Monatshefte für Chemie Chemical Monthly Printed in Austria

Synthesis of Substituted 1-Thiocyanatobutadienes and their Application in a *Diels-Alder*/[3,3] Sigmatropic Rearrangement Tandem Reaction

Sébastien Lanaspèze and Reinhard Neier*

Institute of Chemistry, University of Neuchâtel, CH-2007 Neuchâtel, Switzerland

Received August 9, 2004; accepted August 30, 2004 Published online January 24, 2005 © Springer-Verlag 2005

Summary. The retrosynthetic analysis of *Ibogamine*, a natural psychotropic alkaloid with exceptional anti-addictive properties found in both enantiomeric forms, requires an efficient access to a racemic cyclohexene. This cyclohexene can be obtained *via* the sequence *Diels-Alder*/[3,3] sigmatropic rearrangement reaction starting from substituted 1-thiocyanatobutadienes. An efficient synthesis of the enone, a stable precursor of 1-thiocyanatobutadienes, is reported. Enolisation of this enone was studied to find the optimal conditions to get the desired 1-thiocyanatobutadienes with good *Z*-selectivity.

Keywords. Enols; Ibogamine; Rearrangements; Tandem reactions; Total synthesis.

Introduction

The number of synthetic steps in a total synthesis strongly influences the efficiency. An attractive way to reduce the length of a synthetic pathway is to combine several transformations into one-pot reactions so called tandem, domino, or cascade reaction [1–3]. The tandem process *Diels-Alder* reaction/[3,3] sigmatropic rearrangement combines two well-known pericyclic reactions in one single synthetic step, which allows to obtain interesting synthetic building blocks efficiently [4]. Starting from (*E*)-1-thiocyanatobuta-1,3-diene (1) and acryloyl chloride led to a *cis/trans* 85:15 mixture of diastereoisomers of 1,4-substituted cyclohexene 3 in 84% yield [5] (Scheme 1). The cyclohexene 3 is a precursor for the preparation of the skeleton of *Iboga*-type alkaloids. The isothiocyanate 3 can be easily transformed in three steps to the 2-azabicyclo[2.2.2]oct-5-ene 4 [5] (Scheme 1). This

^{*} Corresponding author. E-mail: reinhard.neier@unine.ch



Scheme 1

bicyclic compound has been used by *Trost et al.* for the synthesis of *desethylibo-gamine* (5) [6, 7].

Ibogamine (**6a**), as other *Iboga*-type alkaloids, has been isolated from an African plant *Tabernanthe iboga* [8], which has well-known psychotropic activities. Studies on rodents have proven that **6a** shows exceptional multi-drug anti-addictive properties [8–10]. Unfortunately, despite the encouraging results described by *Lotsof* [11] about treatment of opioid withdrawal with *ibogaine* (**6b**) on human patients, systematic clinical studies have not been carried out. Since the first total synthesis of **6a** by *Büchi* [12] in 1965, eleven other total synthesis have been reported [7, 13–21] but none of them is cost- and material-efficient enough to allow the production of this compound or of derivatives of this compound in large scale industrial production.



5 (+/-)-Desethylibogamine

6a *R* = H: (+/-)-*Ibogamine* **6b** *R* = OMe: (+/-)-*Ibogaine*

In most of the reported total syntheses of *ibogamine* (**6a**), the crucial step is the formation of the C–C bond between the carbons C(2) corresponding to the α -position of the indole ring and C(16) one of the α -positions of the bridgehead carbon connected to the N-atom. In the total synthesis reported by *Trost et al.* this crucial C–C bond was formed using a palladium-catalysed *Heck*-type C–C coupling, in the presence of stoechiometric amounts of a silver salt [7]. The goal of our studies is to develop an alternative procedure for the synthesis of the strategic C–C bond avoiding use of stoechiometric quantities of a silver salt. Introducing a triflate protected enolate on carbon C(16) should allow to test different catalytic C–C

Synthesis of Substituted 1-Thiocyanatobutadienes



coupling conditions. *Ibogamine* (6a), contrary to the model compound *desethylibogamine* (5), possesses an additional ethyl group in a pseudo-equatorial position on carbon C(20). The retrosynthetic analysis of 6a based on the formation of the strategic C–C bond described above leads to two major building blocks of similar size: the indole derivative 7 (M=B(OH)₂, ZnCl) and the isoquinuclidine ring rac-8. We propose to obtain the isoquinuclidine ring rac-8 via our tandem process *Diels-Alder* reaction/[3,3] sigmatropic rearrangement starting from the 2,4-substituted 1-thiocyanatobuta-1,3-diene 10. In this paper the synthesis of diene 10 via a *Z*-enolisation of the enone 11 is reported and the stereoselectivity of this enolisation is studied.

Results and Discussions

The initial step of the synthesis of enone **11** is the preparation of the phosphorane **13**. According to the procedure described by *Hudson et al.* [22], we started our synthesis from the highly toxic, commercially available 1,3-dichloroacetone **12**. According to the literature procedure the intermediate salt formed from the S_N^2 reaction between the dichloroketone and triphenylphosphine was isolated in 85% yield. The salt was then deprotonated in a separate step to get the ylide **13**. To improve the yield and to reduce the time of exposure to toxic α -chloroketones, we developed a one-pot



Reagents and conditions: (i) PPh₃, THF, reflux, 4h; (ii) Na₂CO₃, MeOH, rt, 30 min, 97%; (iii) KSCN, KI_{cat}, EtOH, 55°C, 5h, 96%; (iv) propionaldehyde, 40°C, 6h, 70%



process. Using this one-pot procedure the ylide **13** could be isolated in almost quantitative yield. Displacement of the first chlorine atom of 1,3-dichloroacetone **12** by triphenylphosphine was monitored by the disappearance of the ³¹P signal of triphenylphosphine ($\delta_P = -4.4 \text{ ppm}$) in its ³¹P NMR (Scheme 3). The second S_N2 reaction replacing the chloride by nucleophilic attack of the sulphur atom of the thiocyanate anion was carried out in ethanol. In order to accelerate the reaction, the presence of a catalytic quantity of iodide was needed, as has been reported by *Sapi et al.* [23]. Thus, the thiocyanate **14** was obtained in very good yield and satisfactory purity so that the raw material could be used in the following step without further purification.

The *Wittig* reaction was assumed to be *E*-selective due to the stabilisation of ylide **14** by the adjacent carbonyl group. As assumed the *E*-enone **11** was obtained as a consequence of the thermodynamic equilibrium between the more stable *trans*-oxaphosphetane **15** and its *cis*-diastereoisomer during the *Wittig* reaction (Scheme 4).

Propionaldehyde was both the solvent and the reactant of our *Wittig* reaction. The progress of the reaction was monitored by the growth of the ³¹P signal of triphenylphosphine oxide ($\delta_P = 30.1 \text{ ppm}$) in ³¹P NMR. The α,β -unsaturated ketone **11** is probably undergoing unwanted polymerisations, which make the purification difficult. Thus, after filtration on a short silica gel column, the enone **11** was purified by a bulb-to-bulb distillation (100°C, 10⁻¹ mbar). By this straight forward three-steps process enone **11** could be obtained in 65% total yield starting from 1,3-dichloroacetone **12**.

In order to optimize the reaction conditions for the enolisation model **16b** was chosen. Synthesis of thiocyanate **16b** was carried out using a similar procedure as for the synthesis of **14** starting from commercially available phenacyl chloride **16a** (Scheme 5).



Reagents and conditions: (i) KSCN, KI_{cat}, EtOH, rt, 4h, 86%; (ii) MsCl, Et₃N, CH₂Cl₂, rt, 45 min, 40%

600

Synthesis of Substituted 1-Thiocyanatobutadienes



The enolisation of α -halogenoketones usually leads to the kinetic (Z)-diastereoisomer [24]. The α -thiocyanatoketones can be viewed as pseudohalogen analogues of α -halogenoketones. The stereoselectivity of the enolisation can be explained using τ -bonds, also called "banana" bonds or bent bonds [25–28]. Using a *Newman* projection to describe the most stable ground state conformation of the α -halogenoketone, the strongest electron withdrawing substituent in the α -position of the carbonyl group (the halogen or the pseudohalogen) has to be in a *syn*-coplanar position relative to the C–O τ -bond (Scheme 6). In this conformation all bonds are staggered, the two hydrogen atoms, which are the best donor atoms, are in a *anti*-periplanar position compared to the two strongly electron attracting C–O τ -bonds (Scheme 6).

Having the precursors for our enolates in hand, we studied the stereoselectivity of the enolisation process of our enone 11 under thermodynamic conditions, using triethylamine or *Hünig*'s base. With our model compound 16b, an excellent stereoselectivity of the enolisation was observed under our experimental conditions. Compound 17 was obtained as a (Z)/(E)-mixture with a diastereoselectivity of

	SCN Electrophile, weak base					
		11	10a	-10e (<i>ZE</i>)/(<i>EE</i>)		
10	R	Procedure	$(ZE)/(EE)^{d}$	Yield		
a	TMS	A^{a}	87:13	78%		
b	TBDMS	A^{a}	62:38	_e		
c	Ms	B^{b}	100:0	71%		
d	Ts	B^{b}	85:15	_f		
e	Tf	C^{c}	65:35 ^g	72% ^h		

Table 1. Study of stereoselectivity of the enolisation of enone 11 under thermodynamic conditions

^a Procedure A: 2.2 eq triethylamine + 2 eq electrophile + 1 eq enone **11** in ether as solvent at rt for 1 h; ^b procedure B: 2–10 eq triethylamine + 1.2–5 eq electrophile + 1 eq enone **11** in CH₂Cl₂ as solvent at 0°C for 30 min; ^c procedure C: 1.4 eq diisopropylethylamine + 1.25 eq electrophile + 1 eq enone **11** in CH₂Cl₂ as solvent at 0°C for 30 min; ^d stereoselectivity estimated by ¹H NMR analysing the product of the *Diels-Alder* reaction; only the (*ZE*)-diastereoisomer can react in a *Diels-Alder* reaction; ^e under the conditions of procedure A, the diene **10b** could not be isolated, due to hydrolysis; the stereoselectivity was tentatively assigned based on the analysis of the raw material; ^f diene **10c** was isolated but not purified; ^g both diastereoisomers (*ZE*) and (*EE*) could be separated by chromatography on a silica gel column and the diasteriomeric ratio was estimated by ¹⁹F NMR; ^h the synthesis of diene **10e** was also carried out under kinetic conditions resulting in a better stereoselectivity



99:1 (Scheme 5). As described in Table 1, four different electrophiles were used to trap the enol, applying three different experimental procedures (Table 1).

As predicted for the formation of the 1(Z),3(E)-diastereoisomers was favoured compared to the formation of the 1(E),3(E)-diastereoisomers. Using the procedure *B* and mesyl chloride as an electrophile the best diastereoselectivity was observed (*de* > 99%). A NOESY experiment showed a crosspeak between H-C(1) and H-C(4) (Scheme 7) proving the structure of compound **10c**. The observed NOESY crosspeak is only compatible with the *s*-*cis* conformation of the 1(Z), 3(E)-diastereoisomer.

Applying the conditions D for the enolisation of enone **11** under kinetic control the dienes **10b** and **10e** were obtained. The lithium enolate **17** is formed under low-temperature conditions. The enolate is then quenched by various electrophiles to get the desired substituted 1-thiocyanatobuta-1,3-diene **10b** and **10e**. As indicated in Table 2 very good yields were obtained under these conditions. A satisfactory diastereoselectivity was observed with *TBDMSCl* as an electrophile (de = 60% in favour of 1(Z),3(E)-diastereoisomer of the silylated 1-thiocyanato-hexa-1,3-diene **10b**). Using triffic anhydride as an electrophile, the influence of addition of *DMPU* as a co-solvent on diastereoselectivity was studied. A good diastereoselectivity (de = 78%) was observed without the use of the co-solvent. Addition of *DMPU* (20% in *THF*) improved the diastereoisomeric excess to de = 95%.

	\sim	$ \begin{array}{c} $		$\frac{\text{SCN}}{10} \xrightarrow{\text{Electrophile}} 10$	R = TBDMS b : $R = Tf$
10	R	Procedure	THF/DMPU	$(ZE)/(EE)^{\rm b}$	Yield
b	TBDMS	D^{a}	10:0	80:20	90%
e e	1j Tf	$D D^{a}$	8:2	97.5:2.5°	50% 85%

Table 2. Stereoselectivity of the enolisation of enone 11 under kinetically controlled condition

^a Procedure D: 1.1 eq LiHMDS in THF/DMPU as solvent at -80° C for 45 min + 1.1 eq electrophile + 1 eq enone **11** in THF as solvent at 0°C for 1 h; ^b stereoselectivity estimated by ¹H NMR analysing the product of the *Diels-Alder* reaction; only the (ZE)-diastereoisomer can react in a *Diels-Alder* reaction; ^c the diastereomeric ratio was estimated by ¹⁹F NMR; ^d non optimised

Conclusions

An efficient three-step synthesis of enone **11** has been developed giving **11** in 65% total yield. The stereoselectivity of the enolisation of **11** was studied under thermodynamically controlled conditions using different electrophiles to trap the enolate. Excellent diastereoisomeric excesses were observed and the best diastereoselectivity was found using mesyl chloride as an electrophile (de > 99%) yielding the 1(Z),3(E)-1-thiocyanatohexa-1,3-dien-2-yl methanesulfonate (**10c**). Under kinetically controlled conditions the 1-thiocyanatobuta-1,3-dienes **10b** and **10e** could be obtained. The addition of *DMPU* as a co-solvent was studied in the case of the 1-thiocyanatobuta-1,3-diene **10e**. The diastereoselectivity (de = 78%) could be improved using *DMPU* (de = 95%). Efficient synthesis of five different substituted 1-thiocyanatobuta-1,3-dienes **10a**-**10e** are reported in good total yields. Their efficiency in our *Diels-Alder*/[3,3] signatropic rearrangement tandem reaction will be studied in view of the development of a new total synthesis of (*rac*)-*Ibogamine* (**6a**).

Experimental Part

All moisture-sensitive reactions were carried out under Ar and N₂ using oven-dried glassware. All reagents were of commercial quality if not specifically mentioned. Solvents were freshly distilled prior to use. Flash chromatography (FC): Brunschwig silica gel 60, 0.032–0.063 mm; under positive pressure. TLC: Merck precoated silica gel thin-layer sheets 60 F 254, detection by UV and treatment with basic KMnO₄ sol. Mp: Gallenkamp MFB-595. IR spectra: Perkin Elmer Spectrum One FT-IR, in cm⁻¹. NMR spectra: Bruker Avance-400 (400 MHz (¹H), 162 MHz (³¹P), and 100 MHz (¹³C)) and Varian Gemini-2000 (188 MHz (¹⁹F)), at rt, chemical shifts δ in ppm rel. to Si*Me*₄ (=0 ppm) as internal reference, coupling constants *J* in Hz. ESI-MS Finnigan LCQ. Elemental analyses of novel compounds agreed favourably with calculated values.

1-Chloro-3-(triphenyl-5 λ -phosphanylidene)propan-2-one (13)

Triphenylphosphine (52.90 g, 200.9 mmol) in 75 cm³ of dry *THF* was added to a solution of 25.12 g of 1,3-dichloroacetone (197.8 mmol) in 25 cm³ of dry *THF*. The mixture was heated to reflux for 4 h and the solvent was then removed. The resulting white salt was dissolved in 100 cm³ of methanol, and an aqueous solution (1.55 *M*) of 10.48 g of Na₂CO₃ (98.9 mmol) was added. The mixture was allowed to stand for 30 min and the white precipitate was dried in air to provide a white powder of pure **13** (67.58 g, 191.6 mmol, 97%). Ylide **13** has been described by *Hudson et al.* [22]. Mp 179–180°C (Ref. [22] 178–179°C); $R_f = 0.25$ (CH₂Cl₂/*Me*OH 97/3); ³¹P NMR (162 MHz, CDCl₃): $\delta = 16.9$ ppm; ¹H NMR (400 MHz, CDCl₃): $\delta = 7.68-7.62$ (m, 6H, H-5²), 7.60–7.54 (m, 3H, H-5⁴)), 7.50–7.45 (m, 6H, H-5³), 4.28 (d, ²*J*(3-P)=23.9 Hz, 1H, H-3), 4.02 (s, 2H, H-1) ppm; ¹³C NMR (100 MHz, CDCl₃): $\delta = 185.1$ (s, 1C, C-2), 133.1 (s, 6C, C-5³), 133.0 (s, 3C, C-5⁴), 128.9 (d, ³*J*(5²-P)=12.5 Hz, 6C, C-5²), 126.2 (d, ⁻¹*J*(5¹-P)=91.3 Hz, 3C, C-5¹), 51.4 (d, ⁻¹*J*(3-P)=114.5 Hz, 1C, C-3), 47.3 (d, ³*J*(1-P)= 16.1 Hz, 1C, C-1) ppm.

1-Thiocyanato-3-(triphenyl-5\lambda-phosphanylidene)propan-2-one (14, C₂₂H₁₈NOPS)

Ylide **13** (64.42 g, 182.6 mmol) in 300 cm³ of ethanol was added to a solution of 21.30 g of potassium thiocyanate (219.1 mmol) in 350 cm³ of hot ethanol and 3.03 g of KJ (18.3 mmol) were added. The mixture was allowed to stir at 55°C for 5 h and cooled to rt. The potassium salts were filtered off, most

of the solvent was evaporated in the rotavap, and 400 cm³ of CH₂Cl₂ were added. The potassium salts were filtered off again and washed with cold CH₂Cl₂. The resulting solution was concentrated to dryness to provide pure **14** (65.75 g, 175.1 mmol, 96%), mp 146–148°C. $R_f = 0.39$ (CH₂Cl₂/*Me*OH 97/3); ³¹P NMR (162 MHz, CDCl₃): $\delta = 16.1$ ppm; ¹H NMR (400 MHz, CDCl₃): $\delta = 7.70-7.52$ (m, 9H, H-5², H-5⁴), 7.51–7.42 (m, 6H, H-5³), 3.96 (d, ²*J*(3-P)=24.0 Hz, 1H, H-3), 3.92 (s, 2H, H-1) ppm; ¹³C NMR (100 MHz, CDCl₃): $\delta = 181.5$ (d, ²*J*(2-P)=3.1 Hz, 1C, C=O), 133.1 (d, ⁴*J*(5³-P)=9.9 Hz, 6C, C-5³), 132.6 (d, ³*J*(5⁴-P)=3.0 Hz, 3C, C-5⁴), 129.2 (d, ³*J*(5²-P)=12.7 Hz, 6C, C-5²), 125.7 (d, ¹*J*(5¹-P)=91.2 Hz, 3C, C-5¹), 114.1 (SCN), 52.9 (d, ¹*J*(3-P)=108.5 Hz, 1C, C-3), 43.1 (d, ³*J*(1-P)=19.1 Hz, 1C, C-1) ppm; IR (KBr): $\bar{\nu} = 2994$, 2153, 1577, 1548, 1482, 1438, 1228, 1108, 764, 747, 719, 692, 518, 507 cm⁻¹.

(E)-1-Thiocyanatohex-3-en-2-one (11, C₇H₉NOS)

A solution of 44.35 g of ylide **14** (118.1 mmol) in 86 cm³ of freshly distilled propanal is allowed to stir at 40°C for 6 h under Ar and cooled to rt. The mixture was concentrated and quickly filtered on a short silica gel column (200 g; *n*-hexane/CH₂Cl₂ 1/1). The resulting brown oil was purified by bulb-to-bulb distillation (100°C at 10⁻¹ mbar) to provide a malodorous pale yellow oil (12.80 g, 82.5 mmol, 70%). $R_f = 0.42$ (CH₂Cl₂); ¹H NMR (400 MHz, CDCl₃): $\delta = 7.03$ (dt, ³*J*(4-3)=16.0 Hz, ³*J*(4-5)=6.3 Hz, 1H, H-4), 6.19 (dt, ³*J*(3-4)=16.0 Hz, ⁴*J*(3-5)=1.7 Hz, 1H, H-3), 4.24 (s, 2H, H-1), 2.31 (qdd, ³*J*(5-6)=7.4 Hz, ³*J*(5-4)=6.3 Hz, ⁴*J*(5-3)=1.7 Hz, 2H, H-5), 1.10 (t, ³*J*(6-5)=7.4 Hz, 3H, H-6) ppm; ¹³C-NMR (100 MHz, CDCl₃): $\delta = 190.5$ (C=O), 153.6 (C-4), 126.5 (C-3), 111.9 (SCN), 42.7 (C-1), 25.9 (C-5), 11.9 (C-6) ppm; IR (film): $\bar{\nu} = 2972$, 2935, 2878, 2158, 1687, 1626, 1460, 1392, 1341, 1297, 1245, 1180, 1109, 1081, 1015, 977, 909 cm⁻¹.

1-Phenyl-2-thiocyanatoethanone (16b)

According to the procedure described for **14**, 5.96 g of phenacyl chloride (38.5 mmol) were treated with 4.70 g of KSCN (48.4 mmol) and 0.66 g of KJ (4.0 mmol) giving 5.90 g of **16b** (33.3 mmol, 86%) as brown crystals. Thiocyanate **16b** has been already isolated by *Prakash et al.* [29] as an oil. Recrystallisation in *n*-hexane/*AcOEt* 2/1 gave needle shaped yellow crystals. Mp 81–82°C; R_f =0.38 (*n*-hexane/*AcOEt* 3/1); ¹H NMR (400 MHz, CDCl₃): δ = 7.96 (ddm, ³*J*(3²-3³)=8.4 Hz, ⁴*J*(3²-3⁴)= 1.3 Hz, 2H, H-3²), 7.69 (tt, ³*J*(3⁴-3³)=7.4 Hz, ⁴*J*(3⁴-3²)=1.3 Hz, 1H, H-3⁴), 7.55 (dd, ³*J*(3³,3²)=8.4 Hz, ³*J*(3³,3⁴)=7.4 Hz, 2H, H-3³), 4.76 (s, 2H, H-2) ppm; ¹³C NMR (100 MHz, CDCl₃): δ = 190.9 (C=O), 134.9 (C-3⁴), 134.1 (C-3¹), 129.3 (C-3³), 128.6 (C-3²), 112.0 (SCN), 43.1 (C-2) ppm; IR (KBr): $\bar{\nu}$ = 2937, 2156, 1678, 1593, 1448, 1379, 1326, 1296, 1202, 998, 757, 687, 630 cm⁻¹.

(Z)-1-Phenyl-2-thiocyanatovinyl methanesulfonate (17, C₁₀H₉NO₃S₂)

Triethylamine (141 mg, 1.4 mmol) was added to a solution of 177 mg of **16b** (1.0 mmol) in 5 cm³ of dry CH₂Cl₂ under Ar. After cooling to 0°C, 143 mg of methanesulfonyl chloride (1.25 mmol) were slowly introduced at 0°C (10 min). After warming to rt, the resulting mixture was stirred for additional 45 min, diluted in 25 cm³ of CH₂Cl₂, and washed with 25 cm³ of H₂O whose *pH* was adjusted to 3–5 with aqueous HCl (1*M*). The organic layer was then washed with brine, dried (MgSO₄), and the solvent was evaporated to dryness to provide a green powder of a mixture of the two stereoisomers (*Z*)/(*E*) (99/1) **17** (102 mg, 0.4 mmol, 40%). $R_f = 0.21$ (*n*-hexane/*AcOEt* 3/1); ¹H NMR (400 MHz, CDCl₃): $\delta = 7.56-7.50$ (m, 2H, H-3³), 7.50–7.42 (m, 3H, H-3², H-3⁴), 6.50 (s, 1H, H-2(*E*)), 6.44 (s, 1H, H-2(*Z*)), 3.15 (s, 3H, H-3(*Z*)), 3.01 (s, 3H, H-3(*E*)) ppm; ¹³C NMR (100 MHz, CDCl₃): $\delta = 151.0$ (C-1), 132.2 (C-3¹), 131.2 (C-3⁴), 129.3 (C-3²), 126.2 (C-3³), 109.5 (SCN), 105.9 (C-2), 39.9 (C-3) ppm; IR (KBr): $\bar{\nu} = 3068$, 3036, 2934, 2164, 1681, 1622, 1354, 1264, 1172, 1008, 961, 864, 801, 779, 718, 524, 505 cm⁻¹.

General Procedures for the Synthesis of the Substituted 1-Thiocyanatobutadienes 10a–10e

Procedure A: To a stirred 0.6M solution of 2.2 eq of triethylamine in diethylether at rt under Ar were added dropwise 2.0 eq of the corresponding electrophile. 1.0 eq of **11** was then added dropwise to the solution for 5 min. The resulting mixture was magnetically stirred at rt for 1 h and the solvent was evaporated. Freshly distilled *n*-pentane was added to the crude product, the salts were filtered off and washed with freshly distilled *n*-pentane again. After evaporating the *n*-pentane *in vacuo*, the product was isolated without further purification.

Procedure B: To a stirred 0.8M solution of 1.0 eq of **11** in CH₂Cl₂ at 0°C under Ar were added 2.0–10.0 eq of triethylamine. A 3.0*M* solution of 1.2–5.0 eq of the corresponding electrophile in CH₂Cl₂ was then added dropwise to the solution for 30 min. The resulting mixture was magnetically stirred at 0°C for 30 min and then diluted in CH₂Cl₂. The organic phase was then washed with H₂O, dried (MgSO₄), and concentrated *in vacuo* to obtain the raw product that was then purified by chromatography on a silica gel column.

Procedure C: To a stirred 0.2 *M* solution of 1.0 eq of **11** in CH₂Cl₂ at rt under Ar were added 1.4 eq of diisopropylethylamine. After cooling to 0°C, 1.25 eq of the corresponding electrophile were added dropwise to the solution during 5 min. The resulting mixture was magnetically stirred at 0°C for 30 min and then diluted in CH₂Cl₂. The organic phase was washed with saturated aqueous NH₄Cl, H₂O, dried (MgSO₄), and concentrated *in vacuo* to get the product that was purified by chromatography on a silica gel column.

Procedure D: A solution of 1.1 eq of Li *bis*(trimethyldisilyl)amide (1*M* in *THF*) was diluted 3 times in a *THF/DMPU* mixture to get a 0.33*M* solution, which was cooled to -90° C. A 1.0*M* solution of 1.0 eq of **11** in freshly distilled *THF* was then added dropwise while maintaining the temperature of the solution below -80° C. The orange solution was then magnetically stirred at -85° C for 45 min and 1.1 eq of the corresponding electrophile were added dropwise maintaining the temperature of the solution below -80° C. The temperature was then allowed to warm slowly to 0° C and the mixture was stirred for additional 1 h at 0° C. After diluting three times the reactions mixture with *n*-hexane, the solution was poured onto saturated aqueous NH₄Cl. The organic phase was washed with H₂O and brine, dried (MgSO₄), and concentrated *in vacuo* to get the product that was purified by chromatography on a silica gel column.

Trimethyl((*3E*)-1-thiocyanatohexa-1,3-dien-2-yloxy)silane (**10a**, C₁₀H₁₇NOSSi)

Procedure A: Freshly distilled enone **11** (0.40 g, 2.58 mmol) was reacted with 0.56 g of trimethylchlorosilane (5.15 mmol) to obtain **10a** as a yellow oil (0.46 g, 2.02 mmol, 78%) and (*ZE*)/(*EE*) mixture (87/13). Compound **10a** was extremely sensitive towards hydrolysis. ¹H NMR (200 MHz, CDCl₃): $\delta = 6.65$ (dt, ³*J*(3-4)=15.4 Hz, ⁴*J*(3-5)=1.1 Hz, 1H, H-3(*EE*)), 6.37 (dt, ³*J*(4-3)=15.3 Hz, ³*J*(4-5)=6.2 Hz, 1H, H-4(*EE*), 6.08 (dt, ³*J*(4-3)=15.3 Hz, ³*J*(4-5)=6.2 Hz, 1H, H-4(*ZE*), 5.89 (dt, ³*J*(3-4)=15.4 Hz, ⁴*J*(3-5)=1.1 Hz, 1H, H-3(*ZE*)), 5.25 (s, 1H, H-1(*ZE*)), 5.12 (s, 1H, H-1(*EE*)), 2.16 (qdd, ³*J*(5-6)=7.4 Hz, ³*J*(5-4)=6.3 Hz, ⁴*J*(5-3)=1.1 Hz, 2H, H-5), 1.03 (t, ³*J*(6-5)=7.4 Hz, 3H, H-6), 0.29 (s, 9H, H-2¹) ppm.

tert-Butyldimethyl((3E)-1-thiocyanatohexa-1,3-dien-2-yloxy)silane (10b, C13H23NOSSi)

Procedure *D*: Enone **11** (0.47 g, 3.00 mmol) was reacted with a 1.0*M* solution of 0.50 g of *tert*butyldimethylchlorosilane (3.31 mmol) in freshly distilled *THF*. After purification by quick filtration on silica gel with *n*-hexane/*AcOEt* 95/5, compound **10b** was obtained as yellow oil (0.73 g, 2.71 mmol, 90%) and (*ZE*)/(*EE*) mixture (80/20) that could not be separated. R_f (*ZE*) = 0.79, R_f (*EE*) = 0.89 (*n*-hexane/*AcOEt* 10/1); ¹H NMR (400 MHz, CDCl₃): δ = 6.63 (dt, ³*J*(3-4)=15.2 Hz, ⁴*J*(3-5)=1.6 Hz, 1H, H-3(*EE*)), 6.42 (dt, ³*J*(4-3)=15.2 Hz, ³*J*(4-5)=6.5 Hz, 1H, H-4(*EE*), 6.07 (dt, ${}^{3}J(4-3)=15.5$ Hz, ${}^{3}J(4-5)=6.5$ Hz, 1H, H-4(ZE)), 5.85 (dt, ${}^{3}J(3-4)=15.5$ Hz, ${}^{4}J(3-5)=1.6$ Hz, 1H, H-3(ZE)), 5.23 (s, 1H, H-1(ZE)), 5.10 (s, 1H, H-1(EE)), 2.23 (qdd, ${}^{3}J(5-6)=7.4$ Hz, ${}^{3}J(5-4)=6.5$ Hz, ${}^{4}J(5-3)=1.6$ Hz, 2H, H-5(ZE)), 2.13 (qdd, ${}^{3}J(5-6)=7.4$ Hz, ${}^{3}J(5-4)=6.5$ Hz, ${}^{4}J(5-3)=1.6$ Hz, 2H, H-5(ZE)), 1.07 (t, ${}^{3}J(6-5)=7.4$ Hz, 3H, H-6(ZE)), 1.02 (t, ${}^{3}J(6-5)=7.4$ Hz, 3H, H-6(ZE)), 1.00 (s, 9H, H-2 ${}^{3}(ZE)$), 0.96 (s, 9H, H-2 ${}^{3}(ZE)$), 0.20 (s, 6H, H-2 ${}^{1}(EE)$), 0.19 (s, 6H, H-2 ${}^{1}(ZE)$) ppm; ${}^{13}C$ NMR (100 MHz, CDCl₃): $\delta = 161.7$ (C-2(ZE)), 157.7 (C-2(ZE)), 141.4 (C-4(EE)), 137.8 (C-4(ZE)), 124.5 (C-3(ZE)), 120.7 (C-3(ZE)), 112.0 (SCN(EE)), 111.5 (SCN(ZE)), 90.0 (C-1(ZE)), 85.8 (C-1(EE)), 25.8 (C-2 ${}^{3}(ZE)$), 25.7 (C-5(ZE)), 25.6 (C-2 ${}^{3}(EE)$), 25.4 (C-5(EE)), 18.5 (C-2 ${}^{2}(EE)$), 18.3 (C-2 ${}^{2}(ZE)$), 13.0 (C-6(ZE)), 13.0 (C-6(ZE)), -3.4 (C-2 ${}^{1}(ZE)$), -4.4 (C-2 ${}^{1}(ZE)$) ppm.

According to procedure A using a 1.0 M solution of *tert*-butyldimethylchlorosilane in diethylether, a (ZE)/(EE) mixture (63/37) of impure **10b** was obtained.

(1Z,3E)-1-Thiocyanatohexa-1,3-dien-2-yl methanesulfonate (10c, C₈H₁₁NO₃S₂)

Procedure *B*: Enone **11** (0.54 g, 3.48 mmol) in 35 cm³ of CH₂Cl₂ was reacted with a 0.5*M* solution of 0.50 g of methanesulfonylchloride (4.35 mmol) in CH₂Cl₂ and 1.34 g of triethylamine (13.23 mmol). Isolation according to *A* to get **10c** as yellow oil (0.58 g, 2.49 mmol, 71%). $R_f = 0.20$ (CH₂Cl₂); ¹H NMR (400 MHz, CDCl₃): $\delta = 6.36$ (dtd, ³*J*(4-3)=15.8 Hz, ³*J*(4-5)=6.6 Hz, ⁵*J*(4-1)=0.5 Hz, 1H, H-4), 6.18 (dd, ⁵*J*(1-4)=0.5 Hz, ³*J*(1-3)=0.5 Hz, 1H, H-1), 5.97 (dtd, ³*J*(3-4)=15.8 Hz, ⁴*J*(3-5)=1.3 Hz, ⁴*J*(3-1)=0.5 Hz, 1H, H-3), 3.05 (s, 3H, H-2¹), 2.19 (qdd, ³*J*(5-6)=7.4 Hz, ³*J*(5-4)=6.3 Hz, ⁴*J*(5-3)=1.3 Hz, 2H, H-5), 1.03 (t, ³*J*(6-5)=7.4 Hz, 3H, H-6) ppm; ¹³C-NMR (100 MHz, CDCl₃): $\delta = 175.9$ (C-2), 148.2 (SCN), 139.0 (C-4), 114.4 (C-3), 41.7 (C-2¹), 25.7 (C-5), 12.8 (C-6) ppm; ¹H NOESY: cross peak between $\delta = 6.36$ and 6.18 ppm.

(3E)-1-Thiocyanatohexa-1,3-dien-2-yl 4-methylbenzenesulfonate (10d, C₁₄H₁₅NO₃S₂)

Procedure *B*: Enone **11** (0.31 g, 2.00 mmol) was reacted with a 0.4*M* solution of 0.76 g of *p*-toluenesulfonyl chloride (4.00 mmol) in CH₂Cl₂ and 2.02 g of triethylamine (20.00 mmol). A (*ZE*)/(*EE*) mixture (65/35) of impure **10d** (0.50 g) was obtained by this procedure. $R_f = 0.14$ (14% *AcOEt* in *n*-hexane). ESI-MS: m/z = 332.1 [M + Na]⁺.

(3E)-1-Thiocyanatohexa-1,3-dien-2-yl trifluoromethanesulfonate (10e, C₈H₈F₃NO₃S₂)

Procedure *D*: Enone **11** (130 mg, 0.84 mmol) was reacted with a 1.0 cm³ of a 1 *M* LiHMDS solution diluted with 2 cm³ of a 7/3 mixture of *THF/DMPU*. Triflic anhydride (0.16 cm³, 0.94 mmol) was added. Compound **10e** was obtained as yellow oil (205 mg, 0.71 mmol, 85%) and (*ZE*)/(*EE*) mixture (97.5/2.5). Both diastereoisomers could be separated by chromatography on a silica gel column using *n*-hexane/*AcOEt* 95/5 as an eluant. ESI-MS: $m/z = 310.0 [M + Na]^+$.

I(Z),3(E)-Diastereoisomer: $R_f = 0.34$ (n-hexane/AcOEt 95/5); ¹⁹F NMR (188 MHz, CDCl₃): $\delta = -73.16$ ppm; ¹H NMR (400 MHz, CDCl₃): $\delta = 6.32$ (dt, ³J(4-3)=15.6 Hz, ³J(4-5)=6.5 Hz, 1H, H-4), 6.09 (s, 1H, H-1), 6.02 (dt, ³J(3-4)=15.6 Hz, ⁴J(3-5)=1.6 Hz, 1H, H-3), 2.24 (qdd, ³J(5-6)=7.4 Hz, ³J(5-4)=6.5 Hz, ⁴J(5-3)=1.6 Hz, 2H, H-5), 1.07 (t, ³J(6-5)=7.4 Hz, 3H, H-6) ppm; ¹³C NMR (100 MHz, CDCl₃): $\delta = 150.3$ (C-2), 141.7 (C-4), 120.1 (C-3), 116.9 (C(SCN)), 108.4 (C-2¹), 104.9 (C-1), 25.8 (C-5), 12.6 (C-6) ppm.

I(E), 3(E)-Diastereoisomer: $R_f = 0.49$ (n-hexane/AcOEt 95/5); ¹⁹F NMR (188 MHz, CDCl₃): $\delta = -73.65$ ppm; ¹H NMR (400 MHz, CDCl₃): $\delta = 6.49$ (dt, ³J(4-3)=15.4 Hz, ³J(4-5)=6.5 Hz, 1H, H-4), 6.35 (dt, ³J(3-4)=15.4 Hz, ⁴J(3-5)=1.5 Hz, 1H, H-3), 6.10 (s, 1H, H-1), 2.32 (m, ³J(5-6)=7.4 Hz, ³J(5-4)=6.5 Hz, ⁴J(5-3)=1.5 Hz, 2H, H-5), 1.11 (t, ³J(6-5)=7.4 Hz, 3H, H-6) ppm; ¹³C NMR (100 MHz, CDCl₃): $\delta = 151.3$ (C-2), 144.8 (C-4), 126.0 (SCN), 116.9 (C-3), 108.3 (C-2¹), 103.0 (C-1), 26.3 (C-5), 12.6 (C-6) ppm.

Synthesis of Substituted 1-Thiocyanatobutadienes

The same procedure without the use of DMPU led to a (ZE)/(EE) mixture (89/11) of **10e** in 50% yield which was not optimised.

Procedure C: Enone 11 (0.40 g, 2.57 mmol) was reacted with triflic anhydride. Compound **10e** was then purified by chromatography on a silica gel column by using CH_2Cl_2 as an eluant to get a (ZE)/(EE) mixture (65/35) (0.53 g, 1.85 mmol) in 72% yield which was not optimised.

Acknowledgements

NMR Spectra (400 MHz) were measured by *H. Bursian* and Dr. *C. Saturnin*, mass spectra by *N. Mottier* and *J. Jean-Denis*, CHN analyses were made in the *Ecole d'ingénieurs et d'architectes de Fribourg*. We thank the University of Neuchâtel and the Swiss National Science Foundation for financial support.

References

- [1] Tietze LF (1996) Chem Rev 96: 115
- [2] Ho TL (1993) Tandem Organic Reactions. Wiley, New York
- [3] Nicolaou KC, Montagnon T, Snyder SA (2003) Chem Comm 551
- [4] Neuschütz K, Velker J, Neier R (1998) Synthesis 227
- [5] Schöpfer J, Marquis C, Pasquier C, Neier R (1994) J Chem Soc Chem Comm 1001
- [6] Trost BM, Genêt JP (1976) J Am Chem Soc 98: 8516
- [7] Trost BM, Godleski SA, Genêt JP (1978) J Am Chem Soc 100: 3930
- [8] Popik P, Skolnick P (1998) Pharmacology of Ibogaïne and Ibogaïne-Related Alkaloïds. In: Cordell GA (ed) The Alkaloïds, vol 52. Academic Press, San Diego, p 197
- [9] Cappendijk SLT, Dzoljic MR (1993) Eur J Pharmacol 241: 261
- [10] Glick SD, Kuehne ME, Raucci J, Wilson TE, Larson D, Keller RW, Carlson JN (1994) Brain Res657: 14
- [11] Alper KR, Lotsof HS, Frenken GMN, Luciano DJ, Bastiaans J (1999) Am J Addictions 8: 234
- [12] Büchi G, Coffen DL, Kocsis K, Sonnet PE, Ziegler FE (1965) J Am Chem Soc 87: 2073
- [13] Nagata W, Hirai S, Kawata K, Okumura T (1968) J Am Chem Soc 90: 1650
- [14] Ikezaki M, Wakamatsu T, Ban Y (1968) Chem Comm 88
- [15] Rosenmund P, Haase WH, Bauer J, Frische R (1975) Chem Ber 108: 1871
- [16] Rahman A-U, Beisler JA, Harley-Mason J (1980) Tetrahedron 36: 1063
- [17] Huffmann JW, Shanmugasundaram G, Sawdaye R, Raveendranath PC, Desai RC (1985) J Org Chem 50: 1460
- [18] Kuehne ME, Reider PJ (1985) J Org Chem 50: 1464
- [19] Imanishi T, Yagi N, Hanaoka M (1985) Chem Pharm Bull 33: 4202
- [20] Henry KJ Jr, Grieco PA, DuBay WJ (1996) Tetrahedron Lett 37: 8289
- [21] White JD, Choi Y (2000) Org Lett 2: 2373
- [22] Hudson RF, Chopard PA (1963) J Org Chem 28: 2446
- [23] Sapi A, Fetter J, Lempert K, Kajtar-Peredy M, Czira G (1997) Tetrahedron 53: 12729
- [24] Tanigushi M, Takeyama Y, Fugami K, Oshima K, Utimoto K (1991) Bull Chem Soc Japan 64: 2593
- [25] Robinson EA, Gillespie JR (1980) J Chem Educ 57: 329
- [26] Rassat A (2004) Phys Chem Chem Phys 6: 232
- [27] Wintner CE (1987) J Chem Educ 64: 587
- [28] Vogel E, Caravatti G, Franck P, Aristoff P, Moody C, Becker A-M, Felix D, Eschenmoser A (1987) Chem Lett 219
- [29] Prakash O, Harpreet K, Batra H, Rani N, Singh SP, Moriarty RM (2001) J Org Chem 66: 2019