## Chemical Reactivity and Charge Densities in Certain 6a-Thiathiophthens

By R. J. S. Beer,\* D. Cartwright, R. J. Gait, R. A. W. Johnstone, and S. D. Ward (The Robert Robinson Laboratories, University of Liverpool, Liverpool 7)

The formation of monosubstitution products by the action of bromine on 2,5-diphenyl- and 2,5-dimethyl-6a-thiathiophthen has been reported. We have examined the reactivity of some unsymmetrically substituted thiathiophthens and compared their observed behaviour with that predicted by molecular orbital theory.

Bromination of 2-methyl-5-phenyl-6a-thiathiophthen (I; R = H) gives a monobromo-derivative, believed to be 3-bromo-2-methyl-5-phenyl-6a-thiathiophthen (I; R = Br) m.p. 113°;  $\lambda_{max}$  251, 273, 334, and 504 m $\mu$  (log  $\epsilon$  4·68, 4·51, 4·04, and 4·03);  $\tau$  7·27 (3H, s),  $\tau$  2·14—2·68 (5H, m),  $\tau$  1·51 (1H, s). The position of bromination is inferred from a comparison of n.m.r. data in the series (see Table).

TABLE

Substituents		Chemical shifts in $CDCl_3$ ( $\tau$ )	
		3-position	4-position
2,5-Diphenyl		1.82	1.82
3-Bromo-2,5-diphenyl			1.26
2,5-Dimethyl		$2 \cdot 43$	$2 \cdot 43$
3-Bromo-2,5-dimethyl			2.06
2-Methyl-5-phenyl		$2 \cdot 31$	1.97
3-Bromo-2-methyl-5-phenyl			1.51

2-Methylthio-5-phenyl-6a-thiathiophthen<sup>1,2</sup> (II; R=H) also brominates smoothly, but its nitration (with concentrated nitric acid in hot acetic acid) is of greater interest, since the structure of the product (II;  $R=NO_2$ ) has been confirmed by an unambiguous synthesis from 3-methylthio-5-phenyl-1,2-dithiolium methosulphate (III) and methyl nitrodithioacetate (IV). The nitro-compound, m.p.  $158-159^\circ$ , has the characteristic3 thiathiophthen absorption spectrum;  $\lambda_{max}$  254, 322, and 480 m $\mu$  (log  $\epsilon$  4·53, 4·26, and 4·04).

The 3-position is also attacked on nitrosation of 2-methylthio-5-phenyl-6a-thiathiophthen with nitrous acid in acetic acid at 5°, but the product, m.p. 138—139° ( $\lambda_{max}$  324 and 422 m $\mu$ ; log  $\epsilon$  4·28 and 4·04), probably exists in the isomeric form V;  $R = CS_2Me$ ).

The methylthio-group in 2-methylthio-5-phenyl-6a-thiathiophthen is slowly replaceable by nucleophiles. Thus prolonged reaction with sodium

ethoxide in ethanol under reflux gave 2-ethoxy-5-phenyl-6a-thiathiophthen, (70%) m.p. 90—91°;  $\lambda_{\rm max}$  258, [280—290], and 478 m $\mu$  (log  $\epsilon$  4·48, [4·28], and 4·02). Primary aliphatic amines reacted similarly with 2-methylthio-5-phenyl-6a-thiathiophthen, giving 2-alkylamino-derivatives.

Attempted nitration and nitrosation of 2,5diphenyl-6a-thiathiophthen yielded the same nitroso-compound (VI; R = NO) or (V; R =COPh), m.p. 172—173°;  $\lambda_{\text{max}}$  252, 298, [330], and 416 m $\mu$  (log  $\epsilon$  4·30, 4·01, [3·94], and 4·01). This product, also obtained by nitrosation of 5-phenyl-1,2-dithiol-3-ylidene-acetophenone, is not intensely coloured, as one might expect for structure (VI; R = NO) and its i.r. spectrum contains a carbonyl-stretching band at 1640 cm.<sup>-1</sup>. Dithiolylidene ketones in which the carbonyl group is adjacent to, and interacts with, the disulphide linkage do not show normal C=O stretching bands.4 We suggest that the nitroso-compound exists in the form (V; R = COPh), with some interaction between the oxygen atom of the nitroso-group and the adjacent sulphur atom.

The nitroso-derivative obtained from 2-methyl-thio-5-phenyl-6a-thiathiophthen, in contrast to the nitro-derivative (II;  $R = NO_2$ ), does not have the electronic absorption spectrum of a thiathiophthen, but resembles the nitroso-ketone (V; R = COPh) and thus should probably be formulated as (V;  $R = CS_2Me$ ). Conversion to the simpler nitroso-compound (V; R = H), m.p.  $131-132^\circ$  ( $\lambda_{max}$  327 and 420 m $\mu$ ; log  $\epsilon$  4·00 and 4·04) was achieved by the action of mercuric acetate.†

Whatever the precise structure of the nitrosocompounds, it is clear that 2-methylthio-5-phenyl-6a-thiathiophthen is attacked at the 3-position by electrophiles and at the 2-position by nucleophiles (with displacement of the methylthio-group). Charge densities for the non-alternant 6a-thiathiophthen system have been calculated both by the Wheland–Mann  $\omega$ -technique<sup>5</sup> and by a PPP-SCF method, using the Maeda model in which the three sulphur atoms are bonded through  $3p-3d_{xz}$  hybridisation of the central sulphur atom. Values of the calculated charge densities and localisation energies for 2-methyl-5-phenyl-6a-thiathiophthen and charge densities for 2-methylthio-5-phenyl-6a-thiathiophthen, obtained by the  $\omega$ -method, are

<sup>†</sup> For similar desulphurisation reactions with mercuric acetate, see ref. 2.

Ph 
$$\stackrel{S}{\longrightarrow}$$
 Me  $\stackrel{S}{\longrightarrow}$  SMe  $\stackrel{S}{\longrightarrow}$  SMe  $\stackrel{MeSO_4}{\longrightarrow}$  O<sub>2</sub>N·CH<sub>2</sub>·CS<sub>2</sub>Me  $\stackrel{S}{\longrightarrow}$  (III) (IV)

$$\stackrel{S}{\longrightarrow}$$
 S O  $\stackrel{S}{\longrightarrow}$  S

shown in diagrams (VII) and (VIII). On the basis of the charge-density values, the 3-position in both molecules should be the position of attack by electrophiles, in agreement with our observed results. Consideration of localisation energies, on the other hand, suggests that the 4-position should be the more reactive (the non-crossing rule9 is broken), but it has been noted10 that charge densities in sulphur-containing non-alternants usually give a better correlation with reactivity than do localisation energies.

The positive charge density at the 2-position in 2-methylthio-5-phenyl-6a-thiathiophthen (VIII) is in accordance with the susceptibility of this compound to attack by nucleophiles. Much other evidence could be adduced to support the view that the 2-position in 6a-thiathiophthens is electron-deficient.

(Received, April 16th, 1968; Com. 470.)

<sup>1</sup> R. J. S. Beer, D. Cartwright, and D. Harris, Tetrahedron Letters, 1967, 953.

<sup>2</sup> R. J. S. Beer, R. P. Carr, D. Cartwright, D. Harris, and R. A. Slater, J. Chem. Soc. (C), in the press.

<sup>3</sup> H. Behringer and A. Grimm, Annalen, 1965, 682, 188; H. Behringer, M. Ruff, and R. Wiedenmann, Chem. Ber., 1964, 97, 1732.

<sup>4</sup> N. Lozac'h, "Organosulfur Chemistry", Interscience, New York, 1967, p. 196. <sup>5</sup> G. W. Wheland and D. E. Mann, J. Chem. Phys., 1949, 17, 264.

<sup>6</sup> J. A. Pople, Trans. Faraday Soc., 1953, 49, 1375.

- K. Maeda, Bull. Chem. Soc. Japan, 1960, 33, 1466; 1961, 34, 785, 1166.
   R. A. W. Johnstone and S. D. Ward, to be published.
- 9 A. Streitwieser, "Molecular Orbital Theory for Organic Chemists", Wiley, London, 1961, p. 347.

<sup>10</sup> R. Zahradnik, Adv. Heterocyclic Chem., 1965, 5, 57.