A fascinating multifaceted redox-active chelating ligand: introducing the $N,N'$-dimethyl-3,3'-biquinoxalinium “methylbiquinoxen” platform†

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To intimately combine a chelating ligand function with the numerous properties of a viologen-like redox-active centre would offer a rare possibility to design controllable multi-redox states, whose properties arise from strongly correlated phenomena between the organic ligand as well as with any metalloid coordinated centres. Such a concept previously proved to be feasible, however is not widely applicable owing to challenges in terms of synthesis, isolation, and aerial sensitivity of both the ligand and its metal complexes. Here we report the first stable example of such a redox-active molecule, $N,N'$-dimethyl-3,3'-biquinoxalinium$^{3+/4+}$ “methylbiquinoxen”, MBqn$^{3+/4+}$, which shows a rich redox chemistry and chelates a metal ion in the case of the metal complex [CdCl$_2$(MBqn)$^0$]. This goes beyond what is possible to achieve using viologens, which are limited by not providing chelation as well as having no accessible biradicaloid state, corresponding to the neutral direduced MBqn$^0$ open-shell behaviour we observe here.

Introduction

Organic molecules with multiple oxidation levels accessed by electro-, photo-, or chemical means are of extreme importance in many biological, chemical and physical processes.1–4 Among these, viologens, or $N,N'$-disubstituted-4,4'-bipyridinium ($V^{2+}$) species, have attracted most attention over the last 80 years. Acting as strong electron-acceptors, viologens undergo two reversible one electron reduction steps leading to a radical cation intermediate ($V^{+}$) and neutral form ($V^0$).5,6

The interest in viologen derivatives is motivated, amongst other things, by the ease of synthesis and tunability, the high degree of control concerning their reversible oxidation-reduction process, the range of colours from colourless ($V^0$) to dark green/blue/purple ($V^{+}$), the paramagnetic nature of $V^{+}$, and their strong structural templating effects ($V^{2+}$).7,8 These possibilities have led to their incorporation in a range of donor-acceptor complexes developed for various applications such as photo- and electrochromic displays,9–11 charge accumulation,12–14 fluorescence,15–17 supramolecular host–guest complexes and molecular machines18–20 as well as ferroelectrics.17

Research on viologens is mainly focused on two aspects. The first of these is the diversification of their properties achieved through incorporation of additional functions. The second aims to improve their longevity, in particular through improving the chemical and air stability of the radical intermediates. In this context, Kaim’s group have replaced some carbon atoms with nitrogen atoms within the aromatic heterocycles, leading to a new type of redox-active ligand that intimately combines a viologen-like centre with the chelating function of 2,2'-bipyridine.21–23 Despite an improved π-electron deficiency and an interesting coordination function that provides strong electronic coupling between the coordinated site (typically a metal ion) and the organic redox centre, synthetic challenges and persistent chemical and air instability, render these conceptually interesting molecules currently not useful for applications.

In the context of the potential for radical-based systems and the challenges to be addressed, we report here on the synthesis and characterization of a new stable reversible organic redox-active chelating ligand $N,N'$-dimethyl-3,3'-biquinoxalinium (MBqn$^{3+}$, 1$^{2+}$, Fig. 1 – grey), “methylbiquinoxen”. In particular the dication 1$^{2+}$ and its monoprotonated direduced form (MBqn$^{0}$H$^+$, 1$^{+}$H$^+$, Fig. 1 – purple) are thermally stable ($1^{2+}$ and $1^{+}$H$^+$ decomposes at $T \approx 500 \text{ K}$ and $T \approx 440 \text{ K}$ respectively, Fig. S2 and S4†), and neither moisture nor air-sensitive in the solid state even without complexation to a metal ion. As a consequence, BF$_4^-$ salts of these can be prepared and handled.

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as simple crystalline starting pro-ligands. A seemingly facile and reversible chemical reduction at room temperature leads to a doublet radical intermediate which is stable in solution under inert atmosphere (Fig. 1 – blue).

Using solvothermal conditions under inert atmosphere, the doubly reduced form MBqno (10) has been isolated in two crystalline forms, firstly as a cation in the monoprotonated species [10-H+]BF4 and secondly as a neutral biradicaloid chelating ligand in the metal complex [CdCl2(I0)]. Noteworthy is that dissolution of [10-H+]BF4 (Fig. 1 – purple) in DMF under inert atmosphere generates the same doublet radical species in solution (Fig. 1 – blue), thus granting the methylbiquinoxalen three different accessible redox states which are chemically reversible (vide infra).

Both 12+ and 10-H+ have been characterized by X-ray diffraction analysis (single crystal, powder), UV-Vis-NIR, IR and mass spectrometry, thermogravimetric analysis, cyclic voltammetry, spectroelectrochemistry measurements, as well as DFT and TDDFT (Time Dependent Density Functional Theory) calculation. Additional SQUID, NMR and EPR studies of 10-H+ in the solid state (SQUID) and dissolved in DMF (SQUID, NMR, EPR) support the presence of the doublet radical intermediate species in solution.

Results and discussion

Synthetic procedures

General procedure for synthesis of compounds [12+][BF4]2, [10-H+]BF4, and [CdCl2(I0)] (Fig. 1). Compound [12+][BF4]2 was obtained from Meerwein alkylation by mixing biquinoxalinine with an excess of trimethylammonium tetrafluoroborate in acetonitrile. After 5 d stirring at room temperature, the resulting solid was isolated by filtration, washed with Et2O, and recrystallized from water to give pale yellow/off-white crystals in 52% yield. Crystals of the direreduced monoprotonated species [10-H+]BF4 as well as of the metal complex [CdCl2(I0)] were obtained from solvothermal reaction using the solvent (here methanol) as the reducing agent. In a Schlenk tube, equipped with a hermetic Teflon stopper, [12+][BF4]2 were mixed with a mixture of MeOH/DMF. The Schlenk was purged with nitrogen and inserted in a solvothermal oven to follow a heat/flat/cool temperature programme, resulting in a pure phase of dark purple crystals of [10-H+]BF4 in 68% yield. Phase purity of [12+][BF4]2 and [10-H+]BF4 was checked by XRPD (Fig. S1 and S3†). For the synthesis of the cadmium complex the same precedent procedure was followed by adding a stoichiometric amount of CdCl2 in the Schlenk tube. A violet crystal of [CdCl2(I0)] was selected from a mixture of phases which always result from the reaction. In spite of many attempts it was only possible to generate the intermediate doublet radical species in solution, but not to isolate it as a crystalline salt.

Crystallography

Single crystals suitable for X-ray diffraction analysis were isolated for the dication [12+][BF4]2 (Fig. 1 – grey) for the direreduced monoprotonated form [10-H+]BF4 (Fig. 1 – purple) as well as for the metal complex [CdCl2(I0)]. A summary of the crystallographic
can be assigned as originating from the molecular entities
Salient C
the 16 H-atoms and 17 H-atoms (MALDI) (Fig. S5 and S6
one H-atom either in solution (FAB) or in the solid state
BF4 contains one independent [MBqn0-H+]([10-H+]
(FFS)0.6 e Å−3, respectively, that are comparable to those found in the literature.23 The pyrazinium structure is particularly clear on the 1,2,3,4 atomic sites, as well as a on the nitrogen on the N‡z position for 10-H+ each
data and refinement results are listed Table 1. Datasets for both structures were collected at 180 K. They were refined in the space groups P1 for [12+][BF4]2 and [CdCl2(I1)], and P21/n for [10-H+]BF4. All atoms including hydrogens were identified from Fourier difference maps. All atoms occupy general positions.
The crystallographic asymmetric units of the structures of [12+][BF4]2 and [10-H+]BF4 are shown in Fig. 2. The asymmetric unit of [12+][BF4]2 contains one independent half [MBqn1−] (12+) (N-methylquinoxalinium MQ+ type) and one independent ordered BF4−. On the other hand, the asymmetric unit of [10-H+]BF4 contains one independent [MBqn10-H+] (10-H+) as well as one independent ordered BF4− per compound.
In the case of 10-H+, after several attempts with different crystals and from different batches, the refinement is such that equal and significant residual electron density peaks (0.4–0.6 e Å−3; background 0.18 e Å−3) are clearly and systematically located on all aromatic and methyl hydrogen sites, as for 12+, but with an additional peak close to the nitrogen on 2′ position (Fig. 3, full explanation in the ESI Tables S1–S3†).
Thus for 10-H+, because 17 H-atoms are clearly present in the molecule, the latter has to be doubly reduced in order to match the singly positive charge balanced by one BF4−. This can be also viewed as the monoprotonated dication (leading to a 16 + 1 = 17H atoms trication) which was further reduced by two electrons to give the so-called direduced monoprotonated cation 10-H+.
Complementary FAB and MALDI-TOF mass spectrometry measurement performed on crystals of [10-H+]BF4 are reproducible and in agreement with a single species, differing only by one H-atom either in solution (FAB) or in the solid state (MALDI) (Fig. S5 and S6†). In particular the MALDI spectra in SM matrix shows a clear difference in peak intensity between the 16 H-atoms and 17 H-atoms (m/z = 288.3 and 289.3) which can be assigned as originating from the molecular entities 12+ and 10-H+.
As a result of the rotational freedom around the central C3–C3 bond, 12+ and 10-H+ could adopt any geometry between planar trans- (ϕ = 180°) and planar cis- (ϕ = 0°), where the dihedral angle is defined as ϕ = C2–C3–C3–C2. What we observe is that for 12+ the geometry is exactly planar trans, whereas for 10-H+ it is nearly planar cis (ϕ = 2.5°).
In the dication 12+; constituted of two equivalent MQ+ moieties, the structure reveals an aromatic π-conjugated system. The average C-C bond length in the benzene ring [1.392(3) Å] corresponds to the intermediate C-C distance of 1.39–1.40 Å observed in a benzene molecule with electron delocalization.22 In the pyrazinium moiety, the aromaticity is confirmed by the values of the C-C and C-N bond distances, in the range 1.410(2)-1.416(2) Å and 1.311(2)-1.383(2) Å respectively, that are comparable to those found in the literature.23 The pyrazinium structure is particularly clear on the 1,2,3,4 atomic positions, as shown by the two shortest distances N1–C2

Fig. 2 Crystallographic structures of 12+/10-H+: view of the asymmetric unit of [MBqn2+][BF4]2, [12+][BF4]2 (a) and [MBqn10-H+][10-H+]BF4, [10-H+]BF4 (b). Salient C–N and C–C bond lengths are indicated.

Fig. 3 Scheme of the C,N skeleton of the molecules overlaid with the residual 3D-Fourier difference map of 12+ (left) and 10-H+ (right). For each 12+ and 10-H+ an equal and significant residual electron density is clearly and systematically located on all aromatic and methyl hydrogen sites, as well as a on the nitrogen on the N‡z position for 10-H+ (highlighted with the yellow circle).
[1.311(2) Å] and N2–C3 [1.315(2) Å], alternating with an intermediate length for C2–C3 [1.410(2) Å] (Fig. 2).

In contrast, in the dimerized monoprotonated cation 10-H+ the different redox state of the species and the protonation of only one side of the molecule gives rise to an asymmetric environment. On one hand, the non-protonated MQ moiety remains essentially similar to 12+, where the bond lengths of both benzene rings and pyrazinium core are comparable [N1–C2 [1.316(2) Å], N2–C3 [1.344(2) Å], C2–C3 [1.413(2) Å]]. On the other hand, in the protonated half molecule, although the benzene ring remains aromatic (C–C ~ 1.39 Å), significant changes occur mostly within the pyrazinium core. The structure is such that the N1–C2, N1–C3, N2–C4 and C3–C4 bond lengths increase by 0.038 Å, 0.038 Å, and 0.075 Å respectively with a concomitant 0.04 Å decrease in the C2–C3 bond distance. The central C3–C3 bond also changes dramatically with a net decrease by 0.052 Å, with 1.424(2) Å for 10-H+ and 1.476(2) Å for 12+.

These structural features of 12+ and 10-H+ have been investigated by quantum chemical calculations using Turbomole24 (for details see Experimental section in ESIT). For both compounds geometrical parameters were fully optimized in the gas phase at the PBE0/def2-TZVPP level of theory (Tables S4, S5 and S8†). The PBE0 hybrid functional being known to provide the best estimates for such π-conjugated systems.25,26 The calculated bond lengths fit well with the experimental data with mean absolute errors (MAE) of 0.0068 Å (12+) and 0.0071 Å (10-H+), whereas the twist angle of the molecule differs only very slightly for 10-H+ (φexp = 2.5°, φcalc = 0°). For 12+ and 10-H+ the frequencies calculations, in addition with a scan of the relaxed potential energy surface along the twist angle (φ) for 12+, confirmed that the described structures correspond to the ground state of the molecules (Fig. S9 and S13, Tables S6 and S9†). The HOMO and LUMO of 12+, as well as the HOMO of 10-H+ are represented in Fig. 4.

As previously stated by A. di Matteo, “a reduction process increasing or decreasing the electron population, in the bonding or antibonding regions, strengthens or weakens the bonds, which become significantly shorter or longer”.27 Here, by populating the LUMO of 12+ (reduction process), a decrease and increase of the bond length is expected between atoms with bonding and antibonding characteristics respectively. This effect will be even more obvious between atoms where the HOMO and LUMO orbitals exhibit the opposite bonding and antibonding features. Thus focusing on the bipyrazinium core, the experimental parameters clearly show that the chemical reduction of 12+ will lead to a shortening of the C2–C3 (C′2–C′3) and C2–C3 bond lengths and to an increase of the N1–C2 (N′1–C′2), N1–C3 (N′1–C′3) and N3–C1 (N′3–C′1) bond lengths, both having respectively bonding and antibonding characteristic in the LUMO orbital.

The HOMO of 10-H+ and the LUMO of 12+ both exhibit comparable bonds with bonding and antibonding characteristics, which is in line with the above reasoning and the observed bond lengths in 10-H+. This also shows that the trans–cis isomerization which exists between 12+ (trans) and 10-H+ (cis) has no strong influence on the way the π-orbitals overlap.

However, the N′2–H situation in 10-H+ creates a dissymmetry within the molecule leading to a strengthening of the bonding and antibonding features in the pyrazinium core of the protonated MQ. This results in a partial stiffening of the system, in the sense that the electron density can be rather described as localized and alternate single and double bonds, like in polyene-like structure such as dipyran or dithiopyran.27,28 The N′2–H system also stabilizes the preferential cis-form in the solid. It creates a strong intramolecular hydrogen bond between the protonated and non-protonated amines (N2⋯H–N′2 ~ 2.25 Å), which is also confirmed by the strong bonding characteristic in the N2⋯H–N′2 region (HOMO of 10-H+ Fig. 4, detail of H-bonds Fig. S7†). This apparently stabilizes 10-H+ and might also act as a template by constraining the cis-form, i.e. the chelate function of the 2,2′-bipyridine.

**Spectroscopy, electrochemistry, magnetic susceptibility, spectroelectrochemistry and TDDFT**

Because of the poor solubility of [10-H+]BF4 and in order to get comparative data, both EPR, UV-Vis-NIR, cyclic voltammetry, spectro-electrochemistry, NMR and SQUID experiments have been performed in DMF. Nevertheless, a complementary UV-Vis experiment has been performed in MeCN for [12+][BF4]2 in order to fully identify its electronic transitions below 400 nm.

**Electron paramagnetic resonance (EPR)** (Fig. 5). A sample of [10-H+]BF4 dissolved in DMF yielded a 9-line spectrum at ambient temperature. This implies the presence of a paramagnetic species arising from either an impurity or a new redox state in solution. The rich hyperfine pattern could be modelled by coupling the single electron of the doublet radical to the I = 1 nuclear spin of all four nitrogen atoms (99.7% natural abundance). This is in line with what is expected for a doublet radical in solution. Simulation yielded spin Hamiltonian parameters giso = 2.0069 and Aiso = 6.05 × 10−4 cm−1 with a magnitude of the 14N hyperfine coupling comparable to that found for N4N′-dialkylated quinoxaline radical cations.29 The g-value is similar to those reported for phosphine-based cation radicals,30–33 but is slightly higher compared to the methylviologen radical cation (g ~ 2.003).7,34,35 This is consistent with the extra heteroatom in the six-membered ring hosting the unpaired spin. Consequently, the 14N hyperfine coupling is larger than for the archetypal viologens.29,34,35
The absence of additional hyperfine structure from coupling to the methyl and aromatic protons of the organic radical confirms these couplings are less than the isotopic linewidth $\sim 2 \times 10^{-4}$ cm$^{-1}$, which is invariant irrespective of the measurement temperature. Nevertheless this does not show conclusively whether the paramagnetic species is an impurity or not.

**Cyclic voltammetry (CV) (Fig. 6a).** In order to gain further insights into the redox process leading to the formation of the paramagnetic species, CV measurements were performed. Furthermore, these were done quantitatively by using the well-known ferrocene/ferrocenium system as internal reference. Regarding its reduction potential ($V$ vs. Fe/Fe$^+$), $1^{2+}$ ($E_{1/2} = -0.54$ V) is an intermediate but strong electron-acceptor between the prototype 3,3$'$-diazamethyliovologen (DMAV$^{+}$) ($E_{1/2} = -0.33$ V$^{39,20,28}$ and the methyliyilogen ($E_{1/2} = -0.89$ V).$^7$ Thus, in two experiments, either the dication $[1^{2+}][BF_4]$ (Fig. 1 - purple) or the doubly reduced monoprotonated cation $[1^{+}H]BF_4$ (Fig. 1 - blue) were dissolved in DMF. Both CVs are perfectly reversible and superposable for all investigated scan rates (50-5000 mV s$^{-1}$).

The deconvolution of the CV allows us to circumvent diffusion problems and to conclude that there is a single-electronic process by comparing with the current peak of the ferrocene/ferrocenium system as internal reference. This rules out the presence of any impurity and points to the oxidation of the doubly reduced monoprotonated cation $[1^{+}H]BF_4$ to give the doubly radical. Furthermore, the voltage separation between current peaks of ca. 200 mV is significantly higher than the theoretical value of 59 mV. This is in line with a redox process inducing a significant structural rearrangement such as a trans–cis isomerization.$^{37}$

**Magnetic susceptibility measurement using SQUID and NMR Evans’ method.** The magnetic behaviour of the redox states in solid and solution states should also show differences due to the diamagnetic nature of the solid state species $[1^{0}H]^+BF_4$, versus the paramagnetic nature of the doublet radical species in solution.

The magnetic susceptibility of $[1^{0}H]^+BF_4$ (Fig. 1 - purple) was measured on a polycrystalline sample over the 4–380 K temperature range under an applied dc field of 1 T. As shown in Fig. S10, the product of magnetic susceptibility with temperature is very close to zero ($<0.007$ cm$^3$ K mol$^{-1}$), which confirms the diamagnetic nature of $[1^{0}H]^+BF_4$ in the solid state.

Furthermore, crystals of $[1^{0}H]^+BF_4$ dissolved in DMF-d$^7$ under inert atmosphere exhibit a very strong diamagnetism at $-5000 \times 10^{-6}$ cm$^3$ mol$^{-1}$, even after correcting with the expected diamagnetic contribution calculated from Pascal constants. Given that we were expecting to see a paramagnetic signature, which would be expected for isolated paramagnetic species, this behaviour suggests the presence of superdiamagnetism. This has been previously observed in 3D inorganic compounds and was coined superdiamagnetism, in an obvious reference to the Meissner effect and the perfect diamagnetism observed in superconductors.$^{38,39}$ More recently superdiamagnetism was induced at low temperatures by the application of a magnetic field in a one-dimensional Pt$^{II}$ complex.$^{40}$ To the best of our knowledge this has never been observed in a purely organic based compound. In our case the diamagnetic susceptibility reaches 6% of the 1/4$\pi$ value expected for complete diamagnetism. This unexpected superdiamagnetism was confirmed using the Evans’$^{1}$H-NMR method$^{41}$ at room temperature from the strong negative chemical shift of the reference tert-butyl alcohol signal at $-27.3$ Hz in DMSO-d$^6$ and $-25.5$ Hz in DMF-d$^7$ in the presence of our studied molecule (Fig. S11 and S12). This is typical of an additional diamagnetic environment around the protons of the reference. The calculated magnetic susceptibilities, respectively $\chi = -962 \times 10^{-6}$ cm$^3$ mol$^{-1}$ and $\chi = -3510 \times 10^{-6}$ cm$^3$ mol$^{-1}$ are in line with the SQUID measurement, though the effect seems much weaker in DMSO.

This allows us to conclude from the CV, EPR, SQUID and NMR results that a single radical intermediate species is present in solution. Due to the perfect planarity of the molecule (no steric hindrance) and the fact that the spin density is located in $\pi$ orbitals, the apparent superdiamagnetic behaviour of such a radical in solution could be tentatively explained from the formation of extended stacks of radicals, in analogy to the unidimensional Pt$^{II}$ complex.$^{40}$ Strongly spin-paired $\pi$-dimers, as already observed in other $\pi$-radicals such as viologen and TTF,$^{34,42,43,45}$ would only provide a “normal” diamagnetic behaviour.

However the question remains: what is the nature of the radical species in solution? In their studies on the prototype $N,N'$-dimethyl-3,3$'$-bipyrazinium dication,$^{20}$ and the $N$-methylpyrazinium,$^{45,46}$ $N$-methylquinoxalinium$^{47}$ and $N$-methylphenazinium$^{48}$ cations, Kaim et al., Rouiller & Laviron, Eaton and Takagi et al. showed that under mild acidic or neutral conditions the chemical reduction of such kinds of cations is accompanied by protonation thus leading to 4-hydro-1-methyl-diazinium $7\pi$ electron radical cations in solution. We thus see two possibilities here. Firstly, the dication $[1^{2+}]$ is reduced and isomerises through the addition of an electron to give the cis-radical cation ($[1^-]$. Secondly, this cis-radical cation is formed and further protonated to give the cis-protonated radical cation ($1^{+}H^+$).

In order to investigate this, additional spectroelectrochemical (SE), UV-Vis-NIR and TDDFT experiments were performed. The purpose was to identify a possible structure/ optical signature relationship for the three different redox
Fig. 6  (a) Deconvoluted cyclic voltammogram (left) \([\text{[10-H}^+]\text{BF}_4\text{]} (0.56 \text{ mM})\) and \(\text{Fc} (0.65 \text{ mM})\) in DMF, with \(\text{TBAP (0.1 M)}\) as supporting electrolyte on Pt at 200 mV s\(^{-1}\). Evolution of the voltammogram (middle) and the current peak of each redox couple (right) with the variation of the scan rate. (b) Spectroelectrochemistry experiments of \(\text{[2+]}\) and \(\text{[10-H}^+]\text{(b)}\), 0.5 mM in 0.1 M TBAPF\(_6\)/DMF on platinum electrode, in thin layer conditions (close to 50 \(\mu\)m) under argon at 10 mV s\(^{-1}\) and 293 K. (b1 and b1\(_0\)) CVs in current vs. potential representation. (b2 and b2\(_0\)) 3D representation: (wavelength, time) vs. \(\Delta\text{absorbance}\). (b3 and b3\(_0\)) 3D representation viewed along the absorbance axis: (wavelength, time) vs. \(\Delta\text{absorbance}\). (b4 and b4\(_0\)) CV and \(\Delta\text{absorbance}/\Delta t\) (at 850 nm) vs. time, showing a good agreement between the optical and electrochemical data. (c) Overlay of UV-Vis-NIR and TDDFT spectra of \(\text{[2+]}\) [crystalline state (c3), dissolved in MeCN (c2)], \(\text{[10-H}^+]\) [solid state (c4)] and of the intermediate species \(\text{[1+}^+\text-H}^+\text{]}\) in DMF solution (c1).
states. Optical spectra have been computed using TDDFT at the
PBE0/def2-TZVPP level of theory in a closed shell (1\(^2\) + and 1\(^4\)-H\(^+\))
and unrestricted open-shell (1\(^1\) + and 1\(^3\)-H\(^+\)) configuration. To
reproduce the behaviour of the molecules in solution, the conductor-like
screening model COSMO has been applied, with \( \varepsilon = 37 \) as a mean value for a mixture of DMF and MeCN.

**Spectroelectrochemistry** (Fig. 6(b and b')). Known to be
a powerful tool for analysing the spectral signature of involved
species during an electrochemical process, time resolved
spectroelectrochemical experiments (SE) were performed, using the
already described home-made cell,\(^{45,47}\) between 350 and 900 nm
donolved 1\(^2\) (Fig. 1 - grey) and 1\(^4\)-H\(^+\) (Fig. 1 - purple) (0.5 mM
in 0.1 M TBAPF\(_6\)/DMF), in order to correlate the changes in their
spectra during reduction and oxidation processes.

In Fig. 6(b and b'), we see that both voltabsorptograms
(Δ(Abs) vs. initial species) are perfect mirror images of each
other, indicating that the two species are part of the same redox
couple. Furthermore, the spectra measured during the SE
experiments are quite similar to those obtained by dissolving crystals of 1\(^2\)-(BF\(_4\))\(_2\) and 1\(^4\)-H\(^+\)BF\(_4\) (Fig. 6c2 and 6c1). The results match perfectly those from the CV in terms of the
exclusive presence of one irreversible and stable 1\(^2\)/1\(^4\)redox
couple in solution.

**Solid and solution state UV-Vis-NIR optical properties of the
dication 1\(^2\).** Crystals of 1\(^2\)-(BF\(_4\))\(_2\) are yellow/white in the
solid state with no significant absorption in the visible range.
The band at \( \lambda_{\text{max}} = 380 \) nm corresponds to the HOMO → LUMO
transition (Fig. 6c3, Table S7\(^{46}\)). Crystals of 1\(^2\)-(BF\(_4\))\(_2\) dissolved
in MeCN give a colourless solution. The SE spectrum of 1\(^2\)
shows no absorption in the visible region (Fig. 6b), which is
confirmed from the UV-Vis spectrum with the lowest energy
absorption band slightly shifted compared with the solid state
spectrum at \( \lambda_{\text{max}} = 360 \) nm (Fig. 6c2).

**Solid state UV-Vis-NIR optical properties of the doubly
reduced monoprotonated cation 1\(^0\)-H\(^+.** Crystals of 1\(^0\)-H\(^+\)BF\(_4\) are
deeper purple and absorb across the whole UV-Vis range and
well into the near-infrared. There are broad absorption bands
centred at 270, 350, 570, 820, and 1000 nm (shouler \( \approx 1300 \) nm)
(Fig. 6c4). The TDDFT spectrum of 1\(^0\)-H\(^+\) matches perfectly
with the electronic transitions observed experimentally in the
range 200-950 nm, with \( \lambda_{\text{max}} = 820 \) nm corresponding to the
HOMO → LUMO transitions (Table S10\(^{f}\)). Analogous to the
behaviour of the methylviologen radical cation either in solution
or in the solid state,\(^{7,8}\) the additional transition present at
lower energy is most probably explained due to the formation of
extended stacks of 1\(^0\)-H\(^+\) species in the solid. This is
actually confirmed by the crystal packing which shows an infinite
1D stack of dimers of 1\(^0\)-H\(^+\) (Fig. S8\(^{f}\)).

**Solution state UV-Vis-NIR optical properties of the doublet
dradical.** In the electronic spectrum in solution for a concentra-
tion of \( \approx 2.4 \times 10^{-4} \) M the formation of the doublet radical
redox state is accompanied by a colour change from the deep
purple of the solid to blue with absorption bands centred at 430,
470, 580, 820 and 920 nm (shoulder \( \approx 1100 \) nm) in the Vis-NIR
region (Fig. 6c1). The SE spectrum of the intermediate radical
species looks very similar, although the position of the bands is
slightly shifted to higher energy by ca. 40 nm (0.2 eV) (Fig. 6b).
To investigate the two previously proposed hypothesis, namely
(a) the species in solution is a cis-radical 1\(^c\) or (b) the
species in solution is the cis-protonated radical dication 1\(^c\)-H\(^+\),
two different TDDFT spectra were simulated from fully
optimized geometries (Tables S12 and S13\(^{f}\)).

The TDDFT spectrum corresponding to hypothesis (a) cis-1\(^c\)
is shown with green lines in Fig. 6c1. This is clearly different
from the experimental spectrum in that it exhibits a single
broad transition at \( \lambda_{\text{max}} = 1200 \) nm and no transitions between
500 and 1100 nm.

The TDDFT spectrum corresponding to hypothesis (b) cis-
1\(^c\)-H\(^+\) is shown with violet lines in Fig. 6c1. The main electronic
transitions observed experimentally in the range 400 to 850 nm
in the SE spectrum are perfectly reproduced theoretically. The
lowest energy band \( \lambda_{\text{max}} = 820 \) nm corresponds to the "SOMO → LUMO transition (Fig. 6c1, Table S13\(^{f}\)).

This leads us to conclude that the radical intermediate
described species present in solution is intrinsically composed of the
cis-protonated radical dication 1\(^c\)-H\(^+\) (Fig. 1 - blue). Furthermore
the formation of extended stacks of radicals 1\(^c\)-H\(^+\) in solution
is substantiated by the presence of an additional band at lower
energy (\( \lambda = 920 \) nm, Fig. 6c1), analogous to the low energy
bands observed in aggregated viologens in solution.\(^{7,8}\)

We also note that although at a first glance the spectra of
1\(^c\)-H\(^+\) (Fig. 6c1) and 1\(^0\)-H\(^+\) (Fig. 6c4) look broadly similar, the
fact that the spectrum of 1\(^0\)-H\(^+\) exhibits no transitions between
400 and 500 nm (Table S10\(^{f}\)), provides a rapid fingerprint for
the identity of the redox state.

**Case of the metal complex [CdCl\(_2\)(1\(^0\))]: chelation and non-
Kekulé structure aspect of 1\(^0\).**

We provide a first example of a metal complex [CdCl\(_2\)(1\(^0\))] which
illustrates the ability of such a ligand to chelate metal ions
(Fig. 7). This proves that the neutral diradical species 1\(^0\) can be
obtained without being protonated unlike in the case of 1\(^0\)-H\(^+\),
which reinforces the suggestion that the additional proton on
N\(_2\) is labile in solution. In addition, the crystals obtained are air
stable, which is important for the further development of robust
compounds.

The fact that we were unable to isolate a pure phase of the
cadmium complex precluded further characterization.
Nevertheless, the reliable crystallographic dataset obtained for this compound has been used as a reference structure for further studies, especially regarding the investigation of the neutral diradical methylbiquinoxalen 1 and its electronic structure.

The asymmetric unit is made of one entire complex [CdCl2(1)]. The cadmium adopts a distorted tetrahedral coordination environment of D3d symmetry.

The 1 chelating ligand is cis-planar ($\phi = 1.3^\circ$) and very symmetric in terms of the relative bond distances (Table S15†). As for 1H+, the multiple bond character of the central C3-C3 bond ($d = 1.449 \text{Å}$) indicates a fully π-conjugated molecule, resulting in a homogenous distribution of the electron density.

Unrestricted DFT calculations for the singlet and triplet state of [CdCl2(1)] have been performed. The energy of the singlet state is slightly lower than the energy of the triplet state. The energy difference ($\Delta E_{ST}$) amounts to 0.14 eV (Table S14†). However, the ground state shows a significant spin contamination. The analysis of the natural orbitals shows that there is a pair of partly occupied orbitals, one with bonding character between C3 and C3′, the other antibonding with occupation numbers of 1.5 and 0.5, respectively, showing partial occupation of the LUMO (Fig. S14†). In addition, the spin density (Fig. S15†) indicates a weakening of the bonding character similar to what is seen in a broken symmetry state. This behaviour is in line with the biradicaloids observed, for example, for under coordinated Sn atoms.52

Regarding 1 in terms of a symmetric and non-protonated structure, not to be confused with the disymmetric closed-shell monoprotonated cation 1H+, such an open-shell behaviour with a small $\Delta E_{ST}$ gap was predictable due to the non-classical (non-Kekulé) electronic structure of 1, as previously identified by Matheis, et al.18,19 in the prototype molecule, 1,1′-dimethyl-3,3′-bipyrazinium (Fig. S16†). By definition, non-Kekulé means a molecule which is “fully conjugated, but each of whose Kekulé structures contains at least two atoms that are not π-bonded”.53,54 Furthermore, the $\Delta E_{ST}$ gap can apparently be significantly reduced by simply twisting the molecule ($\Delta E_{ST} = f(\phi)$ Fig. 7), thus giving a means to thermally access the triplet state.

Conclusions and Outlook

As is already known for viologens, the reversible and controlable structural, electronic, and electrostatic changes between the dication and radical species formed in solution invest the MBqn+ redox couple with electron mediation properties. This should be useful for many radical-assisted reactions (photocatalytic H2 production, C-C bond cleavage, artificial photosynthesis, electrolyte, electrochromism etc.).5,7,37,38

Surprisingly, chelation to a cadmium centre allowed us to access the diradical solid state biradicaloid species MBqn0 as ligand. Additionally, as expected for such a biradicaloid species, the calculated small singlet to triplet energy gap opens perspectives for a range of exotic electronic, optical, magnetic, dielectric etc. properties within a single system composed of purely organic moieties or incorporating metalloid centres.

Intriguingly, the superdiamagnetic behaviour observed for the doublet radical species in solution is such an unexpected property for an organic molecule that this demands further investigation.

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