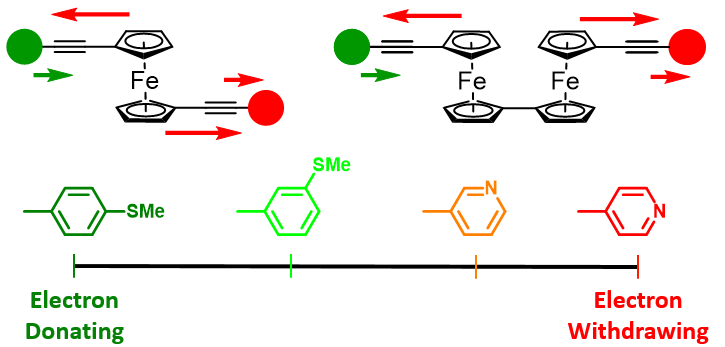
Synthesis, Electrochemistry and Optical Properties of Highly Conjugated Alkynyl-Ferrocenes and -Biferrocenes

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ABSTRACT: Sonogashira reactions are utilized herein to react iodo-ferrocenes and -biferrocenes with terminal alkyne ligands, functionalized with both pyridine and thioanisole groups. High yielding reactions generate both mono-alkynyl and di-alkynyl derivatives, the ratio of which can be altered through changes in the reaction stoichiometry. This methodology allowed us to synthesise a large family of derivatives, comprising four symmetrical derivatives (3**xx,** where **x** represents a phenyl substituted terminal alkynes) and six less-studied asymmetrical derivatives (3**xy**, where **x** and **y** represent two different phenyl substituted terminal alkynes), as well as a number of their biferrocenyl analogues (**6x**, **7xx** and **7xy**), including the first known examples of asymmetrically di-substituted biferrocenes. We examined the electrochemical behavior of all the systems in solution through the use of cyclic voltammetry and demonstrate that these highly conjugated alkynyl ligands impart delicate redox control over the central ferrocene motif. We also note that these substituents display some control over the mixed-valence character present in biferrocene mono-cations, with thioanisole substituents imparting almost an order of magnitude higher *K*com than their pyridyl analogues, and asymmetric systems displaying rare characteristic properties of mixed valence isomers. The electronic structure of these systems was further elucidated through a combination of UV/Vis spectroscopy and DFT calculations. Our methodology provides a facile and adaptable route towards the isolation of a number of novel ferrocene and biferrocene derivatives. From our perspective, the asymmetric nature of these systems, along with the delicate and predictable redox control that this imparts on the central ferrocene unit(s) could lead to applications in molecular electronics, where these properties have previously shown promise in the fabrication of diodes and rectifiers, as well as in the synthesis of donor-π-acceptor systems.

Introduction

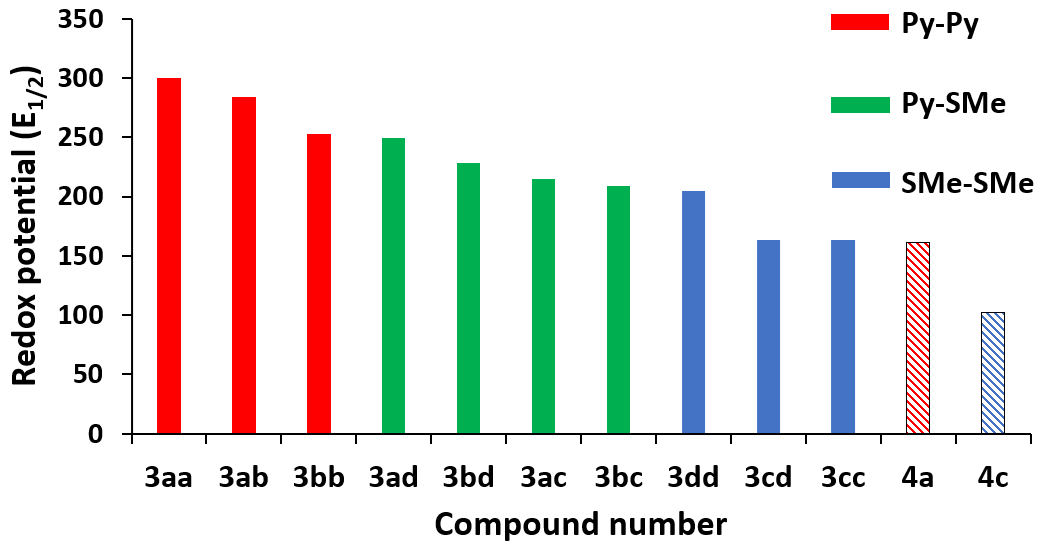
The desire to introduce a well-defined redox centre, such as ferrocene, into a wide-ranging array of molecules, has led to the development of a rich library of new synthetic methods.1,2 This abundance of techniques has resulted in ferrocene derivatives being incorporated into a wide variety of fields including catalysis, materials science, medicine and a large number of different electrochemical applications.3–9 One relatively recent addition to this synthetic tool-kit was the synthesis and isolation of 1,1’-diiodoferrocene in high yields, which has been demonstrated within our group, through the use of an ‘oxidative purification’ as well as by Nijhuis *et al.* whom utilized a sublimation focused route.10,11 Both of these methods generate the highly versatile synthon and its biferrocene equivalent (1,1’’’-diiodobiferrocene) in high yields and with a high degree of purity. The realization of these synthons has opened the door to the use of classical cross-coupling methodologies. In particular, Sonogashira cross-couplings of iodoferrocenes present a valuable opportunity for the synthesis of highly-conjugated redox active molecules through reactions with terminal alkynes. Through substitution of alkynyl moieties onto a ferrocene center we can create a robust, highly conductive structure with the ability to rotate around a central node. These attractive properties have led to the application of these systems in the fabrication of highly-ordered macromolecular assemblies and more prominently in the design and fabrication of electronic materials and single-molecule wires.12–20 However, to date, most studies of ferrocene in these contexts have focused on the synthesis and study of small families of monosubstituted and symmetrically-disubstituted systems, potentially owing to the relative ease of their synthesis and purification.21–28 Although impressive, these studies have often been limited in scope and ignore the potential that asymmetry could impart on the electronic and structural properties of these species. For instance, research in the field of molecular electronics has provided a number of examples where asymmetric design has been applied in the fabrication of systems for molecular rectification and has also been

demonstrated as an effective tool for tailoring different ends of a molecule to different electrode materials.29–33 In an effort to bridge this gap between synthesis and design we designed an adaptable synthetic route that could be applied in the isolation of a large number of new asymmetric ferrocenyl derivatives. For this we selected four different ‘ligands’ with subtly different chemical, and by extension, electronic structures, these being 4-ethynylpyridine (**a**), 3-ethynylpyridine (**b**), 4-ethynylthioanisole (**c**), and 3-ethynylthioanisole (**d**). Herein we demonstrate a facile and scalable synthesis of a series of novel symmetric and asymmetric alkynyl-ferrocenes and –biferrocenes. We examined the properties of these systems through a combination of electrochemistry, UV/Vis and computational methods, demonstrating a fine, predictable redox tunability within the family, as well as uncovering rare examples of mixed valence isomers.

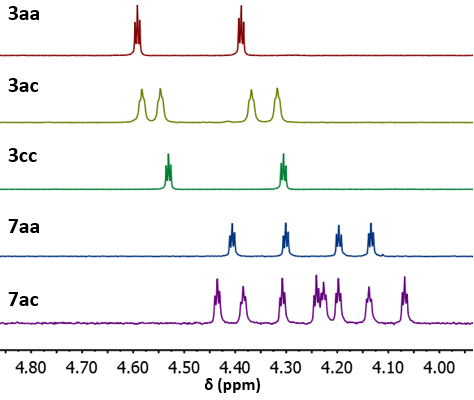
***Scheme 1.*** *Synthetic pathway used in the assembly of 10 ferrocene compounds, disubstituted in each case with two ethynyl-arene substitutents. ‘****Step 1****’ denotes the formation of the ethynyl-arene ligands, ‘****Step 2****’ denotes the initial reactions with 1,1’-diiodoferrocene and 1,1’’’-diiodobiferrocene to form a statistical mixture of mono- and di-substituted ferrocenes and ‘****Step 3****’ shows subsequent reactions that were utilised to introduce asymmetry across the central ferrocenyl units.*

**Synthesis**

Synthesis of each compound was achieved via an iterative Sonogashira coupling process, using a methodology that has previously found success in our group.34 The arylbromide (ArBr, where Ar = 4-pyridyl (**a**), 3-pyridyl (**b**), 4-thioanisole (**c**) or 3-thioanisole (**d**)) was coupled to trimethylsilylacetylene and subsequently deprotected to yield the terminal alkynes in high yields. From here, it was possible to couple the functionalised alkynes to 1,1’-diiodoferrocene (**1**). In this step (Scheme 1, step 2) access to both the monosubstituted products (**2x**) and the symmetrically disubstituted products (**3xx**) can be achieved through careful control of reaction stoichiometry. To preferentially obtain the monosubstituted products (**2x**), an excess of diiodoferrocenes must be employed (this is easily recovered after the reaction through column chromatography) and the converse is true when aiming for the symmetrical doubly-substituted products (**3xx**). Compounds **2x** can then be taken forward in an additional coupling step to obtain the asymmetric disubstituted products (e.g. **3xy**). Conveniently, multiple derivatives can be obtained in a one-pot reaction of a monosubstituted ferrocene (**2x**) with two different alkynes, although this leads to a statistical mixture of products, as is seen in the synthesis of **3bc** and **3cd** (see ESI). Purification is largely facile, however the pyridyl-based analogues require separation on a short alumina(V) column due to their propensity to streak on silica. For this reason, it was beneficial to prepare the asymmetric variants in a stepwise manner, starting from the thioanisole precursors (**2c** or **2d**) in order to maximise yield and ease of purification. For comparative purposes, the ferrocenyl derivatives containing only one arm (i.e. CpFe(C5H4CCAr) (**4**) where Ar = 4-pyridyl or 4-thioanisole) were synthesised from ethynyl-ferrocene according to Scheme 1, Step 2 to generate **4a** and **4c** respectively. This methodology was extended to the synthesis of a select number of biferrocenyl derivatives through reaction of 1,1’’’-diiodobiferrocene (**5**) with both (**a**)and (**c**). This once again produced a mixture of mono- (**6x**) and di-substituted (**7xx**) products, which could be isolated through the use of column chromatography. The yields of the biferrocene reactions were somewhat reduced in comparison to their monoferrocene counterparts which we attribute to decreased reactivity of the biferrocenyl centres. Molecules (**6x**) were then substituted with different alkynes to produce asymmetric biferrocene derivatives (**7xy**), which would have been difficult to isolate through currently available biferrocene functionalization methods.34 All compounds have been characterized by mass spectrometry and the bulk purity was determined through elemental analysis where possible. This synthesis was aided throughout by excellent resolution within the 1H NMR spectra, whereby all compounds show the expected number of pseudo-triplet signals associated with the ferrocene Cp protons at 4.0-4.8 ppm. For the symmetrical mono-ferrocene species (**3aa-dd**), these present as two pseudo-triplets, whereas four distinct pseudo-triplets are observed for the asymmetric variants (e.g. **3ac**). When extending this analysis to the biferrocene family, once again very distinct profiles can be observed in the 1H NMR spectra with the symmetrical biferrocenes producing 4 pseudo-triplets and the asymmetric systems producing 8 (Figure 1).



***Figure 2****: The redox potentials (mV) of compounds* ***3xx*** *(solid) and* ***4x*** *(dashed).*

***Figure 1.*** *Comparative**1H-NMR spectra of a number of different compounds, illustrating the various Cp-H environments that arise from both asymmetry and the inclusion of multiple ferrocene units.*

***Table 1:*** *The CV data obtained for complexes* ***3xx*** *and* ***4x****, in solution (0.1 M NnBu4PF6 in CH2Cl2 taken at 100 mVs-1 with a glassy carbon working electrode and Pt wire for reference and counter. All potentials (mV) are reported vs Fc/Fc+ and have been corrected for iRs with values obtained from AC impedance measurements).*

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| [Fc] | Ep | Ec | E1/2 | ΔE | ipa/ipc |
| 3aa | 336 | 264 | 300 | 72 | 1.23 |
| 3ab | 319 | 249 | 284 | 70 | 1.06 |
| 3ac | 247 | 182 | 215 | 65 | 1.08 |
| 3ad | 284 | 213 | 249 | 71 | 1.11 |
| 3bb | 296 | 209 | 253 | 87 | 1.05 |
| 3bc | 246 | 172 | 209 | 74 | 1.02 |
| 3bd | 268 | 188 | 228 | 80 | 0.89 |
| 3cc | 196 | 130 | 163 | 66 | 0.93 |
| 3cd | 196 | 130 | 163 | 66 | 0.86 |
| 3dd | 237 | 173 | 205 | 64 | 1.00 |
| 4a | 195 | 127 | 161 | 68 | 0.98 |
| 4c | 135 | 69 | 102 | 66 | 0.94 |

Electrochemistry

Cyclic voltammetry experiments were performed on all molecules in dichloromethane solutions with 0.1 M NnBu4PF6 as the supporting electrolyte, and the relevant data are reported in Table 1 (monoferrocenes) and Table 2 (biferrocenes). Looking first at the monoferrocene compounds, these all displayed a characteristic one-electron redox process, associated with the ferrocene motif, at potentials which are more positive than that of standard ferrocene.35,36 This is a consequence of the ethynylarene substituents, which withdraw electron density from the redox-centre by extending the π-system; a fact that is nicely evidenced by comparing the potentials of, for example, **3cc** and **4c**. Additionally, the substituents on the arene were found to exert a significant influence on the overall potential and reversibility of the redox process. The molecules that incorporate only thioanisole groups (**Xc/d**) undergo redox chemistry that is both chemically (ipa/ipc ≈ 1) and electrochemically reversible (ΔE ≈ 65 mV over multiple scan rates), with diffusion-controlled electron transfer rates (ip ∝ ν1/2).37 Changes in redox potential can be observed between the 3- (**3dd**) and 4-substituted (**3cc**) thioanisole compounds, with 3-thioanisoles occurring at a higher potential than their 4-substitued analogues. A rationale for this stems from the ability of the thiomethyl moiety to donate electrons into the π-system and stabilise a positive charge. A qualitative analysis of the relevant resonance forms indicates that the positive charge is delocalized more effectively in the presence of the 4-thioanisole than the 3-thioanisole, thus reflecting the trend observed in the data. A similar, yet opposite trend is observed for the series containing only pyridyl groups (**3a/b**). The 3-pyridyl moiety is able to stabilise the positive charge to a greater degree than the corresponding 4-pyridyls, with the analogue containing both groups (**3ab**) displaying an intermediate potential. The trend in the redox potentials of this series reflects those in both the pyridyl- and thioanisole-based analogues, such that a continuum of potentials can be observed across the whole series of molecules (Figure 2). Thus, while the net effect of the arylalkynes is electron withdrawing, the redox properties of the ferrocene can be tuned further by varying the electron donating/withdrawing properties of the arene. Turning now to examine the biferrocene systems **6x** and **7xx/xy.** All show two distinct redox processes and we attribute these redox features to the sequential oxidation of the two Fe2+/Fe3+ couples present in each system.21 Compounds **7aa** and **7cc** are fully symmetrical systems so the separation between the two redox processes (ΔE1/2) can be attributed to charge stabilisation via electronic coupling between the two ferrocene centres in the mixed valence ion (e.g. [FeII-FeIII]+). The stability of this mixed valence ion can be inferred from the comproportionation constant (KC) for the equilibrium between the mixed valence ion, and its neutral and doubly-oxidised forms, and can be derived from the Nernst equation, according to equation 1.

Using the potentials obtained from the CV of **7aa** and **7cc**, we can determine Kc as 1.3x105 and 9.3x105 respectively, suggesting that they belong in class III of the Robin and Day series. For compounds **6x**, **7ab** and **7ac**, these calculations become more complicated due to the asymmetric structure of the molecules. The monoferrocenes described previously (**3xx/xy** and **4x**) clearly show that the redox properties of an individual ferrocene centre can be influenced by substituents on the Cp ring. Therefore, it is incorrect to consider these biferrocene units as simple mixed valence systems, rather they are rare examples of mixed valence isomers.38 In such systems, ΔE1/2 is no longer purely a consequence of coupling between redox centres, but is also affected by the intrinsic potential difference between the two independent redox sites.39 A consequence of this is that ΔE1/2 values obtained from the CV of mixed valence isomers can provide an overestimation of the electronic coupling present within the system.. To illustrate this, the redox profile of biferrocene **7ac** can be compared to that of monoferrocenes **4a** and **4c**. **4a** is oxidised at 161 mV while **4c** is oxidised at 102 mV, which gives ΔE4a/4c = 59 mV. ΔE1/2 of **7ac** is 370 mV and approximately 59 mV of that must come from the intrinsic potential difference between the two independent ferrocene centres. This means that only around 311 mV (= 370-59) can be attributed to stabilisation via electronic coupling. Gratifyingly, this value of 311 mV is intermediate between those of the analogous symmetrical biferrocenes **7aa** (303 mV) and **7cc** (353 mV). While care must be taken to compare like-for-like systems, 34 Kc is often used as an indicator of coupling strength in mixed valence systems. However, due to the redox asymmetry present in **6x**, **7ab** and **7ac**, Kc cannot be employed in this way. A more in-depth investigation into the electronics of these mixed valence isomers is currently underway in our laboratory.

***Table 2:*** *The CV data obtained for complexes* ***6x*** *and* ***7xx****, in solution (0.1 M NnBu4PF6 in CH2Cl2 taken at 100 mVs-1 with a glassy carbon working electrode and Pt wire for reference and counter. All potentials (mV) are reported vs Fc/Fc+ and have been corrected for iRs with values obtained from AC impedance measurements).*

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| [Fc2] | Ep | Ec | E1/2 | ΔE | ipa/ipc | ΔE1E2 | *Kcom* |
| 6a | 64 415 | -23 279 | 44 347 | 87  136 | 1.15 | 303 | 105.12 |
| 6c | 13  378 | -71 295 | -29 337 | 84 83 | 0.95 | 366 | 106.19 |
| 7aa\* | 3 322 | -57 230 | -27 276 | 60  92 | 0.51 | 303 | 105.12 |
| 7ab\* | 4 347 | -66 241 | -31 294 | 70 106 | 0.62 | 325 | 105.50 |
| 7ac | 49 391 | -56 287 | -34 336 | 105 109 | 1.04 | 370 | 106.26 |
| 7cc | -21 330 | -103 253 | -62 291 | 82 78 | 0.95 | 353 | 105.97 |

**INVESTIGATING ELECTRONIC STRUCTURE**

To investigate the electronic structure of this series, a combination of theoretical calculations and UV/Vis spectroscopy were employed. For this, the most stable ground-state geometry of each compound was determined by DFT calculations in the gas phase, using the B3LYP functional and the 6-311+G(d,p) basis set. In addition, TD-DFT calculations, used to model the UV/Vis spectra, employed the aforementioned optimised geometries and were performed using a dichloromethane conductor-like solvent polarisation continuum model (CPCM) with the PBE1 functional and 6-31+G(d,p) basis set to generate results that could be compared to experimental spectra obtained in the same solvent (see ESI). A few initial conclusions could be drawn from the experimental UV/Vis spectra. For instance, it has previously been reported that unsubstituted ferrocene displays a transition at approximately 442 nm, which relates to two overlapping spin-allowed, Laporte forbidden transitions.40 We see that, in all cases, our introduction of alkynyl ligands causes a bathochromic shift of this peak to higher wavelengths (between 445-460 nm), coinciding with previous work in the literature which suggests that inclusion of an alkyne substituent acts to decrease the HOMO-LUMO gap.41–43 This is clearly demonstrated when comparing molecules **4a** and **4c**, where the more electron withdrawing pyridyl causes a larger shift, though the trend becomes far less distinct upon introduction of a second alkynyl substituent. Trends relating to other ferrocene-associated transitions are harder to quantify as these are often obscured by large intense absorptions that occur between 250-325 nm. Analysis of the frontier molecular orbital (FMO) energies and their iso-surfaces (as determined by DFT calculations, alongside predicted absorption spectra (as modeled by TD-DFT calculations) provided further useful insights into the nature of these optical transitions. Our calculations suggest that the lower energy signals that appear around 450 nm are primarily MLCT in character while those around 350 nm result from transitions between orbitals that are more delocalized across the whole molecule, and are essentially π🡪π\* in character. The absorption bands observed at lower wavelengths can be assigned to higher-energy π🡪π\* transitions, though it should be noted that those molecules containing two different arylalkyne substituents display a significant degree of inter-ligand charge transfer (ILCT) that also contributes to the overall signals below 350 nm. In most cases, the HOMO and LUMO are somewhat delocalised across the whole molecule, although in pyridyl-based analogues there is a clear bias towards a ferrocene-based HOMO and a LUMO situated on the ethynylpyridine. In addition to this, the FMOs of those systems containing both a pyridine and a thioanisole substituent seem to show particular favorability for the HOMO to be localized on the thioanisole fragment and the LUMO to be localized on the pyridyl fragment as is particularly evident in the FMO’s of **3ac** and **7ac** (Figure 3). When considering the biferrocene systems, the UV/Vis analysis demonstrates that these show similar transitions to their monoferrocene analogues, with the primary difference being a general increase in their intensity. The TD-DFT data suggests that the majority of these transitions are dominated by complex MLCT processes (see for example Table S14). Finally, it is also worth noting that an analysis of the frontier molecular orbital energies of these systems generated a trend in the energy of the HOMOs which mirrored that which was observed in the redox potentials of our systems as reported above (Figure S83), suggesting that a predictable redox tunability exists within this family. Energy level diagrams, iso-surface depictions of the FMOs and a list of predicted transitions, with their full assignments, for each compound, are provided in the supporting information.

Conclusion

Herein we have shown a facile synthesis of a large library of novel, highly conjugated, alkynyl-substituted ferrocene and biferrocene derivatives that have been characterised in full. Cyclic voltammetry was used to examine the electronic structure of these systems and it was shown that the differing substitution patterns impart fine redox control over the central ferrocene unit(s). The biferrocene molecules display subtly different mixed valence character, with **7ac**, and likely **7ab**, displaying the characteristics of rare mixed valence isomers. A combination of UV/Vis spectroscopy and theoretical calculations, derived from DFT, were used to further elucidate the electronic structure of these compounds, providing clear justification for the redox tuning as a function of HOMO energy, occurring in a highly-predictable manner. This analysis also allowed us to analyse the optical transitions observed for these compounds, noting a variety of complex MLCT processes, with those asymmetric systems containing both pyridyl and thioanisole fragments displaying more IVCT characteristics than their symmetrically substituted counterparts. Broadly speaking, this work demonstrates the ease and practicality of introducing asymmetry across a redox-centre and how this can be used to subtly alter said center’s electronic structure. Asymmetric molecules have previously shown promise in the field of molecular electronics in devising new systems for diodes, rectifiers and in the tuning of optical band gaps.44–51 The stepwise nature of our synthesis provides a routine method of incorporating an organometallic redox-active fragment into these designs and also provides an opportunity to study how their properties can be changed by externally controlling the ferrocene redox processes through the use of electrochemical gating.52–55 In addition, this fine control of electronic structure provides a further parameter for the predictable tuning of FMO energies, which has large implications in both electronic and thermal conductivity as both are highly dependent on the alignment of the energy levels of electrodes and analytes.56–61 We are currently undertaking further work to examine how these molecules can be assembled onto a surface and how they differ in terms of their conductivity when studied as either single-molecules or as parallel-arrays of molecules, such as those seen in a self-assembled monolayer.

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*Fig 3. Frontier molecular orbitals for a number of compounds that illustrate the differences in the formation of the HOMOs and LUMOs that result from the addition of an additional ferrocene motif and also from introducing asymmetry across the ferrocene unit.*

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at \*\*\*

Experimental details including full synthetic procedures, spectroscopic data, cyclic voltammetry data, ultraviolet-visible spectroscopy data as well as full molecular orbital depictions and energy levels for all molecules (\*\*\*)

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**Notes**

The authors declare no competing financial interest.

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