Mono-, di- and tetrarhenium Fischer carbene complexes with thienothiophene substituents

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The oxidative cleaving of a rhenium-rhenium bond by bromine in binuclear Fischer carbene complexes proves to be an effective method to prepare mononuclear bromido-carbene complexes. The reaction of mono- and dilithiated thieno[2,3-b]thiophene (2,3-b-TTH 2) and thieno[3,2-b]thiophene (3,2-b-TTH 2) with [Re2(CO)10] afford dirhenium nonacarbonyl ethoxycarbene complexes, [Re2(CO)9(C(OEt)2,3-b-TTH)] (1a) and [Re2(CO)9(C(OEt)3,2-b-TTH)] (1b), and the tetrarhenium bis[ethoxycarbene] complexes from the dilithiated thiophene substrates, ([Re2(CO)9(C(OEt))2{μ-2,3-b-TT}] (2a) and ([Re2(CO)9(C(OEt))2{μ-3,2-b-TT}] (2b) featuring bridging thiophene linkers. Rhenium-rhenium bond cleavage by bromine of the mononuclear complexes yielded the scarce class of monorhenium bromido-carbene complexes, cis-[Re(CO)4(C(OEt)2,3-b-TTH)Br] (3a) and cis-[Re(CO)4(C(OEt)3,2-b-TTH)Br] (3b), while the corresponding reaction of the biscarbene tetrarhenium carbonyl complex of thieno[2,3-b]thiophene afforded the cleaving of both metal-metal bonds to give the novel dirhenium biscarbene dibromido complex with a thienothiophene spacer, ([Re(CO)Br(C(OEt))2{μ-2,3-b-TT}] (4a). A new indirect aminolysis route is described to prepare the chlorido dimethylamino-carbene complex cis-[Re(CO)4Br(NMe2)2,3-b-TTH]Cl], with unexpected cleavage of the Re-Re bond. Single crystal X-ray diffraction studies were performed on 1a, 2a, 3a, 5a, 1b and 3b. Spectroscopic and electrochemical methods are employed to investigate the electronic effect of the different conjugation pathways in the different thienothiophenyl carbene substituents, and the replacement of the rhenium pentacarbonyl fragment with a bromido ligand.

Introduction

The syntheses and applications of neutral rhenium carbonyl complexes of the type, [Re(CO)4(carbene)X], containing both a Fischer alkoxycarbene and halogen ligand, are under-explored, 1 in contrast to the more prevalent dirhenium nonacarbonyl Fischer carbene complex (FCC) analogues. 2 This can be ascribed to the challenging synthetic procedure that result as a consequence of the competitive nature of the nucleophilic attack on carbonyl ligands and lithium-halogen exchange reactions with [M(CO)3X] (M = Mn, Re; X = Br, I).
Scheme 1 illustrates the fate of two Re(CO)₅Br precursors being modified during the lithiation process to give monocarbene dirhenium nonacarbonyl complex A after alklylation, and suggests the elimination of Br₂.⁴⁻⁵ Alternatively, a radical mechanism could also account for the metal-metal bond formation. In addition, low yields (less than 10%) of the resulting dirhenium nonacarbonyl FCCs, are reported. The preferred approach for the synthesis of A is the direct reaction of thienyllithium (RLi) with [Re₂(CO)₁₀] which affords [Re₂(CO)₅(C(OEt)R)] in almost quantitative yields.⁶ Monometal carbene bromido complexes of rhenium are not accessible by the direct reaction of lithiated thiophene with [Re₂(CO)₅]Br. However, unlike for the analogous [Mn(CO)₅]Br, it is possible to synthesize rhenium carbene bromido complexes in reasonable yields, when the lithiated thiophene is π-coordinated to a chromium tricarbonyl fragment acting as an electron-sink.⁷ The corresponding reaction of manganese pentacarbonyl bromide by lithium-halogen exchange proceeded to yield novel α,π-Cr/Mn complexes with bridging thiienyl and carboxythienyl ligands.

Alternatively, reversal of the reaction sequence shown in Scheme 1, by oxidative cleaving of the metal-metal bond in pre-prepared dimetal nonacarbonyl ethoxycarbene complexes with bromine, yields the desired monometal carbene bromido complexes.⁸

The interest in thiophene rings as useful Fischer carbene substrates lie in their planar structure, their electron excessive nature and ability to move charge by electron delocalization as a result of π-conjugation.⁹ Condensed thiophene ring derivatives, thienv[2,3-b]thiophene (2,3-b-TTH) and thienv[3,2-b]thiophene (3,2-b-TTH), (Fig. 1) bear the same general properties of thiophene, yet these substrates display subtle differences in their structural and electronic features. Inter-ring π-resonance effects are possible in 3,2-b-TTH (Fig. 1 (ii)), while electron delocalization in 2,3-b-TTH is predominantly limited to the separate rings only (Fig. 1 (i)). In this study, the preparation of dirhenium(I) nonacarbonyl FCCs containing condensed thiophene substituents are reported, as well as the oxidative cleaving of the Re-Re bonds to yield the mononuclear bromido analogues. The effect of the replacement of a −Re(CO)₅ fragment with a bromido ligand on the steric and electronic properties of the complexes is investigated using spectroscopic and electrochemical techniques, while a new synthetic route to access a novel tetracarbonyl rhenium(I) chlorido aminocarbene complex is reported.

Results and discussion

Synthesis of complexes 1–5

For both condensed thienothiophene substrates, 2,3-b-TTH and 3,2-b-TTH₂, lithiation favours the α-positions of thiophene.⁰ Facile mono- and di-lithiation of 2,3-b-TTH and 3,2-b-TTH₂ followed by reaction of the dirhenium decacarbonyl precursor and alklylation of the resultant metal acylates yield the targeted mono- (1) and biscarbene (2) complexes of dirhenium nonacarbonyl (Scheme 2) in the classic synthetic methodology for Fischer carbene preparation using metal carbonyl precursors. However, low yields of the tetrarhenium biscarbene complexes 2a and 2b were obtained, in contrast to the group 6 metal carbonyl analogues.¹ This low yield is ascribed to the steric congestion caused by the cis- coordination of the two Re(CO)₅–fragments relative to the thienothiophene substrate during the second lithiation, rather than incomplete lithiation. Unlike manganese nonacarbonyl FCCs, where bulky carbene ligands readily occupy axial positions due to steric constraints,¹² regardless of the electronic requirements,⁶ axial coordination of the carbene ligand trans to the Re-Re bond is rarely encountered.¹⁷ Previous studies in our group have shown that linking two dirhenium nonacarbonyl fragments by a 2,5-bis(ethoxycarbene)thiylene spacer yields the eq,eq-isomer only, with marked deviation from planarity of the thiophene linker and a yield of only 5%.¹⁹ Using an extended 5,2'-bithienyl linker results in less strain and higher yields of up to 60%.

In an earlier study on the effect of oxidative cleavage of the metal-metal bonds of dimanganese nonacarbonyl FCCs with bromine, we have shown that the carbene ligands remain intact and the procedure affords a synthetic route to obtain the very scarce class of [Mn(CO)₅(C(OEt)R)Br] (R = heteroaryl) complexes.¹⁰ Applying this methodology to rhenium FCCs results in good conversions when...
reacting 1 with a slight excess of bromine in solvent hexane to yield the desired carbene bromido complexes 3a and 3b (Scheme 3). In the case of the tetrarhenium biscarbene complex 2a, it was possible to cleave both Re–Re bonds with two equivalents of bromine affording novel mononuclear rhenium bromido tetracarbonyl fragments linked by a biscarbene thiophene unit (4a) in 34% yield.

The electronic carbene-stabilization effect of replacing the alkoxy carbene substituent with an amino-group is well-documented, but for comparative reasons it was decided to aminolyse FCC 1a with dimethylamine through a simple, yet effective method of carbene substituent with an amino-group is well-documented, and the electronic carbene-stabilization effect of replacing the alkoxy fragments linked by a biscarbene thiophene[2,3-b]thiophene unit (4a) in 34% yield.

The use of hydrochloric acid in a one-pot carbonyl ligand substitution reaction with Re₂(CO)₉ has been reported to result in Re–Re bond cleavage, although the reaction mechanism remains unexplored. The ethoxycarbene complex 1a, dimethylamine hydrochloride (dimethylammonium chloride) and sodium hydroxide are suspended in THF (Scheme 4). A small amount of distilled water is added to the solution to ensure dissolution of all reagents. Upon solvation, dimethylamine is produced in situ in an acid-base reaction which leads to the rapid aminolysis of the ethoxycarbene complexes with dimethylamine. However, cleavage of the Re–Re bond occurs during the reaction to yield the tetracarbonyl chloride amino-FCC 5a in 56% yield, as well as the pentacarbonyl rhenium chloride cleavage product. Previous reaction with Re₂(CO)₉ has been reported to result in Re–Re bond cleavage, although the reaction mechanism remains unexplored.

Spectroscopic characterization
The carbene complexes were studied by NMR, IR and UV-Vis

Table 1

<table>
<thead>
<tr>
<th>Compound</th>
<th>1a</th>
<th>1b</th>
<th>2a</th>
<th>3a</th>
<th>3b</th>
<th>5a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bond lengths</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>Re–Re (1,2)/Br (3)/Cl (5)</td>
<td>3.1067(3)</td>
<td>3.1020(2)</td>
<td>3.0612(4)</td>
<td>2.6322(4)</td>
<td>2.6268(8)</td>
<td>2.490(3)</td>
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<tr>
<td>Re–C_carb</td>
<td>2.127(4)</td>
<td>2.118(3)</td>
<td>2.099(5)</td>
<td>2.163(3)</td>
<td>2.185(7)</td>
<td>2.205(11)</td>
</tr>
<tr>
<td>C_carb–OEt/NMe₂</td>
<td>1.332(5)</td>
<td>1.335(4)</td>
<td>1.313(6)</td>
<td>1.319(4)</td>
<td>1.289(8)</td>
<td>1.290(13)</td>
</tr>
<tr>
<td>Re–CO₄ trans to Carb</td>
<td>1.976(4)</td>
<td>1.981(4)</td>
<td>1.973(6)</td>
<td>1.991(3)</td>
<td>1.987(8)</td>
<td>1.963(15)</td>
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<tr>
<td>Re–CO₄ trans to Br/Cl</td>
<td></td>
<td></td>
<td></td>
<td>1.925(4)</td>
<td>1.912(9)</td>
<td>1.897(15)</td>
</tr>
<tr>
<td>Re–CO₄ trans to Re–Re</td>
<td></td>
<td></td>
<td></td>
<td>1.922(4)</td>
<td>1.936(4)</td>
<td>1.928(6)</td>
</tr>
<tr>
<td>Re–CO₄ trans to CO₂</td>
<td>1.990(4)</td>
<td>1.991(4)</td>
<td>1.988(6)</td>
<td>2.008(3)</td>
<td>2.015(8)</td>
<td>2.007(14)</td>
</tr>
<tr>
<td>Re–CO₄ trans to CO</td>
<td>2.000(4)</td>
<td>1.998(4)</td>
<td>1.994(6)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Re–CO₄ trans to Re–Re</td>
<td>1.934(4)</td>
<td>1.938(4)</td>
<td>1.927(6)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>C_carb–C₂</td>
<td>1.457(6)</td>
<td>1.459(5)</td>
<td>1.464(7)</td>
<td>1.441(4)</td>
<td>1.439(10)</td>
<td>1.485(13)</td>
</tr>
</tbody>
</table>

| Bond angles (°) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Re–C_carb–O/N | 131.0(3) | 130.2(2) | 123.4(4) | 127.6(2) | 129.3(5) | 128.6(8) |
| Re–C_carb–C₂ | 123.8(3) | 125.5(2) | 125.3(4) | 125.1(2) | 123.0(5) | 115.4(7) |
| O/N–C_carb–C₂ | 105.0(3) | 104.0(3) | 111.3(5) | 107.3(3) | 107.3(6) | 116.0(9) |

| Torsion angle (°) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Re–C_carb–C₂–C₃ | -30.1(6) | 31.2(6) | 27.9(12) | 4.4(5) | -11.7(11) | 69(2) |
| O/N–C_carb–C₂–C₃ | 154.9(4) | -154.5(4) | -153.8(9) | -176.8(3) | 175.3(7) | -113(2) |
| C₉ trans to carb–Re–Re–C_2 | -30.6(2) | 31.8(2) | -38.7(3) | | | |
| C₉–Re–Re–C_2 | -23.0(2) | 35.4(2) | -37.1(3) | | | |

| Angle between two mean planes (°) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 31.79 | 31.45 | 36.02 | 6.64 | 11.02 | 70.80 |

*Averaged bond length. †Smallest torsion angle. ‡The first mean plane drawn through C₂, C₃, C₄ and C₅, and the second through Re, C_carb and O/N.
spectroscopy, as well as mass spectrometry. A pattern of nine bands for the carbonyl stretching vibrations in the infrared spectra of 1 and 2 corresponds to equatorially substituted \([\text{Re}_2\{\text{CO}\}_9\text{L}]\). Four bands of approximately equal intensity correspond to the vibrational modes of a cis-M(\text{CO})_4L moiety and was observed for 3, 4a and 5a. The coordination geometry assignments are supported by the molecular structures obtained from the X-ray crystal structure determinations of 1a, 1b, 2a, 3a, 3b and 5a, respectively (vide infra). Coordination of a Fischer carbene carbon to a dirhenium nonacarbonyl fragment has a significant impact on the chemical shifts of \(H_\beta\) of the thiophene substituents in the \(^1\text{H}\) NMR spectra. This proton resonance is shifted downfield due to the strong electron-withdrawing properties of the metal carbonyl carbene moiety. In addition, compared to non-condensed thiophene Re-FCC substituents previously reported, the \(H_\beta\)-protons are shifted to lower field, in the order thiényl < dithiényl < thiêno(3,2-b)thiényl \((1b) < \text{thiêno}(2,3-b)\text{thiényl} \((1a) \delta 7.65 < 7.73 < 7.98 = 7.99 \text{ ppm})\).\(^{25}\) The order correlates with the electron delocalization ability of the heteroaryl rings as a result of \(\pi\)-resonance effects of the thiényl substituents. For the biscarbene complexes the same trend in \(H_\beta\) resonances was observed for the thiényl substituents (\(\delta 7.47\) (thiényl) < 7.71 (dithiényl) < 7.84 (2,3-b-TT, 2b) < 8.02 (3,2-b-TT, 2a) ppm).

Replacing a \(-\text{Re}(\text{CO})_2\) fragment (1) by a \(-\text{Br}\) ligand (3) markedly affects the electronic features of the complex and shifts all the proton resonances in the \(^1\text{H}\) NMR spectra to lower field (ESI Fig. S2, S7, S15 and S18). As for 1 and 2, increased ring delocalization – perceived in the increasingly downfield \(H_\beta\) resonances compared to non-condensed thiophene carbene substituents – is still observed.

\(^{25}\) The \(^{13}\text{C}\) NMR spectra display carbene carbon chemical shifts for both the mono- and biscarbene complexes 1 and 2 and the analogous thienyl and dithienyl Re-FCCs in the range of \(6 \text{ 289–296 ppm, whereas the corresponding carbene bromido complexes 3a, b,}\) display carbene carbon signals shifted significantly upfield (\(\delta 260–274 \text{ ppm). The carbonyl carbons of the nonacarbonyl compounds 1 and 2 resonate as one broad peak in the \(^{13}\text{C}\) NMR spectra and the chemical shifts are found more downfield (ca. \(\delta 193 \text{ ppm}) than those of the tetracarbonyl complexes, 3. For the tetracarbonyl ligands, three carbon signals are observed. The two cis-CO ligands’ carbon atoms (trans with respect to each other) resonate as a single chemical shift of higher intensity, downfield relative to the other carbonyl C-atoms. The two lower intensity peaks observed more upfield represents the carbonyl carbon resonances \(\text{trans}\) to the carbene and the bromido ligands, respectively. The resonances of the thienothiophene ipso-carbons (\(C_\text{w}\)-atoms) directly attached to the carbene carbons are similar, with values for the dirhenium carbene complexes 1 and 2 being shifted (\(\pm \delta 3 \text{ ppm}) further downfield compared to the corresponding carbene bromido complexes, 3. Comparison of the 2D \(^1\text{H}, \ ^{13}\text{C}\) HSQC spectra of 1 and 3, reveals that the most downfield thienothiophene proton \(H_\beta\) in 1 is bonded to the second most downfield aromatic carbon, \(C_p\). However, for 3, both \(H_\beta\) and \(C_p\) are the most downfield thienothiophene resonances (ESI, Fig. S4, S9, S17 and S20), in all probability due to a H–Br interaction also observed in the crystal structures of 3a and 3b (vide infra).

Comparing the amino- and ethoxycarbene complexes 5a and 3a, respectively, a significant upfield shift of \(H_\beta\) (\(6.670 \text{ ppm}) and the carbene carbon resonance (\(6.242 \text{.1 ppm}) of 5a is observed, in contrast to \(H_\beta\) (\(6.811 \text{ ppm}) and \(C_\text{arbene}\) (\(6.270.2 \text{ ppm}) of 3a. In addition, the steric bulk of the dimethyl-substituted amino-moiety results in the observance of four individual carbonyl carbon resonances observed in the \(^{13}\text{C}\) NMR spectrum of 5a, as well as the significant broadening of both the \(H_\beta\) and \(C_p\) resonances in both the \(^1\text{H}\) and \(^{13}\text{C}\) NMR spectra (Fig. S12 and S13).

Structural studies

Single crystals suitable for X-ray diffraction studies of the dirhenium monocarbene complexes \(eq-[\text{Re}_2\{\text{CO}\}_9\{\text{carbene}\}]\) (1a and 1b), the tetrarhenium biscarbene complex \(eq-[\text{Re}_4\{\text{CO}\}_{12}\{\mu\text{-biscarbene}\}]\) (2a) and monorhenium halido-carbene complexes cis-[Re(CO)\(_4\){carbene}] (3a, 3b \((X = \text{Br})\) and 5a (X = Cl)) were grown from dichloromethane \((\text{CH}_2\text{Cl}_2)\) solutions layered with hexanes. The
Fig. 3 ORTEP (ellipsoids at 55% probability) diagrams of 2a (hydrogen atoms omitted for clarity).

(a)

(b)

(c)

Fig. 4 ORTEP (ellipsoids at 55% probability) diagrams of (a) 3a, (b) 3b and (c) 5a

molecular structures are shown in Fig. 2–4 and selected bond lengths, angles and torsion angles are listed in Table 1. The structures of 1a and 1b consist of two Re fragments, –Re(CO)5 and –Re(carbene)(CO)4, held together by Re-Re bonds, while in 3 the –Re(CO)5 moiety is replaced by a –Br ligand. In 2, the thienothiophene substituent acts as a linker between two carbene functionalities forming a bridging ligand that combine two dirhenium units of 1 in a tetrarhenium complex assembly. For the dirhenium nonacarbonyl complexes 1 and 2, equatorial coordination of the carbene ligand (cis to the Re-Re bond) is confirmed, as well as cis disposition of the carbene ligand with respect to the bromido in the mononuclear complexes 3a, b. The equatorial carbonyl ligands around the Re-Re bond, in the dirhenium complexes’ structures, are in a staggered conformation. The torsion angles are larger than 23° and increase from 1a < 1b < 2a. The ethoxy substituents are orientated on the sulphur side of the rings (Z-isomer), except for 2a and 5a where these species lie across a centre of inversion and thus the thienothiophene rings are disordered by 50%, with 50% rotated by 180° with respect to the metal-carbene moieties. Longer Re-C_Carbene bond lengths are observed when the –Re(CO)5 fragment is replaced with a bromido ligand (compare 3a, 2.163(3) Å and 3b, 2.185(7) Å with 1a, 2.127(4) Å and 1b, 2.118(3) Å, respectively). This observation seems to indicate an overall less electron-rich metal centre with the substitution of –Re(CO)5 with –Br, due to the greater electronegativity of the Br atom, and a formal +1 oxidation state of the rhenium metal. As a consequence, compensation with regards to electrophilic carbene carbon stabilization by the carbene heteroatom substituent results in the shortening of C_Carbene–OEt bond distance in 3a and 3b, compared to 1a and 1b (1.319(4) Å, 1.289(8) Å vs. 1.332(5) Å, 1.335(4) Å). Similar structural consequences were observed for the oxidative cleavage of the Mn-Mn bonds in dimanganese nonacarbonyl FCCs with halogens.10

The torsion angles, Re–C_Carbene–C–C3 and O–C_Carbene–C–C3, serve as indicators that show how far the ethoxy and metal fragment is pushed out of the thienyl mean plane. In the case of 1a, 1b and 2a these fragments are pushed out of the plane by approximately 30°, each in opposite directions, while for the monorhenium carbene bromido complexes 3a and 3b the angle never exceeds 12°, because of the much smaller and less bulky metalcarbonyl fragment.

For both 3a and 3b, Br···H3 (the Hβ thienothiophene hydrogen atom) intramolecular interactions are observed in the packing of the crystal lattices (ESI, Fig. S29); 2.809 Å and 2.787 Å for 3a and 3b, respectively. This could account for the downfield NMR shifts observed for these atoms in the corresponding NMR spectra (vide supra).

Similarly, as observed in the NMR spectra of 5a, the X-ray structural studies demonstrate the increased electron-donating/handcarbene stabilization effect of the amino-group compared to the ethoxy-group of 3a. A longer Re–C_Carbene bond length of 2.205(11) Å for 5a is observed than the corresponding Re–C_Carbene bond length of 2.163(3) Å for 3a. This longer Re–C_Carbene bond length is accompanied by a N–C_Carbene bond length of 1.290(13) Å, indicative of a bond order greater than 1 in 5a.
The characteristic A moiety bound to the metal carbonyl fragment. A greater shift of ca. 80 cm⁻¹ compared to the carbonyl ligand (in which case a much smaller shift is expected), but on a distal moiety bound to the metal carbonyl fragment. This is supported by the spin density calculations as visualized in Fig. S32 and Fig. S35. Spin density analyses of the oxidized forms of 1a and 1b, [1a]⁺ and [1b]⁺, respectively, and the reduced forms [1a]⁻ and [1b]⁻, calculated with DFT (Fig S30-35) were used to confirm the redox events.

Cyclic voltammetry, UV-Vis-NIR and IR spectrophotometry and DFT calculations

The electrochemical properties of 1a, 1b, 2a, 3a, 5a and 2b were evaluated by cyclic voltammetry. In all cases, the cyclic voltammograms (measured at a glassy carbon electrode in CH₂Cl₂) show irreversible oxidation waves assigned to the rhenium metal centres ranging from ca. 0.62–0.77 V for the di- and tetranuclear FCCs 1 and 2a, to ca. 1.33–1.51 V for the mononuclear bromido FCCs 3 (Fig. 5 and ESI, Fig. S21–S26), (vs, Ag/Ag⁺, referenced to the ferrocene/ferrocinium couple at 0 V). The values obtained for 1–3 and 5a are summarized in Table 2. Single quasi-reversible reduction waves at negative potentials were observed for 1a, 1b, 3a and 3b, while two overlapping, irreversible reduction events were observed for the biscarbene 2a. As expected, the CVs of 5a display less positive oxidation potentials (0.47 V) compared to the ethoxycarbene bromido analogue 3a, as anticipated for the more electron-donating aminocarbene ligand. In addition, no reduction event was detectable within the workable potential window of CH₂Cl₂ as previously observed for other metal carbonyl aminocarbene FCCs.¹⁰

The IR stretching frequencies of the CO ligands of 1a and 3a in the neutral and reduced state were measured by IR spectrophotometry, and revealed a shift of the characteristic 9-band pattern for 1a, and 4-band pattern for 3a to lower energies. For example, the A₁' and A₄' vCO bands of 1a at 2102 cm⁻¹ and 1996 cm⁻¹, shift to 2089 cm⁻¹ and 1973 cm⁻¹, respectively, for [1a]⁺; while the characteristic A₁'' band of 3a is shifted from 2002 cm⁻¹ to 1973 cm⁻¹ for [3a]⁻ (Fig S27 and S28). This shift of 13–29 cm⁻¹ is consistent with the localization of the negative charge not on the Re–centre bonded directly to the carbonyl ligands (in which case a much greater shift of ca. 80–100 cm⁻¹ is expected),¹³ but on a distal moiety bound to the metal carbonyl fragment.¹⁴ This is supported by the spin density calculations as visualized in Fig. S32 and Fig. S35.

| Table 2 Cyclic voltammetry data (potentials (V) and currents (µA)) for the two redox processes observed for 1–3 and 5a |
|---|---|---|---|---|---|---|---|
| 1a | 1b | 2a | 3a | 3b | 5a |
| [1a]⁺ | [Re=C/Re-C] | [1b]⁺ | [Re=C/Re-C] | [2a]⁺ | [Re=C/Re-C] | [3a]⁺ | [Re=C/Re-C] | [3b]⁺ | [Re=C/Re-C] | [5a]⁺ | [Re=C/Re-C] |
| E° (V) | 0.63 | 0.62 | 0.69 | 0.77 | 1.33 | 1.47 | 1.47 | 1.47 | 0.47 |
| Epa (V) | – | – | – | – | – | – | – | – | – |
| Epoc (V) | – | – | – | – | – | – | – | – | – |
| ΔEpa (V) | 0.12 | 0.09 | 0.09 | 0.09 | 0.09 | – | – | – | – |
| ΔEpoc (V) | – | – | – | – | – | – | – | – | – |
| jpa (µA) | 26.5 | 6.5 | 14 | 4 | 38, 25⁺ | – | 58, 76⁺ | 5 | 24 |
| jpoc (µA) | – | – | 10 | – | 11, 4 | – | 13 | – | 10 | – | – |

* Overlapping waves

The electrochemical properties of 1a, 1b, 2a, 3a, 5a and 2b were evaluated by cyclic voltammetry. In all cases, the cyclic voltammograms (measured at a glassy carbon electrode in CH₂Cl₂) show irreversible oxidation waves assigned to the rhenium metal centres ranging from ca. 0.62–0.77 V for the di- and tetranuclear FCCs 1 and 2a, to ca. 1.33–1.51 V for the mononuclear bromido FCCs 3 (Fig. 5 and ESI, Fig. S21–S26), (vs, Ag/Ag⁺, referenced to the ferrocene/ferrocinium couple at 0 V). The values obtained for 1–3 and 5a are summarized in Table 2. Single quasi-reversible reduction waves at negative potentials were observed for 1a, 1b, 3a and 3b, while two overlapping, irreversible reduction events were observed for the biscarbene 2a. As expected, the CVs of 5a display less positive oxidation potentials (0.47 V) compared to the ethoxycarbene bromido analogue 3a, as anticipated for the more electron-donating aminocarbene ligand. In addition, no reduction event was detectable within the workable potential window of CH₂Cl₂ as previously observed for other metal carbonyl aminocarbene FCCs.¹⁰

The IR stretching frequencies of the CO ligands of 1a and 3a in the neutral and reduced state were measured by IR spectrophotometry, and revealed a shift of the characteristic 9-band pattern for 1a, and 4-band pattern for 3a to lower energies. For example, the A₁' and A₄' vCO bands of 1a at 2102 cm⁻¹ and 1996 cm⁻¹, shift to 2089 cm⁻¹ and 1973 cm⁻¹, respectively, for [1a]⁺; while the characteristic A₁'' band of 3a is shifted from 2002 cm⁻¹ to 1973 cm⁻¹ for [3a]⁻ (Fig S27 and S28). This shift of 13–29 cm⁻¹ is consistent with the localization of the negative charge not on the Re–centre bonded directly to the carbonyl ligands (in which case a much greater shift of ca. 80–100 cm⁻¹ is expected),¹³ but on a distal moiety bound to the metal carbonyl fragment.¹⁴ This is supported by the spin density calculations as visualized in Fig. S32 and Fig. S35.
localisation of the HOMO on the Re-carbonyl moieties, while the LUMO spreads across the Re=C double bond and the conjugated heteroaryl (TTH) carbene substituent.

The use of the metal-based oxidation potentials as a probe for determining the electronic environment surrounding the central rhenium(I) metal is limited as a comparative tool, due to the irreversibility of the oxidation waves observed. However, it is clear that higher positive potentials are required for the Re(I) oxidation in the bromido FCCs 3a and 3b (1.33 V, 1.51 and 1.47 V), where the rhenium is formally in oxidation state +1, compared to the FCCs 1a and 1b (0.63 V and 0.62 V) containing the Re-Re bond with an – Re(CO) fragment instead of a halide ligand. For the tetrarhenium biscarbene complex 2a, the observance of both, two oxidation waves (\( E_{pa} = 0.69 \) V and 0.77 V), and two reduction waves (\( E_{pc} = -1.48 \) V and -1.66 V) could be indicative of successive redox reactions leading to mixed-valent redox-active intermediates (for example, \([2a]^+\) and \([2a]^2+\), respectively). This observation may on the whole indicate the through-bond electronic communication between the molecular metal-carbene fragments, as supported by our spectroelectrochemistry studies (vide infra), rather than a split of the individual irreversible, oxidation/reduction waves due merely to an electrostatic effect.10c

Redox potentials may not directly provide information on the electron-donating capacity of individual ligands, but is a good measure of the tendency of a system to acquire electrons and thus be reduced. Therefore, in our study, in the observed reduction potentials of complexes 1 and 3, more negative potentials are required for the one electron reduction of the Re=C carbene bond in the case of the dirhenium(0) nonacarbonyl FCCs 1a and 1b (\( E^{\text{in}} = -1.61 \) V and -1.58 V, respectively), than for the bromido rhenium(I) tetracarbonyl FCCs 3a and 3b (\( E^{\text{in}} = -1.48 \) V and -1.43 V, respectively). The more subtle effect of the carbene ligand condensed heteroaryl substituent, namely 2,3-b-TH for FCCs 1a and 3a, and 3,2-b-TH for 1b and 3b, can be deduced by analysing not only the reduction potentials (Table 2), but also the HOMO–LUMO energy levels (band gaps, Fig. 6) of the corresponding compounds. In the case of 1a and 3a, greater negative potentials are required for reduction than for 1b and 3b, respectively. In the same way, for 1a compared to 1b, a higher MO energy gap (3.039 eV compared to 2.976 eV, ESI, Table S2) is calculated, which is also the case for 3a compared to 3b (3.104 eV vs. 2.980 eV). These relatively small energy differences were ascribed to the greater ability for the b series FCCs to stabilise the resultant radical anion following reduction. This is mainly caused by increased conjugation path length and inter-ring π-resonance which is not as prominent in the a series FCCs where electron delocalisation is predominantly restricted to one thiophene ring of the condensed rings (Fig. 7).

The electrochemical and MO analyses results prompted us to use UV-Vis-NIR spectroelectrochemistry (SEC) measurements for compounds 1a, 1b, 2a, 3a and 3b to further investigate the effect of the condensed thienothiophene carbene substituents on the nature of the reduction of these FCCs. SEC is a hybrid analytical technique which can directly indicate the formation of charge transfer states in the molecule under study. The SEC measurements, upon reduction of molecules 1a and 1b (Fig. 8) at an applied potential range of -1.1 to -1.4 V, showed a weak absorption band centred around 540 nm with extinction coefficients below 5000 M$^{-1}$ cm$^{-1}$ clearly indicating the formation of charge transfer species. Furthermore, the weak 540 nm feature appears with the disappearance of 450 nm band in 1a and rise of 420 nm band in 1b. This is also evidenced from our time-dependent density functional theoretical (TD/DFT) calculations. The natural transition orbital (NTO) visualisations as obtained from the TD/DFT calculations (Fig. S30, S31) indicate that the 460 nm (493 nm as calculated) band corresponds to the HOMO–LUMO transition with 98% contribution to the absorption (Table S3) where the hole is spread on the Re-C end and the electron is delocalised on the TT side. The NTO and absorption analysis of the [1b]$^-$ also indicates the rise of 420 nm...
band in 1b (Fig. S33, S34). The TD/DFT calculations reveal the new transitions at ≈540 nm to be a ligand-to-metal charge transfer (LMCT) with a minor σ−π transition within the TT moiety. In the case of 3a, a weak absorption at 650 nm for 3a and 540 nm for 3b has been observed upon reduction of voltage range -1.1 to -1.4 V (Fig. S37, S38). Similarly, for 2a, a weak 560 nm band appears under reductive conditions (Fig. S36) at an expense of 450 nm absorption and a very weak absorption of 1600 nm with an absorption coefficient of ≈200 M⁻¹ cm⁻¹. This rather weak transition can be attributed to the intercalation charge transfer (IVCT) between the Re centres as explained in the electrochemistry section. It is worth mentioning here that all of the compounds under study, underwent a charge transfer state via an isosbestic point (Fig. 8, S36, S37, S38) clearly indicating a direct electron transfer process. Nevertheless, the results obtained from cyclic voltammetry, MO calculations and spectroelectrochemistry suggest that the compounds under investigation undergo Re centred oxidations, while reduction results in the delocalisation of the negative charge on the Re\text{C}_{\text{carbene}}-TT moiety.

**Experimental**

**General**

Thieno[2,3-b]thiophene, thieno[3,2-b]thiophene, bromine, n-BuLi (1.6 M solution in hexane) and dirhenium decacarbonyl were purchased from Sigma Aldrich and Strem Chemicals and used as received. Triethyltetrafluoroborate was prepared according to literature procedure. All operations were carried out using standard Schlenk techniques under an inert atmosphere of nitrogen or argon. Silica gel 60 (particle size 0.063–0.20 mm) was used as resin for all column chromatography separations. Anhydrous THF (tetrahydrofuran) and hexane were distilled over sodium metal and benzophenone, and CH₂Cl₂ over CaH₂ under N₂ atmosphere. NMR spectra were recorded on Bruker AVANCE 500, Ultrashield Plus 400 AVANCE 3 and Ultrashield 300 AVANCE 3 spectrometers, at 25°C, using CDCl₃ as solvent. The 1H NMR spectra were recorded at 500.139, 400.13 or 300.13 MHz, and the 13C NMR spectra at 125.75, 100.613 or 75.468 MHz. The chemical shifts were referenced to deuterated chloroform (CDCl₃) signals at 7.26 ppm for δH and 77.00 ppm for δC. Coupling constants are reported in Hz. Numerical numbering of atoms for NMR spectral assignments is indicated in the ESI, Figure S1. Infrared spectroscopy was performed on a Bruker ALPHA FT-IR spectrophotometer with a NaCl cell, using hexane as solvent. Mass spectral analyses were performed on a Synapt G2 HDMS, by direct infusion at 10 μL/min, using hexane as solvent. Mass spectral analyses were performed on a Synapt G2 HDMS, by direct infusion at 10 μL/min.

**Crystallography**

Single crystal diffraction data for 1a, 1b, 2a, 3a, 3b and 5a were collected at 150 K on a Bruker D8 Venture diffractometer with a kappa geometry goniometer and a Photon 100 CMOS detector using a Mo-Kα λ=1.54Å micro focus source. Data were reduced and scaled using SAINT. Absorption corrections were performed using SADABS. The structures were solved by a novel duel-space algorithm using SHELXTL and were refined by full-matrix least-squares methods based on F² using SHELXL. All non-hydrogen atoms were refined anisotropically. All hydrogen atoms were placed in idealized positions and refined using riding models. The CIF files for complexes 1a, 1b, 2a, 3a, 3b and 5a are available as Supporting Information.

**Cyclic voltammetry**

The CVs at a glassy carbon electrode in CH₂Cl₂ were recorded using a three-electrode cell. The reference electrode was a non-aqueous Ag/Ag⁺ electrode, separated by the test solution by a frit with fine porosity. Glassy carbon disc (3.0 mm diameter) was the working electrode and the counter electrode was a platinum wire. Measurement was done by a Metrohm µAutolab type III potentiostat, using NOVA 2.0 electrochemistry software at scan rates of 100 mV.s⁻¹.

To a deoxygenated solution of 0.1 M [N'⁵Bu₄][PF₆] (supporting electrolyte) in HPLC grade CH₂Cl₂, the test compound (1.0 mM) and the internal standard, 1.0 mM [Fe(n²-C₅H₅)₂]⁺ was added (vs Ag/Ag⁺, E° = 0 V for the ferrocene/ferrocenium couple).

**UV-Vis-NIR Spectroelectrochemistry**

Measurements were performed in CH₂Cl₂ containing 0.1 M N'⁵Bu₄[B(C₆F₅)₄] at 2 mM analyte concentrations within a OTTLE (optical transparent thin-layer electrochemistry) cell at 25 °C in and with a Varian Cary 5000 spectrophotometer. The values obtained by deconvolution could be reproduced within εmax = 100 L·mol⁻¹·cm⁻¹, νmax = 50 cm⁻¹, and Δν/Δν = 50 cm⁻¹. Between the spectroscopic measurements the applied potentials have been increased step-wisely using step heights of 100, 50, 25 or 20 mV. The spectral simulations were calculated using the least amount of gaussian functions to still achieve a reasonable fit of the simulation with the experimental spectrum.

**IR Spectroelectrochemistry**

Measurements were performed in CH₂Cl₂ containing 0.1 M N'⁵Bu₄[B(C₆F₅)₄] at 2 mM analyte concentrations within a OTTLE (optical transparent thin-layer electrochemistry) cell at 25 °C with a Varian Cary 5000 spectrophotometer. Between the spectroscopic measurements the applied potentials have been increased step-wisely using step heights of 100, 50, 25 or 20 mV.

**Computational details**

The geometry optimisation and time dependent DFT calculation were done on Gaussian 09 program. The geometry optimizations, single point and TD/DFT calculations were performed using B3LYP/LANL2DZ basis sets. The molecular orbitals were visualized using Gaussview 5. The UV-Vis spectral visualisations and the spin populations were analysed using Gaussian 3.0 and Chemissian 4.43 programs.

**Preparation of** [Re₂(CO)₉(C(OEt)₂-2,3-b-TTH)] (1a), [(Re₂(CO)₉(C(OEt)₂)|μ-2,3-b-TTH)] (2a), cis-[Re(CO)₉Br(C(OEt)₂-2,3-b-TTH)] (3a), and [(Re(CO)₉Br(C(OEt)₂)|μ-2,3-b-TTH)] (4a)
A solution of 0.07 mL (0.7 mmol) of thieno[2,3-b]thiophene in 15.0 mL of THF was treated with 0.75 mL (1.2 mmol) of n-BuLi at -78°C. After stirring for 30 min, Re(CO)₅(C₅H₅) (0.65 g, 1.0 mmol) was added to the solution and stirred for a further 30 min in the cold and then allowed to reach room temperature over 40 min. The solvent was removed under reduced pressure and the residue dissolved in 10 mL of CH₂Cl₂ and cooled to -20°C. The reaction mixture was treated with 0.23 g (1.2 mmol) of [Et₂P][BF₄] in CH₂Cl₂ and subsequently allowed to reach room temperature. The reaction mixture was filtered through a small silica gel plug using CH₂Cl₂ and the solvent was removed under reduced pressure. After absorbing the reaction mixture on silica, it was dry loaded onto a silica gel column. The respective products (1a and 2a) were separated by column chromatography using gradient elution with hexane and CH₂Cl₂. To synthesize 3a and 4a, 0.20 g (0.2 mmol) 1a and 0.10 g (0.1 mmol) 2a were each dissolved in hexane. Excess bromine dissolved in hexane were added dropwise to the solution while stirring, and allowed to react until no more colour change was observed. The solvent was removed in vacuo. Column chromatography was used to purify the products (3a and 4a, respectively). Hexane was used as initial eluent and the polarity was gradually increased using CH₂Cl₂.

1a: Yield: 0.37 g (0.5 mmol, 68%), red-orange crystals. λmax(CH₂Cl₂)/nm 450 (ε/dm³ mol⁻¹ cm⁻¹ 14835), 390 (12265), 320 (20995). υ(ν(CO)) cm⁻¹ 2104 (A''), 2041s (A''), 2026w (A''), 1956 (A''), 1979 (A''), 1976 (A''), 1955s (A''), 1946m (A''). δH(400.13 MHz; CDCl₃; MeSi) 7.99 (1 H, s, H4), 7.36 (1 H, d, J₆,7 = 5.3, H5'), 7.30 (1 H, d, J₆,7 = 5.3, H4'), 4.58 (2 H, q, J = 7.0, CH₃). 1.63 (3 H, t, J = 7.0, CH₃). 1.23 (10 H, t, J = 7.0, CH₃). 1.35 (100 H, m, CH₂). m/z: 2083 (A''), 1933s (A''), 1984s (A''), 1929m (B'), 1929m (B'). δH(400.13 MHz; CDCl₃; MeSi) 7.31 (1 H, d, J₆,7 = 5.2, H5'), 7.19 (1 H, d, J₆,7 = 5.2, H4'), 6.70 (1 H, s, br, H4), 3.72 (3 H, s, NCH₃), 3.27 (3 H, s, NCH₃). δC(100.613 MHz; CDCl₃; MeSi) 242.1 (C₂ar), 197.8, 194.9, 193.7, 189.1 (Re(CO)₅), 157.0 (C₅), 112.3 (C₄), 145.8 (C₃), 136.4 (C₂), 128.4 (C₃'), 120.2 (C''), 53.5 and 46.9 (CH₃), m/z (C₂H₃O)²⁺, 258.98 g/mol 493.9702 (18%, [M+Cl]⁻), 497.0697 (14%, [M+CH₃Cl]⁻), 465.9721 (24%, [M+Cl]⁻). Column chromatography was used to purify the products. The solution stirred for 15 minutes at -78°C and was allowed to stir at room temperature for 40 minutes. The solvent was removed in vacuo. The residue was dissolved in 10 mL of CH₂Cl₂ and cooled to -30°C, followed by treatment with 0.95 g (5.0 mmol) of [Et₂O][BF₄] in CH₂Cl₂. The reaction mixture was allowed to reach room temperature. Most of the solvent was removed in vacuo, and the sample was wet loaded. Separation was done with a silica column, starting with hexane and gradually increasing the polarity of the solvent by adding CH₂Cl₂, resulting in the isolation of the respective products 1b and 2b. To synthesize 3b, 0.36 g (0.4 mmol) of 1b was dissolved in hexane. Excess bromine dissolved in hexane was added dropwise to the solution while the mixture was stirred. The reaction was terminated as soon as the solution lightened by removing the solvent in vacuo. Column chromatography was used to purify 3b, with hexane as initial eluent and the polarity was gradually increased using CH₂Cl₂.

1b: Yield: 0.73 g (0.89 mmol, 59%), red-orange crystals. λmax(CH₂Cl₂)/nm 455 (ε/dm³ mol⁻¹ cm⁻¹ 14165), 395 (12546), 322 (18907). υ(ν(CO)) cm⁻¹ 2103m (A''), 2041s (A''), 2025w (A''), 2023m (A''), 1994vs (A''), 1978m (A''), 1976m (A''), 1955s (A''), 1945m (A'). δH(400.13 MHz; CDCl₃; MeSi) 7.98 (1 H, s, H4), 7.61 (1 H, d, J₆,7 = 5.3, H5'), 7.24 (1 H, d, J₆,7 = 5.3, H4'), 4.57 (2 H, q, J = 7.0, CH₃), 1.62 (3 H, t, J = 7.0, CH₃). δC(100.613 MHz; CDCl₃; MeSi) 272.3 (C₂ar), 193.3 (br, Re(CO)₅), 159.6 (C₅), 126.2 (C₄), 144.7 (C₃), 139.4 (C₂), 132.8(C₅), 119.8 (C₅), 77.6 (CH₃), 14.6 (CH₃).
m/z(C₃H₅O₃S₅Re₂, 821.87 g/mol) 820.8818 (39%, [M+H]), 792.8371 (100%, [M-H CO]), 764.8448 (47%, [M-H-CO]).

2b: Yield: 0.039 g (0.025 mmol, 2%), purple crystals. δ_H(300.13 MHz; CDCl₃)H4'), 7.86 (1 H, d, J = 6.0 Hz, H5'), 7.28 (1 H, dd, J = 5.3, 0.5 Hz, H4'), 5.31 (2 H, q, J = 7.0 Hz, CH₂), 1.64 (6 H, t, J = 7.0 Hz, CH₃).

3b: Yield: 0.14 g (0.24 mmol, 60%), yellow-orange crystals. λmax(CHCl₃)/nm 442 (ε/1000 mol−1 cm−1 7685), 395 (19827), 343 (7265). v(Re=O)(hexane) cm−1 2101 (A′(1)), 2026 (A′(2)), 2001 (B′), 1943 (B′). δ_H(400.13 MHz; CDCl₃)H4'), 8.17 (1 H, d, J = 5.3 Hz, H5'), 7.28 (1 H, dd, J = 5.3, 0.5 Hz, H4'), 5.31 (2 H, q, J = 7.0 Hz, CH₂), 1.69 (3 H, t, J = 7.0 Hz, CH₃). δ_C(13C)100.613 MHz; CDCl₃)H4'), 7.86 (1 H, d, J = 6.0 Hz, H5'), 7.28 (1 H, dd, J = 5.3, 0.5 Hz, H4'), 5.31 (2 H, q, J = 7.0 Hz, CH₂), 1.69 (3 H, t, J = 7.0 Hz, CH₃). δ_C13C100.613 MHz; CDCl₃)H4'), 8.17 (1 H, d, J = 5.3 Hz, H5'), 7.28 (1 H, dd, J = 5.3, 0.5 Hz, H4'), 5.31 (2 H, q, J = 7.0 Hz, CH₂), 1.69 (3 H, t, J = 7.0 Hz, CH₃).

Conclusions

The preparation of new dirhenium FCCs 1, tetrarhenium biscarbene complexes 2, and subsequent efficient Re-Re bond cleavage of these FCCs by bromine resulting in the FCCs 3 and 4, are reported. Aminolysis of 1a with dimethylamine hydrochloride in a novel synthetic route resulted not only in the desired substitution of the alkoxycarbene substituent with a dimethylamino-group, but also in the cleavage of the Re-Re bond to give the monorhenium(I) tetracarbonyl chlorido amino-FCC 5a. Spectroscopic techniques (NMR, FT-IR, HRMS) and single crystal XRD were employed to confirm the structures of the compounds and to illustrate the electronic and structural effects of (i) the oxidative Re-Re bond cleavage with bromine on the metal-carbene ligand interaction, (ii) alkoxycarbene ligand substitution with aminocarbene ligand (3a vs 5a), and (iii) the increased ring-delocalization pathway available in the FCCs (1b–3b) with the thieno[3,2-b]thienyl carbene substituent, compared to the thieno[2,3-b]thienyl-substituted FCCs (1a–3a). FT-IR and UV-Vis-NIR SEC measurements, and MO DFT calculations supported these findings, and confirmed the localization of the HOMOs on the rhenium carbonyl moieties; LUMOs delocalized across the carbene ligand; with larger band gaps for the Re(I) tetracarbonyl bromido monocarbene complexes than observed for the dirhenium(0) nonacarbonyl monocarbene complexes. As anticipated, electrochemical reduction yields radical anions with the negative charge delocalized across the carbene-PT ligand (more efficiently stabilized by the thieno[3,2-b]thienyl carbene ligands (series b) than by the thieno[2,3-b]thienyl carbene ligands (series a)), and in the case of the biscarbene FCCs 2, UV-Vis-NIR SEC also indicated the presence of through-bond intramolecular electronic communication between the terminal rhenium-carbene moieties.

Conflict of interest

There are no conflicts of interest to declare.

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