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Detection of Water Molecules on the Radical Transfer Pathway of Ribonucleotide Reductase by ¹⁷O Electron–Nuclear Double **Resonance Spectroscopy**

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ABSTRACT: The role of water in biological proton-coupled electron transfer (PCET) is emerging as a key for understanding mechanistic details at atomic resolution. Here we demonstrate ¹⁷O high-frequency electron-nuclear double resonance (ENDOR) in conjunction with H_2^{17} O-labeled protein buffer to establish the presence of ordered water molecules at three radical intermediates in an active enzyme complex, the $\alpha_2\beta_2$ E. coli ribonucleotide reductase. Our data give unambiguous evidence that all three, individually trapped, intermediates are hyperfine coupled to one water molecule with Tyr-O···¹⁷O distances in the range 2.8-3.1 Å. The availability of this structural information will allow for quantitative models of PCET in this prototype enzyme. The results also provide a spectroscopic signature for water H-bonded to a tyrosyl radical.

Water is no longer known as just the solvent in which biochemical reactions take place but has been recognized as an essential player in these reactions.¹ Of particular interest is water involvement in electron transfer processes,²⁻⁵ its action as a proton wire⁶⁻⁸ or its role in proton-coupled electron transfer (PCET).⁹⁻¹² The identification of internal water in proteins can be achieved by X-ray diffraction.¹³⁻¹⁵ However, the crystallization of transient protein complexes is difficult. One key approach for detection of water in biological systems has been the use of ¹⁷O-enriched water in conjunction with magnetic resonance spectroscopy.¹⁶⁻²¹ Among these methods, electron paramagnetic resonance (EPR) can take advantage of high selectivity, as it detects nuclei only in the ligand sphere $(r \leq 1.5 \text{ nm})^{22}$ of paramagnetic centers.

EPR-based ¹⁷O hyperfine (hf) spectroscopy has been established for the detection of water binding to transitionmetal ions, where the oxygen usually coordinates to the ion and large hyperfine couplings (several MHz) can be observed.^{23–25} However, the most common water coordination motif to biological radicals occurs via H-bond interactions. The hf coupling to ¹⁷O is diminished in comparison to the metal ion coordination, due to a longer interspin distance. In addition, the small $^{17}{\rm O}$ gyromagnetic ratio $(\gamma_{\rm H}/\gamma_{^{17}{\rm O}}\approx7.4)^{26}$ and high nuclear spin (I = 5/2) have rendered the ¹⁷O hf coupling difficult to resolve.

Here we illustrate that high-frequency (94 and 263 GHz) electron-nuclear double resonance (ENDOR) spectroscopy can detect the ¹⁷O signal of ordered water molecules at an Hbond distance to radical intermediates in E. coli ribonucleotide reductase (RNR). The enzyme uses a long-range (32 Å) radical transfer (RT) to initiate nucleotide reduction (Scheme 1).²⁷ Three tyrosines $(Y_{356}, Y_{731}, \text{ and } Y_{730})$ are essential pathway residues, which form transient intermediates in the active complex $\alpha_2\beta_2$, consisting of the two homodimeric

Scheme 1. Current PCET Model of the 32 Å (Y_{122} to C_{439}) RT in *E. coli* RNR^{27,32,a}



^aRedox-active tyrosines 356, 731, and 730 are shown in cyan, electron transfer steps as red arrows, and proton transfer steps as blue arrows. Water molecules revealed in this study in respective site-selective mutants are shown in boldface.

subunits α_2 and β_2 .^{13,27} Water has been observed only crystallographically in inactive α_2 s without β_2 .^{13,14,28,29} Using site-selectively inserted tyrosine analogues to trap Y intermediates, 30 our previous $^{1}H/^{2}H$ ENDOR and DFT studies^{10,11,31} revealed H-bonds attributed to water molecules and proposed a role of water in RT. However, all active sites of proteins have exchangeable protons, and thus alternative interpretations to our water proposal were possible. Recently, a cryo-EM structure of $\alpha_2\beta_2$ was reported but the resolution

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was insufficient for water observation.³² Since our original proposals, studies using photo-RNRs and MD simulations implying waters in $\alpha_2\beta_2$ have appeared.^{33,34} However, water has never been directly detected.

Therefore, we explored the capability of $H_2^{17}O$ ENDOR spectroscopy by exchanging the RNR buffer with $H_2^{17}O$. $\alpha_2\beta_2$ - Y_{356}^{\bullet} was generated by a 2,3,5- $F_3Y_{122}^{\bullet}$ mutation in β_2^{35} whereas radicals at Y_{731} and Y_{730} were trapped by replacing the respective residue with 3-aminotyrosine (NH₂Y),³⁶ leading to $\alpha_2\beta_2$ -NH₂Y₇₃₁^{\bullet} and $\alpha_2\beta_2$ -NH₂Y₇₃₀^{\bullet}. The individual variants were mixed with the complementary α_2 or β_2 protein, CDP as a substrate, and ATP as an effector. The reaction was then quenched after a few seconds inside EPR tubes. Details on the sample preparation are given in sections SI1 and SI2.

Figure 1 displays representative 94 GHz ¹⁷O Mims³⁷ ENDOR spectra of the radical intermediates.

Each spectrum shows a sharp doublet centered on the ¹⁷O Larmor frequency (19.3 MHz at 3.4 T), which can be assigned to the central spin transition (m_I (¹⁷O) = +1/2 \rightarrow -1/2) of one coupled ¹⁷O nucleus. As ¹⁷O is contained only in the water of



Figure 1. 94 GHz ¹⁷O Mims³⁷ ENDOR spectra of (A) the intermediate Y_{356}^{\bullet} and (B) NH₂Y₇₃₁ \bullet and NH₂Y₇₃₀ \bullet at $B_0||g_y$ in the EPR line (T = 50 K, $\tau_{\text{Mims}} = 390$ ns). Acquisition time: 46 h (Y_{356}^{\bullet}), 40 h (NH₂Y₇₃₁ \bullet), and 18 h (NH₂Y₇₃₀ \bullet). Y_{356}^{\bullet} is from β_2 -F₃Y₁₂₂ \cdot/α_2 - $Y_{730}F$, which gives the highest radical yield (section SI2). Experimental spectra are shown in gray, with a Savitzky–Golay filter (fourth-order polynomial, 20-point window) shown in color. Simulations used Easyspin³⁸ (section S11.6) with parameters given in Table 1 and section S3. Solid lines (teal) represent transitions among $m_1 > 0$ manifolds and dashed lines (orange) those among $m_1 < 0$ manifolds. The simulation does not distinguish between dihedral $\theta = 0^{\circ}$ or $\theta = 180^{\circ}$.

the protein buffer, these sharp signals must arise from water molecules coupled to the radicals. Control experiments with only β_2 protein confirmed that the signal is associated to the radicals generated in $\alpha_2\beta_2$ (section SI3). The broad resonances at ± 2.5 MHz are attributed to other nuclear transitions of the I = 5/2 spin system, broadened by nuclear quadrupole coupling (Figure 1A). Additionally, we note asymmetry of the doublet, which arises from second-order effects of the quadrupole coupling (section SI4). A comparison of the ENDOR spectra at the low $(B_0||g_x)$ and high-field $(B_0||g_z)$ edges of the EPR line (section SI5) indicates an almost isotropic hf coupling, with the dipolar contribution dominating the line width of the central doublet. The ¹⁷O ENDOR spectra could be simulated with one ¹⁷O nucleus, from which the asymmetry of the central peaks resulted using full diagonalization of the spin Hamiltonian (Figure 1 and section SI8). Parameters are given in Table 1 and section SI3. The spectra of Y₃₅₆ and

Table 1. Simulation and DFT Parameters for 17 O and 1 H hf Couplings of Water in RNR Intermediates^{*a*}

	Y ₃₅₆ • sim/ DFT _{small}	$\underset{sim}{\overset{NH_{2}Y_{731}}{\bullet}}$	$\frac{\rm NH_2Y_{730} \bullet sim}{\rm DFT_{large}} /$
A_{x} (¹⁷ O)	0.43/0.19	0.70	0.65/0.24
A_{y} (¹⁷ O)	0.66/0.59	0.84	0.80/0.6
A_{z} (¹⁷ O)	0.70/0.65	0.89	0.89/0.6
$A(H_1)$	$6.2^{31}/7.4$	$\leq 2.5^{b}$	2.7 ^b /4.2
$ ho(^{17}{ m O})^{c}$ (%)	0.05		0.03
$r_{\rm O^{17}O}$ (Å)	2.9 ± 0.1	~3.0	~3.0

^{*a*}Except as noted, values are in MHz. Simulated quadrupole values for ¹⁷O were { $P_{xi}P_{yi}P_{z}$ } = {-0.02;-0.32;0.34} MHz with $e^2qQ/h = 6.8$ MHz and $\eta = 0.93$.⁴¹ ^{*b*}Values from ²H couplings in refs 11 and 10 using $\gamma_{^{1}H}/\gamma_{^{2}H}^{2} \approx 6.5$.²⁶ ^{*c*}Loewdin spin density⁴² from DFT. Uncertainties in coupling constants are less than 10% for simulations and up to 20% for DFT.

 $\rm NH_2Y_{731}^{\bullet}$ additionally contain signals close to the Larmor frequency not reproduced in the simulations, which likely originate from second-sphere water molecules at the subunit interface. Additional broadening is also observed, particularly at $\rm NH_2Y_{731}^{\bullet}$. It might be caused by conformational distribution of this residue, which was found to have flexibility.^{39,40,33}

To rationalize the coupling, we began with a DFT-optimized small model (25 atoms, details in section SI1) of Y_{356}^{\bullet} , as previous ENDOR spectra revealed ¹H couplings consistent with one water at the H-bond distance $r_{O-H} \approx 1.8$ Å.³¹ The ¹⁷O coupling from this model was $A_{max}(^{17}O) \approx 1$ MHz, slightly exceeding the present experimental value of 0.6 ± 0.05 MHz. To optimize the model, we computed dihedral θ (C₃-C₄-O··· H) and distance scans for ¹⁷O couplings, including the quadrupole tensor and the relative energies (section SI6). The DFT equilibrium distance always resulted in $r_{\rm O-H} \approx 1.8$ Å. We found that hf couplings and energies vary significantly with θ , while the quadrupole coupling is less affected (Figure S9A-C). A_{xyz} values of $\lesssim 1$ MHz are found for θ in the range $\lesssim \pm 30^{\circ}$ (or equivalently $150^\circ \leq \theta \leq 240^\circ$): i.e., close to the ring plane. Water coordination in the ring plane also results in minimal relative energies (Figure S9B). Importantly, predicted spin densities on ¹⁷O are <0.1% but are sufficient for producing a marked ¹⁷O isotropic splitting. The spin density transfer or spin polarization is likely related to the H-bond nature. A distance scan for the optimized dihedral of +20° predicts $A_{\text{max}}(^{17}\text{O})$ in the range 0.75–0.56 MHz (Figure S10A) for $r_{\rm O-H} \approx 1.8-2.0$ Å. Consideration of the DFT-predicted ¹H couplings (Figure S10B) and comparison with the experimental values³¹ of ~6.2 (H₁) and ~1.6 MHz (H₂) indicates that the water is located at $r_{\rm Tyr-O...^{17}O} = 2.9 \pm 0.1$ Å, corresponding to an $r_{\rm O-H}$ value of 1.9 ± 0.1 Å. Notably, the DFT-predicted dipolar coupling ($T_{\rm II} \approx 0.3$ MHz, section S16) is consistent with the point-dipole model and the aforementioned broadening of the sharp peaks.

Analogous DFT calculations were performed on the isolated amino tyrosyl NH₂Y[•].^{10,11,36} We observed a trend for the 17 O hf coupling in the dihedral and distance scans (section SI7) very similar to the Y[•] model. The calculation predicts that $A_{iso}(^{17}\text{O})$ of NH₂Y[•] is slightly larger (10–15%) than that of Y[•] at similar Tyr-O...¹⁷O distances and orientations, which could explain the experimental observation. The amino group introduces an asymmetry in the radical, and the energetically most favored water orientation is found at the opposite side of the amino group (Figure S11B). Nevertheless, this small model could not account simultaneously for the ¹⁷O and ¹H couplings observed for these two intermediates (Figure S12). As noted in a previous g_x calculation,¹⁰ the coordination of the water molecule to NH₂Y[•]s is influenced by the surrounding secondsphere residues, as these two intermediates are buried in $\alpha_{2}\beta_{2}$ (Scheme 1).

Having established that at least one water molecule is hfcoupled to each of the three intermediates, we examine their current molecular models in light of this finding. First, we consider the radical site Y_{356}^{\bullet} (Scheme 1). To explain the unprecedented g_x value of Y_{356}^{\bullet} ($g_x = 2.0062$), we previously proposed that two almost equivalent waters might be simultaneously bonded to Y_{356}^{\bullet} .³¹ While the present results are most consistent with the distance and orientation proposed for one water, the 94 GHz ¹⁷O ENDOR spectra (Figure 1A) cannot resolve a second water. We note that the spectral line shape and ¹⁷O hf coupling in Figure 1A are conserved in other RNR constructs that generate Y_{356}^{\bullet} (section SI8), including the $F_3Y_{122}^{\bullet}/E_{52}Q_{-}\beta_2$ double mutant used to solve a recent cryo-EM structure.⁴³

To gain spectral resolution, we recorded ${}^{17}O$ ENDOR spectra of Y_{356}^{\bullet} at 263 GHz/9.4 T (Figure 2).

The results illustrate that the line width of the central doublet substantially narrows, particularly at $B_0 || g_z$ (Figure 2). Despite the narrowing, a factor of approximately 2 from 94 to 263 GHz, we cannot discern two distinct ¹⁷O contributions. Simulations of the 263 GHz spectra with the same parameters used at 94 GHz reproduce the line narrowing and support the analysis at 94 GHz. The lack of evidence for a second, almost equivalent water H-bonded to Y_{356}^{\bullet} strongly suggests that the two-water model has become very unlikely and alternative explanations for the shifted g_x value of Y_{356}^{\bullet} will have to be examined. The precise location of second-sphere residues might play a role, ¹² which will require further experimental and computational investigation.

For the radical intermediates in the subunit α , a previous combined ENDOR/DFT model of NH₂Y₇₃₀ • proposed a water molecule coordinated in plane at a distance $r_{\rm NH_2Y_{730}-O...} \approx 3.0$ Å.¹⁰ The present results are consistent with this model and provide direct evidence for this postulated water in the enzyme complex $\alpha_2\beta_2$ -NH₂Y₇₃₀•. The DFT-predicted hf parameters (DFT_{large}) for this large model (140 atoms) are reported in Table 1, and the model is displayed in section SI9.



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Figure 2. Comparison of 94 and 263 GHz Mims ENDOR of Y_{356}^{\bullet} at the three canonical positions in the EPR line. Total acquisition time for 263 GHz (T = 20 K): 18 h ($B_0 || g_x$), 10 h ($B_0 || g_y$), and 11 h ($B_0 || g_z$). Experimental spectra are shown in gray, with a Savitzky–Golay filter (fourth-order polynomial, 10 point window) in color. Simulations of 263 GHz spectra are in black with parameters as for 94 GHz (see Table 1 and Table S4).

Finally, for $\alpha_2\beta_2$ -NH₂Y₇₃₁, large-scale (215 atoms) DFT calculations previously proposed three models of the trapped intermediate (section SI10). Among these models, only one (model 3, Figure S15) contained a water molecule at an H-bond distance. The DFT-predicted ¹⁷O hf couplings of model 3 (~2.5 MHz), however, largely exceed the present experimental values (Table S5). However, this DFT model did not include residues from the β subunit, which we now know are close to this residue in the active complex.⁴³ Therefore, the model will require further refinement. Nevertheless, the present results give evidence for a water molecule coordinated almost in the plane of NH₂Y₇₃₁.

In conclusion, we have reported the capability of ¹⁷O highfrequency ENDOR to detect water H-bonded to tyrosyl radicals. The spectroscopic approach led to the first detection of ordered water molecules at three trapped radicals proposed to be representative of Y[•] intermediates in the PCET of *E. coli* RNR. These results verify previous hypotheses on the presence and role of water in the RNR mechanism and provide a new starting point for computational studies. Knowledge of this ¹⁷O signature will also be generally useful for many other biological systems, in which tyrosyl radicals are involved.

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ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c01359.

Experimental procedure, radical yield determination, ENDOR spectra of Y_{122}^{\bullet} and $F_3Y_{122}^{\bullet}$, ENDOR spectra of I = 5/2 nuclei, orientation-selective ENDOR spectra, DFT models of Y[•] and NH₂Y[•], ¹⁷O Y₃₅₆[•] spectra of different mutants, previous large DFT models of $\alpha_2\beta_2$ -NH₂Y₇₃₀[•] and $\alpha_2\beta_2$ -NH₂Y₇₃₁[•] (PDF)

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Notes

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