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## Redox-active tetraruthenium metallacycles: reversible release of up to eight electrons resulting in strong electrochromism†

Daniel Fink, Bernhard Weibert and Rainer F. Winter\*

**Tetraruthenium macrocycles with 1,4-divinylphenylene and diarylamine-substituted isophthalic acids as the sides display up to eight one-electron redox steps and rich electrochromic behaviour with strong absorptions of the dications in the near infrared and of the tetra- and hexacations at low energies in the visible.**

Metallacycles are typically constructed from metal coligand fragments as nodes and two kinds of ditopic bridging ligands, so-called linkers as the sides. The shapes and sizes of such structures are determined by the preferred coordination geometries of the metal ions and the topologies of the linkers. Hence, a great number of metallacycles with vastly different architectures and astounding levels of complexity have been realized.<sup>1–6</sup> Considering that many of these structures contain redox-active metal ions or linkers, or even both, only relatively few studies were specifically devoted to investigating or capitalizing on that property. Inherent perspectives such as triggering changes in guest-binding inside the cavities of metallacycles, manipulating their optical and magnetic properties, or studying intracage charge transfer phenomena have only sporadically been tackled. The elegant work of Hupp and coworkers, who explored charge transfer between diimine or porphyrinic linkers in mixed-valent tetraruthenium rectangles with ligand-based redox activity,<sup>7–10</sup> and of Kaim, Stang and Therrien on tetraplatinum, -rhenium or -ruthenium metallacycles with oxidizable or reducible linkers stand as instructive examples.<sup>11–16</sup>

We here report on tetraruthenium metallacycles which are constructed from two pairs of diruthenium 1,4-divinylphenylene- and triarylamine-derived ditopic linkers as two different kinds of redox-active entities that allow for the pairwise release of up to eight electrons per macrocycle resulting in strong polyelectrochromism.<sup>17–19</sup> Discrete divinylphenylene-bridged

diruthenium complexes  $\{\text{Ru}(\text{CO})(\text{L})(\text{P}^i\text{Pr}_3)_2\}_2(\mu\text{-CH=CH-C}_6\text{H}_4\text{-CH=CH-1,4})$  (L = mono- or bidentate monoanionic ligand) are generally oxidized in two consecutive, reversible one-electron steps. Their mixed-valent radical cations exhibit complete charge and spin delocalization as is indicated by the observation of just one Ru(CO) IR-band (the observed small band splitting arises from the non-degeneracy of the symmetrical and anti-symmetrical Ru(CO) stretching vibrations) and resolved EPR hyperfine splitting of the unpaired spin to four equivalent <sup>31</sup>P nuclei of the P<sup>i</sup>Pr<sub>3</sub> coligands.<sup>20,21</sup> Triaryl amines constitute paradigmatic organic redox-systems and have found great use as selective oxidants and in redox catalysis.<sup>22–26</sup> More electron-rich representatives can be further oxidized to persistent dications.<sup>26</sup> A macrocycle constructed from two of these entities each may thus reversibly release six or even eight electrons per macrocycle.<sup>27,28</sup>

When the 1,4-divinylphenylene-bridged diruthenium complex **1-Cl** as 180° building block was combined with diphenyl- or (di-4-anisylamino)benzene-3,5-dicarboxylic acid (**2-H** or **2-OMe**) as a 0° building block<sup>4</sup> with K<sub>2</sub>CO<sub>3</sub> base in MeOH/CH<sub>2</sub>Cl 1 : 1, a complex mixture of products was initially formed as indicated by the presence of several sets of vinyl proton and <sup>31</sup>P NMR signals.

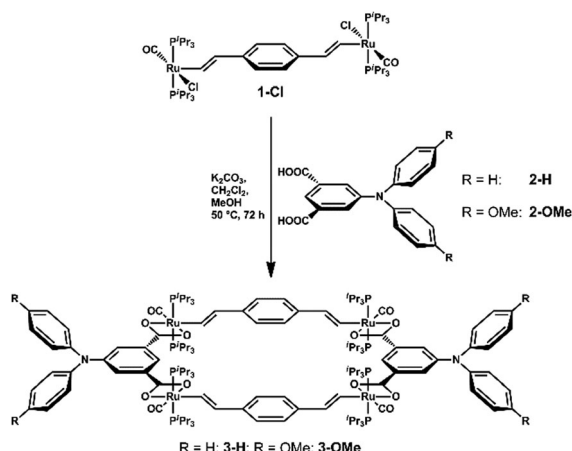
On further warming to 50 °C for 48 h that mixture became uniform, showing just one set of proton NMR signals for the divinylphenylene and dicarboxylate building blocks and a single <sup>31</sup>P NMR resonance. Target macrocycles **3-H** and **3-OMe** (Scheme 1) were isolated in yields of ca. 60% as pale yellow microcrystalline solids and were characterized by <sup>1</sup>H, <sup>13</sup>C{<sup>1</sup>H} and <sup>31</sup>P{<sup>1</sup>H} NMR spectroscopy, ESI-MS and combustion analysis (ESI†).

Crystals of **3-OMe** that lent themselves for X-ray crystallography were grown from benzene/MeOH and CH<sub>2</sub>Cl<sub>2</sub>/MeOH, respectively, and the results are shown in Fig. 1. In both structures, the divinylphenylene diruthenium moieties adopt a *cisoid* conformation and both vinyl groups point outward the inner cavity of the metallamacrocyclic ring. This generates an oval cavity of approximate dimensions of 7.9 Å × 12.4 Å or 7.8 Å × 12.4 Å as measured between opposing hydrogen atoms. Forcing the divinylphenylene units into a metallamacrocyclic ring causes some minor torsional strain. In the benzene solvate **3-OMe**·7C<sub>6</sub>H<sub>6</sub> this strain is evident from the torsion

Fachbereich Chemie, Universität Konstanz, Universitätsstraße 10, D-78453 Konstanz, Germany. E-mail: rainer.winter@uni-konstanz.de

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Scheme 1 Synthesis of the tetraruthenium metallamacrocycles.

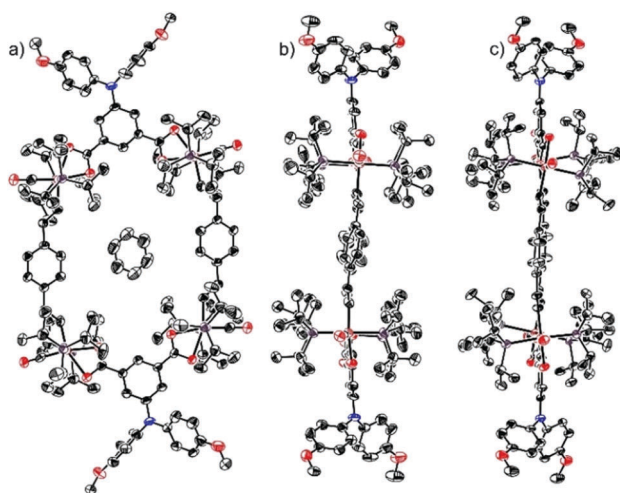


Fig. 1 Structures of **3-OMe** as the benzene solvate (a) top view, (b) side view and (c) side view of the  $\text{CH}_2\text{Cl}_2$  solvate. Solvent molecules other than the benzene molecule in the cavity and hydrogen atoms are removed for reasons of clarity. The ellipsoids are displayed at a 50% probability level.

of one ruthenium alkenyl moiety of each divinylphenylene linker by  $\phi = 27.3^\circ$  with respect to the adjacent phenylene ring whereas the other remains in a more coplanar arrangement with  $\phi = 6.6^\circ$ . Torsional angles O–C–C–C at the benzene-3,5-dicarboxylates of  $7.6$  and  $4.5^\circ$  and at the Ru–C=C–C linkages of  $3.5$  and  $4.8^\circ$  are nearly ideal and the P–Ru–P vectors of opposing nodes are almost parallel as indicated by P–Ru–Ru–P torsion angles of  $3.5$  and  $4.1^\circ$ . In the dichloromethane solvate **3-OMe**· $8\text{CH}_2\text{Cl}_2$  the divinylphenylene moieties achieve a somewhat higher degree of coplanarity with the central phenylene rings with torsion angles  $\phi$  of  $10.8$  and  $13.3^\circ$  while the torsional angles O–C–C–C and Ru–C=C–C are larger at  $7.2^\circ$  and  $14.9^\circ$  or  $7.2^\circ$  and  $8.4^\circ$ . The overall result is a twist of the P–Ru–P vectors as measured by P–Ru–Ru–P torsions of  $19.0^\circ$  and  $21.9^\circ$ , which in turn induces a bending of one divinylphenylene moiety to above and the other one to below the mean plane of the macrocycle (Fig. 1c). The two structures can thus be viewed as snapshots demonstrating the conformational degrees of freedom inherent to these macrocycles.

In **3-OMe**· $7\text{C}_6\text{H}_6$ , one molecule of benzene stacks in a coplanar, but laterally offset fashion with respect to the divinylphenylene sides, whereas the cavity of the  $\text{CH}_2\text{Cl}_2$  solvate accommodates two solvent molecules that show no H-bonding interactions or short contacts with the cavity walls (see ESI†). This indicates that the tetraruthenium macrocycles, like other structures of this type, might also exhibit interesting host–guest chemistry.<sup>29,30</sup>

The electrochemical properties of macrocycles **3-H** and **3-OMe** are best compared to those of precursor **1-Cl** and its bis(benzoato) substitution product,  $\{\text{Ru}(\text{CO})(\text{P}^i\text{Pr}_3)_2(\kappa^2\text{O}, \text{O}'\text{-OOCPh})\}_2(\mu\text{-CH=CH-C}_6\text{H}_4\text{-CH=CH-1,4})$  (**1-OOCPh**). Both complexes display two consecutive, reversible one-electron waves at half-wave potentials  $E_{1/2}$  of  $-0.09$  and  $+0.17$  V<sup>20</sup> or  $-0.25$  and  $+0.02$  V, respectively.<sup>31</sup> The cathodic shift of  $\sim 150$  mV on substitution of the chloro by the benzoato ligands follows the increase in valence electron count at the Ru atoms from 16 to 18 and the concomitant raise in energy of the redox-orbital(s), which receive major contributions from the  $\pi$ -conjugated bridging ligand.<sup>20,21,32–37</sup> For **3-H**, three consecutive waves are apparent in the cyclic voltammograms ( $\text{CH}_2\text{Cl}_2/\text{NBu}_4\text{PF}_6$ , Fig. 2). On close inspection one notes a small splitting of the first wave into two closely spaced one-electron processes. No such effect is however seen for the second and third waves.  $E_{1/2}$  values under these conditions are summarized in Table 1. Nearly identical results were obtained with the even less ion-pairing  $\text{B}\{\text{C}_6\text{H}_3(\text{CF}_3)_2-3,5\}_4^-$  counterion (ESI†). Since all waves are associated with similar peak currents and based on the overall composition and potentials we can safely assign the first two composite waves as the oxidations of the divinylphenylene sides and the

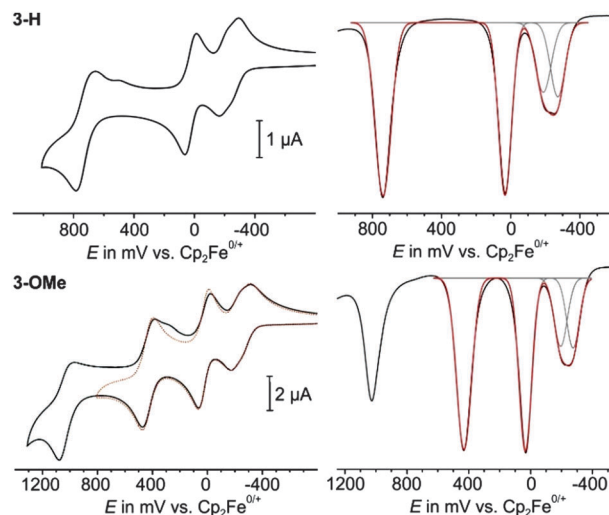


Fig. 2 Cyclic voltammograms ( $\text{CH}_2\text{Cl}_2/0.1 \text{ M NBu}_4\text{PF}_6$ , r.t. at  $v = 0.1 \text{ V s}^{-1}$ ) (left) and square wave voltammograms (right) of **3-H** and **3-OMe** along with deconvolutions.

Table 1 Half-wave potentials of metallamacrocycles **3-H** and **3-OMe** in mV vs.  $\text{Cp}_2\text{Fe}^{0/+}$

	$E_{1/2}^{0/+}$	$E_{1/2}^{+/2+}$	$E_{1/2}^{2+/4+}$	$E_{1/2}^{4+/6+}$	$E_{1/2}^{6+/8+}$
<b>3-H</b>	−272	−188	33	739	—
<b>3-OMe</b>	−275	−194	34	431	1026



third wave as the concurrent oxidations of the amine appendices. Thus, all three waves correspond to two closely spaced or coincident one-electron processes.

In the case of **3-OMe**, a fourth such wave is seen close to the anodic discharge limit of the electrolyte, which obviously corresponds to the second oxidations of the peripheral triarylamine sites (Fig. 2). The above assignments are reconfirmed by the fact that substitution of the phenyl by anisyl substituents only affects the redox potentials past the 4+ states (Table 1).

Bis(alkenylarylene)-bridged diruthenium complexes with extended conjugated bridging ligands and [2.2]paracyclophanyl-bridged analogs with cofacial and coplanar arranged styryl decks were found to exhibit rather substantial degrees of ground-state delocalization in the mixed-valent states despite potential splittings  $\Delta E_{1/2}$  near the purely statistical limit.<sup>32,38</sup> This is, however, not the case here as shown by the results of IR and UV/Vis/NIR (NIR = near infrared) spectroelectrochemistry. Spectroscopic changes on the first composite two-electron oxidation, where one electron is removed from each divinylphenylene diruthenium entity, are essentially identical to the first oxidation of **1-Cl** or **1-OOCPh**. Thus, the Ru(CO) IR band blue-shifts by 34 cm<sup>-1</sup> with some discernible splitting into a less intense band at higher and a more intense one at lower energy (Fig. 3 and ESI†). That splitting is due to a non-degeneracy of the symmetrical and antisymmetrical combinations of Ru(CO) stretches. Further spectroscopic changes include the growth of bands in the region of ring stretching and bending vibrations and of an intense absorption in the NIR at ca. 7700 cm<sup>-1</sup> (ESI†). That band also shows up as part of an intense, structured NIR absorption in the UV/Vis/NIR experiments along with an equally intense one in the Vis (see Table 2, Fig. 3 and ESI†). Characteristic spectroscopic data are collected in Table 2.

The relation between  $\Delta E_{1/2}$  (in Volts) and  $K_c$ , the equilibrium constant of the comproportionation reaction  $3\text{-H}^{2+} + 3\text{-H} \rightleftharpoons 2\text{3-H}^+$ ,  $K_c = \exp[n \cdot F \cdot \Delta E_{1/2} / (R \cdot T)] = \exp(39.33 \cdot \Delta E_{1/2})$  at  $T = 295$  K dictates that, after release of one equivalent of charge, the mixed-valent (MV) forms **3-H**<sup>•+</sup> or **3-OMe**<sup>•+</sup> are the dominant species in the equilibrated solutions, even when  $\Delta E_{1/2}$  is vanishingly small. Still, no spectrum recorded during electrolysis has features other than those corresponding to the respective dications and the neutral precursors and, in particular, none that could be ascribed to charge transfer from the remaining reduced divinylphenylene side to the already oxidized one. This means that the radical cations are mixed-valent species of Class I with non-interacting redox sites.<sup>39,40</sup> Obviously the dicarboxylic linkers, which do not contribute to the redox orbitals HOMO and HOMO-1,<sup>31,41</sup> and the lateral positioning of the divinylphenylene moieties of the macrocycles mutually insulate the two divinylphenylene sides from each other. As a consequence, **3-H**<sup>2+</sup> and **3-OMe**<sup>2+</sup> are EPR active and display a somewhat broad isotropic signal at  $g = 2.039$  (ESI†).

The same overall behaviour repeats on the second composite oxidation encompassing the concurrent 2+/3+ and the 3+/4+ waves, corresponding to the second oxidation of the divinylphenylene sides (Fig. 3 and ESI†). Thus, the overall spectroscopic changes with a further blue-shift of the Ru(CO) band by ca. 40 cm<sup>-1</sup>, the collapse of the NIR band and the further growth

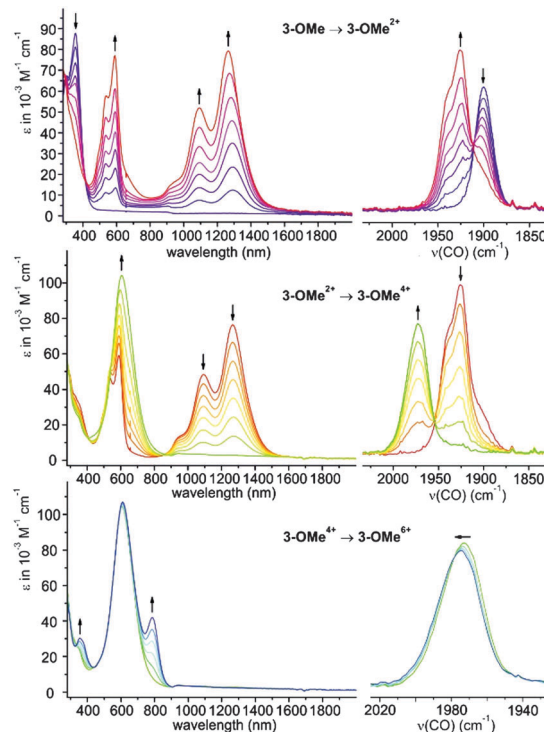


Fig. 3 Changes in the UV/Vis/NIR (left) and in the  $\nu(\text{CO})$  region of the IR spectra (right) of macrocycle **3-OMe** upon oxidation to **3-OMe**<sup>2+</sup> (top), **3-OMe**<sup>4+</sup> (middle) and **3-OMe**<sup>6+</sup> (bottom) (CH<sub>2</sub>Cl<sub>2</sub>/NBu<sub>4</sub>PF<sub>6</sub>, r.t.).

Table 2 Characteristic spectroscopic data for macrocycles **3-H** and **3-OMe** in all their accessible oxidation states

	$\nu(\text{CO})$ (cm <sup>-1</sup> )	$\lambda$ (nm) ( $\epsilon \times 10^{-3}$ (M <sup>-1</sup> cm <sup>-1</sup> ))
<b>3-H</b>	1901	306 (60.3), 357 (75.6)
<b>3-H</b> <sup>2+</sup>	1943, 1927	293 (49.8), 538 (42.4), 590 (60.4), 1093 (36.0), 1267 (59.9)
<b>3-H</b> <sup>4+</sup>	1975	276 (53.1), 605 (101.9)
<b>3-OMe</b>	1900	304 (67.4), 355 (87.8)
<b>3-OMe</b> <sup>2+</sup>	1942, 1925	282 (70.3), 537 (57.8), 590 (76.9), 1093 (52.0), 1267 (79.4)
<b>3-OMe</b> <sup>4+</sup>	1973	276 (60.2), 606 (104.3)
<b>3-OMe</b> <sup>6+</sup>	1975	275 (54.5), 354 (30.4), 610 (107.0), 787 (42.1)

of the Vis absorption are very similar or virtually identical to those observed on the second oxidation of diruthenium complexes **1-Cl** and **1-OOCPh**. Again, no absorption bands other than those of the di- and tetracations are seen at intermediate stages of the electrolysis where **3-H**<sup>3+</sup> or **3-OMe**<sup>3+</sup> constitute the major species in solution. Hence, the trications are also valence localized MV species of Class I. The divinylphenylene diruthenium entities thus behave as two independent electrophoric units and the only consequence of forging two of them together in a metallacyclic structure is a predictable doubling of molar extinction coefficients to rather impressive values of ca. 80 000 for the prominent NIR absorption of **3-OMe**<sup>2+</sup> and >100 000 for the Vis band of **3-OMe**<sup>4+</sup> (neutral **3-OMe** is transparent in these regimes).

Owing to the inherent instability of **3-H**<sup>6+</sup>, the spectral consequences of further oxidizing the triarylamine appendices





could only be monitored for **3-OMe**. They include a very modest shift of the Ru(CO) IR band from 1973 to 1975 cm<sup>-1</sup> and of the prominent Vis band from 606 nm in **3-OMe**<sup>4+</sup> to 610 nm in **3-OMe**<sup>6+</sup> (see Table 2). Additional bands at 354 nm and at 787 nm with extinction coefficients of 30 400 and 42 100 M<sup>-1</sup> cm<sup>-1</sup> closely resemble those of the radical cation of **2-OMe** ( $\lambda_{\text{max}} = 365$ , 793 nm).<sup>42</sup> Moreover, chemically generated **3-OMe**<sup>6+</sup> displays a non-binomial triplet at  $g = 2.014$  and a <sup>14</sup>N hyperfine splitting of 9.6 G in its EPR spectrum as it is typical of aminium-type radicals,<sup>43</sup> thereby reconfirming the above assignments of the individual redox processes (ESI<sup>†</sup>). The only negligible impact of amine oxidation on the position of the  $\pi \rightarrow \pi^*$  absorption band of the bis(oxidized) divinylphenylene diruthenium entities and the Ru(CO) band shifts further emphasizes the lack of electronic interaction between the two different kinds of redox sites. Further oxidation to **3-OMe**<sup>8+</sup> was, however, not possible under the conditions of spectroelectrochemistry, probably due to the proximity of that wave to the solvent background and a poor solubility of this polycation.

In conclusion, we have prepared and investigated amine-functionalized tetraruthenium metallamacrocycles in good yields from divinylphenylene-diruthenium complexes and amine functionalized isophthalic acids. These compounds can be oxidized by up to eight electrons with stepwise charge loss from the divinylphenylene linkers and the triarylamine appendices. Spectroscopic consequences of oxidation up to **3-OMe**<sup>6+</sup> were monitored by IR and UV/Vis/NIR spectroelectrochemistry and revealed strongly electrochromic behaviour.<sup>44</sup> The lateral placement of the redox-active divinylphenylene entities and the insulating nature of the dicarboxylate linkers shut down intramolecular charge transfer between the individual redox sites and lead to paramagnetic behaviour.

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