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Hyperfine Coupling to the Bridging ¹⁷O in the Di-µ-oxo Core of a Mn^{III}–Mn^{IV} Model Significant to the Core Electronic Structure of the O₂-Evolving Complex in Photosystem II

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It is generally accepted that Mn in the S₂ state of the oxygenevolving center (OEC) is in the +3 and +4 oxidation states with μ -oxo linkages.^{1,2} X-ray crystallography has provided the resolution to model the OEC as a Mn₃CaO₄ cube connected to a fourth Mn via a μ -oxo bridge.^{3,4} ⁵⁵Mn electron nuclear double resonance (ENDOR) is highly consistent with the cuboidal structure.^{5,6}

Di- μ -oxo-bridged Mn^{III}-Mn^{IV} compounds are widely studied models for this S₂ state.^{1,7,8} Such models present a manganese multiline EPR pattern from S = 2 Mn^{III} and $S = \frac{3}{2}$ Mn^{IV}, antiferrromagnetically coupled to each other to yield a net spin $S = \frac{1}{2}$.⁶ Electron spin echo envelope modulation (ESEEM) and ENDOR of such Mn^{III}-Mn^{IV} models have resolved hyperfine couplings for Mn^{IV} and Mn^{III},^{9,10} protons,⁹ and liganding nitrogen.¹¹ However, the ubiquitous μ -oxygens which physically couple the Mn^{III} and Mn^{IV} have yet to have their electronic structure and electron-spin density elucidated.

Recently Tagore et al.1 showed incorporation of isotopically enriched oxygen into the di-µ-oxo bridges of Mn^{III}–Mn^{IV} models, simply by slow exchange from trace water in dry CH₃CN. For our work the di-µ-oxo Mn^{III}-Mn^{IV} bipyridyl dimer [(bpy)₄Mn₂^{III/IV}(µ- $O_{2}[(ClO_{4})_{3} (bpy = 2,2'' bipyridine) was synthesized according to$ literature methods.^{1,12} A CH₃CN (HPLC grade, Fisher) solution 2.5 mM in Mn^{III}-Mn^{IV} bipyridyl dimer was prepared, and trace H₂O, either as H₂¹⁶O or as isotopically enriched H₂¹⁷O (84% atomic enrichment in ¹⁷O, Isotec.), was added at 1 μ L of water to 200 μ L of CH₃CN. The exchange time of the water oxygen into the μ -oxo cross bridges at room temperature is about 20 min.¹ An equal volume of CH₂Cl₂ (Fisher, reagent grade) was added, and the sample precooled at -80 °C for several hours. The precooled 70 µL sample, in a 2.0 mm i.d., 2.4 mm o.d. quartz EPR tube, was glassed by plunging into liquid nitrogen. A glass inhibits paramagnetic species from aggregating upon freezing to prevent these aggregates from interfering with ENDOR. (CH₃CN-DMF also provided an even better glass. However, the DMF contained reductants that produced Mn^{II} artifacts but did not hamper di-uoxo ¹⁷O hyperfine measurements (see Supporting Information).)

X-band EPR (9.525 GHz) was carried out at 15 K as previously described.¹³ CW Q-band (34.1 GHz) ENDOR was performed under dispersion (χ') and rapid passage field-modulated conditions at 2 K.¹³ A nucleus, *Z*, with $I \ge 1$, namely, ¹⁷O ($I = 5/_2$) or ¹⁴N (I = 1), will have first-order ENDOR frequencies given as ${}^{Z}\nu^{\pm}_{ENDOR} = |{}^{Z}A/2 \pm {}^{Z}\nu + 3{}^{Z}P(2m-1)/2|$, where $-I + 1 \le m \le I$, ^ZA, and ^ZP are hyperfine and quadrupole coupling constants and ${}^{Z}\nu$ is the nuclear Zeeman frequency.¹⁴ At 12200 G ${}^{17}\nu = 7.03$ MHz and ${}^{14}\nu = 3.76$ MHz. For the ¹⁷O features here, $|{}^{17}A/2| \approx {}^{17}\nu$. The ${}^{17}\nu^{-}_{ENDOR}$ branch is close to zero frequency and is not resolved because $|{}^{17}A/2|$ and

¹⁷ ν cancel. The ¹⁷ $\nu^+_{\rm ENDOR}$ branch occurs at a frequency of approximately $|^{17}A/2 + ^{17}\nu|$ because as elsewhere, ¹⁷O quadrupolar splittings contribute only to line broadening.^{14,15} For ¹⁴N, the $^{14}\nu^+_{\rm ENDOR}$ branch, like the ¹⁷ $\nu^+_{\rm ENDOR}$ branch, is the one observable by rapid passage CW Q-band ENDOR.¹³

The X-band EPR signal from the di- μ -oxo Mn^{III}—Mn^{IV} bipyridyl dimer (Figure 1A) was similar to that reported by Cooper et al.¹² The outer features 300–600 G above the center (at g = 1.99 and \sim 3400 G) of the multiline pattern showed the most well-resolved structure. There was significant broadening of this structure brought on by the H₂¹⁷O. In Figure 1B, we compare second-derivative X-band features, which show significant ¹⁷O-induced broadening.

A comparison (Figure 2) of ENDOR signals from the Mn^{III–} Mn^{IV} bipyridyl dimers, respectively, exchanged with H₂¹⁶O and with H₂¹⁷O, showed a new feature from the ¹⁷O sample near 13.5 ± 1.0 MHz. This feature was best resolved ~300 to 600 G above and below the Q-band EPR line center (which occurs at g = 1.99 or 12240 G). The hyperfine coupling, derived from ¹⁷ ν ⁺_{ENDOR} = |¹⁷A/2 ± ¹⁷ ν |, was |¹⁷A| = 12.8 ± 1.0 MHz. A feature near 10.5 MHz occurred from all samples. We assign this as the liganding bipyridyl ¹⁴N nitrogen with an approximate hyperfine coupling of ||¹⁴A| = 13.5 ± 0.3 MHz; corresponding ||¹⁴A| couplings of the Mn^{III–}Mn^{IV} CYCLAM and TMPA complexes were, respectively, 9.2 and 11.2 MHz.¹¹

Correlating EPR Line Broadening with ¹⁷O Hyperfine **Coupling.** For *two* equivalent $I = \frac{5}{2}$ ¹⁷O-di- μ -oxo nuclei, an elevenline pattern is expected with peaks in the ratio of 1:2:3:4:5:6:5:4: 3:2:1 and a separation between peaks of $|^{17}A|$ (in Gauss). [Note that $2.79|^{17}A|$ (in Gauss) = $|^{17}A|$ (in MHz).] This packet shape is well approximated by a Gaussian function whose peak width between derivative extrema is $4.84 \cdot |^{17}A|$ (in Gauss).¹⁶ To replicate EPR line broadening, we convoluted the narrower second-derivative spectrum of the ¹⁶O-di-µ-oxo Mn^{III}-Mn^{IV} bipyridyl dimer with a Gaussian broadening function and, in the Supporting Information, with an exact 1:2:3:4:5:6:5:4:3:2:1 distribution. The convolutionally broadened EPR spectra were compared (Figure 1B) with the broadened spectrum of the ¹⁷O-di-u-oxo Mn^{III}-Mn^{IV} bipyridyl dimer. Best agreement was obtained with a Gaussian broadening function having a 22 \pm 3 G peak width between derivative extrema. This width corresponded to an intrinsic di- μ -oxo ¹⁷O coupling of $|^{17}A| = 4.6 \pm 0.6$ (in Gauss) = 12.9 ± 1.8 (in MHz). The coupling estimated from the EPR line width compares favorably with the hyperfine coupling of $|{}^{17}A| = 12.8 \pm 1.0$ MHz from ENDOR.

There has been little direct experimental hyperfine evidence on the oxygen hyperfine structure at μ -oxo cross bridges. There happens to be ENDOR hyperfine information from the bridging μ -oxygen between Fe(III) ($S = \frac{5}{2}$) and Fe(IV) (S = 2) in the di-Fe cluster of ribonucleotide reductase.¹⁵ There the hyperfine

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Figure 1. (Spectra A) First-derivative X-band EPR spectra of di-µ-oxo Mn^{III}-Mn^{IV} bipyridyl dimers exchanged with H₂¹⁶O (black) and H₂¹⁷O (red) in CH₃CN-CH₂Cl₂, recorded at T = 15 K, 6 G field modulation, 100 s signal averaging with a 2000 G field sweep, 2 mW microwave power, EPR frequency = 9.525 GHz. (Spectra B) Experimental second-derivative X-band EPR spectra of the same dimers exchanged with H₂¹⁶O (black) and H₂¹⁷O (red) and recorded in the 3700-4100 G range using 3 G field modulation; the blue overlay shows that the EPR spectrum from the di- μ -oxo-¹⁷O dimer can be obtained from the narrower line di- μ -oxo ¹⁶O dimer by convolution of that narrower spectrum with a Gaussian broadening function (of 22 G width between derivative extrema) using the Origin 7.0 data analysis program.



Figure 2. We present ENDOR of Mn^{III}-Mn^{IV} bipyridyl dimers exchanged with H216O (black) and H217O (red) in CH3CN-CH2Cl2 and in CH3CN-DMF glasses. The fields in the figure from top to bottom are approximately 500 and 600 G below g = 1.99 and approximately 300, 400, and 500 G above g = 1.99. ENDOR conditions: adiabatic rapid passage, T = 2 K, microwave power = $0.2 \,\mu$ W, 100 kHz mod = 5 G ptp, time constant = 90 ms, radio frequency power ≈ 20 W, radio frequency sweep rate = 2 MHz/ s, averaging time/spectrum = 1000 s, v_{EPR} = 34.10 GHz.

coupling of \sim 23 MHz is nearly double that measured here for the $Mn^{III}\text{--}Mn^{IV}$ bipyridyl dimer. The di-Fe cluster couplings should be larger because Fe tends to be more covalent than Mn and because the $S = \frac{5}{2}$ ferric ion has a spin-containing $d(x^2 - y^2)$ orbital directed for σ bonding toward the oxygen 2s orbital. This σ bonding should lead to sizable ¹⁷O Fermi hyperfine coupling, whereas, the di-Mn system has no such spin-containing $d(x^2-y^2)$ orbital. Antiferromagnetic coupling between paramagnetic metals depends on covalent electron-spin transfer through bridging ligands.¹⁷ DFT (density functional theory) computations on di-µ-oxo-Mn^{III}–Mn^{IV} systems indirectly utilize the μ -oxo covalent spin transfer to predict Mn^{III}-Mn^{IV} antiferromagnetic coupling.^{18,19} The present work provides experimental underpinnings for testing future high level DFT calculations that give a comprehensive prediction of spin density, di-µ-oxo hyperfine coupling, and Mn^{III}-Mn^{IV} antiferromagnetic coupling.

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Supporting Information Available: X-band EPR spectra are provided from the Mn^{III}-Mn^{IV} bipyridyl dimer in CH₃CN-DMF glassing solvent. A comparison is provided of line broadening simulations due to a Gaussian packet, a di-17O 1:2:3:4:5:6:5:4:3:2:1 packet, and a mono-17O 1:1:1:1:1 packet. This material is available free of charge via the Internet at http://pubs.acs.org.

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