,7-O-isopropylidenedioxy-1,2,3,4,4a,5,6,8aoctahydro-endo-1,4-methanonaphthalen-8-one (32). Iodotrimethylsilane (0.10 ml, 0.70 mmol) was added dropwise to a stirred solution of **31** (120 mg, 0.36 mmol) in carbon tetrachloride (4 ml) at -20 °C under argon, and stirring was continued for 3 h at -10 °C. The reaction was quenched with 20% aqueous sodium thiosulfate (2 ml) at -10 °C, and then the mixture was diluted with ether (40 ml). The organic layer was washed successively with 20% aqueous sodium thiosulfate $(2 \times 20 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 20 \text{ ml})$, and brine $(2 \times 20 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 4:1) to give **32** (147 mg, 89%) as a colorless viscous oil. $[\alpha]_D^{20} + 62.7$ (*c* 0.96, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.89 (3H, s, C-Me), 1.32 (3H, s, C-Me), 1.84 (1H, d, J=11.3 Hz, C9–H), 2.24 (1H, d, J=11.3 Hz, C9-H), 2.88-2.97 (4H, m, C1-H, C4-H, C4a-H, $CH_{a}H_{b}Ph$), 3.07 (1H, dd, J=4.9, 10.1 Hz, C8a–H), 3.35 (1H, d, J=14.1 Hz, CH_aH_bPh), 3.61 (1H, d, J=2.4 Hz, C2-H), 4.25 (1H, t, J=3.7 Hz, C5-H), 4.36 (1H, d, J= 4.3 Hz, C6-H), 4.80 (1H, d, J=5.4 Hz, C3-H), 7.22-7.32 (5H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 26.5, 28.3, 33.0, 36.5, 38.7, 39.7, 47.0, 47.7, 48.2, 75.7, 76.1, 82.1, 89.4, 110.3, 126.8, 127.9 (2 carbons), 132.0 (2 carbons), 135.3, 210.6; IR (neat) 520, 610, 660, 700, 730, 770, 850, 890, 910, 940, 970, 1050, 1060, 1120, 1160, 1220, 1230, 1260, 1380, 1450, 1490, 1710, 2890, 2930, 2990 cm⁻ EIMS (m/z) 466 (M^+) , 451 $[(M-Me)^+]$, 408 $[(M-Me)^+]$ $Me_2CO)^+$], 375 [(M – PhCH₂)⁺]; HREIMS (*m*/*z*) calcd for $C_{21}H_{23}IO_4$ (M⁺): 466.0641, found 466.0636.

4.1.19. (1*R*,4*S*,4*aR*,5*R*,6*S*,7*S*,8*aS*)-7-Benzyl-5-hydroxy-6,7-*O*-isopropylidenedioxy-1,4,4*a*,5,6,7,8,8*a*-octahydro*endo*-1,4-methanonaphthalen-8-one (33). Zinc powder (272 mg, 4.2 mmol) and acetic acid (0.24 ml, 4.2 mmol) were successively added to a stirred solution of 32 (130 mg, 0.28 mmol) in methanol (5 ml) at room temperature. The mixture was gradually warmed to 60 °C and stirred for 1 h at the same temperature. After cooling, the mixture was diluted with ether (80 ml) and filtrated. The filtrate was washed with saturated aqueous sodium hydrogen carbonate $(2 \times 30 \text{ ml})$, and brine $(2 \times 30 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ ethyl acetate, 4:1) to give 33 (86 mg, 91%) as a white solid. Recrystallization from hexane/dichloromethane (10:1) afforded colorless prisms, mp 169–170 °C; $[\alpha]_D^{20}$ +161.0 (c 0.99, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 0.79 (3H, s, C-Me), 1.41 (1H, d, J=8.4 Hz, C9-H), 1.47 (3H, s, C-Me), 1.54 (1H, d, J=8.4 Hz, C9-H), 1.58 (1H, d, J=3.2 Hz, OH), 2.73 (1H, d, J = 14.4 Hz, CH_aH_bPh), 2.99 (1H, s, C1-H), 3.21 (2H, m, C4-H, C8a-H), 3.41 (1H, d, J= 14.4 Hz, CH_aH_bPh), 3.46 (1H, dd, J=3.7, 11.6 Hz, C4a–H), 4.40 (1H, d, J=4.6 Hz, C6–H), 4.45 (1H, q, J=3.9 Hz, C5–H), 6.20 (1H, dd, J=2.9, 5.4 Hz, C2–H), 6.48 (1H, dd, J=3.1, 5.4 Hz, C3–H), 7.20 (1H, m, Ph), 7.28–7.26 (4H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 26.5, 27.5, 38.8, 43.3, 43.5, 45.3, 47.2, 51.0, 68.1, 80.1, 83.2, 111.4, 126.3, 127.7 (2 carbons), 132.0 (2 carbons), 133.0, 136.6, 139.1, 207.6; IR (KBr) 540, 620, 660, 700, 730, 750, 820, 850, 910, 980, 1060, 1080, 1150, 1170, 1220, 1240, 1260, 1340, 1380, 1450, 1500, 1710, 2940, 2990, 3520 cm⁻¹; EIMS (*m/z*) 340 (M^+) , 325 $[(M - Me)^+]$, 282 $[(M - Me_2CO)^+]$, 249 $[(M - Me_2CO)^+]$ $PhCH_2$)⁺]. Anal. Calcd for $C_{21}H_{24}O_4$: C, 74.09, H, 7.11. Found: C, 74.01, H, 7.17.

4.1.20. (4R,5S,6S)-6-Benzyl-4-hydroxy-5,6-O-isopropylidenedioxy-2-cyclohexen-1-one (34). A stirred solution of 33 (60 mg, 0.18 mmol) in diphenyl ether (4 ml) was heated at 230 °C for 4 h. After cooling, the mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, $1:0 \rightarrow 2:1$) to give **34** (39.5 mg, 81%) as a colorless viscous oil. $[\alpha]_D^{20} = 0.7$ (c 0.97, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.08 (3H, s, C-Me), 1.29 (3H, s, C-Me), 1.70 (1H, d, J = 6.2 Hz, OH), 3.04 (1H, d, J = 14.1 Hz, CH_aH_bPh), $3.17 (1H, d, J = 14.1 \text{ Hz}, CH_aH_bPh), 4.13 (1H, t, J = 1.8 \text{ Hz},$ C5-H), 4.61 (1H, m, C4-H), 6.16 (1H, d, J=10.1 Hz, C2-H), 6.81 (1H, ddd, J = 2.0, 4.9, 10.1 Hz, C3-H), 7.25-7.34 (5H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 26.1, 27.4, 38.9, 64.6, 78.5, 81.8, 108.6, 127.1, 127.8 (2 carbons), 128.3, 131.5 (2 carbons), 134.7, 144.4, 199.6; IR (neat) 530, 590, 620, 670, 700, 730, 760, 800, 830, 860, 900, 920, 970, 1030, 1060, 1080, 1110, 1140, 1170, 1230, 1240, 1380, 1440, 1450, 1500, 1680, 2930, 2990, 3030, 3060, 3450 cm^{-1} ; EIMS (*m*/*z*) 274 (M⁺), 259 [(M-Me)⁺]; HREIMS (m/z) calcd for C₁₅H₁₅O₄ [$(M - Me)^+$]: 259.0970, found 259.0992.

4.1.21. (1*R*,5*S*,6*S*)-5-Benzyl-5,6-*O*-isopropylidenedioxy-**4-oxo-2-cyclohexenyl methanesulfonate** (4). Methanesulfonyl chloride (0.17 ml, 2.1 mmol) was added to a stirred solution of **34** (58.8 mg, 0.21 mmol) in dichloromethane (5 ml) containing triethylamine (0.42 ml, 3.0 mmol) and 4-dimethylaminopyridine (DMAP) (24 mg, 0.21 mmol) at 0 °C, and stirring was continued for 3 h at room temperature. The reaction was quenched with saturated aqueous sodium hydrogen carbonate (2 ml) at 0 °C, and the mixture was diluted with ether (80 ml). The organic layer was successively washed with 3% aqueous hydrochloric acid $(2 \times 30 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 30 \text{ ml})$, and brine $(2 \times 30 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 1:1) to give 4 (63 mg, 85%) as a colorless viscous oil. $[\alpha]_{\rm D}^{20}$ -62.6 (c 1.19, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.08 (3H, s, C-Me), 1.30 (3H, s, C-Me), 2.86 (1H, d, J = 14.5 Hz, $CH_{a}H_{b}Ph$), 3.10 (3H, s, Ms), 3.27 (1H, d, J=14.5 Hz, CH_aH_bPh), 4.14 (1H, t, J = 1.7 Hz, C5–H), 5.45 (1H, dd, J =1.8, 4.8 Hz, C4–H), 6.30 (1H, d, J=10.1 Hz, C2–H), 6.85 (1H, ddd, J=1.8, 4.8, 10.1 Hz, C3-H), 7.25-7.34 (5H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 26.3, 27.4, 38.1, 38.8, 71.2, 76.3, 81.6, 109.7, 127.3, 128.1 (2 carbons), 130.8, 131.4 (2 carbons), 134.1, 138.9, 197.8; IR (KBr) 530, 620, 700, 760, 790, 850, 900, 950, 980, 1070, 1090, 1120, 1150, 1180, 1230, 1370, 1450, 1500, 1690, 1740, 2940, 2990, 3030 cm^{-1} ; EIMS (*m/z*) 352 (M⁺), 337 [(M-Me)⁺], 294 $[(M-Me_2CO)^+]$; HREIMS (m/z) calcd for $C_{17}H_{20}O_6S$ (M⁺): 352.0981, found 352.0982.

4.1.22. (1R,5S,6S)-5-Benzyl-5,6-dihydroxy-4-oxo-2cyclohexenyl methanesulfonate (35). A solution of 4 (60 mg, 0.17 mmol) in trifluoroacetic acid/water (6:1) (1 ml) was stirred at 0 °C for 30 min. The mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, 1:1) to give **35** (45 mg, 85%) as a colorless viscous oil. $[\alpha]_D^{20}$ $-49.8 (c 0.95, \text{CHCl}_3);$ ¹H NMR (500 MHz, CDCl₃) δ 2.99 (1H, d, J=3.6 Hz, C3–OH), 3.16 (3H, s, Ms), 3.18 (2H, s, CH₂Ph), 3.35 (1H, s, C6–OH), 4.14 (1H, dt, J = 1.1, 3.7 Hz, C5-H), 5.47 (1H, dt, J=1.1, 3.9 Hz, C4-H), 6.24 (1H, dd, J=1.1, 10.2 Hz, C2-H), 6.85 (1H, ddd, J=1.1, 3.7, 10.2 Hz, C3-H), 7.14-7.19 (2H, m, Ph), 7.22-7.31 (3H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 38.6, 40.9, 73.3, 75.7, 78.1, 127.3, 128.4 (2 carbons), 129.0, 130.6 (2 carbons), 134.1, 141.7, 197.1; IR (neat) 530, 700, 730, 780, 850, 880, 940, 980, 1060, 1110, 1170, 1360, 1440, 1450, 1490, 1690, 2930, 3030, 3480 cm⁻¹; EIMS (*m/z*) 312 (M^+) , 294 $[(M-H_2O)^+]$; HREIMS (m/z) calcd for $C_{14}H_{16}O_6S$ (M⁺): 312.0668, found 312.0656.

4.1.23. (4S,5S,6S)-6-Benzyl-4,5-epoxy-6-hydroxy-2cyclohexen-1-one (2). 0.2 M Sodium hydroxide (0.7 ml, 0.14 mmol) was added dropwise to a stirred solution of 35 (30 mg, 96 µmol) in ether (8 ml) at 0 °C. After 10 min, the mixture was extracted with ether $(2 \times 30 \text{ ml})$. The combined extracts were washed with brine $(3 \times 20 \text{ ml})$ and dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 2:1) to give 2 (19 mg, 90%) as a white solid. Recrystallization from hexane/ether (5:1) afforded colorless prisms, mp 96–97 °C; $[\alpha]_{D}^{20}$ +45.6 (c 0.80, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 2.93 (1H, d, J= 13.6 Hz, CH_aH_bPh), 3.01 (1H, d, J=13.6 Hz, CH_aH_bPh), 3.60 (1H, dt, J=1.6, 3.9 Hz, C4–H), 3.65 (1H, s, OH), 3.77 (1H, d, J=3.9 Hz, C5-H), 6.16 (1H, dd, J=1.5, 9.9 Hz,C2-H), 7.09-7.15 (3H, m, Ph, C3-H), 7.22-7.32 (3H, m, Ph); ¹³C NMR (125 MHz, CDCl₃) δ 44.4 (CH₂Ph), 47.9 (C4 or C5), 56.0 (C4 or C5), 77.7 (C6), 127.3 (Ph or C3), 128.4 (2 carbons, Ph), 130.2 (Ph or C3), 130.3 (2 carbons, Ph), 133.6 (Ph), 145.1 (C2), 197.5 (C1); IR (KBr) 500, 540, 580, 630, 670, 700, 750, 790, 840, 860, 900, 960, 1030, 1090, 1130, 1150, 1200, 1240, 1250, 1300, 1380, 1450, 1490, 1600, 1690, 2850, 2920, 3030, 3060, 3480 cm⁻¹; HREIMS (*m*/*z*) calcd for $C_{13}H_{12}O_3$ (M⁺): 216.0786, found 216.0806. Anal. Calcd for $C_{13}H_{12}O_3$: C, 72.21; H, 5.59. Found: C, 71.81; H, 5.58.

4.1.24. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4,5-O-isopropylidenedioxy-4-[[(4R)-2,2-dimethyl-3-(ptoluenesulfonyl)oxazolidin-4-yl]hydroxymethyl]tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one (24). Lithium bis(trimethylsilyl)amide in THF (1.0 M solution, 5.30 ml, 5.3 mmol) was added dropwise to a stirred solution of 25 (800 mg, 2.4 mmol) in dry THF (40 ml) at -78 °C under argon. After 30 min, a solution of (R)-N-(p-toluenesulfonyl)-N,O-isopropylidene serinal (9) (1.72 g, 6.0 mmol) in dry THF (20 ml) was added slowly at -78 °C, and the resulting mixture was further stirred for 1 h at the same temperature. The reaction was quenched with saturated aqueous ammonium chloride (3 ml) at -78 °C, and the mixture was diluted with ether (300 ml). The organic layer was washed successively with saturated aqueous ammonium chloride (2×100 ml), saturated aqueous sodium hydrogen carbonate $(2 \times 100 \text{ ml})$, and brine $(2 \times 100 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, $2:1 \rightarrow 1:1$) to give 24 (1.27 g, 98%) (inseparable mixture, major/minor=9:1) as a colorless viscous oil. ¹H NMR (500 MHz, CDCl₃) δ 1.38 (3H, s, C-Me), 1.39 (3H, s, C-Me), 1.44 (3H, s, C-Me), 1.68 (3H, s, C-Me), 1.79 (1H, d, J=11.3 Hz, C11–H), 1.85 (1H, d, J = 11.3 Hz, C11–H), 2.25 (1H, d, J = 5.2 Hz, C1–H), 2.35 (1H, dd, J=1.8, 5.2 Hz, C2–H), 2.41 (3H, s, Me of Ts), 2.45 (1H, s, C8–H), 2.70 (1H, d, J=6.8 Hz, OH), 2.90 (1H, t, J=2.3 Hz, C7–H), 3.76 (1H, d, J=6.4, 9.8 Hz, C4^{\prime}–H), 4.20 (1H, d, J=6.5 Hz, C5'-H), 4.39 (1H, d, J=7.0 Hz, CH-OH), 4.51 (1H, dd, *J*=1.3, 9.8 Hz, C5'–H), 4.55 (1H, t, J = 2.3 Hz, C9–H), 4.59 (1H, t, J = 2.8 Hz, C7–H), 5.05 (1H, d, J = 3.1 Hz, C6–H), 7.30 (2H, d, J = 8.2 Hz, Ar), 7.68 (2H, d, J = 8.2 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 20.6, 21.5, 24.4, 26.1, 27.8, 28.8, 29.2, 30.5, 32.6, 41.2, 47.2, 59.6, 64.7, 73.0, 76.2, 78.9, 83.2, 88.1, 96.9, 109.6, 127.8 (2 carbons), 129.7 (2 carbons), 137.8, 143.5, 206.7; IR (neat) 550, 590, 680, 730, 820, 830, 880, 940, 1030, 1100, 1150, 1230, 1250, 1340, 1370, 1380, 1460, 1710, 2880, 2940, 2990, 3440 cm⁻¹; HREIMS (m/z) calcd for C₂₇H₃₃NO₈S (M⁺): 531.1927, found 531.1903.

4.1.25. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4,5-Oisopropylidenedioxy-4-[[(4R)-2,2-dimethyl-3-(p-toluenesulfonyl)oxazolidin-4-yl](methyldithiocarbonyloxy)methyl]tetracyclo[$6.2.1.0^{2,7}.0^{2,10}$]undecan-3-one (36). Sodium bis(trimethylsilyl)amide in THF (1.0 M solution, 0.85 ml, 0.85 mmol) was added dropwise to a stirred solution of 24 (378 mg, 0.71 mmol) in dry THF (20 ml) at -78 °C under argon. After 30 min, carbon disulfide (0.43 ml, 7.1 mmol) was added slowly to the mixture at -78 °C, and stirring was continued for 1 h at the same temperature. The resulting mixture was gradually warmed to $-50 \,^{\circ}\text{C}$ over 1 h, and then iodomethane (0.54 ml, 7.1 mmol) was added slowly to the above mixture at -78 °C. After 1 h, the mixture was gradually warmed to -50 °C over 1 h. The reaction was quenched with saturated aqueous ammonium chloride (3 ml) at 0 °C, and then the mixture was diluted with ether (200 ml). The organic layer was washed successively with saturated aqueous sodium thiosulfate $(2 \times 80 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 80 \text{ ml})$, and brine $(2 \times 80 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 2:1) to give 36 (389 mg, 88%) as a white solid. Recrystallization from hexane/ether (5:1) afforded colorless prisms, mp 252–253 °C; $[\alpha]_D^{20}$ –2.0 (c 1.12, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.19 (3H, s, C-Me), 1.41 (3H, s, C-Me), 1.47 (3H, s, C-Me), 1.51 (3H, s, C-Me), 1.79 (1H, d, J=11.4 Hz, C11-H), 1.84 (1H, d, J= 11.4 Hz, C11–H), 2.25 (1H, d, J=5.2 Hz, C1–H), 2.42 (4H, s, Me of Ts, C8-H), 2.65 (3H, s, S-Me), 2.72 (1H, dd, J= 1.9, 5.2 Hz, C10–H), 2.88 (1H, t, J=2.4 Hz, C7–H), 3.77 (1H, dd, J=7.0, 9.5 Hz, C4'-H), 4.46 (1H, dd, J=1.4, 9.6 Hz, C5'–H), 4.50 (1H, t, J=2.7 Hz, C6–H), 4.52 (1H, t, J=2.2 Hz, C9–H), 4.57 (1H, d, J=2.8 Hz, C5–H), 4.65 $(1H, d, J = 5.9 \text{ Hz}, C5' - H), 6.88 (1H, s, CH - OCS_2Me), 7.31$ (2H, d, J=8.2 Hz, Ar), 7.69 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 19.3, 20.3, 21.5, 24.1, 25.8, 27.7, 28.7, 29.3, 30.5, 32.5, 41.4, 47.5, 58.6, 64.7, 75.6, 79.5, 80.9, 82.9, 87.4, 96.8, 109.4, 128.2 (2 carbons), 129.6 (2 carbons), 137.5, 143.4, 205.6, 214.0; IR (neat) 520, 550, 590, 650, 680, 730, 820, 880, 910, 940, 1060, 1100, 1150, 1180, 1210, 1250, 1350, 1370, 1460, 1710, 2880, 2940, 2990 cm⁻¹; EIMS (m/z) 621 (M^+) , 606 $[(M-Me)^+]$. Anal. Calcd for C₂₉H₃₅NO₈S₃: C, 56.02; H, 5.67; N, 2.25; S, 15.47. Found: C, 55.85; H, 5.69; N, 2.29; S, 15.31.

4.1.26. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4,5-Oisopropylidenedioxy-4-[[(4R)-2,2-dimethyl-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl]tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one (37). Tri-n-butyltin hydride (0.33 ml, 1.2 mmol) and triethylborane in hexane (1.0 M solution, 0.63 ml, 0.63 mmol) were added successively to a stirred solution of 36 (384 mg, 0.62 mmol) in dry toluene (24 ml) at room temperature. After 1 h, the mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, $1:0 \rightarrow 2:1$) to give 37 (303 mg, 95%) as a white solid. Recrystallization from hexane/ether (5:1) afforded colorless prisms, mp 207–208 °C; $[\alpha]_{D}^{20}$ +49.1 (c 1.04, CHCl₃); ¹H NMR $(500 \text{ MHz}, \text{ CDCl}_3) \delta 1.34 (3H, s, C2'-Me), 1.37 (3H, s, s)$ C-Me), 1.40 (3H, s, C-Me), 1.67 (3H, s, C2'-Me), 1.79 (1H, d, J=11.4 Hz, C11-H), 1.84 (1H, d, J=11.4 Hz, C11-H), 2.20 (1H, dd, J = 10.9, 14.7 Hz, C4–CH_aH_b–C4'), 2.29 (1H, d, J=5.3 Hz, C1-H), 2.41 (3H, s, Me of Ts), 2.46 (1H, s, C8–H), 2.51 (1H, dd, J=1.8, 5.2 Hz, C10–H), 2.57 (1H, d, $J = 14.6 \text{ Hz}, \text{ C4-CH}_{a}H_{b}-\text{C4}'), 2.89 \text{ (1H, t, } J = 2.3 \text{ Hz},$ C7-H), 3.67 (1H, m, C5'-H), 4.12-4.18 (2H, m, C4'-H, C5'-H), 4.34 (1H, d, J=2.9 Hz, C5-H), 4.57 (2H, d, J=2.6 Hz, C6–H, C9–H), 7.28 (2H, d, J=8.1 Hz, Ar), 7.65 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 20.0, 21.5, 24.0, 26.8, 27.8, 30.3, 30.5, 31.7, 41.4, 44.5, 47.0, 55.7, 69.2, 76.30, 77.2, 83.5, 84.9, 84.9, 96.6, 109.7, 127.5 (2 carbons), 129.6 (2 carbons), 138.0, 143.2, 207.2; IR (neat) 520, 550, 600, 650, 680, 710, 840, 880, 920, 940, 1030, 1060, 1240, 1300, 1340, 1370, 1450, 1710, 2880, 2940, 2990 cm⁻¹; EIMS (m/z) 500 $[(M-Me)^+]$; CIMS (m/z) 516 $[(M+H)^+]$. Anal. Calcd for C₂₇H₃₃NO₇S: C, 62.89; H, 6.45; N, 2.72; S, 6.22. Found: C, 62.59; H, 6.49; N, 2.74; S, 6.25.

4.1.27. (1S,2S,3S,4R,4aR,5R,6S,7S,8aR)-3,5-Epoxy-2iodo-6,7-O-isopropylidenedioxy-7-[(2S)-2-p-toluenesulfonylamino-3-hydroxypropyl]-1,2,3,4,4a,5,6,8a-octahydro-endo-1,4-methanonaphthalen-8-one (38). Iodotrimethylsilane (82 µl, 0.46 mmol) was added dropwise to a stirred solution of 37 (98 mg, 0.15 mmol) in carbon tetrachloride (10 ml) at -20 °C under argon, and stirring was continued at -10 °C for 3 h. The reaction mixture was quenched with saturated aqueous sodium thiosulfate (2 ml) at 0 °C, and then the mixture was diluted with chloroform (80 ml). The organic layer was washed successively with 20% aqueous sodium thiosulfate $(2 \times 30 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2×30 ml), and brine $(2 \times 30 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 1:1) to give **38** (104 mg, 91%) as a white amorphous solid. $[\alpha]_{D}^{20} + 44.3$ $(c 1.09, CHCl_3)$; ¹H NMR (500 MHz, CDCl₃) δ 1.31 (6H, s, C-Me), 1.74 (1H, d, J = 11.2 Hz, C9–H), 1.86 (1H, dd, J =3.5, 15.5 Hz, C1'-H), 2.20-2.32 (3H, m, C9-H, C1'-H, OH), 2.42 (3H, s, Me of Ts), 2.61 (1H, dd, J = 4.8, 10.5 Hz, C8a-H), 2.75 (1H, br, C1-H), 2.87 (1H, br, C4-H), 2.91 (1H, dt, J=3.8, 10.5 Hz, C4a–H), 3.43 (1H, m, C2'–H), 3.67 (2H, t, J = 4.7 Hz, $C3'H_2OH$), 3.75 (1H, d, J = 1.9 Hz, C2–H), 3.80 (1H, t, J=2.6 Hz, C5–H), 4.17 (1H, d, J=2.4 Hz, C6–H), 4.60 (1H, d, J=11.2 Hz, C3–H), 5.84 (1H, d, J=4.6 Hz, N–H), 7.27 (2H, d, J=8.0 Hz, Ar), 7.70 (2H, d, J=8.2 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.5, 28.2, 28.5, 32.4, 35.8, 36.5, 42.0, 44.3, 46.6, 48.1, 51.4, 66.3, 77.0, 82.8, 85.5, 88.2, 112.8, 127.6 (2 carbons), 129.3 (2 carbons), 137.42, 143.0, 210.6; IR (neat) 550, 670, 730, 810, 850, 920, 940, 1050, 1090, 1130, 1160, 1220, 1230, 1330, 1380, 1420, 1600, 1710, 2890, 2930, 2980, 3280, 3510 cm^{-1} ; HREIMS *m*/*z* for C₂₃H₂₇INO₆S [(M-CH₂OH)⁺]: 572.0604, found 572.0604.

4.1.28. (1S,2S,3S,4R,4aR,5R,6S,7S,8aR)-3,5-Epoxy-2-iodo-6,7-O-isopropylidenedioxy-7-[(4S)-2,2-dimethyl-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-1,2,3,4,4a,5,6,8a-octahydro-endo-1,4-methanonaphthalen-8-one (**39**). *p*-Toluenesulfonic acid (6 mg, 34 µmol) was added to a stirred solution of 38 (100 mg, 0.17 mmol) in benzene (6 ml) containing 2,2-dimethoxypropane (0.20 ml, 1.7 mmol) at room temperature. The mixture was gradually warmed to 60 °C and stirred for 1 h at the same temperature. After cooling, the mixture was diluted with ether (80 ml). The organic layer was washed with saturated aqueous sodium hydrogen carbonate $(2 \times 30 \text{ ml})$ and brine $(2 \times$ 30 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo gave a residue, which was purified by column chromatography (hexane/ethyl acetate, 2:1) to give **39** (88 mg, 83%) as a colorless viscous oil. $[\alpha]_D^{20} + 138.4$ (*c* 0.97, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.37 (3H, s, C-Me), 1.43 (3H, s, C-Me), 1.44 (3H, s, C-Me), 1.70 (3H, s, C-Me), 1.83 (1H, d, J=11.2 Hz, C9-H), 2.13 (1H, dd, J= 10.7,14.8 Hz, C7–C H_2 –C4'), 2.27 (1H, d, J=11.2 Hz, C9–H), 2.43 (3H, s, Me of Ts), 2.48 (1H, d, J=4.6 Hz, $C7-CH_2-C4'$), 2.89–2.98 (3H, m, C1–H, C4–H, C4a–H), 3.02 (1H, dd, J=4.9, 10.2 Hz, C8a–H), 3.67 (1H, dd, J=5.5, 8.1 Hz, C5'–H), 3.93 (1H, d, J=2.1 Hz, C2–H), 4.16– 4.22 (2H, m, C5-H, C5'-H), 4.40-4.46 (2H, m, C4'-H, C6-H), 4.80 (1H, d, J=5.1 Hz, C3-H), 7.32 (2H, d, J= 8.0 Hz, Ar), 7.83 (2H, d, J=8.3 Hz, Ar); ¹³C NMR

(125 MHz, CDCl₃) δ 21.5, 24.2, 27.3, 28.2, 30.5, 33.0, 36.8, 40.2, 41.0, 46.6, 47.7, 48.3, 55.6, 68.7, 75.9, 81.4, 81.9, 89.2, 96.9, 111.3, 127.8 (2 carbons), 129.5 (2 carbons), 138.0, 143.2, 209.9; IR (neat) 510, 550, 590, 650, 680, 710, 730, 820, 840, 920, 940, 1100, 1160, 1230, 1340, 1370, 1460, 1600, 1710, 1890, 2930, 2990 cm⁻¹; HREIMS (*m*/*z*) calcd for C₂₆H₃₁INO₇S [(M-Me)⁺]: 628.0866, found 628.0853.

4.1.29. (1R,4S,4aR,5R,6S,7S,8aS)-5-Hydroxy-6,7-O-isopropylidenedioxy-7-[(4S)-2,2-dimethyl-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-1,4,4a,5,6,7,8,8a-octahydro-endo-1,4-methanonaphthalen-8-one (40). Zinc powder (123 mg, 1.9 mmol) and acetic acid (0.11 ml, 1.9 mmol) were successively added to a stirred solution of **39** (81 mg, 0.13 mmol) in methanol (6 ml) at room temperature. The mixture was gradually warmed to 60 °C and stirred for 1 h at the same temperature. After cooling, the mixture was diluted with ether (50 ml) and filtrated. The filtrate was washed with saturated aqueous sodium hydrogen carbonate $(2 \times 20 \text{ ml})$, and brine $(2 \times 20 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 2:1) to give **40** (64 mg, 98%) as a colorless viscous oil. $[\alpha]_D^{20} + 127.1$ (*c* 1.14, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.39 (1H, m, C9-H), 1.45 (3H, s, C-Me), 1.47 (3H, s, C-Me), 1.53 (4H, s, C-Me, C9–H), 1.67 (3H, s, C-Me), 1.81 (1H, d, J=2.3 Hz, OH), 2.17 (1H, dd, J = 10.5, 14.0 Hz, C7–C H_aH_b –C4'), 2.25 $(1H, dd, J = 1.8, 14.4 \text{ Hz}, C7-CH_aH_b-C4'), 2.41 (3H, s, Me$ of Ts), 2.98 (1H, s, C4-H), 3.13 (1H, s, C1-H), 3.21 (1H, dt, J=3.4, 11.7 Hz, C4a–H), 3.43 (1H, dd, J=3.6, 11.7 Hz, C8a–H), 3.60 (1H, dd, J = 5.5, 9.0 Hz, C5'–H), 4.12 (1H, dd, J = 1.8, 9.0 Hz, C5'-H), 4.37 (1H, br, C5-H), 4.43 (1H, d, J = 4.3 Hz, C6–H), 4.45 (1H, m, C4'–H), 6.19 (1H, dd, J =3.0, 5.5 Hz, C3–H), 6.50 (1H, dd, J=3.1, 5.6 Hz, C2–H), 7.27 (2H, d, J=8.2 Hz, Ar), 7.85 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.5, 24.4, 26.5, 27.3, 30.1, 41.8, 43.1, 43.6, 45.5, 46.4, 51.2, 56.4, 68.1, 68.7, 83.0, 85.2, 96.9, 112.2, 127.7 (2 carbons), 129.4 (2 carbons), 132.7, 138.5, 140.0, 142.9, 207.6; IR (neat) 550, 600, 650, 680, 710, 730, 780, 830, 920, 1050, 1100, 1160, 1210, 1240, 1340, 1380, 1450, 1600, 1720, 1880, 2940, 2990, 3530 cm⁻¹; HREIMS (m/z) calcd for C₂₆H₃₂NO₇S $[(M - Me)^+]$: 502.1900, found 502.1869.

4.1.30. (4*R*,5*S*,6*S*)-4-Hydroxy-5,6-*O*-isopropylidenedioxy-7-[(4S)-2,2-dimethyl-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-2-cyclohexen-1-one (41). A stirred solution of 40 (23.0 mg, 44 µmol) in diphenyl ether (5 ml) was heated at 230 °C for 2 h. After cooling, the mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, $1:0 \rightarrow 1:1$) to give 41 (5.0 mg, 25%) as a colorless viscous oil. $[\alpha]_D^{20}$ +58.1 (c 0.46, CHCl₃); ¹H NMR (500 MHz, CDCl₃) & 1.27 (3H, s, C-Me), 1.39 (6H, s, C-Me, C-Me), 1.68 (3H, s, C-Me), 2.13 (1H, dd, J=10.8, 14.5 Hz, C6–C $H_{a}H_{b}$ –C4'), 2.42 (3H, s, Me of Ts), 2.43 (1H, d, J= 14.5 Hz, C6–CH_a H_{b} –C4'), 2.52 (1H, br, OH), 3.74 (1H, ddd, J=1.3, 5.4, 9.2 Hz, C5'-H), 4.07 (1H, dd, J=5.3, 10.7 Hz, C4^{\prime}-H), 4.09 (1H, d, J=9.0 Hz, C5^{\prime}-H), 4.16 (1H, t, J = 1.7 Hz, C5–H), 4.66 (1H, br, C4–H), 6.17 (1H, d, J =10.2 Hz, C2–H), 6.84 (1H, ddd, J=2.0, 4.6, 10.1 Hz, C3–H), 7.29 (2H, d, J=8.0 Hz, Ar), 7.69 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.5, 24.1, 26.7, 27.2, 30.3, 39.5, 55.3, 64.7, 69.2, 80.7, 83.8, 97.0, 109.5, 127.6 (2 carbons), 128.2, 129.6 (2 carbons), 137.7, 143.4, 143.6, 198.8; IR (neat) 550, 590, 650, 680, 710, 750, 830, 880, 910, 1040, 1100, 1160, 1230, 1340, 1370, 1460, 1490, 1600, 1680, 2880, 2940, 2990, 3470 cm⁻¹; HREIMS (*m*/*z*) calcd for C₂₁H₂₆NO₇S [(M-Me)⁺]: 436.1430, found 436.1403.

4.1.31. (1R,2S,4S,5S,6R,7R,8R,9S,10S)-6,9-Epoxy-4,5-Oisopropylidenedioxy-4-[[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl]tetracyclo[6.2.1.0^{2,7}.0^{2,10}]undecan-3-one (43). 1.0 M Hydrochloric acid (1.36 ml, 1.4 mmol) was added to a stirred solution of **37** (320 mg, 0.62 mmol) in THF (15 ml) at room temperature, and the mixture was heated at 55 °C for 6 h. After cooling, the mixture was neutralized with saturated aqueous sodium hydrogen carbonate (ca. 20 ml), and the resulting mixture was extracted with ether $(3 \times 50 \text{ ml})$. The combined extracts were washed with brine $(2 \times 50 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 2:1) to give an equilibrium mixture (290 mg) of **42a** and **42b** $(1:1 \text{ by } 500 \text{ MHz}^{-1}\text{H NMR})$ as a colorless viscous oil. This equilibrium mixture was directly used for the following reaction without separation.

Trichloromethyl chloroformate (1.23 ml, 6.2 mmol) was added dropwise to a stirred solution of the above equilibrium mixture of 42a and 42b (290 mg, 0.61 mmol) in dry THF (30 ml) containing pyridine (1.96 ml, 24 mmol) at 0 °C, and the mixture was gradually warmed to room temperature. After 2 h, the reaction was quenched with saturated aqueous sodium hydrogen carbonate (5 ml) at 0 °C, and the mixture was diluted with ether (200 ml). The organic layer was washed successively with 3% aqueous hydrochloric acid $(2 \times 80 \text{ ml})$, saturated aqueous sodium hydrogen carbonate (2×80 ml), and brine (2×80 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 3:2) to give 43 (209 mg, 67% in two steps) as a white solid. Recrystallization from hexane/ether (5:1) afforded colorless needles, mp 174–175 °C; $[\alpha]_{\rm D}^{20}$ +27.5 (c 1.16, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.36 (3H, s, C-Me), 1.39 (3H, s, C-Me), 1.82 (1H, d, J=11.6 Hz, C11-H), 1.87 (1H, d, J= 11.6 Hz, C11–H), 2.02–2.09 (1H, m, C4– $CH_{a}H_{b}$ –C4'), 2.25 (1H, dd, J=1.8, 5.2 Hz, C10-H), 2.34 (1H, d, J=5.1 Hz)C1-H), 2.45 (3H, s, Me of Ts), 2.50 (1H, s, C8-H), 2.90 (1H, t, J=2.2 Hz, C7-H), 2.92 (1H, d, J=14.6 Hz, C4–CH_a H_{b} –C4'), 4.33 (1H, d, J=3.0 Hz, C5–H), 4.37– 4.44 (2H, m, C4'=H, C5'=H), 4.57 (1H, d, J=5.4 Hz, C5' = H), 4.60 (1H, t, J = 2.8 Hz, C6–H), 4.62 (1H, t, J =2.2 Hz, C9–H), 7.33 (2H, d, J=8.4 Hz, Ar), 7.86 (2H, d, J= 8.4 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 20.6, 21.7, 26.8, 27.8, 30.3, 30.5, 32.1, 41.3, 43.3, 46.6, 54.6, 68.6, 76.2, 83.3, 83.9, 84.1, 110.3, 128.5 (2 carbons), 129.8 (2 carbons), 134.7, 145.7, 152.3, 205.8; IR (KBr) 540, 580, 620, 670, 750, 810, 840, 880, 920, 940, 990, 1030, 1070, 1090, 1140, 1170, 1220, 1250, 1300, 1370, 1600, 1710, 1790, 2880, 2940, 2990 cm⁻¹; EIMS (*m/z*) 501 (M⁺), 486 $[(M-Me)^+]$; HREIMS (*m/z*) calcd for C₂₅H₂₇NO₈S (M⁺): 501.1457, found 501.1481.

4.1.32. (1S,2S,3S,4R,4aR,5R,6S,7S,8aR)-3,5-Epoxy-2-iodo-6,7-O-isopropylidenedioxy-7-[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-1,2,3,4,4a,5,6,8a-octahydroendo-1.4-methanonaphthalen-8-one (44). Iodotrimethylsilane (68 µl, 0.48 mmol) was added dropwise to a stirred solution of 43 (210 mg, 0.42 mmol) in carbon tetrachloride (20 ml) at -20 °C under argon. After 1 h, the reaction was quenched with saturated aqueous sodium thiosulfate (2 ml) at -20 °C, and then the mixture was diluted with ether (150 ml). The organic layer was washed successively with 20% aqueous sodium thiosulfate $(2 \times 80 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 80 \text{ ml})$, and brine $(2 \times 80 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 3:2) to give 44 (195 mg, 74%) as a white solid. Recrystallization from hexane/ether (5:1) afforded colorless needles, mp 233-234 °C; $[\alpha]_{D}^{20}$ + 152.8 (c 1.04, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.35 (3H, s, C-Me), 1.41 (3H, s, C-Me), 1.85 (1H, d, J = 11.3 Hz, C9–H_a), 2.05 (1H, dd, J = 10.7, 14.6 Hz, $C7-CH_{a}H_{b}-C4'$), 2.29 (1H, d, J=11.1 Hz, C9-H_b), 2.45 (3H, s, Me of Ts), 2.89-2.95 (3H, m, C1-H, C4a-H, C7-CH_aH_b-C4'), 2.97-3.04 (2H, m, C4-H, C8a-H), 3.80 (1H, d, J=2.4 Hz, C2–H), 4.29 (1H, t, J=3.7 Hz, C5–H), 4.42 (1H, t, J=9.0 Hz, C5'-H), 4.45 (1H, d, J=4.1 Hz, C6–H), 4.56 (1H, dd, J=4.4, 9.3 Hz, C5^{\prime}–H), 4.80 (1H, m, C4'-H), 4.85 (1H, d, J=5.3 Hz, C3-H), 7.37 (2H, d, J=8.3 Hz, Ar), 7.96 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) & 21.7, 27.1, 28.0, 32.2, 36.7, 39.9, 40.7, 46.9, 47.9, 48.3, 54.5, 68.8, 75.6, 80.8, 81.0, 89.3, 111.5, 128.5 (2 carbons), 129.8 (2 carbons), 134.8, 145.5, 152.5, 210.3; IR (KBr) 540, 570, 610, 670, 760, 820, 920, 1050, 1090, 1130, 1170, 1310, 1370, 1450, 1600, 1700, 1790, 2990 cm⁻¹; EIMS (m/z) 629 (M^+) , 614 $[(M - 1)^2]$ Me)⁺]; HRCIMS (m/z) calcd for C₂₅H₂₉INO₈S [$(M+H)^+$]: 630.0659, found 630.0693. Anal. Calcd for C₂₅H₂₈INO₈S: C, 47.70; H, 4.48; N, 2.23; S, 5.09. Found: C, 47.78; H, 4.47; N, 2.30; S, 4.82.

4.1.33. (1R,4S,4aS,5R,6S,7S,8aS)-5-Hydroxy-6,7-O-isopropylidenedioxy-7-[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-1,4,4a,5,6,7,8,8a-octahydro-endo-**1.4-methanonaphthalen-8-one** (45). Zinc powder (300 mg, 4.6 mmol) and acetic acid (0.26 ml, 4.6 mmol) were successively added to a stirred solution of 44 (194 mg, 0.31 mmol) in THF/methanol (1:1) (20 ml) at room temperature. The mixture was gradually warmed to 60 °C and stirred for 1 h at the same temperature. After cooling, the mixture was diluted with ether (150 ml) and filtrated. The filtrate was washed with saturated aqueous sodium hydrogen carbonate (2×70 ml), and brine (2×70 ml), then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 3:2) to give 45 (148 mg, 95%) as a colorless viscous oil. $[\alpha]_D^{20}$ +155.9 (c 1.00, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.41 (1H, d, J=8.4 Hz, C9–H), 1.50 (3H, s, C-Me), 1.53 (3H, s, C-Me), 1.56 (1H, d, J=8.4 Hz, C9–H), 1.94 (1H, s, OH), 2.30 (1H, dd, J = 10.5, 14.2 Hz, C7–C H_aH_b –C4'), 2.44 (3H, s, Me of Ts), 2.72 (1H, dd, J=2.1, 14.1 Hz, C7–CH_aH_b–C4'), 3.01 (1H, s, C4-H), 3.14 (1H, s, C1-H), 3.25 (1H, dt, J=3.2)11.6 Hz, C4a–H), 3.44 (1H, dd, J=3.6, 11.6 Hz, C8a–H), 4.17 (1H, t, J=8.5 Hz, C5'-H), 4.36 (1H, dd, J=5.1,

9.3 Hz, C5'–H), 4.44 (1H, s, C5–H), 4.59 (1H, d, J=4.3 Hz, C6–H), 4.92 (1H, m, C4'–H), 6.20 (1H, dd, J=3.0, 5.3 Hz, C3–H), 6.49 (1H, dd, J=3.1, 5.5 Hz, C2–H), 7.34 (2H, d, J=8.3 Hz, Ar), 7.95 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.7, 26.3, 27.2, 41.1, 43.1, 43.7, 45.5, 46.1, 51.2, 54.8, 67.9, 69.2, 82.9, 84.0, 112.5, 128.5 (2 carbons), 129.8 (2 carbons), 132.9, 134.9, 139.6, 145.5, 152.8, 207.7; IR (neat) 540, 580, 610, 670, 700, 760, 820, 850, 920, 1050, 1090, 1120, 1170, 1210, 1250, 1310, 1380, 1450, 1600, 1720, 1780, 2940, 2990, 3540 cm⁻¹; HREIMS (m/z) calcd for C₂₅H₂₉NO₈S (M⁺): 503.1614, found 503.1595.

4.1.34. (4*R*,5*S*,6*S*)-4-Hydroxy-5,6-*O*-isopropylidenedioxy-6-[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4yl]methyl-2-cyclohexen-1-one (46). A stirred solution of 45 (148 mg, 0.29 mmol) in diphenyl ether (15 ml) was heated at 230 °C for 2 h. After cooling, the mixture was concentrated in vacuo to afford a residue, which was purified by column chromatography (hexane/ethyl acetate, $1:0 \rightarrow 1:1$) to give 46 (75.9 mg, 59%) as a colorless viscous oil. $[\alpha]_D^{20}$ + 80.9 (c 1.05, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.26 (3H, s, C-Me), 1.40 (3H, s, C-Me), 2.09 (1H, dd, J=11.0, 14.5 Hz, C6–CH₂H_b–C4'), 2.20 (1H, d, J=5.3 Hz, OH), 2.44 (3H, s, Me of Ts), 2.92 (1H, dd, J=2.2, 14.4 Hz, C6–CH_a H_{b} –C4[']), 4.19 (1H, t, J=1.7 Hz, C5–H), 4.45 (2H, d, J = 6.2 Hz, C5'-H₂), 4.59 (1H, m, C4'-H), 4.72 (1H, t, *J*=4.7 Hz, C4–H), 6.19 (1H, d, *J*=10.2 Hz, C2–H), 6.90 (1H, ddd, J=1.9, 4.8, 10.1 Hz, C3–H), 7.34 (2H, d, J= 8.2 Hz, Ar), 7.87 (2H, d, J=8.2 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.7, 26.6, 27.2, 39.0, 54.3, 64.1, 69.2, 80.0, 83.2, 109.9, 128.0, 128.4 (2 carbons), 129.9 (2 carbons), 134.5, 144.7, 145.8, 152.5, 198.0; IR (neat) 540, 570, 600, 670, 760, 820, 910, 1040, 1090, 1130, 1170, 1230, 1380, 1490, 1600, 1680, 1790, 2930, 3480 cm⁻¹; CIMS (*m*/ z) 438 $[(M+H)^+]$; HREIMS (*m/z*) calcd for C₁₉H₂₀NO₈S $[(M-Me)^+]$: 422.0910, found 422.0926.

4.1.35. (1R,5S,6S)-5,6-O-Isopropylidenedioxy-4-oxo-5-[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-2-cyclohexenyl methanesulfonate (5). Methanesulfonyl chloride (73 µl, 0.93 mmol) was added to a stirred solution of 46 (68.3 mg, 0.16 mmol) in dichloromethane (7 ml) containing triethylamine (0.17 ml, 1.2 mmol) and 4-dimethylaminopyridine (38.0 mg, 0.31 mmol) at 0 °C, and stirring was continued for 4 h at room temperature. The reaction was quenched with saturated aqueous sodium hydrogen carbonate (1 ml) at 0 °C, and the mixture was diluted with ether (70 ml). The organic layer was successively washed with 3% aqueous hydrochloric acid $(2 \times 30 \text{ ml})$, saturated aqueous sodium hydrogen carbonate $(2 \times 30 \text{ ml})$, and brine $(2 \times 30 \text{ ml})$, then dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ ethyl acetate, 1:1) to give 5 (66.8 mg, 83%) as a colorless viscous oil. $[\alpha]_{D}^{20}$ +47.3 (c 1.07, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 1.27 (3H, s, C-Me), 1.42 (3H, s, C-Me), 2.10 (1H, dd, J = 10.8, 14.3 Hz, C6–CH_aH_b–C4'), 2.45 (3H, s, Me of Ts), 2.91 (1H, dd, J=2.2, 14.3 Hz, C6–CH_a H_{b} –C4[']), 3.19 (3H, s, Me of Ms), 4.34 (1H, t, J =1.8 Hz, C5–H), 4.43 (1H, dd, J=4.8, 9.4 Hz, C5'–H), 4.46 (1H, t, J=9.3 Hz, C5'-H), 4.58 (1H, m, C4'-H), 5.58 (1H, m)dd, J=1.7 Hz, C4-H), 6.34 (1H, d, J=10.1 Hz, C2-H),

6.89 (1H, ddd, J=1.9, 5.0, 10.2 Hz, C3–H), 7.36 (2H, d, J=8.2 Hz, Ar), 7.88 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.7, 26.8, 27.1, 38.5, 39.1, 54.0, 69.0, 69.7, 79.8, 80.9, 111.0, 128.4 (2 carbons), 129.9 (2 carbons), 130.9, 134.4, 138.9, 145.8, 152.2, 196.5; IR (neat) 540, 570, 600, 620, 670, 730, 760, 820, 860, 950, 990, 1060, 1090, 1130, 1170, 1230, 1370, 1600, 1690, 1790, 2930, 2990 cm⁻¹; CIMS (*m*/*z*) 516 [(M+H)⁺]; HREIMS (*m*/*z*) calcd for C₂₀H₂₂NO₁₀S₂ [(M−Me)⁺]: 500.0685, found 500.0696.

4.1.36. (1R,5S,6S)-5,6-Dihydroxy-4-oxo-5-[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4-yl]methyl-2-cyclohexenyl methanesulfonate (47). A solution of 5 (59.2 mg, 0.11 mmol) in trifluoroacetic acid/water (6:1) (3 ml) was stirred at 0 °C for 30 min. The mixture was concentrated in vacuo to give 47 (54.6 mg, quant.) as a colorless viscous oil. $[\alpha]_{D}^{20}$ + 25.9 (c 1.22, CHCl₃); ¹H NMR (500 MHz, CDCl₃) δ 2.25 (1H, dd, J = 10.0, 14.7 Hz, C6–C H_aH_b –C4'), 2.30–2.40 (1H, br, OH), 2.45 (3H, s, Me of Ts), 2.50-2.90 (1H, br, OH), 2.98 (1H, d, J = 14.6 Hz, C6–CH_aH_b–C4'), 3.18 (3H, s, Me of Ms), 4.20-4.27 (2H, m, C5-H, C4'-H), 4.30 (1H, dd, J=4.4, 9.3 Hz, C5'–H), 4.45 (1H, t, J=8.8 Hz, C5'–H), 5.47 (1H, t, J=3.5 Hz, C4–H), 6.38 (1H, d, J=10.2 Hz, C2-H), 6.89 (1H, ddd, J=1.3, 4.1, 10.2 Hz, C3-H), 7.37 (2H, d, J=8.2 Hz, Ar), 7.87 (2H, d, J=8.3 Hz, Ar); ¹³C NMR (125 MHz, CDCl₃) δ 21.7, 38.8, 40.2, 53.6, 70.4, 74.8, 75.0, 77.0, 128.5 (2 carbons), 129.3, 130.0 (2 carbons), 134.2, 140.6, 146.1, 152.4, 198.0; IR (neat) 540, 570, 670, 730, 760, 820, 850, 940, 1090, 1170, 1360, 1600, 1700, 1780, 2360, 2930, 3480 cm⁻¹; HRCIMS (m/z) calcd for $C_{18}H_{22}NO_{10}S_2$ [(M+H)⁺]: 476.0685, found 476.0658.

4.1.37. (4S,5S,6S)-4,5-Epoxy-6-hydroxy-6-[(4S)-2-oxo-3-(p-toluenesulfonyl)oxazolidin-4-yl]methy-2-cyclohexen-1-one (3b). 0.2 M Sodium hydroxide (1.5 ml, 0.30 mmol) was added dropwise to a stirred solution of 47 (54.5 mg, 0.11 mmol) in ether (5 ml) at 0 °C. After 20 min, the mixture was extracted with ether $(3 \times 30 \text{ ml})$. The combined extracts were washed with brine $(3 \times 30 \text{ ml})$ and dried over Na₂SO₄. Concentration of the solvent in vacuo afforded a residue, which was purified by column chromatography (hexane/ethyl acetate, 1:1) to give 3b (32.3 mg, 75%) as a white solid. Recrystallization from hexane/dichloromethane (3:1) afforded colorless prisms, mp 224–225 °C; $[\alpha]_D^{20}$ +153.3 (*c* 0.99, CHCl₃); ¹H NMR $(500 \text{ MHz}, \text{ CD}_2\text{Cl}_2) \delta 2.34 (1\text{H}, \text{dd}, J=10.5, 14.2 \text{ Hz},$ $C6-CH_{a}H_{b}-C4'$, 2.45 (3H, s, Me of Ts), 2.51 (1H, dd, $J=1.1, 14.2 \text{ Hz}, \text{ C6-CH}_{a}H_{b}-\text{C4}'), 3.49 (1\text{H}, \text{br}, \text{OH}),$ 3.67 (2H, m, C4-H, C5-H), 4.16 (1H, m, C4'-H), 4.34 (1H, dd, J=4.6, 9.4 Hz, C5'-H), 4.44 (1H, t, J=9.1 Hz, C5'-H), 6.35 (1H, m, C2-H), 7.27 (1H, m, C3-H), 7.38 (2H, m, Ar), 7.80 (2H, m, Ar); ¹³C NMR (125 MHz, CD_2Cl_2) δ 21.9 (Me of Ts), 40.7 (C6– CH_2 –C4'), 48.6 (C4'), 53.7 (C4), 56.5 (C5), 70.5 (C5'), 77.3 (C6), 128.7 (2 carbons, Ar), 130.3 (3 carbons, Ar, C2), 134.7 (C3), 145.9 (Ar), 146.6 (Ar), 152.4 (C2'), 197.9 (C1); IR (KBr) 540, 570, 600, 670, 760, 840, 990, 1090, 1170, 1370, 1690, 1780, 2360, 2930, 3460 cm⁻¹; HRCIMS (m/z)calcd for $C_{17}H_{17}NO_7S$ [(M+H)⁺]: 380.0804, found 380.0786.

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Tetrahedron

Tetrahedron 62 (2006) 1609

Erratum

Erratum to "A general strategy for the synthesis of azapeptidomimetic lactams" [Tetrahedron 61 (2005) 10277]

Robert L. Broadrup, Bei Wang and William P. Malachowski*

Department of Chemistry, Bryn Mawr College, Bryn Mawr, PA 19010-2899, USA

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The publisher apologises for the following error:

'room temperature' that followed the gas chromatography (GC) and high-pressure liquid chromatography (HPLC) information in the experimental section should have read 'retention time'. This error occurred on p 10282, sections 4.1.12, 4.1.13, 4.1.16 and p 10283, sections 4.1.17, 4.1.18, 4.1.19, 4.1.20, and 4.1.21.

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* Corresponding author. Tel.: +1 610 526 5016; fax: +1 610 526 5086; e-mail: wmalacho@brynmawr.edu



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Tetrahedron

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Corrigendum

Corrigendum to "5-Hydroxy-3-phenyl-5-vinyl-2-isoxazoline and 3-phenyl-5-vinylisoxazole: synthesis and reactivity" [Tetrahedron 61 (2005) 11270]

Leonardo Di Nunno,* Antonio Scilimati and Paola Vitale

Dipartimento Farmaco-Chimico, Università degli Studi di Bari, Via E. Orabona, 4, 70125 Bari, Italy

Available online 20 December 2005

At the end of the Abstract '...3-hydroxy-5-phenyl-5-vinyl-2-isoxazoline' should read '5-hydroxy-3-phenyl-5-vinyl-2-isoxazoline'.

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* Corresponding author. Tel.: + 39 080 5442734; fax: + 39 080 5442231; e-mail: dinunno@farmchim.uniba.it