SYNTHETIC STUDIES ON ERIOLANIN: 1,2-CARBONYL TRANSPOSITIONS OF CIS-BICYCLO[4.2.0] OCTANONE VIA ENOL THIOETHER FORMATION

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Abstract - 1,2-Carbonyl transposition of cis-fused cyclobutanone is demonstrated by means of two procedures(method A and method Bl. This relatively short sequencelmethod Bl provides **e** new and convenient method for conversion of carbonyi group in the entitled rlng system.

Dichiorocyciobutanones, readily available by dichioroketene cycioadditions to olefins¹ are extremely useful intermediates in organic synthesis.² In a continuation of our work directed toward the synthesis of an antitumor sesquiterpene, eriolanin 1^3 , we required a concise and general method for $1, 2$ carbonyl transposition $\begin{bmatrix} 2 & -3 \end{bmatrix}$ in cis-bicyclo[4.2.0] octanone system. While previous works **an various** aspects of csrbonyi transposltlon **have** been reported, methods to transpose the carbonyl group of ketones **are** iacking4in the entitled ring fusion.

We now report the two different methods A and **B on** 1,2-carbonyl transposition, which provided enoi thloethers **8 or** 14 respectively **as** intermediates for cyclobutanones.

Treatment of 4 with lithium bis(trimethyIsilyI)amide in tetrahydrofuran(THF) at -78° C to 0° C for 1.5 h followed by addition of diphenyldisulfide(2.5 equiv)³ in hexamethyiphosphoric triamide(HMPA) at 0° C for 14 h gave bis(phenylthio)cyclobutanones **5a** and 5b in a 4:l ratio in 90% yield.

It is difficult to control the regioselectivity for monosulfenylation of 4 **at C-2 position via depratonation with an amide base. Enollration usually** occurs at C-8 to yield the corresponding cyclobutanone enolate.^{2a, 6} Reduction of 5a⁷ with sodium borohydride afforded smoothly a 1:6.6 mixture of 6a' ['H NMR(CLXJ₃)^o 3.34(dd, J_{bc}=7.8 and J_{ab}=10.0 Hz, IH_b),3.93(d, J_{bc}=7.6Hz, $H_{\rm H}$),4.02(d, $H_{\rm H}$),4.02(d, **Jbc=4.8Hz,IHcl] in quantitative yield, of which the stereochemistry was I determined from J values of H NvR, the observation of NOE and chemical transformations as shown in Scheme 1. A similar reduction of 4:1 mixtures 5e and 5b afforded 6a, 6b, and 6c in a rsitio** of **5:40:1 in 91% yield, of which compound 6c arose from 5b. Desulfurizatton of 6a and 6b with Raney NI(W-2) in ethanol led to their cyciobutanois 7a and 7b respectively, while 7a was available from the reduction of cyclobutanone 4.' Hence, the reduction products were found to possess the stereochemistry indicated. Treatment of 6a and 66 with methanesulfonyi chloride in pyridine and subsequent** elimination of the resulting methanesulfonates with potassium tert-butoxide **in dmethyl sufoxide at room temperature produced effectively the same enoi** thioether 8^7 [mp 47°C; ¹H NMR(CDCl₃) δ 5.93(s, 1H); Mass m/z 324(M⁺)] in good yields.
Hydrolysis of 8 with mercuric chloride in aqueous acetonitrile at room **Hydrolysis of 8 with mercuric chloride in aqueous acetanitrile at room 7** ~. **temperature5 spontaneously cyclobutenone 9 in 45% isolated yield. Scheme** 1 **(method A)**

^a**LIN(TI!S12(3 €~uiv)/THF/-78~C-O'Clfhen PhSSPh(2.5 e~utv)lHMPA/O'C,9O%** ; **b** NOBH₄/MeOH,99X from <u>5a</u>, 90X from <u>5b</u> , c Raney Ni(W-Z)/EtOH/rt ; d MsCl/pyr.
b NaBH₄/MeOH,99X from <u>5a</u>, 90X from <u>5b</u> , c Raney Ni(W-Z)/EtOH/rt ; d MsCl/pyr. I NaBH₄/MeOH,99% from <u>5a</u>, 90% from <u>5b</u> ; c Raney Ni(W-2)/EtOH/rt ; d MsCl/pyr.
/rt ; e t-BuOK/DMSO/rt,76% from <u>6a</u>, 84% from <u>6b</u> ; f H9Cl₂/CH₃CN-H₂O(3:l)/rt, **451 i 9 H2/PtlACOEt** ; **h PCClMolecuior Sleves 3A/CH2CI2.33% fran 9**

Catalytic hydrogenation of double bond in 9 with platinum under a hydrogen atmosphere gave rise to the overreduced cyclobutanol, which was immediately **oxidized to cyclobutanone 4 with pyridlnlum chlorochramate in dichlaramethane** in 33% overall yield. **Scheme 2** (method B) **Scheme** 2 (method B)

t3b **NOE** 13a 14 NQE **12b** 13_b

⁰ZnINH4Cl-W/O'C13Cmln,61% from cyclohexene ; **b PhSNolMeOH/O-C-rt.86I** ; **C NaBH4/MeOH.80% from** I.&. **90% from** UP ; **d NsCl/DYr./rt 1 e t-BuOK/DmOlrt, 87% from 130, 100% from 130** ; f HgCl₂/CH₃CN-H₂0(3:1)/50°C,62% ; g Raney Ni(W-2)

To improve the later stage in method A, we next turned our attention to the following method B which provided a more convenient and easy operation rather than method A. Dichlorocyclobutanone 10 was carefully reduced with zinc in **methanol ic amnonium chlorldeg at O'C for 30 mln to afford monochlorocyclobutanone 11**¹H **NNR**(CDC₁₃)^{δ} 4.95(dd, J=2.2 and 8.8 Hz, IH)] as the sole product in 61% **yield from cyclohexene. When 11 was allowed to react with sodium thiophenolate** *in methanol, an 86% yield of a 1:1 mixture of diastereoisomers 12a⁷[¹H NMR* $[CDC1₃]$ ⁵ 4.16(dd, J=2.2 and 5.4 Hz, 1H)] and $12b⁷[¹H N/R(CDC1₃)$ ⁵ 4.65(dd, J=2.2 **and 8.5Hr,lH)I was obtained. These diastereoisomers were easily separated by chromatography on silica gel. Reduction of 12a and 12b with sodium borohydride** gave their cyclobutanols 13a ['H NWR(CDCl₃)⁰ 3.54(dd, J_{bc}=7.6 and J_{ab}=10.0Hz,
,1H_b),3.99(t, J_{be}=J_{ed}=7.6Hz,1H_c)] and 13b⁷[¹H NWR(CDCl₃)δ 4.16(ddd,J_{bd}=2.2,J_{be}= **6.1, and Jab=8.3Hz,lHbl.4.50(dt, J,,=2.0 and Jbc=JCd=6.1 Hz,1Hc11 respectively in good yields. Stereochemistry of 13a and 13b obtained in this way was established in the similar manner mentioned above and results were indi**cared in Scheme 2. The enol thioether 14^{T} [H NMR(CDC1₄) δ 5.90(d, J=0.7 Hz, lH); Mass m/z 216(M⁺)] was obtained both from 13a and 13b in two steps (I) MsCl/py**ridine (11) t-BuCK/IMSO.**

Fjnaily, hydrolysis of the resulting enol thioether 14 with mercuric chloride at 50°C produced the desired cyclobutanone 4 in 62% yield. All reaction mixtures produced from either method A or B are not required their separations since they yield the same enol thioethers 8 or 14. This method should provide ready **access** to 1.2-carbonyl transposition of **cis-blcyclo[4.2.0loctanone** derivatives. Stereochemistry during reduction of cyclobutanones 5a, 5b, 12a, and 12b with sodium borohydride resulted in steric control as indicated in Scheme 3. We are currently exploiting this methodology in **our** approaches to the synthesis of erioianin I. **Scheme 3**

**12a R'=H, R²=SPh
ACKNOWLEDGMENTS 12b R'=SPh, R²=H**

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- For earlier leading references, **see; (a)** j. C. Stevens, D. A. Reich. D. 1. R. Brandt, K. R. Fountain, and E. J. Gaughan, J. Am. Chem. Soc.. **87,** 5257 (1965); (b) R. Montalgue and L. Ghasez, Angew. Chem. Int. Ed. 5257 (1965); (b) R. Montaigue and L. Ghosez, <u>Angew. Chem. Int. Ed.
Engl., 7</u>, 221 (1968); (c) W. T. Brady and O. H. Waters, <u>J. Org. Chem.,</u>
32, 3707 1967); (d) A. Hassner, V. R. Fletcher and D. E. O. Harmon,
1. Am. Cham
- J. Am. Chem. Soc., 93, 264 (1971).
- (a) C. R. Clark, J. Lin, and M. Nikaido, <u>Tetrahedron Lett.,</u> 2645
(1984); (b) A. E. Greene, M. J. Luche, and J. P. Depres, <u>J. Am. Chem.</u>
<u>Soc., 105</u>, 2435 (1983); (c) R. F. Newton and S. M. Roberts, $2.$ (19841: (b) A. E. **Greene,** M. J. Luche, and J. P. Depres, J. **Am.** Chem. Soc., 105, 2435 (1983); (c) R. F. Newton and S. M. Robe
Tetrahedron, 36, 2163 (1980); (d) B. M. Trost, Top. Curr. Chem., 43, 563 (1975); **(el** A. Hassner, H. W. Pinnick, and J. M. Ansell, Chem., 43, 1774 (1978); (f) W. T. Brady, Tetrahedron, 37, 2949 (1981).
- Isolation; S. M. Kupchan, **R.** L. Baxter, C.-K. Chiang, C. 1. Gllmore, and $3.$ R. F. Bryan, J. Chem. Soc. Chemm. Commun., 842 (1973). Total Synthesis; (a) P. A. Grieco, T. Oguri, S. Gilman, and G. T. DeTitta J. Am. Chem. Soc., 100, 1617 (1978); (b) M. R. Roberts and R. H. Schlessinger, J. Am. Chem. Soc., 103, 724 (1981).
- The chemistry of 1.2-carbonyl transpos~tion, **see;** V. V. **Kane,** V. Singh, $4.$ A. Martin, and D. L. Doyle, Tetrahedron, **39,** 345 (1983).
- 8. M. Trost, K. Himi, end S. Kurorumi, J. Am. Chem. **Soc.. 97,** ⁴³⁸ $5.$ (1975).
- T. Cohen, L.-C. Yu, and W. M. Daniewski, J. Org. Chem., *50,* 4596 (1985). 6.
- 7. A₁₁ new compounds gave satisfactory ¹H NMR, IR, and mass spectra in accord with indicated structures.
- 8. K. B. Wiberg and J. G. Pfeiffer, J. Am. Chem. Soc., 92, 553(1970).
- Reductive dedichlorination, see; P. W. Jeffs, G. Molina, M. W. Cass, and 9. N. A. Cortese, **J.** Org. Chem., 47, 3871 (1982).

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