Synthesis of Furan-, Thiophene- and Pyrrole-fused Sultines and their Application in Diels–Alder Reactions†

Wen-Sheng Chung,* Wen-Ju Lin, Wen-Dar Liu and Liang-Gyi Chen

Department of Applied Chemistry, National Chiao Tung University, Hsinchu, Taiwan 30050, R. O. C.

The synthesis of 1,4-dihydrofurano[3,4-d]-3,2-oxathiine 2-oxide **8**, 5,7-dimethyl-1,4-dihydrothieno[3,4-d]-3,2-oxathiine 2-oxide **9** and 1,4-dihydro-6-tosylpyrrolo[3,4-d]-3,2-oxathiine 2-oxide **10**, precursors for nonclassical heteroaromatic *o*-quinodimethanes, and their application in the Diels–Alder reactions are reported.

The chemistry of heterocyclic analogues of *o*-quinodimethane (*o*-QDM, **1**) has attracted a great deal of attention recently, and several reports of the generation of the furan and pyrrole derivatives have appeared.¹ The compounds 3,4-dimethylidenefuran **2**, 3,4-dimethylidenethiophene **3** and 3,4-dimethylidenepyrrole **4** are π -conjugated non-Kekulé molecules that have aroused theoretical and synthetic interest.^{1,2} These 3,4-dimethylidene-heteroaromatics **2–4** were generated from the corresponding diazenes² and have been detected by EPR, UV and NMR spectroscopy. They have also been shown by Berson and coworkers² to react with a series of alkenes to form two types of cycloadducts: fused and bridged. The results are synthetically useful and have been elaborated upon by Takayama and coworkers in syntheses of multicyclic compounds.^{1b,c}

Diazene precursors are usually unstable at room temperature, therefore the search for possible substitutes becomes important. Various methods for generation of these highly reactive dienes have been developed.^{3,4} Among the many known methods for their preparation, that involving cheletropic elimination of SO₂ from heteroaromatic-fused 3-sulfolenes 5-7 has drawn the most attention.^{1,4} Durst *et al.* were the first to generate *o*-QDM by thermal elimination of SO₂ from a sultine, 1,4-dihydro-2,3-benzoxathiine 3-oxide.⁵ A significant advantage of using sultines is that their thermolysis occurs at a much lower temperature than that of corresponding sulfolenes (80 vs. 170-220 °C). Recently other papers on using sultines as o-QDM precursors have appeared,6,7 however, the use of sultines in heteroaromatic analogues is still rare. We report here our work on the synthesis of furanosultine 8, thienosultine 9 and pyrrolosultine 10 and their applications in Diels-Alder reactions with alkenes and alkynes.

Previously unknown furanosultine 8 and thienosultine 9 are synthesized in two steps with good yield, as shown in Scheme 1. The 3,4-bis(chloromethyl)furan 11 was prepared by the known method.^{2b,8} The bis(chloromethyl)thiophene 12 was obtained by chloromethylation of 2,5-dimethylthiophene adapted according to the procedures developed by Wynberg *et al.* in the synthesis of corresponding sulfones $6.^9$ The 3,4-bis(chloromethyl)-*N*-tosylpyrrole 13 was synthesized in four steps (41%)

overall yield) as described by Berson and co-workers in the syntheses of corresponding diazenes.² The key step in the syntheses of sultines **8–10** is the use of Rongalite^{6,7a} (sodium formaldehyde sulfoxylate) with the corresponding dichlorides **11–13**.[‡]

The Diels-Alder reactions of furan-fused sultine 8 with several dienophiles are presented in Table 1 (entries 1-5) and Scheme 2. Heating 8 with 3 equivalents of dimethyl acetylenedicarboxvlate (DMAD) in benzene at 120-123 °C in a sealed tube for 1 h produced 5,6-dimethylidene-7-oxanorbornene 14a in 38% yield plus some polymer by-products (Table 1, entry 1). Essentially the same types of reactions were observed with diethyl fumarate (DEF), dimethyl maleate (DMM) and fumaronitrile (FN) (Table 1, entries 2-4). With N-phenylmaleimide (NPM), thermolysis at 134-140 °C gave a new type of product 15e in 24% yield. The low to medium yields of these reactions of furanosultine 8 with various dienophiles are disappointing, however, as their reaction products are different from those reported by Takayama and co-workers for the reactions of furan-fused sulfolene $5, 1^{1b,c}$ where two types of products 14 and 15 were formed. Furthermore, the high yield of intractable polymers in all cases also implies that radical or biradical-initiated polymerization is involved.

Thermal Diels-Alder reaction of 2,5-dimethylthiophenosultine 9 with several dienophiles was studied next (Table 1, entries 6–9). In the absence of a dienophile, sultine 9 underwent thermolysis to give the sulfolene 6 in excellent yield (90%, entry 6). Heating 9 with DMAD, DMF or NPM at 180 °C gave both the isomerized sulfolene 6 and fused adducts 16g and 16h in 89--97% yields. Unexpectedly, no adducts were formed when the thiophene-fused sulfolene $\mathbf{6}$ was heated in the presence of DMAD or DMF at the same reaction temperature (Table 1, entries 10-11). About 10% of the fused adduct 16h was obtained only when a strong dienophile such as NPM was used (entries 9, 12). Wynberg *et al.*^{9*a*} were the first to synthesize thiophene-fused sulfolene **6b** ($R = CO_2Me$), however, they did not detail its chemical reactivities other than pyrolysis. Our results show that thiophene-fused sulfolene 6 has very low reactivity compared to the corresponding sultine 9 in the Diels-



5 X = O 6 X = S 7 X = NR 8 R = H, X = O 9 R = Me, X = S 7 X = NR 10 R = H, X = N-SO₂C₇H₇

11 R = H, X = O 12 R = Me, X = S 13 R = H, X = N-SO₂C₇H₇



Scheme 1 Reagents and conditions: i, AlCl₃, diethyl ether, room temp.; ii, Rongalite, TBAB, DMF, 25 °C; iii, ClCH₂OMe, $SnCl_4$ –CS₂

2538

Alder reaction. Thus, in the reaction conditions studied here, the fused Diels-Alder adducts 16g-h can be obtained from the reaction of sultine 9 with dienophiles, but not from sulfolene 6. It is interesting to observe the great similarity between our results with sultines with those with diazenes reported by Berson *et al.*² For example, in the thiophene-biradical 3 trapping experiments, the sole adducts found (85–100% yields) had the fused structure 16 rather than the bridged structure 14.

Reactions of N-tosylpyrrolosultine 10 with a series of dienophiles (3 equiv.) at 150–170 °C run smoothly to give three types of products: sulfolene 7 (20–40%), 1:2 Diels–Alder adducts 15 (0–42%) and fused adducts 16 (5–73%) (Table 1, entries 13–18). Basically the same reaction products are obtained even at 110 °C (Table 1, entry 15), however to our surprise, only 7 and the fused adduct 16i were obtained when

DMAD was reduced to 1 equiv. (entry 14). The *N*-tosylpyrrolosulfolene **7** had been shown to be unreactive¹⁰ with dimethyl fumarate (DMF) even at 240 °C, however the *N*-tosylpyrrolosultine **10** reacted with DMF at 170 °C to give a fused adduct **16j** (63%) and the isomerized **7** (32%) (Table 1, entry 16 vs. 20). It is also of interest to isolate the fused adduct **16i** (5%) in the reaction of sultine **10** with DMAD compared with that of sulfolene **7**,^{*ib.c*} where **15i** was the only product formed (entry 13 vs. 19).

These results can be explained by two mechanisms. The most obvious possibility is the formation of non-Kekulé biradicals (2–4), followed by Diels–Alder reaction with a dienophile to form either bridged (14) or fused adducts (16).² Both adducts can add another dienophile to form the 1:2 adducts (15). Alternatively, a Diels–Alder reaction may first occur on the aromatic moieties of sultines 8–10 to give 17, from which SO₂

Table 1 The Diels-Alder reactions of sultines 8-10 and sulfolenes 6-7 with dienophiles^a

Entry	Diene	Dienophile	T/°C	t/h	Products (yield %) ^c	Total yield (%)	Ref.
1	8	DMAD	120-123	1	14a (38)	38	
2	8	DEF	140-145	6	14b (46)	46	
3	8	DMM	130-138	12	14c (38)	38	
4	8	FN	135142	4	14d (53)	53	
5	8	NPM	134-140	3	15e (24)	24	
6	9	_	180	24	6 (90)	90	
7	9	DMAD	180	24	6(44) + unknown	>44	
8	9	DMF	180	24	6 (49) + 16g (40)	89	
9	9	NPM	180	24	6(6) + 16h(91)	97	
10	6	DMF	180	24	6 (99)	99	
11	6	FN	180	24	6 (99)	99	
12	6	NPM	180	24	6 (89) + 16h (10)	99	
13	10	DMAD	165-170	12	7 (40) + 15i (42) + 16i (5)	87	
14	10	DMAD (1 equiv.)	155	12	7 (40) + 16i (9)	49	
15	10	DMAD	110	12	7 (30) + 15i (38) + 16i (8)	76	
16	10	DMF	170	14	7 (32) + 16j (63)	95	
17	10	FN	170	17	7 (34) + 16k (50)	84	
18	10	NPM (4 equiv.)	156160	22	7(20) + 16I(73)	93	
19	7	DMAD	170	14	15i (97)	97	1b, 1c
20	7	DMF	170–240 ^b	33	7 (73)	73	1b, 1c

^a Reactions were run with 3 equiv. of dienophiles in benzene (sealed tube) unless otherwise specified. ^b Reaction was run in xylene. ^c Isolated yield. For structures of products **14a–16l** see Scheme 2.



Scheme 2 Possible reaction pathways

J. CHEM. SOC., CHEM. COMMUN., 1995

is eliminated instantaneously to give bridged adducts 14. Compound 14 further reacts with another dienophile to give the 1:2 adducts 15, and finally a retro-Diels-Alder reaction of 15 would occur to form 16 (Scheme 2). The latter mechanism was proposed by Takayama to explain results with corresponding sulfolenes 5 and 7. We conclude that the furano-, thieno- and Ntosylpyrrolo-fused sultines 8-10 reacted under milder condition than the corresponding sulfones 5-7 and their reaction products were different in many cases. When generated in the presence of a dienophile, they can provide elegant synthons for the formation of [4 + 2] cycloadducts. If less reactive trapping agents were used, the diene was recaptured by SO₂ to afford the sulfones 5-7 (path 3 in Scheme 2).11

We thank the National Science Council of the Republic of China for financial support (Grant No. NSC 84-2113-M-009-002). W. S. C. would like to thank Professor J. A. Berson for his encouragement and valuable suggestions.

Received, 11th September 1995; Com. 5/05968B

Footnotes

† Presented in the 15th International Congress of Heterocyclic Chemistry, Taipei, R. O. C., August, 1995, Book of Abstracts, PO3-257.

[‡] Dichlorides 11,² 12^{9b} and 13² have been reported in the literature and our samples correspond in all respects with the reported properties. For all products satisfactory spectral data were obtained. Selected data for 8: light yellow oil; ¹H NMR (300 MHz, CDCl₃) & 7.40 (1 H, s), 7.36 (1 H, s), 5.28 $(1~{\rm H},{\rm AB},J~14.4~{\rm Hz}),\,5.02~(1~{\rm H},{\rm AB},J~14.4~{\rm Hz}),\,3.87~(1~{\rm H},{\rm A'B'},J~15.6~{\rm Hz})$ and 3.65 (1 H, A'B', J 15.3 Hz); ¹³C NMR (75.4 MHz, CDCl₃) & 141.01 (CH), 136.38 (CH), 113.92 (C_q), 108.00 (C_q), 55.03 (CH₂) and 46.64 (CH₂). For **9**: white solid, mp 70–71 °C; ¹H NMR, δ 5.16 (1 H, AB, J 14.1 Hz), 4.92 (1 H, AB, J 13.5 Hz), 3.88 (1 H, A'B', J 15.6 Hz), 3.58 (1 H, A'B', J 15.0 Hz) and 2.27 (6 H, s); ¹³C NMR, 133.15 (C_q), 128.72 (C_q), 125.40 (C_q), 119.13 (Cq), 58.73 (CH₂), 50.92 (CH₂) and 12.11 (CH₃); m/z 202 (M+), 138 (100%), 123 and 91; (Found: M+ 202.0116 C₈H₁₀O₂S₂ requires 202.0122). For 10, white solid, mp 152–153 °C; ¹H NMR, δ 7.76 (2 H, d, J 8.8 Hz), 7.31 (2 H, d, J 8.1 Hz), 7.07 (1 H, s), 7.01 (1H, S), 5.20 (1 H, AB, J 14.2 Hz), 4.91 (1 H, AB, J 14.2 Hz), 3.88 (1 H, A'B', J 15.6 Hz), 3.56 (1H, A'B', J 15.6 Hz) and 2.42 (3 H, s); 13 C NMR, 145.50 (C_q), 135.54 (C_q), 130.15 (CH), 127.03 (CH), 119.19 (CH), 117.12 (C_q), 114.44 (CH), 110.97 (C_q),

References

and 92.

- 1 For recent review see; (a) R. A. Aitken, I. Gosney and J. I. G. Cadogan, Prog. Heterocycl. Chem., 1992, 4, 1; 1993, 5, 1; (b) T.-S. Chou, Rev. Heteroatom Chem., 1993, 8, 65; (c) K. Ando and H. Takayama, Heterocycles, 1994, 37, 1417; (d) K. Ando, M. Kankake, T. Suzuki and H. Takayama, Tetrahedron, 1995, 51, 129.
- 2 (a) L. C. Bush, R. B. Heath and J. A. Berson, J. Am. Chem. Soc., 1993, 115, 9830; (b) M. M. Greenberg, S. C. Blackstock, K. J. Stone and J. A. Berson, J. Am. Chem. Soc., 1989, 111, 3659; (c) K. J. Stone, M. M. Greenberg, S. C. Blackstock and J. A. Berson, J. Am. Chem. Soc., 1989, 111, 3671 and references cited therein.
- 3 S. Braverman, Y. Duar and D. Segev, Tetrahedron Lett., 1976, 3181; P. J. Garratt and S. B. Neoh, J. Org. Chem., 1979, 44, 2667
- 4 J. L. Charlton and M. M. Alauddin, Tetrahedron, 1987, 43, 2873; R. L. Funk and K. P. C. Vollhart, Chem. Soc. Rev., 1980, 9, 41; W. Oppolzer, Synthesis, 1978, 793; K. C. Nicolaou, W. E. Barnette and P. Ma, J. Org. Chem., 1980, 45, 1463.
- 5 T. Durst, J. L. Charlton and D. B. Mount, Can. J. Chem., 1986, 64, 246; J. L. Charlton and T. Durst, Tetrahedron Lett., 1984, 25, 5287; F. Jung, M. Molin, R. Van Den Elzen and T. Durst, J. Am. Chem. Soc., 1974, 96, 935.
- 6 For a review of sultines see: D. C. Dittmer and M. D. Hoey, The Chemistry of Sulphinic Acids, Esters and Their Derivatives, Wiley, Chichester, 1990, pp. 239-273.
- 7 (a) M. D. Hoey, D. C. Dittmer, J. Org. Chem., 1991, 56, 1947; (b) W. F. Jarvis, M. D. Hoey, A. L. Finocchio and D. C. Dittmer, J. Org. Chem., 1988, 53, 5750; (c) G. Attardo, W. Wang, J.-L. Kraus and B. Belleau, Tetrahedron lett., 1994, 35, 4743.
- 8 M. P. Cava and A. A. Deana, J. Am. Chem. Soc., 1959, 81, 4266; C. E. Doecke, P. J. Garratt, H. Shahriari-Zavareh and R. Zahler, J. Org. Chem., 1984, 49, 1412.
- 9 (a) H. Wynberg and D. J. Zwanenburg, J. Org. Chem., 1964, 29, 1919; (b) D. J. Zwanenburg and H. Wynberg, J. Org. Chem., 1969, 34, 333.
- 10 K. Ando, M. Kankake, T. Suzuki and H. Takayama, Synlett., 1994, 741; K. Ando, M. Kankake, T. Suzuki and H. Takayama, J. Chem. Soc., Chem. Commun., 1992, 1100.
- 11 Sulfur dioxide appears to be an especially reactive trapping agent which can intercept the furan, thiophene and pyrrole xylenes. See for example: P. M. S. Chauhan, A. P. A. Crew, G. Jenkins, R. C. Storr, S. M. Walker and M. Yelland, Tetrahedron Lett., 1990, 31, 1487; 1491.