Radical Relatives: Facile Oxidation of Hetero-Diarylmethene Anions to Neutral Radicals

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ABSTRACT. Furan and thiophene diarylmethenes are potential redox-active ligands for metal centers that could be exploited in the development of non-traditional, stoichiometric, and catalytic redox reactions. As such, we describe here the selective meso-deprotonations of the dithiophene, difuran, and diimine-difuran diarylmethanes to form the π-conjugated anions, for which only the diimino-difuryl anion is truly isolable and studied by X-ray crystallography. In all cases, facile one-electron oxidation of these anions occurs which allows the isolation of the neutral dithienyl and diimino-difuryl radicals. UV-visible and TD-DFT studies reveal that the oxidation of the dithienyl anion to its radical is associated with an increase in the H(S)OMO-LUMO gap, evident
through a hypsochromic shift of the main absorption band in the electronic spectrum, whereas oxidation of the diimino-difuryl anion causes only minor spectroscopic changes. Electrochemical studies support the stability of the radicals with respect to the anion, showing strongly negative oxidation potentials. The control of the redox activity of these diarylmethene carbanions through variation of the nature of the substituents, donor-atom, and the conjugated π-system and their potential as ligands for redox-inert metal centers makes them intriguing candidates as non-innocent partners for redox reactions.

INTRODUCTION

In conventional redox reactions, the redox process is managed solely by a metal center which varies its formal oxidation state according to gain or loss of electrons. In contrast, an increasingly prevalent approach to redox catalysis is to use the redox ability of ligands to store or supply reducing equivalents, so limiting, or negating very high or low metal oxidation states. This latter approach takes inspiration from biological systems such as enzymes which use, for example, porphyrin or cubane ligands to store or supply electrons for redox reactions. Many metal complexes of redox-active ligands have been developed and studied and display redox chemistry that makes use of both ligand redox activity and the Lewis acidity of the metal center to ensure substrate binding. Important recent advances in this field include the reduction of carbon dioxide at an iron-porphyrin center that occurs only upon double reduction of the porphyrin to a tetra-anion, or the conferment of noble reactivity to non-noble metals such as iron through cooperation with a redox-active pyridinediimine (PDI) ligand.

Diarylmethanes have a rich radical chemistry, with early work showing that radical formation was inherent in the C-C coupling of diarylmethanes to form tetraarylethes.
diaryl methane radicals have been generated by flash photolysis of diarylmethanes through an oxidation/deprotonation mechanism, and related diarylmethane radical anions have been studied by EPR spectroscopy. Importantly, unlike simple diarylmethanes, the heteroatom congeners such as dipyrrins, and dipyridyl-, difuryl-, and dithienylmethenes could potentially act as redox-active chelating ligands to transition metals, having some similarity to the well-exploited β-diketiminate ligand family which have also been shown to exhibit ligand non-innocence.

Simple neutral difurylmethene radicals are accessible and have been studied by EPR spectroscopy, and dipyridylmethene complexes of the p-block elements form ligand-radical complexes of varying stability. Due to their relevance to porphyrins and the fluorescent properties of BODIPY dyes and related complexes, dipyrrins have been extensively studied and shown to exhibit redox activity in their complexes and dipyrrin-fullerene triads. Similarly to these studies, we have shown that a simply prepared anionic, imine-expanded dipyrrin acts as a ligand for iron and uranium complexes, the latter showing ligand redox activity. Due to the ligand-based LUMO, the sequential outer-sphere, one-electron reduction of the uranyl(VI) center to U(V) and U(IV) occurs by initial ligand reduction followed by electron-transfer to the metal. We have also reported the redox-active bis(iminothienyl)methene anion that undergoes facile single-electron oxidation to form the stable and isolable neutral iminothiophene radical. This radical formed the dinuclear Cu(I) ligand-radical complex upon reaction with CuI, and under more oxidizing conditions, the radical cation was generated. The mild potentials at which these redox processes take place and the poor coordinating ability of the sulfur donors of the thiophene heterocycles prompted us to study additional variants of the diarylmethene scaffold. As such, we describe here the influence of structural variations such as the
truncation of the imine functionalities or the use of a stronger coordinating furan heterocycle on
the redox and coordination properties of the hetero-triarylmethe unit.

Figure 1. The hetero-diarylmethene anions investigated in this work and their relationships to the
previously reported imine-expanded dithienylmethene compounds $L^{NS}$.

EXPERIMENTAL SECTION

**General experimental details.** The syntheses of all air- and moisture-sensitive compounds were
carried out using standard Schlenk techniques. Vacuum Atmospheres and MBraun glove boxes
were used to manipulate and store air- and moisture-sensitive compounds under an atmosphere of
dried and deoxygenated dinitrogen. All glassware was dried in an oven at 160 °C, cooled under
$10^{-3}$ mbar vacuum, and then purged with nitrogen. All solvents for use with air- and moisture-
sensitive compounds were stored in ampoules containing pre-dried 4 Å molecular sieves. Solvents
were collected from a Vacuum Atmospheres solvent tower drying system, where they had been
passed over a column of molecular sieves for 24 h prior to collection. They were then degassed
prior to use and subsequent storage.

$^1$H NMR spectra were recorded on a Bruker AVA400 spectrometer operating at 399.90 MHz, a
Bruker AVA500 or Bruker PRO500 operating at 500.12 MHz or a Bruker AVA600 spectrometer
operating at 599.81 MHz. $^{13}$C {$^{1}$H} NMR spectra were recorded on a Bruker AVA500 or Bruker PRO500 operating at 125.76 MHz. $^{1}$H and $^{13}$C {$^{1}$H} NMR spectra are referenced to residual solvent resonances calibrated against SiMe$_{4}$. $^{19}$F {$^{1}$H} NMR spectra were recorded on a Bruker AVA500 spectrometer operating at 470.59 MHz and referenced to CCl$_{3}$F. EPR spectra were recorded on a Bruker ELEXSYS E500 spectrometer and spectral simulations were carried out using Bruker’s Xsophe software package. UV-Vis absorption spectra were recorded on a Jasco V-670 spectrophotometer in a 10 mm quartz cuvette fitted with a Young's tap for air-sensitive compounds.

Electrochemical measurements were made using an Autolab ECO Chemie PGSTAT potentiostat and the data processed using GPES Manager, version 4.9. Experiments were carried out under a flow of N$_{2}$ in a 10 mL cell. The solution employed was 1 mM of the analyte in THF, with 0.1 M [$^{8}$Bu$_{4}$N][PF$_{6}$] as the supporting electrolyte. Cyclic voltammograms were recorded for quiescent solutions at variable scan rates between 100–500 mV s$^{-1}$. The nature of an observed redox process (reduction or oxidation) was determined by linear sweep voltammetry measured for stirred solutions with scan rates between 10–20 mV s$^{-1}$. The working electrode used was glassy carbon or platinum disc (d = 1 mm), with a platinum gauze counter electrode. Ag$^{+}$/Ag pseudo-reference electrode was used with potentials calibrated internally against the ferrocenium/ferrocene couple, Fc$^{+}$/0.

X-ray crystallographic data were collected at 170 K on an Oxford Diffraction Excalibur diffractometer using graphite monochromated Mo-K radiation equipped with an Eos CCD detector ($\lambda = 0.71073 \text{ Å}$). Structures were solved using ShelXT by direct methods or intrinsic phasing and refined using a full-matrix least square refinement on $|F|^2$ using ShelXL. All programs were used within the Olex suite. All non-hydrogen atoms were refined with anisotropic displacement parameters and H atom parameters were constrained to parent atoms and refined using a riding model unless otherwise stated.

DFT calculations were performed using the Gaussian09$^{42}$ package on the Eddie server system at the University of Edinburgh. Initial guess geometries were generated using the Avogadro program (version 1.1.1). All structures discussed in the text were optimized and converged according to the criteria for maximum displacement and maximum force. Frequency calculations were conducted to confirm that the optimized structures represented minimum energy geometries, which were confirmed by having no imaginary frequencies. The "\OPT=NoRaman" and "\FREQ=NoRaman"
options were used to improve computational efficiency. TD-DFT calculations were conducted on the first 40 excited states using the SCRF solvent model (Solvent=Dichloromethane). All optimization, frequency and TD-DFT calculations were carried out using the CAM-B3LYP functional and 6-311G(d,p) basis set. Molecular orbital surfaces were exported as cubefiles, visualized in UCSF-Chimera, and rendered with the Pov-Ray raytracer program.

Elemental analyses were carried out by Mr Stephen Boyer at the London Metropolitan University and were measured in duplicate.

The reagents 2-bromothiophene, furan, benzaldehyde, POCI₃, Amberlyst 15®, DMF, tert-butylamine and CuI were all used as supplied by Sigma-Aldrich, Fisher Scientific or VWR. Compounds 1, 2, and HL⁰ were synthesized according to published procedures.⁴⁴-⁴⁶

**Synthetic Details**

HL⁵ – A reported procedure was adapted.¹¹ Solid pentafluorobenzaldehyde (0.925 g, 4.70 mmol), pentafluorophenyl thiophene 1 (3.54 g, 14.1 mmol) and Fe(acac)₃ (0.332 g, 0.94 mmol) were dissolved in Me₃SiCl (5 mL), purged with argon and stirred at 40 °C for 18 h. The resulting mixture was diluted with CH₂Cl₂, quenched with aqueous NaHCO₃ and washed with water and brine before being dried over MgSO₄. Evaporation of the solvent provided a yellow/brown solid which was washed with hexanes to provide the product as an off-white powder (4.09 g, 87%). ¹H NMR (500 MHz, chloroform-d): δ = 7.42 (d, J = 3.8 Hz, 2H, γ-C₆H), 7.08 (d, J = 3.8 Hz, 2H, β-C₆H), 6.29 (s, 1H, meso-C₆H). ¹³C{¹H} NMR (126 MHz, chloroform-d): δ = 145.1 (m, 2xC), 145.0 (dm, J = 126 Hz), 143.1 (dm, J = 126 Hz), 141.3 (dm, J = 126 Hz), 140.2 (dm, J = 126 Hz), 138.2 (dm, J = 126 Hz), 130.3 (t, J = 5.0 Hz), 127.2 (s), 126.7 (m), 116.1 (t, J = 13.9 Hz), 109.7 (td, J = 15.1 Hz, J = 3.8 Hz), 36.70 (s). ¹⁹F NMR (471 MHz, benzene-d₆) δ = −140.5 (d, J = 15.5 Hz), −140.7 (m), −153.4 (t, J = 21.6 Hz), −155.7 (t, J = 21.6 Hz), −160.6 (m), −162.2 (td, J = 21.5, 5.9 Hz). Anal. Calcd. for [C₂₇H₃₅F₁₅S₂]: C, 47.8; H, 0.74%; found: C, 47.89; H, 0.76%. HRMS (APPI) (m/z): Calcd. for [C₂₇H₃₅S₂F₁₅], 676.95094; found 677.95555 [M-H]⁺.

KL⁵ – A solution of HL⁵ in THF (3 mL) (0.678 g, 1.00 mmol) was added to a suspension of KH (0.048 g, 1.20 mmol) in THF (3 mL) and the resulting mixture was heated at 60 °C for 24 h. Attempts to isolate KL⁵ resulted in its decomposition upon drying, therefore no isolated yield is reported. ¹H NMR (400 MHz, tetrahydrofuran-d₈): δ = 7.31 (d, J = 4.5 Hz, 2H, γ-CH), 5.93 (d, J
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\delta = 151.5 \text{ (2xC), 147.5 (dm, } J = 126 \text{ Hz), 142.9 (dm, } J = 126 \text{ Hz), 140.2 (dm, } J = 126 \text{ Hz), 139.1 (dm, } J = 126 \text{ Hz), 135.2 (dm, } J = 126 \text{ Hz), 132.0 (m), 120.3 (t, } J = 20.2 \text{ Hz), 114.3 (td, } J_1 = 15.1 \text{ Hz, } J_2 = 2.5 \text{ Hz), 110.1 (s), 105.9 (s), 77.9 (s). \]

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\delta = -139.6 \text{ (dd, } J = 25.8, 8.2 \text{ Hz), -146.1 (dt, } J = 22.8, 5.5 \text{ Hz), -163.3 (t, } J = 20.9 \text{ Hz), -166.0 (ddd, } J = 25.8, 20.9, 8.2 \text{ Hz), -167.5 (td, } J = 21.7, 5.3 \text{ Hz), -172.2 (tt, } J = 21.2, 5.4 \text{ Hz). UV-vis (THF): } \lambda_{\text{max}} (\varepsilon) 464 \text{ nm (9500 dm}^3\text{ mol}^{-1}\text{ cm}^{-1}), 636 \text{ nm (54400 dm}^3\text{ mol}^{-1}\text{ cm}^{-1}). \]

**LS•** - A solution of HL₈ (0.339 g, 0.50 mmol) in THF (3 mL) was added to a suspension of CuI (0.105 g, 0.55 mmol) and KH (0.024 g, 0.60 mmol) in THF (3 mL) and the resulting mixture was heated at 60 °C for 24 h, providing a yellow/brown solution. Filtration, solvent removal and washing with hexane provided LS• as a light yellow solid (0.220 g, 65%). UV-vis (THF): \( \lambda_{\text{max}} (\varepsilon) \) 211 (17500), 291 (24700), 456 nm (23000 dm³ mol⁻¹ cm⁻¹). Anal. Calcd. for [C₂₇H₄F₁₅S₂] C, 47.87; H, 0.60%; found: C, 48.01; H, 0.65%. HRMS (APPI): Calcd. for [C₂₇H₄F₁₅S₂], 676.95094; found 676.95084 [M⁺].

**(LO•)(KI)ₙ** - A solution of HL₀ (1.260 g, 5.00 mmol) in THF (5 mL) was added to a suspension of CuI (1.140 g, 5.00 mmol) and KH (0.401 g, 10.00 mmol) in THF (5 mL) and the resulting mixture was heated at 60 °C for 24 h, providing a dark brown solution. The crude mixture was cooled to −30 °C and filtered while cold. The filtrate was evaporated to dryness under vacuum and washed with hexanes to provide (LO•)(KI)ₙ as a dark brown solid (0.944 g, 75% based on LO•). Anal. Calcd. for LO• [C₁₇H₁₅O₂] C, 81.25; H, 6.02%; for (LO•)₂(KI) [C₁₇H₁₅O₂KI] C, 61.08; H, 4.52%; found: C, 64.32; H, 4.91%. HRMS (APPI) \( m/z \): Calcd. for [C₁₇H₁₅O₂], 251.10666; found 251.10743 [M⁺]. Calcd. for [C₃₄H₃₀O₄], 502.21386; found 502.21433 [M+H⁺].

**Bis(5,5′-formyl-2,2′-furan)phenylmethane, 2** – A reported procedure was adapted.⁹ Neat POCl₃ (10.0 mL, 107 mmol) was added at 0 °C to a solution of I (9.6 g, 42.9 mmol) in DMF (100 mL) providing a light brown mixture that was stirred at room temperature for 24 h during which the color changed to red/purple. The reaction was quenched by the addition to NaOAc(aq) at 0 °C followed by stirring for 1 h at room temperature. The crude product was extracted in CH₂Cl₂ (150 mL), washed sequentially with water (3 × 50 mL), HCl(aq) (3 × 50 mL) and NaHCO₃(aq) (3 × 50 mL) and dried over MgSO₄ to obtain a dark red oil. Purification by flash chromatography (hexanes:EtOAc 7:3, Rₓ = 0.4) provided 2 as a light brown oil (4.2 g, 35%). ^1H NMR (500 MHz,
chloroform-d): δ = 9.59 (s, 2H, CHO), 7.40-7.28 (m, 5H, o-CH, m-CH, p-CH), 7.23 (d, J = 3.6, 2H, β-CH), 6.40-6.31 (m, 2H, γ-CH), 5.66 (s, 1H, meso-CH). 13C{1H} NMR (126 MHz, chloroform-d): δ = 177.4 (s), 159.1 (s), 152.5 (s), 136.5 (s), 129.0 (s), 128.2 (s), 128.0 (s), 122.4 (s), 111.3 (s), 45.4 (s).

**HL**NO − Neat tert-butylamine (2.82 mL, 37.2 mmol) was added to a solution of 2 (1.88 g, 9.30 mmol) in toluene (60 mL) and 4 Å molecular sieves (3.0 g) and the resulting mixture was stirred for 36 h at room temperature. The resulting mixture was filtered through Celite and the solvent removed under vacuum, providing a dark brown oil which was dissolved in hexanes (30 mL) and filtered to remove undissolved impurities. Evaporation of the filtrate under vacuum provided HL**NO as a dark brown solid (3.23 g, 90%). 1H NMR (601 MHz, chloroform-d): δ = 8.02 (s, 2H, N=CH), 7.33-7.29 (m, 2H, m-CH), 7.28-7.24 (m, 1H, p-CH), 7.23-7.20 (m, 2H, o-CH), 6.72 (d, J = 3.4, 2H, β-CH), 6.10-6.05 (m, 2H, γ-CH), 5.65 (s, 1H, meso-CH), 1.26 (s, 18H, C3H18). 13C{1H} NMR (126 MHz, chloroform-d): δ = 156.5 (s), 152.5 (s), 145.3 (s), 138.8 (s), 128.8 (s), 128.6 (s), 127.5 (s), 113.6 (s), 110.4 (s), 57.7 (s), 45.3 (s), 29.8 (s). Anal. Calcd. for [C25H30N2O2]: C, 76.89; H, 7.74; N, 7.17%; found: C, 76.49; H, 7.52; N, 7.05%. UV-Vis (toluene): λmax (ε) 284 nm (22000 dm3 mol–1 cm–1). HRMS(ESI) (m/z): calcd for [C25H30N2O2]+, 391.2380; found 391.2356 [M]+.

**KL**NO − A dark brown solution of HL**NO (6.50 mmol, 2.5 g) in THF (15 mL) was added to a suspension of KH (7.80 mmol, 312 mg) in THF (15 mL) and the resulting dark green mixture was stirred at RT for 2 h and vented every 30 min. The mixture was filtered and the solvent evaporated under vacuum to provide KL**NO as a brown/gold solid (2.56 g, 92%). 1H NMR (500 MHz, tetrahydrofuran-d8): δ = 7.61 (s, 2H, N=CH), 7.36 (d, J = 7.9, 2H, o-CH), 7.23 (t, J = 7.5, 2H, m-CH), 7.09-7.04 (m, 1H, p-CH), 6.48 (d, J = 4.0, 2H, β-CH), 5.16 (d, J = 4.0, 2H, γ-CH), 1.27 (s, 18H, CH3). 13C{1H} NMR (126 MHz, tetrahydrofuran-d8): δ = 163.0 (s), 144.3 (s), 143.5 (s), 142.4 (s), 133.6 (s), 128.9 (s), 125.3 (s), 123.8 (s), 96.8 (s), 84.4 (s), 55.7 (s), 31.1 (s). Anal. Calcd. for [C25H30N2O2K]: C, 70.06; H, 6.82; N, 6.54%; found: C, 69.68; H, 6.94; N, 6.42%. UV-Vis (THF): λmax (ε) 269 (28900), 346 (24500), 453 (19400), 630 nm (23100 dm3 mol–1 cm–1).

**(L**NO•)(KI) − A dark green mixture of KL**NO (1.49 mmol, 640 mg) and CuI (1.47 mmol, 280 mg) in toluene (8 mL) was stirred at room temperature for 72 h, during which it turned dark brown. The mixture was filtered, and the toluene evaporated under vacuum to give a brown solid which was washed with hexanes (3 x 5 mL) providing (L**NO•)(KI) as a light brown solid (0.518 g, 64%).
HRMS (APPI) (m/z): calcd for [C$_{25}$H$_{29}$N$_2$O$_2$.], 389.22235; found 389.22181 [M$^+$]. Anal. Calcd. for L$^{\text{LO}}$ [C$_{25}$H$_{29}$N$_2$O$_2$] C, 77.0; H, 7.50; N, 7.19%; for (L$^{\text{LO}}$)(KI) [C$_{25}$H$_{29}$N$_2$O$_2$KI] C, 54.05; H, 5.26; N, 5.04%; found: C, 57.00; H, 4.93; N, 4.53%.

**RESULTS AND DISCUSSION**

**Synthesis of the precursors.** The synthesis of HL$^5$ involves a two-step procedure starting from 2-bromothiophene with an initial arylation protocol to access the intermediate 2-pentafluorophenylthiophene 1, followed by an acid-catalyzed condensation with pentafluorobenzaldehyde to provide the desired precursor as an off-white solid (Scheme 1). Similarly, compound HL$^0$ is obtained in a one-step procedure through condensation of benzaldehyde and the commercially available 2-methylfuran 3; all attempts to prepare pentafluorophenyl-substituted variants of HL$^0$ were unsuccessful. In contrast, the donor-expanded variant HL$^{\text{NO}}$ is obtained straightforwardly as a brown/orange solid in high yield through condensation of the dialdehyde intermediate 2 with tert-butylamine.
Scheme 1. Synthesis of the anionic hetero-diarylmethenes $KLS$, $KLN$O, and $KLO$ and their oxidation reactions to the corresponding radicals $LS^*$, $LNO^*$($KI)_n$, and $LO^*$($KI)_n$ (isolated yields are reported in brackets).

The reactions between $HL^S$ or $HL^NO$ and KH both display color changes that are the consequence of the full conjugation of the carbon framework that would be achieved through meso-H deprotonation; the thiophene anion $KLS$ has a dark blue color, whereas the iminofuran anion $KLN$O displays a dichroic dark green/red color in solution. Further evidence for the meso-selective deprotonation is provided in the $^1H$ NMR spectra of these anions, which show the disappearance of the peaks at 6.58 ppm for $HL^S$ and 5.59 ppm for $HL^NO$ along with concomitant shifts in the heterocyclic peaks (Figure 2).

Figure 2. Portion of the $^1H$ NMR spectra (tetrahydrofuran-$d_8$) for $HL^S/KLS$ (left) and $HL^NO/KLN$O (right).

Dithiophene $HL^S$. In the case of $HL^S$, the choice of pentafluorophenyl as the 5,5'-substituents proved important as alternative alkyl-, aryl- and fused arene-substituted analogs decomposed upon deprotonation with KH. Even so, while the clean formation of $KLS$ is seen in the $^1H$ NMR
spectrum, it remains highly reactive and slowly decomposes in either THF or pyridine solution to an NMR-silent, dark yellow solution, so precluding its isolation. Additionally, HL^S is sufficiently acidic to be deprotonated by the weaker silylamide base KN(SiMe3)2; however, the resulting blue KL^S solution is gradually oxidized to a paramagnetic species and at a faster rate than seen with KH. As such, a targeted synthesis of the paramagnetic compound was carried out by reacting HL^S with KH in the presence of stoichiometric CuI. As the reaction occurs the initial colorless solution turns gradually dark green, then light green and finally dark yellow as the NMR-silent product L^S\textsuperscript{•} is formed. The EPR spectrum of isolated L^S\textsuperscript{•} shows a resonance at g\textsubscript{iso} = 2.0036, consistent with an organic radical (Figure 3). The hyperfine structure is highly complex due to the coupling of the unpaired electron to all four protons from the thienyl moieties and all fifteen fluorine atoms (^19F, I = \frac{1}{2}, 100% abundant) of the pentafluorophenyl substituents.
**Figure 3.** X-band EPR spectrum of radical $L^5\$ in CH$_2$Cl$_2$ at 293 K (experimental conditions: frequency, 9.8562 GHz; power, 2.0 mW; modulation, 0.01 mT). Experimental data are represented by the black line with the simulation depicted by the red trace. $g_{iso} = 2.0036$; $A_{iso} = 3.71 \times 10^{-4}$ cm$^{-1}$ (2 $\times$ 1H); $A_{iso} = 3.05 \times 10^{-4}$ cm$^{-1}$ (2 $\times$ 1H); $A_{iso} = 1.71 \times 10^{-4}$ cm$^{-1}$ (3 $\times$ 19F); $A_{iso} = 1.16 \times 10^{-4}$ cm$^{-1}$ (6 $\times$ 19F); $A_{iso} = 0.47 \times 10^{-4}$ cm$^{-1}$ (2 $\times$ 19F); $A_{iso} = 0.34 \times 10^{-4}$ cm$^{-1}$ (4 $\times$ 19F).

Interestingly, the dithienyl radical $L^5\$ is a stable acyclic radical which, in contrast to its imine-expanded relative, possesses a less extensive $\pi$-system with consequent minimal stabilization by conjugation. In addition, it would be expected that the strongly electron-withdrawing pentafluorophenyl substituents would destabilize the radical in favor of the anion. Due to the redox activity displayed by this simple dithienyl compound, reactions between $KL^5$ and a variety of metal salts were attempted to access metal complexes of this potentially redox-active ligand. Unfortunately, these reactions result in ligand oxidation to form $L^5\$, even in the case of typically redox-inert metals like ZnCl$_2$ and MgCl$_2$. Consequently, direct deprotonation reactions were carried out with metal complexes provided with internal strong bases such as magnesium and zinc alkyls; however, even in these cases only the neutral radical species $L^5\$ is observed. These unsuccessful metalation reactions reinforce both the ease of oxidation of the dithienyl anion $KL^5$ and the poor coordinating properties of the endocyclic sulfur donor, in line with our previous studies.

**Difuran HL.$^0$.** Replacing the thiophene heterocycles with furan causes a dramatic change in stability, and thus reactivity, of the anionic species. While the reaction between the colorless precursor $HL.$ and KH in THF results in a color change to dark brown, the $^1$H NMR spectrum of the reaction mixture is silent and paramagnetic according to qualitative Evans’ method measurements; a similarly silent $^1$H NMR spectrum is seen for the reaction between $HL.$ and...
KN(SiMe$_3$)$_2$. As such, the tendency of the expected anion KL$^O$ to oxidize spontaneously to the radical L$^{O*}$ is more pronounced than for KL$^S$ and the anionic species KL$^O$ is not even observed. The radical nature of L$^{O*}$ is corroborated by its EPR spectrum which is consistent with an organic radical with $g_{iso} = 2.0043$, albeit with unresolved hyperfine structure (Figure 4). In a previous report, L$^{O*}$ was prepared, but not isolated, either by oxidation of HL$^O$ to form the difurylmethene cation, followed by single-electron reduction by Zn mirror, or by a deprotonation/oxidation protocol similar to that used here.$^{24}$ While EPR data were provided for the difuryl complexes in which the meso-substituent is either methylfuryl, ethylfuryl, C$_6$H$_4$(OH-4), C$_6$H$_4$(Bu-3)$_2$(OH-4), or C$_6$H$_4$(OMe-3) and showed extensive hyperfine structure to the $^1$H nuclei, no data were provided for L$^{O*}$. In our case, we were unable to isolate L$^{O*}$ as an analytically pure compound due to non-stoichiometric incorporation of KI and, as such, the purity of L$^{O*}$ remains uncertain, with the lack of hyperfine structure seen in the EPR spectrum of L$^{O*}$ likely a consequence of its coordination to KI. However, the APPI-MS of L$^{O*}$ provides further support for its identity, showing ions at $m/z$ 251 and 252 consistent with the cations [L$^O$]$^+$ and [L$^{O+H}$]$^+$ formed on ionization.
Figure 4. X-band EPR spectrum of radical L^0• in THF at 293 K (experimental conditions: frequency, 9.6709 GHz; power, 10 mW; modulation, 0.1 mT). Experimental data are represented by the black line; simulation is depicted by the red trace composed: $g_{\text{iso}} = 2.0043$; $A_{\text{iso}} = 2.2 \times 10^{-4}$ cm$^{-1}$ (2 × 1H); $A_{\text{iso}} = 1.6 \times 10^{-4}$ cm$^{-1}$ (2 × 1H); $A_{\text{iso}} = 1.5 \times 10^{-4}$ cm$^{-1}$ (2 × 1H); $A_{\text{iso}} = 1.1 \times 10^{-4}$ cm$^{-1}$ (1 × 1H).

Iminofuran HL^NO. In contrast to the spontaneous oxidation chemistry displayed by the anions KL^S and KL^O, the iminofuran KL^NO is straightforwardly isolated as a translucent dark green powder in high yield. Noticeably, KL^NO can be stored under an inert atmosphere in a glove box for months without any appreciable decomposition. Red crystals of KL^NO suitable for crystallographic analysis were obtained by layering hexanes on a concentrated THF solution. Interestingly, in the solid state, KL^NO exists as a 1D coordination polymer in which each unit is
linked to the next through $\pi$-coordination of a furan group to the potassium, with no incorporation of the THF donor solvent (Figure 5).

**Figure 5.** Solid-state structure of the iminofuran anion $\text{KL}^{\text{NO}}$. For clarity, all hydrogen atoms and a second molecule in the unit cell are omitted (displacement ellipsoids are drawn at 50% probability). Right: $\pi$-coordination of the furan moiety from the neighboring unit to the $\text{K}$ cation.

Selected bonds (Å) and angles (°): $\text{K1}–\text{O1}$ 2.740(2), $\text{K1}–\text{O2}$ 2.751(2), $\text{K1}–\text{N1}$ 2.773(3), $\text{K1}–\text{N2}$ 2.748(3), $\text{N1}–\text{C5}$ 1.294(4), $\text{N2}–\text{C21}$ 1.280(5), $\text{O1}–\text{C6}$ 1.400(4), $\text{O1}–\text{C9}$ 1.392(4), $\text{C9}–\text{C10}$ 1.386(5), $\text{C10}–\text{C17}$ 1.418(4), $\text{O2}–\text{C17}$ 1.386(4), $\text{O2}–\text{C20}$ 1.396(4); $\text{C9}–\text{C10}–\text{C17}$ 127.4(3), $\text{O1}–\text{K1–O2}$ 60.31(7), $\text{O1}–\text{K1–N1}$ 62.56(8), $\text{O2}–\text{K1–N2}$ 62.40(8), $\text{N1}–\text{K1–N2}$ 120.15(9).

In the structure, the *meso* carbon $\text{C17}$ is $\text{sp}^2$ hybridized with essentially coplanar iminofuran linkages (torsion angle 8.7°). In contrast to the iminothiophene relative $\text{KL}^{\text{NS}}$ which adopts a dimeric structure, the $\text{N}_2\text{O}_2$ compartment in $\text{KL}^{\text{NO}}$ binds a single potassium cation due to the shorter $\text{N1}–\cdots\text{N2}$ separation between the imine nitrogen atoms of 4.785(4) Å (compared with 7.357(1) Å in $\text{KL}^{\text{NS}}$). Such a feature is also likely responsible for the positioning of the $\text{K}$ cation.
above the N₂O₂ plane (N₂O₂ plane···K1 1.222 Å), although influence by the adjacent π-coordinated furan cannot be excluded (centroid···K1 3.029 Å).

In a similar manner to the related thiophene anion KL₅, reaction between KL̄NO and CuI leads to the toluene-soluble, neutral radical L̄NO• as a brown/orange solid in good yield, and isolated as the salt-incorporated adduct KI(L̄NO•). The radical nature of L̄NO• is supported by the lack of features in its NMR spectrum and is confirmed by its fluid-solution EPR spectrum which shows a resonance at g_{iso} = 2.0035, consistent with an organic radical. A complex hyperfine structure arises from the coupling of the unpaired electron with six hydrogen and two nitrogen (¹⁴N, I = 1, 99.7% abundant) nuclei from the iminofuryl backbone and three additional hydrogen nuclei from the meso-phenyl group (Figure 6).

Figure 6. Fluid-solution X-band EPR spectrum of radical L̄NO• in THF at 293 K. (experimental conditions: frequency, 9.8554 GHz; power, 6.3 mW; modulation, 0.01 mT). Experimental data are represented by the black line; simulation is depicted by the red trace composed: g_{iso} = 2.0035; A_{iso}
= 3.54 × 10^{-4} \text{ cm}^{-1} (2 \times ^1\text{H}); A_{\text{iso}} = 3.22 \times 10^{-4} \text{ cm}^{-1} (2 \times ^1\text{H}); A_{\text{iso}} = 1.60 \times 10^{-4} \text{ cm}^{-1} (3 \times ^1\text{H}); A_{\text{iso}} = 1.57 \times 10^{-4} \text{ cm}^{-1} (2 \times ^1\text{H}); A_{\text{iso}} = 0.53 \times 10^{-4} \text{ cm}^{-1} (2 \times ^{14}\text{N}).

**Electronic spectroscopy and TD-DFT analysis.** Upon deprotonation of the precursor HL^5 to the corresponding potassium salt KL^5, two bands at 636 (HOMO-LUMO) and 465 nm arise in the UV-Vis spectrum (Figure 7). Upon oxidation of KL^5 into L^5\textsuperscript{•} a weak band at 455 nm is seen, similar in shape to that of KL^5, and due to combined $\alpha/\beta$ SOMO-LUMO transitions. As in the related thiophene compounds,\textsuperscript{38,47} the oxidation of the anion to the radical is accompanied by a hypsochromic shift which is indicative of a larger orbital gap in the radical of 5.50 eV ($\alpha$-spin) and 5.31 eV ($\beta$-spin) against 4.24 eV for the anion. The electronic absorption spectra of the furan-containing compounds are different, in which the HOMO-LUMO transition observed at 284 nm for HL\textsuperscript{NO} (Figure S11) is modified to multiple bands on deprotonation to KL\textsuperscript{NO}, with the low-energy band at 630 nm associated with the HOMO-LUMO transition (Figure 7). Compared to the thiophene-based diarylmethenes, oxidation of KL\textsuperscript{NO} to L\textsuperscript{NO\textsuperscript{•}} does not cause major variations in the electronic spectrum and preserves the slightly red-shifted, four-band motif with the low-energy absorption at 632 nm due to combined $\alpha/\beta$ SOMO-LUMO transitions. The electronic spectrum of the difuran radical L\textsuperscript{O\textsuperscript{•}} appears mostly unresolved (Figure S12), with the absorption bands broadened into a large absorption curve and likely caused by the ability of this radical to coordinate non-stoichiometric quantities of KI (see above); precipitation of a colorless solid, presumably KI, is seen during data acquisition.
Electrochemical studies. Analysis of the cyclic voltammograms (CVs) of the compounds provides additional insight into their electronic properties. The precursor \( \text{HL}^8 \) is electrochemically inert over a potential window spanning from +1.5 V to −2.0 V versus \( \text{Fc}^{+}/\text{0} \). Due to the difficulty in isolating the anion \( \text{KL}^5 \), the electrochemical investigation of \( \text{L}^{5\text{r}} \) is therefore crucial to obtain information on the redox properties of this system. The voltammogram for \( \text{L}^{5\text{r}} \) displays two redox waves, an oxidation at +0.27 V and a reduction and −1.09 V (Figure 8), which are best interpreted as forming the diamagnetic cation \( \text{L}^{5\text{c}} \) upon oxidation, while the reduction event most likely refers
to the $L^+/L^-$ couple. No meaningful CV data were acquired for $L^{O-}$ and $L^{O-}$ due to the difficulties encountered in their isolation.

The CV for $HL^{NO}$ comprises three events, an oxidation at $-0.11 \text{ V}$ and an irreversible oxidation at $-0.48 \text{ V}$ associated with an irreversible reduction at $-1.89 \text{ V}$ (Figure S14). On deprotonation to form $KL^{NO}$, these waves are replaced by a quasi-reversible process at $-1.28 \text{ V}$, assigned to the anion-radical redox couple, $KL^{NO}/L^{NO*}$ (Figure 8). Unexpectedly, the CV of isolated $L^{NO*}(\text{KI})_n$ obtained by chemical oxidation of $KL^{NO}$ with CuI does not map on to the wave observed for the anion but instead comprises two oxidations, one irreversible at $+0.20 \text{ V}$ and one quasi-reversible at $+0.53 \text{ V}$, with no trace of the reduction event allied to $KL^{NO}$. It is therefore evident that the radical formed by chemical oxidation of $KL^{NO}$ by CuI or I$_2$ has a different composition to that formed by electrochemical oxidation in a non-coordinating electrolyte. As with $L^{O*}(\text{KI})_n$, the unusual electrochemical features of $L^{NO*}(\text{KI})_n$ could derive from the observation by elemental analysis that one equivalent of KI is incorporated and might therefore alter the redox characteristics of the radical, hampering its reduction to the anion. This is reinforced by the CV of $L^{NO*}$ that is generated in the CV cell by the addition of Cu$^+$(MeCN)$_4$BF$_4$ to $KL^{NO}$ which shows the expected $KL^{NO}/L^{NO*}$ redox couple at $-1.16 \text{ V}$. The single oxidation seen at $0.13 \text{ V}$ is similar to that seen for $L^{NO*}(\text{KI})_n$ at $+0.20 \text{ V}$, suggesting that this is a ligand-based oxidation, i.e. $L^+/L^{NO*}$ and that the second oxidation at $+0.53 \text{ V}$ relates to the $I_2/I^-$ couple. We currently do not fully understand the effect of KI coordination on the redox chemistry of $L^{NO}$ but are exploring this through the synthesis of further examples of complexes of $L^{NO}$ with redox-inert metals.
Figure 8. Comparison of CV data for L^S• (orange), KL^NO (green), L^{NO•} (blue) and L^{NO•}(KI)_n (red); glassy carbon working electrode, platinum gauze counter electrode, silver wire pseudo-reference electrode, 100 mV s\(^{-1}\), referenced to Fc\(^{+/-}\), 1-5 mM analyte, 0.1 M \([\text{Bu}_4\text{N}][\text{PF}_6]\) electrolyte in dry CH\(_2\)Cl\(_2\) under \(\text{N}_2\).

Compared to the parent iminothiophene compound KL^NS, the diarylmethene anions KL^NO and KL^S investigated here display significantly more facile oxidation chemistry, evident from the highly negative potentials at which the waves related to the formation of the radical species are found. These potentials, at \(-1.09\) V and \(-1.28\) V for KL^S and KL^NO, respectively, are indeed over 1.0 V more negative than that seen for KL^NS at \(-0.12\) V and highlight the ease of formation of the
radicals. While the oxidation of the anion KL\textsuperscript{NO} is more thermodynamically facile than for KL\textsuperscript{5}, the poorer coordinating ability of the dithienyl donor set to the potassium cation compared to the N\textsubscript{2}O\textsubscript{2} donor set likely limits the kinetic stability of KL\textsuperscript{5} and so inhibits its isolation.

**CONCLUSIONS**

The hetero-diarylmethene compounds KL\textsuperscript{5}, KL\textsuperscript{O}, and KL\textsuperscript{NO} have been synthesized and studied by means of spectroscopic, crystallographic, and electrochemical techniques. These conjugated carbanions display a pronounced redox activity and provide stable neutral acyclic radicals upon oxidation under mild conditions. Significantly, the radical L\textsuperscript{S\textsuperscript{r}} was isolated despite a reduction in molecular conjugation compared to L\textsuperscript{N\textsuperscript{S\textsuperscript{r}}} and the incorporation of strongly electron-withdrawing pentafluorophenyl substituents, providing a rare example of a non-macrocyclic thiophene-based radical.\textsuperscript{38,48} Spectroscopic investigation highlighted a shift of the main absorption bands as a consequence of the oxidation of the anion to the corresponding radical, and provided evidence of an increased SOMO-LUMO energy gap in the thiophene-derived compound. Electrochemical analysis allowed the identification of the strongly negative potentials at which the oxidation reactions occur, confirming the tendency of these anions to be spontaneously oxidized to the radicals. These results emphasize how the extensive redox activity of diarylmethene carbanions can be controlled by varying the nature of the substituents, donor-atom, and the conjugated \pi-system, and that these variations impact considerably on the coordinating properties of these species; the N\textsubscript{2}O\textsubscript{2} donor set of the iminofuran anion allows the isolation and crystallographic characterization of the anion KL\textsuperscript{NO} and promotes KI incorporation in the radical L\textsuperscript{NO\textsuperscript{r}}. We are currently exploiting these facets in the development of reagents and catalysts for the reduction of small molecules such as carbon dioxide.
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The Supporting Information is available free of charge on the ACS Publications website at http://pubs.acs.org. NMR and UV-vis spectra, electrochemical, crystallographic, and computational data.

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REFERENCES


Difuryl- and dithienymethene anions are potential chelating ligands for metals, and are shown to exhibit marked redox activity, undergoing facile one-electron oxidation to form stable, neutral radicals. The stability of the anions and the ease of formation of the neutral radicals are related to the extent of conjugation, the substituent pattern, and the nature of the heteroatom.