Covalently Attached Ferrocene and Tetrathiafulvalene Redox Systems

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The synthesis and solution redox properties of several covalently attached ferrocene and tetrathiafulvalene units are reported, together with the X-ray crystal structure of a 1,1'-bis(1,3-dithiole-2-ylidene)-substituted ferrocene derivative.

The study of molecules comprising two different redox centres that are covalently linked is a burgeoning topic within supramolecular chemistry, with applications in molecular electronic devices, sensors, electrocatalysis and energy conversion. Current research in this area focuses on gaining control over charge-transfer interactions between combinations of organic and organometallic species, *e.g.*, bipyridinium cations, quinones, metallocenes and metal-coordinated macrocycles.²

The covalent incorporation of tetrathiafulvalene (TTF) units into multistage redox assemblies3 presents a fascinating prospect for the following reasons: (i) oxidation of TTF to the radical cation species occurs reversibly at a relatively low potential (ca. 0.34 V, vs. Ag/AgCl for unsubstituted TTF) which can be finely tuned by the attachment of appropriate substituents; (ii) TTF+ is thermodynamically a very stable species; (iii) oxidised TTF units readily form highly ordered arrays with close intermolecular interactions involving both π - π overlap and S···S contacts. However, the chemistry of functionalised TTF derivatives which are needed to construct such assemblies is virtually unexplored. 4-6 There are two main reasons for this: (i) direct substitution onto the TTF ring is frequently an inefficient process, yielding a mixture of products which are difficult to separate (ii) very few methods are available whereby the TTF system can be built up from functionalised components, and these are invariably lengthy procedures involving a capricious coupling reaction of two 1.3-dithiole units as the final step.8

We report here the synthesis of prototype systems comprising covalently linked ferrocene and TTF moieties. Additionally, we describe the solution electrochemical properties of the new materials, an X-ray crystal structure and preliminary data on complex formation with tetracyano-p-quinodimethane (TCNQ). Two distinct structural combinations of ferrocene and TTF are described, utilising functionalised TTF and 1,3-dithiole building blocks that we have recently developed in the context of studies on organic metals. Ferrocene has been attached both to the periphery of TTF (compounds 1b–1d, 3b, 4b and 5) and incorporated as a spacer unit between two 1,3-dithiole-2-ylidene rings (compounds 6a and 6b).

Systems 1b-1d, comprising one ferrocene and one TTF unit separated by one, two and five atom spacer groups, respectively, were prepared as follows. Ferrocenecarbonyl chloride 1a reacted with tetrathiafulvalenyllithium 2a⁷ (-78°C in diethyl ether) to yield compound 1b (12% yield).† Reaction of compound 1a with TTF-thiolate anion 2b (generated from thiobenzoate derivative 2c and sodium ethoxide in ethanol at -10 °C)⁵ gave thioester derivative 1c (27% yield). The reaction of acid chloride 1a with TTF-alcohol 2d4 (triethylamine, dichloromethane, 20°C) was particularly efficient, affording ester 1d in 80% yield. Directly analogous reactions have provided bis-TTF and bis-ferrocene derivatives 3b and 4b, respectively. Thus the reaction of 1,1'-ferrocenedicarbonyl dichloride 3a and TTF-alcohol 2d gave the product 3b (20% yield, based on 1,1'-ferrocenedicarboxylic acid), and TTF-dialcohol 4a9 and compound 1a gave product 4b (60% yield).

Ferrocene-TTF system 5, with a *trans*-ethene linkage, was obtained (58% yield, after separation from a mixture of *trans*-and *cis*-isomers) from the reaction of TTF-carboxaldehyde **2e**¹⁰ and the Wittig reagent generated from ferrocene-CH₂-PPh₃+ iodide (butyllithium, tetrahydrofuran, -78 °C).¹¹

The incorporation of conjugated acyclic and cyclic units (e.g. ethene, anthracene and thiophene groups) between two 1,3-dithiole rings, to form 'stretched' TTF derivatives with unusual structural and redox properties, has been the subject of extensive studies in our laboratory¹² and elsewhere. ¹³ Hitherto, the use of an organometallic spacer group, which is

[†] Satisfactory elemental analysis, mass spectra and ¹H NMR spectra were obtained for all new compounds.

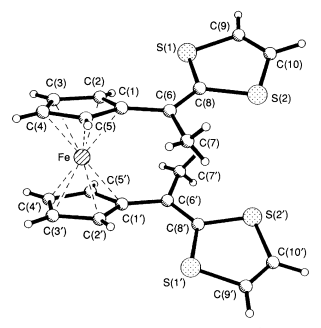


Fig. 1 Molecular structure of $[Fe(C_5H_4CMe=C_3S_2Me_2)_2]$ **6a.** Selected bond lengths (Å): Fe-C(1) 2.073(5), Fe-C(2) 2.049(7), Fe-C(3) 2.042(8), Fe-C(4) 2.049(5), Fe-C(5) 2.046(7), Fe-Cp-plane 1.65, C(1)-C(6) 1.463(6), C(6)-C(8) 1.356(7), C(8)-S(1) 1.760(6), C(8)-S(2) 1.770(4), S(1)-C(9) 1.744(7), S(2)-C(10) 1.723(10), C(9)-C(10) 1.307 (12)

itself a redox species, has not been reported in this context. Compounds **6a** and **6b** have now been prepared in 60-70% yield by the twofold reaction of 1,1'-diacetylferrocene and the 1,3-dithiole Horner–Wittig reagents **7a** and **7b**, 12b respectively (butyllithium, tetrahydrofuran, -78 °C).

The structure of compound **6a** was determined by X-ray diffraction‡ (Fig. 1). The molecule is situated around a twofold crystallographic axis, with the cyclopentadienyl rings parallel to within 1.3° and staggered by 17.6°. The entire ligand is remarkably non-planar, being twisted by 26.0° around the C(1)-C(8) axis and by ca. 3.4° around the C(6)=C(8) double bond. The dithiole ring adopts an envelope conformation, folding by 11° along $S(1)\cdots S(2)$.

The electrochemical redox properties of the ferrocene-TTF systems have been studied by cyclic voltammetry; the data are collated in Table 1 and the cyclic voltammograms of compounds 3b, 5 and 6a are shown in Fig. 2. In all the compounds where ferrocene and TTF are linked by a bridge (viz. 1b-1d, 3b, 4b and 5) three reversible electron transfer oxidations are observed within the range ca. 0.3-0.85 V. These correspound to TTF \rightarrow TTF+, TTF+, \rightarrow TTF2+ and ferrocene \rightarrow

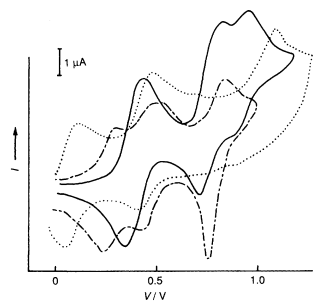


Fig. 2 Cyclic voltammograms of compounds 3b (----), 5 (---) and 6a (-----)

Table 1 Cyclic volatmmetric data^a

Compound	$E_1^{\frac{1}{2}}$	$E_2^{\frac{1}{2}}$	$E_3^{\frac{1}{2}}$
TTF	0.34	0.71	
Ferrocene	0.36		
1b	0.38	0.73	0.88
1c	0.45	0.73^{b}	
1d	0.35	0.61	0.75
3b	0.37^{c}	0.71^{c}	0.85
4b	0.45	0.62^{c}	0.82
5	0.26	0.44	0.77
6a	0.10	0.56	1.15^{d}
6b	0.06	0.51	1.05^{d}

^a Data are in volts, recorded using a Pt working electrode, Pt gauze counter electrode, Ag/AgCl reference electrode, 0.2 mol dm⁻³ Bu¹₄N+PF₆⁻, 10⁻⁴ mol dm⁻³ compound in dry acetonitrile under nitrogen at 20 °C, with iR compensation. All waves represent a reversible, one electron process except where indicated. ^b Broad two electron wave. ^c Two electron wave. ^d Irreversible wave.

ferrocinium. In all cases the first oxidation is assigned to TTF → TTF++, whereas the second oxidation is assigned to TTF++ → TTF2+ for compounds 1d and 3b, and to the ferrocene unit for compound 4b. This interpretation is supported by data obtained for the model compound 89 which does not contain the TTF system. For compounds 1b, 1c and 5 the assignment of the second and third oxidations is unclear. Both compounds 6a and 6b display two, reversible single-electron oxidations and a third irreversible oxidation at a higher potential. A striking feature of the data for 6a and 6b is that the first oxidation occurs at a remarkably low potential (ca. 0.1 V); we, therefore, assign this wave to the oxidation of the entire delocalised system involving significant interaction between the two 1,3-dithiole rings and ferrocene.

Compounds **6a** and **6b** yield charge-transfer complexes with TCNQ (40% yield, from dichloromethane solution) which have high room temperature conductivity values ($\sigma_{rt} = 0.2$ and 0.1 S cm⁻¹, respectively, using two-probe techniques on compressed pellets). The stoichiometry of both complexes, as judged by elemental analysis, is 1 donor: 2 TCNQ.

[‡] Crystal data for **6a**: C₂₀H₁₈FeS₄, M=442.4, monoclinic, space group C2/c (No. 15), a=17.847(2), b=10.191(2), c=12.849(2) Å, $\beta=126.01(1)^\circ$, U=1890.4(6) Å, Z=4, $D_c=1.56$ gcm⁻³, μ (Mo-K α) = 12.4 cm⁻¹, F(000)=912, crystal size $0.15\times0.25\times0.40$ mm. 2170 independent intensity data were collected at 23 °C on a Rigaku AFC-6S diffractometer using graphite-monochromated Mo-K α radiation ($\lambda=0.71073$ Å) and $2\theta/\omega$ scan mode ($2\theta<55^\circ$). The structure was solved by direct methods. All non-hydrogen atoms were refined in anisotropic approximation and all H atoms in isotropic approximation by full-matrix least-squares to R=0.056 ($R_w=0.065$, goodness-of-fit 1.90) for 1592 reflections with $|F|>4\sigma(F)$, corrected for absorption by semi-empirical methods (108 azimuthal scans of 3 reflections used, min. to max. transmission ratio 0.687). Atomic coordinates, bond lengths and angles, and thermal parameters have been depostied at the Cambridge Crystallographic Data Centre. See Notice to Authors, Issue No. 1.

We thank the SERC and ICI FCMO for funding this work, and The Royal Society for financial support (to A. S. B.).

Received, 21st December 1992; Com. 2/06746C

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