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A conjugated diosma-octacyclic complex and its mixed-valence singly reduced state[†]

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In this work two dicationic diosma-octacyclic complexes have successfully been synthesized and fully characterized. One of them, with aromatic osmapentalene termini linked by the non-aromatic osmafuran moiety to the rigid naphthalenediolate bridge, represents the first example of a fused, fully conjugated dimetalla-octacyclic complex. A reference complex with more flexible biphenolate in the bridging position was also prepared. Their redox and electronic properties were investigated by combined methods of cyclic voltammetry and UV-vis-NIR-IR spectroelectrochemistry, supported by density functional theory (DFT) and time-dependent density functional theory (TD-DFT) calculations. The rigid octacyclic complex forms a stable singly reduced mixed-valence species with the spin density localized at one of the osmapentalene termini.

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Introduction

Analysis and understanding of unconventional aromaticity in organometallic compounds (Hückel, Craig and Möbius types) has long been in the center of interest for both experimental chemists and theoreticians. The chemistry of metallaaromatics has been developing rapidly and new challenges have been identified.^{1–10} Recently, polycyclic metallaaromatic compounds such as metalla-naphthalenes,^{11–14} metalla-naphthalynes,^{15,16} metalla-anthracene,^{17,18} metalla-fluoranthenes,¹⁹ metalla-anthracynes,^{20,21} and carbolong

complexes^{22–31} have widely been studied due to excellent thermodynamic stability and outstanding performance involving charge transfer,³² NIR luminescence²² photothermal effect,^{33–35} photovoltaics,^{36,37} photoactive therapy³⁸ and catalysis.³⁹

To date, most of these complexes contain only a single transition metal in the polycyclic skeleton. Whilst abundant transition metal-organic dinuclear complexes have yielded unusually brilliant results in catalysis, magnetism, electronic communication, and optical materials, metallaaromatic complexes containing two metal centres (including dimetallabenzenes) have been less developed.^{40,41}

In 2014, Xia and co-workers synthesised metal-bridged tricyclic aromatic complex 2 (monocationic, a BF_4^- salt) through the nucleophilic addition of phenol to cationic osmapentalyne 1 (Scheme 1).⁴² The aromaticity of the red-highlighted core in 2 obtained support from the calculated negative values of nucleus-independent chemical shifts (NICS) for each of the three fused five-membered metalla-rings. In this work, the tricyclic, reportedly osmaaromatic⁴² moieties in 2 are interconnected by a conjugated bridge to form dicationic dinuclear complexes 3 and 4 (Scheme 1). Flexible 4,4'-biphenol was used as the nucleophilic reagent to form the osmafuran-appended biphenol bridge in 3. Then, 2,6-naphthalenediol was chosen to prepare octacyclic complex 4 with a rigid bridge core. To the best of our knowledge, metallaaromatic compounds with an octacyclic skeleton are unprecedented in the literature, regarding both monometallic and dimetallic systems. Aromaticity of the molecular rings in both 3 and 4 was investigated by the recently developed electron density of delocalized bonds (EDDB) method^{43,44} (instead of the popular NICS calculations^{45–47}) and the anisotropy of induced current

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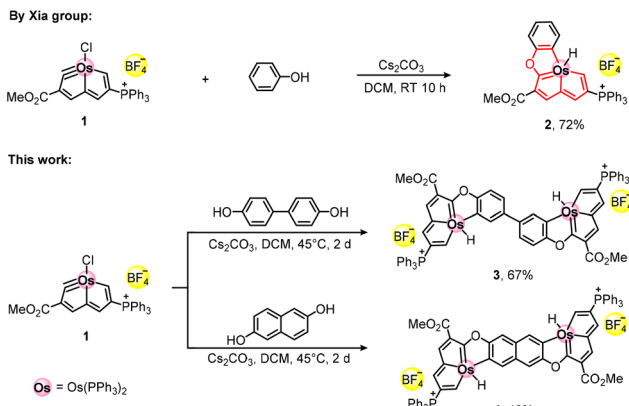
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† Electronic supplementary information (ESI) available: Experimental details and syntheses; NMR spectra and HR-MS data; X-ray crystallographic data; complete cyclic voltammograms; IR spectroelectrochemistry; additional DFT, TD-DFT and EDDB data. CCDC 2024859 and 2024860. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d2qi01523d>

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Scheme 1 Syntheses of 2–4.

density (AICD) analyses.^{48,49} Their redox and electronic properties and the nature of the singly reduced mixed-valence state were explored by cyclic voltammetry, *in situ* UV-vis and IR spectroelectrochemistry at variable temperature, and quantum mechanical calculations.

Results and discussion

The synthetic routes to dinuclear **3** and **4** are shown in Scheme 1. Commercially available 4,4'-biphenol and slightly more than two equivalents of osmapentalyne **1** were refluxed in the presence of Cs_2CO_3 in dichloromethane for 2 days to yield **3** in 67% yield. Complex **4** was synthesized from **1** and 2,6-naphthalenediol under the same reaction conditions and isolated in 40% yield. In the solid state, both **3** and **4** are stable and can be stored in air for one year without any signs of decomposition. The composition and purity of dications **3** and **4** were revealed by their HRMS/ESI⁺ signals (Fig. S17 and S18, ESI†) and elemental analyses, respectively. The calculated mass of each molecular ion complies with the experimental value.

The structures of **3** and **4** are presented in Fig. 1 and 2, respectively, and in Fig. S1 and S2 (ESI†). Crystallographic data are summarized in Tables S1–S3 (ESI†). Single crystals of **3** suited for X-ray diffraction were grown by slow diffusion of *n*-hexane into the dichloromethane solution of the complex. In the crystal, **3** is centrosymmetric and each osmium atom is shared by three five-membered rings. The sum of the angles in the five-membered ring $\text{Os}1-\text{C}1-\text{C}2-\text{C}3-\text{C}4$ is 540.0° , which is an ideal value indicating its perfect planarity. The same applies for the five-membered rings $\text{Os}1-\text{C}4-\text{C}5-\text{C}6-\text{C}7$ and $\text{Os}1-\text{C}7-\text{O}1-\text{C}9-\text{C}8$, where the deviation from the ideal value for ring planarity is very small, *viz.* 539.9° and 539.8° , respectively. The small torsion angles in the three osmacycles are illustrated in Fig. S1(a) (ESI†). The sum of the angles in the phenolate rings ($\text{C}8-\text{C}13$) is 719.5° , which is again close to the ideal value of a planar six-membered ring. The slight deviation from planarity of the polycyclic skeleton in the crystal of **3**

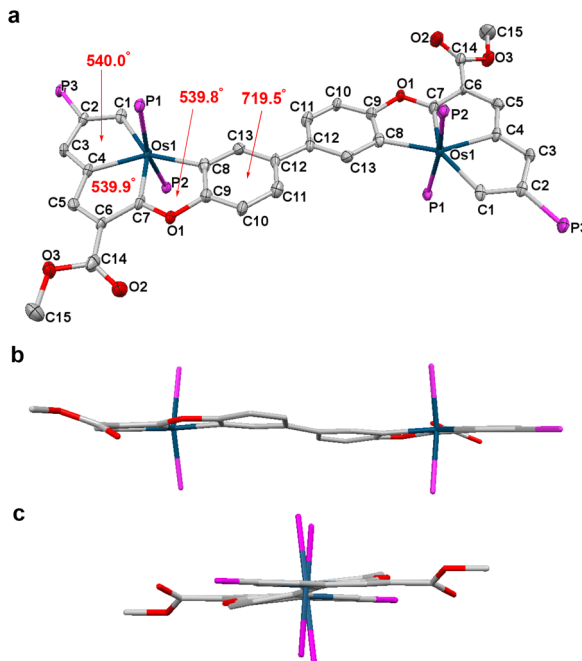


Fig. 1 (a) Crystal structure of **3**- $4\text{CH}_2\text{Cl}_2$ (thermal ellipsoids set at 50% probability); (b) front view; (c) side view. Hydrogen atoms, BF_4^- counterions, CH_2Cl_2 molecules and phenyl substituents in PPh_3 have been omitted for clarity. Selected bond lengths (\AA) and angles ($^\circ$) are presented in Table S2 and Fig. S2(a) (ESI†).

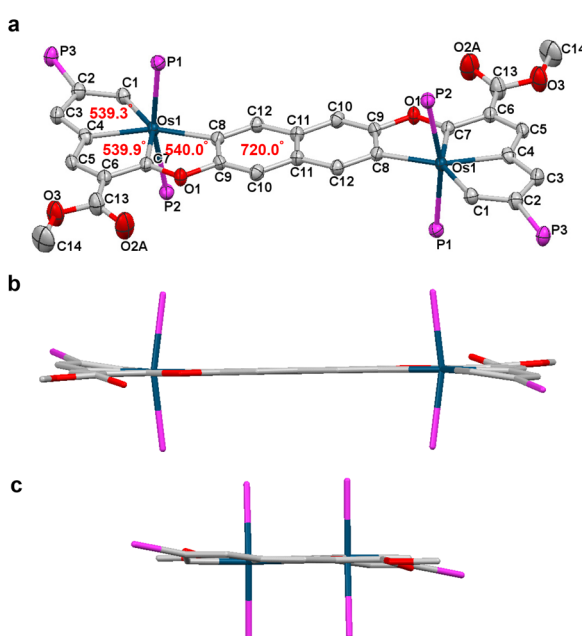


Fig. 2 (a) Crystal structure of **4**- BPh_4^- - $2\text{CH}_2\text{Cl}_2$ (thermal ellipsoids set at 50% probability); (b) front view; (c) side view. Hydrogen atoms, anions, CH_2Cl_2 molecules and phenyl substituents in PPh_3 have been omitted for clarity. Selected bond lengths (\AA) and angles ($^\circ$) are presented in Table S3 and Fig. S2(b) (ESI†).



seen in Fig. 1(b and c) has its origin in the twist and S-shaped distortion of the biphenyl core due to the flexible inter-ring C12–C12' bond. However, the C12–C12' bond length of 1.378 Å is significantly smaller compared to the value of 1.494 Å for the inter-ring C–C σ-bond in biphenyl and comparable with intra-ring distances in benzene (1.397 Å),⁵⁰ proving strong π-conjugation in the biphenolate bridge in 3.

Diverse attempts to grow single crystals of dicationic 4 with BF_4^- counterions failed. Consequently, the counterions had to be exchanged for BPh_4^- by reacting 4 with $\text{Na}[\text{BPh}_4]$ in methanol. Suitable single crystals of **4-BPh₄** were obtained by slow diffusion of *n*-hexane into the dichloromethane solution of the complex. The crystal structure of dication **4-BPh₄** is also centrosymmetric. Each ring of the octacyclic skeleton exhibits excellent planarity, while sum of the ring angles in each half reaching 539.3°, 539.9°, 540.0° and 720.0° (Fig. 2). Differently from 3, the planar octacyclic structure of **4-BPh₄** with the rigid naphthalene core is undistorted (Fig. 2b and c). The values of the small torsion angles in the three osmacycles are given in Fig. S1(b) (ESI†).

The three Os–C bonds in the five-membered rings of 3 (2.060–2.124 Å) are in average slightly longer compared with 2,⁴² but still falling within the range reported for planar Möbius aromatic osmapentalene (1.926–2.139 Å).^{22–30,51} The Os–C8 bond to the biphenolate linker is slightly longer (2.154 Å). Interestingly, all four Os–C bonds in **4-BPh₄** (2.048–2.130 Å) containing the naphthalenediolate linker are shorter compared with 3, which reflects increased conjugation in the rigid octacyclic system, potentially facilitating electronic communication between the redox-active osmapentalene halves (see the Cyclic voltammetry and DFT parts hereinafter).

Complexes 3, 4 and **4-BPh₄** were further characterized by nuclear magnetic resonance (NMR) spectroscopy (Fig. S6–S15, ESI†). In the ¹H NMR spectra, the resonances in the high-field region (δ –3.33 ppm for 3 and δ –2.92 ppm for 4) are assigned to OsH. The low-field signals at δ 12.00 ppm for 3 and δ 12.22 ppm for 4 belong to OsCH, which complies with the chemical shifts of OsCH reported for aromatic osmapentalenes (11.93–12.46 ppm).^{42,51} The ¹³C{¹H} NMR spectra of 3 and 4 exhibit low-field signals at δ 257.5 ppm for 3 and 257.0 ppm for 4, which are assigned to the OsCH resonance. Compared to 4, the ¹³C{¹H} NMR spectrum of **4-BPh₄** reveals a new quartet signal at δ 164.5 ppm assigned to BC in the BPh_4^- counterion.

Density functional theory (DFT) calculations of models 2'', 3'', and 4'' (see Computational details in ESI†) have appreciably reproduced the experimental ¹H NMR chemical shifts of OsCH (see Fig. S16 in ESI† for details). The downfield chemical shifts of the H1, H3 and H5 nuclei, ranging from 8.54 to 12.22 ppm, comply with the aromaticity of the two metallapentalene rings.^{22,23,42} The difference between the experimental and calculated OsH values is about 3 ppm; this is understandable, as the hydride is directly bound to the 5d transition metal, and its magnetic shielding can strongly be affected by spin-orbit (SO) effects.⁴⁸

The planarity, high stability in air and the OsCH downfield shifts in the ¹H NMR spectra of 3 and 4 indicate that these

diosmium complexes are aromatic in the osmapentalene rings. To probe the aromaticity of the whole octacyclic system and compare the results with the data in the previous report by Xia and co-workers,⁴² electron density of delocalized bonds (EDDB)^{43,44} was used to investigate the aromaticity of these compounds instead of the alternative popular indices such as nucleus-independent chemical shifts (NICS),^{45,46} as the latter calculations feature certain limitations in evaluating the aromaticity of metallacycles.⁵² The strong local paramagnetic currents around heavy atoms have been reported to affect the value of NICS. Hence, NICS might not be a reliable criterion in evaluating ring aromaticity in osmapentalene derivatives⁵³ and metal cluster systems.^{54,55} Note that the EDDB analysis is an advanced electronic criterion for aromaticity, proposed by Szczepanik *et al.*, which has shown superiority in the quantification of electronic delocalization in aromatic rings.^{43,44} The EDDB analysis can efficiently separate the σ- and π-contributions from bonds in the molecules, and can also specifically investigate the electronic delocalization of specific loops in complex systems. With the popularity of corresponding computing programs, the EDDB analysis is often performed to quantitatively assess the strength of aromaticity in an increasing number of aromaticity studies.^{56–59} Therefore, the EDDB analysis has been used to evaluate the aromaticity of 2'', 3'' and 4'' (see Computational details in ESI†) in comparison with the tricyclic part of 2'' reported by Xia and co-workers⁴² (Fig. 3). The calculations show that the π-EDDB value for the peripheral 11-membered metal-bridged tricyclic aromatic system is 5.850 *e*. The corresponding values for the whole series 2'', 3'' and 4'' are slightly smaller (Fig. 3a). Detailed analyses of the π-EDDB values for each individual ring are presented in Fig. 3b. The π-EDDB values determined for the two five-membered metalla-rings of the osmapentalene moiety in

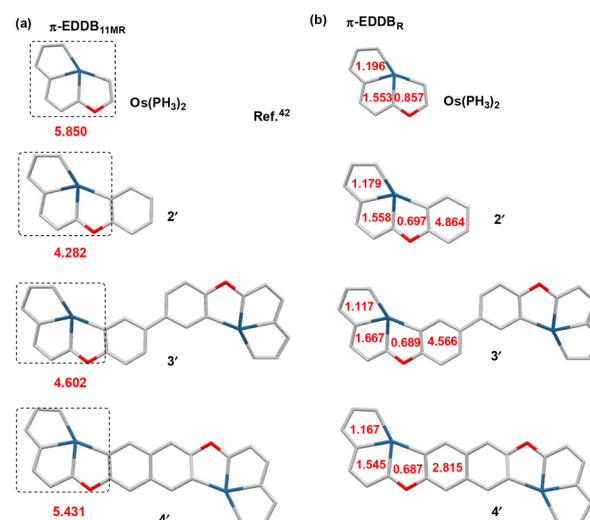


Fig. 3 EDDB analysis of the ring aromaticity in 2'', 3'' and 4''. (a) π-EDDB values for the 11-membered triple ring system. (b) Separate π-EDDB values for each ring, including the additional benzene ring in the bridge fused to the osmafuran moiety.

2', 3' and 4' are very similar to those in the limited tricyclic system, indicating that the osmapentalene rings in 2', 3' and 4' are aromatic. In contrast, the π -EDDB value (0.857 e) calculated for the osmafuran ring in the previously reported tricyclic system⁴² is slightly larger than those (0.687–0.697 e) for the osmafuran ring in 2', 3' and 4' fused with the additional benzene ring of the bridge (Fig. 3b). As the π -EDDB values (0.687–0.857 e) for the osmafuran ring in all these complexes are much smaller than that (2.483 e) of furan (Fig. S20†), together with no distinguishable ring current of the osmafuran ring in the ACID plots (Fig. 4), we can draw the conclusion that the osmafuran rings in all these complexes, including the previously reported⁴² tricyclic species, is nonaromatic. In addition, the π -EDDB value (2.815 e) for the benzene ring in 4' is much smaller than those in 2' and 3'. Similarly, the EDDB and π -EDDB value (3.144 e) for the benzene ring in naphthalene is also significantly reduced in comparison with that (5.327 e) of benzene (Fig. S20†). The much lower π -EDDB value for the naphthalene linker in 4' (Fig. 3) is understandable due to its mobile π -sextet in the Clar structures.^{60–62} For comparison, the overall EDDB values obtained from all contributing molecular orbitals in the 11-membered ring system, and the separate overall EDDB values for each ring are summarised in Fig. S19 (ESI†), leading to the same conclusions.

In line with the EDDB calculations, AICD analyses^{48,49} of both mononuclear 2' (Fig. S21, ESI†) and dinuclear 3' and 4' (Fig. 4) also demonstrate the nonaromaticity of the osmafuran rings and aromaticity of the other rings. Specifically, the clockwise diatropic ring currents are shown in the two osmapentalene rings as well as the central biphenyl a naphthalene rings in 3' and 4', respectively. In contrast, no distinct ring currents could be identified in the inserted osmafuran rings. This also applies for tetracyclic 2'. In summary, fully conjugated 4 cannot be classified as a so far unprecedented fully aromatic dimetalla-octacyclic compound. Different from the previously reported aromatic osmafurans,⁶³ the osmafuran ring in 4 rep-

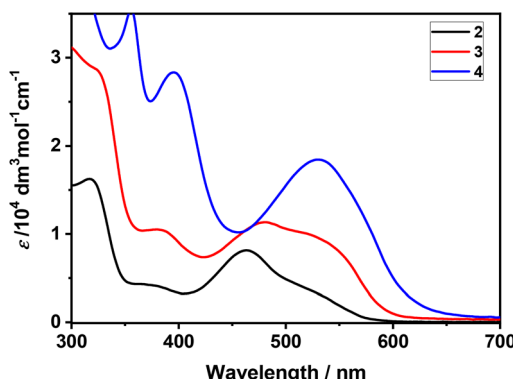


Fig. 5 UV-vis absorption spectra of 2–4 measured in CH_2Cl_2 at $T = 293$ K.

resents a non-aromatic linker between the aromatic osmapentalene termini and the naphthalene bridge.

The electronic absorption spectra of 2–4 are shown in Fig. 5. The absorption maximum of 2 at $\lambda = 464$ nm (with a shoulder at 520 nm) is close to the value of 461 nm reported by Xia and co-workers.⁴² The corresponding visible absorption of dinuclear 3 and 4 is slightly shifted to a lower energy and gains intensity. An absorption band between 350–400 nm is also observed for all three complexes, its intensity following the same trend, from weakly absorbing 2 to strongly absorbing 4.

The electronic absorption of dinuclear 3 and 4 in the visible-near-UV region and the nature of the participating molecular orbitals, including the HOMO and LUMO (Fig. 6) examined in the electrochemical studies (*vide infra*), have been clarified by DFT and TD-DFT calculations conducted on their optimized models. The relevant data are presented in Tables S4–S6 (ESI†). The conjugated tetracyclic halves in 3 manifest a good planarity, but the large dihedral angle (35.8°) ascertained in the model structure further diminishes their electronic communication. In contrast, the diosma-octacyclic skeleton of optimized 4 is fully planar.

The visible electronic absorption of planar 4 has been adequately reproduced by the TD-DFT results. The broad asymmetric band at 530 nm (at $T = 293$ K) encompasses two closely lying intense electronic transitions attributed to combined $\text{HOMO} \rightarrow \text{LUMO}$ and $\text{HOMO-1} \rightarrow \text{LUMO}$ excitations corres-

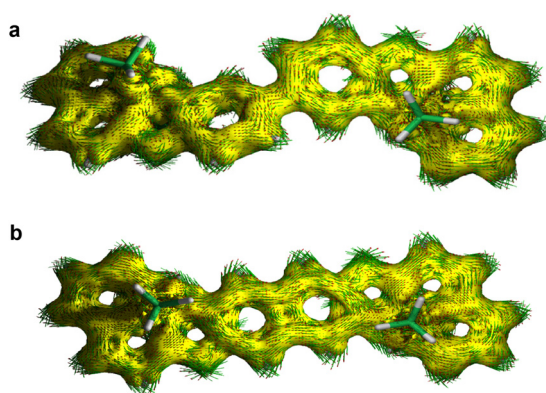


Fig. 4 AICD plots of (a) 3' and (b) 4' visualizing π -ring current contributions. Current density vectors are plotted onto the AICD isosurface of 0.030 to indicate diatropic ring currents. The magnetic field vector is orthogonal with respect to the ring plane and points upwards. The aromatic rings exhibit clockwise diatropic circulations.

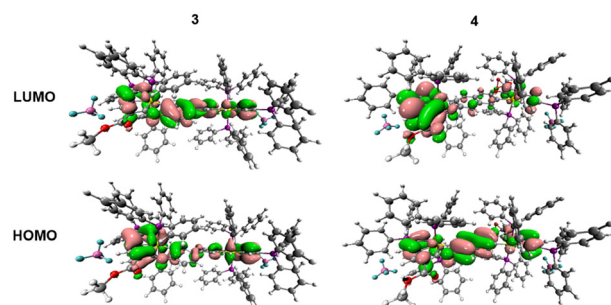


Fig. 6 HOMO and LUMO of 3 (left) and 4 (right).

ponding to transfers of π -electron density from the naphthalenediolate bridge and/or one of the osmapentalene moieties to the other osmapentalene moiety (interpreted as LMCT/MMCT). The strong electronic absorption of **4** at 395 nm is dominated by a charge transfer from one osmapentalene unit to the triphenylphosphonium substituent, HOMO-1 \rightarrow LUMO+2, and a charge transfer within the bis(osmapentalene) and central naphthalene π -system, HOMO-3 \rightarrow LUMO, resembling the HOMO \rightarrow LUMO excitation in the visible region.

For **3** the TD-DFT data have predicted strong absorption at 498 nm assigned as the HOMO \rightarrow LUMO excitation within the bis(osmapentalene)-biphenolate backbone π -system, with a limited charge-transfer character. The higher-lying absorption in the visible region has a similar, even more delocalized nature due to the strong contribution from the HOMO-3 \rightarrow LUMO excitation. On the other hand, the near-UV absorption is dominated by the HOMO-2 \rightarrow LUMO+1 excitation that involves the biphenyl core and one of the osmapentalene moieties and can therefore be interpreted as a charge-transfer transition (LMCT). Although the TD-DFT data comply appreciably with the experimental near-UV-vis absorption of **3**, some prudence needs to be taken, as the biphenolate twist angle in the optimized model structure is larger than unveiled in the crystal structure of the complex (Fig. 1).

The redox behaviour and electron-transfer properties of conjugated complexes **2–4** were essentially explored by cyclic voltammetry (CV) at variable temperature (Fig. 7, Fig. S3 in ESI,† and Table 1). Mononuclear **2** shows reversible reduction at $E_{1/2} = -2.04$ V at ambient temperature (Fig. S3a†), but its oxidation remains totally irreversible even at $T = 233$ K (Fig. 7(a)). Dinuclear **3** and **4** follow this behaviour. Differently from refer-

Table 1 Redox potentials of **2–4** from cyclic voltammetry at a Pt microdisc electrode^a

Osmapentalene	$E_{1/2}$ (red1), Volt	$E_{1/2}$ (red2), Volt	$E_{p,a}$ (oxid.), Volt
2	-2.04	n/a	0.57
3	-2.03	-2.11	0.34
4	-1.96	-2.09	0.43

^a Experimental conditions: dichloromethane/*n*-Bu₄NPF₆ and $T = 233$ K. The electrode potentials are referenced against the standard ferrocenium/ferrocene (Fc⁺/Fc) couple.

ence **2**, however, they exhibit two separate reversible reduction steps (Fig. 7(b) and (c), respectively). The HOMO–LUMO gap (E_g) of 2.07 eV has been estimated⁶⁴ from the cyclic voltammogram of complex **3**. This value is nearly identical with 2.08 eV determined⁶⁴ from the lowest electronic absorption of **3** (in CH₂Cl₂/*n*-Bu₄NPF₆, see below), which is shown by TD-DFT to encompass largely (86%) the HOMO \rightarrow LUMO excitation (Table S4, ESI†). For **4** the difference of the optical band gap from the CV-based value of 2.14 eV remains limited to *ca.* -0.05 eV, which complies with the very close energies of the HOMO-1 and HOMO involved in the lowest-energy optical excitation (Tables S4 and S6, ESI†). The potential difference $\Delta E_{1/2}$ between the successive reduction waves (obtained by deconvolution) increases from 80 mV for **3** to 130 mV for **4**. The LUMO of **4** resides largely on one of the osmapentalene termini while the HOMO is strongly delocalized over the naphthalenediolate bridge and both osmapentalene termini (Table S5, ESI†). This observation is signalling some weak electronic coupling between the Os centres in parent **3**, which increases in rigid diosma-octacyclic **4**. However, the presence of the non-aromatic osmafuran rings in the conjugated dinuclear compounds is the likely reason for the small $\Delta E_{1/2}$ value even in the latter case. The reversibility of the close-lying cathodic steps reveals the formation of stable mixed-valence (M–V) complexes in the 1e⁻-reduced state, which may, however, undergo redox disproportionation. This assumption is further supported by UV-vis spectroelectrochemistry (*vide infra*).

The DFT description of singly reduced radical **4**[–] (with the PPh₃ groups in **4**⁺ replaced by PH₃ for successful geometry optimization) has indeed exposed the α -HOSO and α -LUSO being localized on the opposite osmapentalene units (Table S7, ESI†) and the spin density being distributed mainly over one osmapentalene fragment (Fig. 8). This striking asymmetry (Fig. S5, ESI†), in combination with the moderate $\Delta E_{1/2}$ value (Table 1), permits to assign **4**[–] as a class I rather than class II M–V system, with trapped valences according to the Robin–Day classification,⁶⁵ although, most M–V systems have been based on distinct metal oxidation states.^{66–70} Recording of the electronic absorption spectrum of **4**[–] was then attempted to identify a low-lying intervalence charge-transfer (IVCT) absorption, which would otherwise verify the class II M–V assignment.

UV-vis and IR spectroelectrochemical experiments with **2–4** to monitor their electrochemical reduction were conducted in

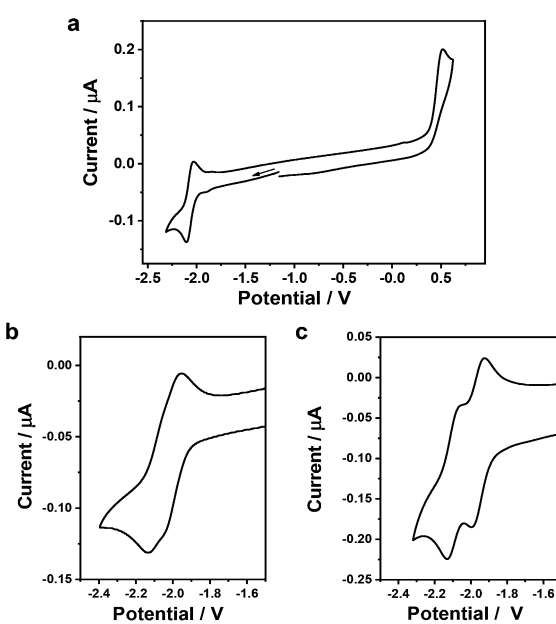


Fig. 7 Cyclic voltammograms of 1 mM **2** (a), **3** (b) and **4** (c) in CH₂Cl₂/*n*-Bu₄NPF₆ at $v = 100$ mV s⁻¹, $T = 233$ K. The potential scales correspond to the standard ferrocenium/ferrocene redox couple.

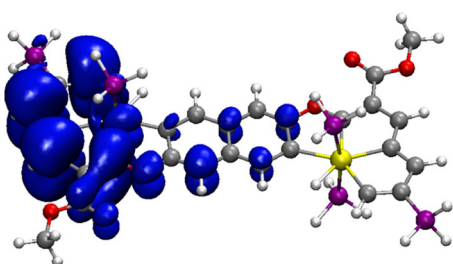


Fig. 8 Spin density distribution in mixed-valence $4''^-$. Contour values: $\pm 0.04 \text{ (e bohr}^{-3})^{1/2}$.

non-coordinating dichloromethane at $T = 233 \text{ K}$, using an OTTLE cell. UV-vis spectral changes recorded during the cathodic electron transfer steps are depicted in Fig. 9.

The 1e^- reduction of 2 in the chilled electrolyte at -2.04 V proved to be reversible only in the initial phase, in contrast to the reversible cathodic CV response (Fig. 7 and Table 1). The prominent absorption band at 463 nm was gradually decreasing, which was accompanied by the growth of new low-energy absorption between $550\text{--}650 \text{ nm}$ tailing to 800 nm . Having reached *ca.* 50% conversion (as shown in Fig. 9(a)), the low-energy absorption of 2^- began to disappear; subsequent re-oxidation at a strongly positively shifted anodic counterwave did not lead to the recovery of parent 2 . The secondary reduction product, denoted as $\{2\}^-$, has not been identified.

On the contrary, the initial 1e^- reduction of 4 to 4^- was fully reversible at $T = 233 \text{ K}$, as manifested by the persistent isosbestic points in the UV-vis absorption spectra (Fig. 9(c)) and smooth recovery of the parent absorption at a relatively

small overpotential during the reverse anodic step. The strong absorption band of 4 at 508 nm was replaced by a half-reduced maximum at 570 nm and a broad absorption of low intensity between $650\text{--}900 \text{ nm}$. These characteristic features resemble the transient absorption observed for mononuclear 2^- (Fig. 9(a)) prior to its irreversible transformation to $\{2\}^-$, which complies with the localized valence of 4^- as manifested by DFT calculations (Fig. 8). The parallel monitoring of the reduction of 4 to 4^- in the IR fingerprint region revealed a small red shift of the $\nu(\text{C=O})$ absorption of the ester group on the osmapentalene skeleton from $1718 \text{ to } 1710 \text{ cm}^{-1}$, accompanied by a significant gain of the absorption intensity (Fig. S4, ESI†). The apparent asymmetry of the new $\nu(\text{C=O})$ band at 1710 cm^{-1} reveals the presence of the underlying, diminished band at 1718 cm^{-1} also in valence-localized 4^- where only one of the osmapentalene termini gets singly reduced. As demonstrated by Fig. 8, the affected ester group in 4^- hardly bears spin density, in line with the limited decrease in the $\nu(\text{C=O})$ wavenumber. The consecutive cathodic step at -2.09 V (Table 1) converted 4^- to a poorly known secondary species, $\{4\}^{2-}$, showing a new absorption band at 492 nm on the foot of the strong near-UV absorption (Fig. 9(d)). Like $\{2\}^-$, the reoxidation of $\{4\}^{2-}$ was found in the thin-layer cyclic voltammogram strongly positively shifted beyond the parent reduction wave and reproduced neither singly reduced 4^- nor parent 4 .

The stepwise electrochemical reduction of 3 was not resolved sufficiently due to the small $\Delta E_{1/2}$ value (Table 1) to permit the complete generation of singly reduced 3^- . As shown in Fig. 9(b), the characteristic electronic absorption of 3^- between $600\text{--}900 \text{ nm}$ (cf. 4^- in Fig. 9(c)) only appears at the onset of the reduction wave at -1.73 V . The concomitant redox disproportionation of 3^- results in the direct observation of the secondary reduction product, $\{3\}^{2-}$ (Fig. 9(b)), showing similar absorption features between $350\text{--}600 \text{ nm}$ as $\{4\}^{2-}$ (Fig. 9(d)). The UV-vis spectroelectrochemical results have revealed that 3^- and 4^- are electronically close and contain the same chromophores. The twisted $4,4'$ -biphenolate and coplanar naphthalenediolate linkers in the parent complexes markedly control the weak to moderate electronic communication between the osmapentalene termini.

The electronic absorption of stable 4^- in the visible region was analysed by TD-DFT calculations performed on its optimized model $4''^-$ (see Fig. S5 in ESI†) having the six PPh_3 groups replaced by PH_3 . The data are summarized in Table 2 and Table S7 (ESI†). The weak absorption recorded between $650\text{--}900 \text{ nm}$ (Fig. 9(c)) is dominated by the $\alpha\text{-HOSO} \rightarrow \alpha\text{-LUSO+2}$ transition within the π -orbital system delocalized over the reduced osmapentalene moiety, also including a charge transfer to the phosphonium ring substituent. There is no evidence for a low-lying intense IVCT transition that would comply with the assignment of model $4''^-$ as a class II M-V complex, as discussed above at the cyclovoltammetric response of 4 . The medium-intensity absorption band of 4^- at 570 nm represents a π -electronic excitation close to the weak lowest-energy absorption, with the central naphthalene rings of

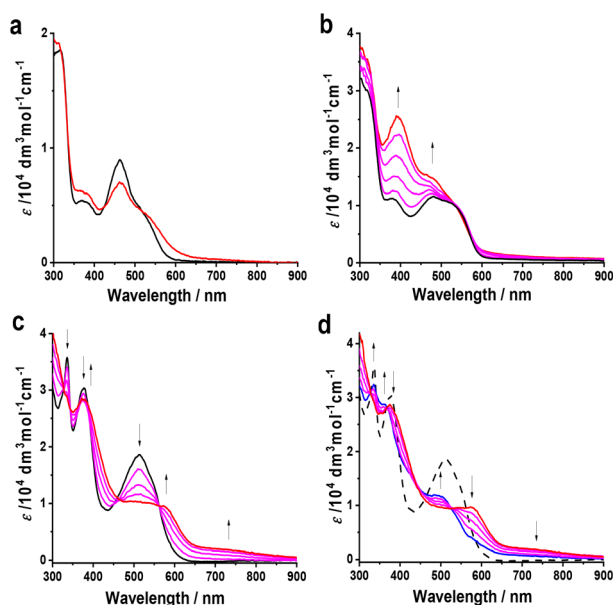


Fig. 9 Spectral changes in the UV-vis-NIR absorption region recorded during the electrochemical reduction in a cryostatted OTTLE cell: $2 \rightarrow 2^-$ (a), $3 \rightarrow \{3\}^{2-}$ (b), $4 \rightarrow 4^-$ (c), $4^- \rightarrow \{4\}^{2-}$ (d). Conditions: CH_2Cl_2 , $n\text{-Bu}_4\text{NPF}_6$, $T = 233 \text{ K}$, $c_{\text{parent}} = 2 \text{ mmol dm}^{-3}$.



Table 2 Experimental (exp.) electronic absorption of singly reduced 4^- ($T = 233$ K, in dichloromethane/ n -Bu₄NPF₆) and TD-DFT calculated (calc.) electronic absorption in model $4''^-$

Excitation (percentage)	<i>f</i>	λ_{\max} (calc.)/nm	λ_{\max} (exp.)/nm ($\epsilon/10^4 \times M^{-1} \text{ cm}^{-1}$)
α -HOSO \rightarrow α -LUSO+2 (82%)	0.001	719	~710 (0.195)
α -HOSO-3 \rightarrow α -LUSO (28%)	0.0001	663	
α -HOSO \rightarrow α -LUSO+6 (19%)	0.092	474	570 (0.955)
α -HOSO \rightarrow α -LUSO+6 (19%)	0.091	453	
β -HOSO \rightarrow β -LUSO+2 (23%)	0.023	427	440 (1.31)
α -HOSO \rightarrow α -LUSO (55%)			
α -HOSO \rightarrow α -LUSO+6 (18%)	0.197	423	
β -HOSO \rightarrow β -LUSO+2 (53%)			
β -HOSO-8 \rightarrow β -LUSO (15%)	0.071	394	377 (2.82)
β -HOSO-2 \rightarrow β -LUSO (15%)			
α -HOSO-9 \rightarrow α -LUSO (16%)	0.028	391	
β -HOSO-9 \rightarrow β -LUSO (16%)			
α -HOSO-1 \rightarrow α -LUSO (11%)	0.094	386	
β -HOSO-2 \rightarrow β -LUSO (11%)			

α -LUSO+6 being more involved. The α -HOSO \rightarrow α -LUSO transition (Table S7†) with an apparent IVCT character is shown to contribute significantly only to the higher-lying optical excitation around 450–500 nm. In general, the absence of a clear IVCT absorption band in the NIR region and the complex nature of the visible electronic absorption of 4^- (represented by model $4''^-$) do not meet the criteria for an intermediate class II M-V system marked by a low activation energy for facile interconversion of the distinct valences (electron-hopping) on the timescale of 10^{-14} s. Even though the rigid central naphthalenediolate linker in **4** may mediate a stronger electronic interaction between the osmapentalene termini compared to slightly twisted dinuclear **3**, the singly reduced M-V radical species with the spin density localized at a distinct osmapentalene site (Fig. 8) can more reliably be classified as a class I valence-trapped complex.

Conclusions

In summary, two dicationic conjugated diosma-octacyclic complexes **3** and **4** were successfully synthesized by a nucleophilic addition reaction of osmapentalyne with 4,4'-biphenol and 2,6-naphthalenediol, respectively. The downfield proton chemical shifts, AICD analysis and calculated EDDB values confirmed the aromaticity of the terminal osmapentalene units and the bridge core. In contrast, the osmafuran ring linking these aromatic regions is shown to be nonaromatic. Complex **4** is the first reported conjugated dimetalla-octacyclic compound. Cyclic voltammetry and DFT calculations reveal a moderate electronic communication between the osmapentalene termini mediated by the rigid naphthalenediolate core. The rare singly reduced mixed-valence species, 4^- , was characterised by UV-vis and IR spectro-electrochemistry as a Robin-Day class I valence-trapped complex. This work has not only enriched the family of metallaaromatic (osmapentalene) complexes by novel structures, but also clarified their peculiar electronic nature, redox behaviour, and ability to undergo electronic communica-

cation between polycyclic redox centres, opening a vast new playground for exploring such phenomena in this field by experiment and theory in the years ahead.

Author contributions

Funding acquisition: Sheng Hua Liu, František Hartl; syntheses: Yu Xuan Hu; investigation: Yu Xuan Hu, Qianqian Deng, Xiaofei Yang, Jing Zhang, Emily K. Garrett, Yaping Ou, František Hartl; supervision: Jun Zhu, Sheng Hua Liu, František Hartl; visualization: Yu Xuan Hu, Jing Zhang, František Hartl; writing – original draft: Yu Xuan Hu, František Hartl; writing – review & editing: Jun Zhu, Sheng Hua Liu, František Hartl.

Conflicts of interest

The authors declare no conflict of interest.

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