



Biophysics Hot Paper



In-Cell Characterization of the Stable Tyrosyl Radical in *E. coli* Ribonucleotide Reductase Using Advanced EPR Spectroscopy

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Abstract: The *E. coli* ribonucleotide reductase (RNR), a paradigm for class Ia enzymes including human RNR, catalyzes the biosynthesis of DNA building blocks and requires a di-iron tyrosyl radical (Y_{122}^{\bullet}) cofactor for activity. The knowledge on the *in vitro* Y_{122}^{\bullet} structure and its radical distribution within the $\beta 2$ subunit has accumulated over the years; yet little information exists on the *in vivo* Y_{122}^{\bullet} . Here, we characterize this essential radical in whole cells. Multi-frequency EPR and electron-nuclear double resonance (ENDOR) demonstrate that the structure and electrostatic environment of Y_{122}^{\bullet} are identical under *in vivo* and *in vitro* conditions. Pulsed dipolar EPR experiments shed light on a distinct *in vivo* Y_{122}^{\bullet} per $\beta 2$ distribution, supporting the key role of Y^{\bullet} concentrations in regulating RNR activity. Additionally, we spectroscopically verify the generation of an unnatural amino acid radical, $F_3Y_{122}^{\bullet}$, in whole cells, providing a crucial step towards unique insights into the RNR catalysis under physiological conditions.

Introduction

Ribonucleotide reductases (RNRs) catalyse the conversion of nucleotides to their corresponding deoxynucleotides in every living cell.^[1] Due to their central role in DNA replication and repair, they have been a target for cancer therapeutics.^[2] *E. coli* class Ia RNR, a paradigm for class Ia enzymes including human RNR, consists of two homodimeric subunits, $\alpha 2$ and $\beta 2$. The active $\alpha 2\beta 2$ complex is formed transiently upon substrate and effector binding.^[3] The catalytic reaction of *E. coli* class Ia RNR relies on at least five redox-active amino acids, which are involved in an unprecedented reversible radical transfer that occurs over $\approx 32 \text{ \AA}$.^[4] β harbours the stable di-iron tyrosyl radical cofactor Y_{122}^{\bullet} and the redox-active tyrosine residue Y_{356} . α contains the active site for nucleotide reduction, two allosteric effector binding sites and redox active residues Y_{731} , Y_{730} , and C_{439} that are a part of the radical transfer pathway. Y_{122}^{\bullet} reversibly oxidizes

C_{439} to a thiyl radical, which in turn initiates the irreversible catalytic reaction.

As di-iron tyrosyl radical cofactor initiates the catalytic reaction, radical generation at this tyrosine residue is essential for RNR activity. Y^{\bullet} is generated by self-assembly requiring Fe^{2+} and O_2 , both in structurally homogenous *E. coli* and *Saccharomyces cerevisiae* (*S. cerevisiae*) class Ia RNRs.^[5] However, radical generation efficiency varies among different organisms and studies (*in vivo* vs. *in vitro*).^[2b,5] *In vitro* studies with *E. coli* RNR showed that usually 1.2 $Y^{\bullet}/wt-\beta 2$ were generated, suggesting $\approx 60\%$ of the $\beta 2$ population is active and $\approx 40\%$ possesses inactive diferric clusters without any Y^{\bullet} s (“two or none” radical distribution model).^[6] *In vivo* Y^{\bullet} levels, however, were substoichiometric compared to that of iron loading.^[7] Furthermore, *in vivo* radical distribution in *E. coli* (one or two radicals per $\beta 2$) remains unknown. In contrast, studies with *S. cerevisiae* revealed stoichiometric amounts of Y^{\bullet} with 1 Y^{\bullet} per $\beta\beta'$ heterodimer both *in vivo* and *in vitro*.^[5,8] These studies together demonstrated that *S. cerevisiae* RNR activity is not regulated by the modulation of Y^{\bullet} concentrations, but Y^{\bullet} concentrations play a key role in regulation of *E. coli* RNR activity in cells.

The rate-limiting step in the *E. coli* RNR catalytic reaction, which is the conformational change(s) upon substrate and allosteric effector binding, has prevented spectroscopic detection of other redox-active amino acids in wild-type enzyme.^[9] The involvement of Y^{\bullet} intermediates in RNR catalytic reaction was evidenced by site-specific incorporation of unnatural amino acids (UAAs) with altered pK_a s and reduction potentials such as DOPA, NH_2Y , NO_2Y , and F_nYs .^[10] Radical formation on these UAAs not only allowed characterization of tight $\alpha 2\beta 2$ complex but also provided valuable insights into proton-coupled electron transfer steps within RNR via transient absorption^[11] and advanced electron paramagnetic resonance (EPR) spectroscopies.^[12] For example, distance measurements conducted using double electron-electron resonance (DEER, also known as PELDOR)^[13] combined with incorporation of UAAs provided structural constraints for redox-active pathway tyrosine residues and displayed an asymmetry within the $\alpha 2\beta 2$ complex.^[10c,14] Same strategy was further used to gain insights into the relative redox potentials of three subsequent pathway tyrosyl radicals,^[10g] and to detect the flexibility of pathway residue Y_{731} in the $\alpha 2\beta 2$ complex for the first time.^[12b] Very recently, near-atomic-resolution cryo-EM structure of the active $\alpha 2\beta 2$ complex, which is asymmetric, was achieved by employing $F_3Y_{122}^{\bullet}$.^[4a]

One of the most challenging tasks for techniques used in structural biology is to achieve high resolution within the native environment of biomolecules, that is, the cells. Among

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the methods employed, EPR spectroscopy is an emerging tool for in vivo studies.^[15] It is the natural choice for investigating biomolecules with intrinsic paramagnetic centres, and it has been central to our understanding of not only RNRs but also other fundamental enzymes such as hydrogenases, nitrogenases and photosystem II.^[16] Here, we have generated Y_{122}^{\cdot} , the stable di-iron tyrosyl radical cofactor of *E. coli* class Ia RNR in whole cells. We characterized its in-cell conformation and H-bonding environment at nanometre-scale resolution using multi-frequency EPR and electron-nuclear double resonance (ENDOR) spectroscopies. Subsequently, we detected in vivo distances between two Y_{122} 's by DEER (Figure 1). These orientation-selective experiments revealed insights into not only the in vivo structure and dynamics of Y_{122}^{\cdot} and $\beta 2$ subunit but also the in vivo radical distribution within $\beta 2$. This first example of distance measurements between two native paramagnetic centres of an enzyme within intact cells where the protein under investigation is expressed adds another dimension to similar attempts where in a spin label is directly introduced in the cells.^[17] Additionally, we have site-specifically incorporated 2,3,5-trifluorotyrosine (F_3Y) at residue 122, generated and identified its radical form $F_3Y_{122}^{\cdot}$ in the cells, and obtained in vivo distances between F_3Y_{122} 's. These results verify the generation of an unnatural amino acid radical in whole cells and show that F_3Y incorporation does not alter protein structure within cells.

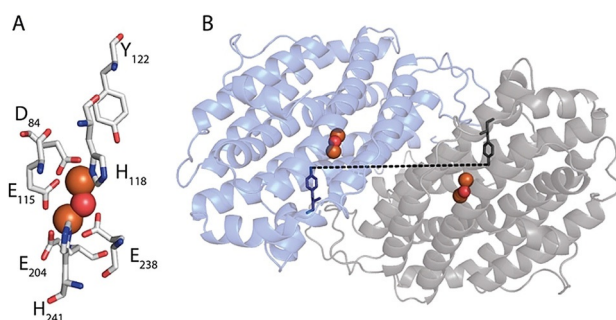


Figure 1. A) Environment of Y_{122} in the $\beta 2$ subunit of *E. coli* RNR. Y_{122} (grey sticks) is located next to a di-iron centre (Fe: orange and O: red-sphere), which is ligated by six amino acids (white sticks). B) The distance measured in this work is shown with a dashed line in the homodimeric $\beta 2$ subunit (PDB 5CI4).

Results and Discussion

Generation, Detection, and Characterization of Y_{122}^{\cdot} in Cells

Presence of the di-iron centre is prerequisite for the radical state of Y_{122} and for the catalytic activity of *E. coli* RNR. Previously, it was shown that in vitro incubation of iron-free, apo- $\beta 2$ with Fe^{II} and O_2 resulted in the reconstitution of the radical state of Y_{122} , and thus led to recovery of the RNR activity.^[18] Here, we extend this technique to in vivo conditions. Apo- $\beta 2$ in *E. coli* was over-expressed because Y_{122} concentrations are below detection limit under normal growth conditions.^[7] Cell cultures were harvested, washed several times and resuspended in a buffer containing Fe^{II} for

10 min. Next, the cell suspension was saturated with O_2 and transferred into EPR tubes (Supporting Information (SI) 1.1). Prior to EPR experiments, the viability of the cells was checked using cell counting experiments (SI 1.3). These proved that the cells used for EPR measurements were intact and alive. In addition, the possibility of cell damage, and thus high amounts of protein leakage from the cells into the suspension buffer was ruled out using polyacrylamide gel electrophoresis and EPR experiments (SI 2.1).

Comparison of 9.6 GHz EPR spectra of treated and untreated whole *E. coli* cells showed that the Fe^{II} and O_2 treatment led to the generation of a radical species whose EPR signature is similar to that of Y_{122}^{\cdot} (SI 2.2). To examine the generated radical, we recorded multi-frequency (9.6, 34 and 94 GHz) EPR spectra of treated whole *E. coli* cells (termed in-cell) and analysed the data using spectral simulations (Figure 2). Principal g values and hyperfine coupling parameters related to ring- and β -methylene protons reported for *E. coli* Y_{122}^{\cdot} under in vitro conditions (Table S3 and Scheme S1) were in excellent agreement with the multi-frequency EPR/ENDOR dataset.^[19] These results showed that the radical generated within cells is Y_{122}^{\cdot} having an average bulk concentration of $18 \pm 5 \mu M$ (see SI 2.3 for concentration determination).

94 GHz EPR spectrum of in vitro Y_{122}^{\cdot} is also shown in Figure 2 for comparison. At high frequencies the EPR line shape is dominated by g - and hyperfine anisotropy and is highly sensitive to molecular environment of tyrosyl radi-

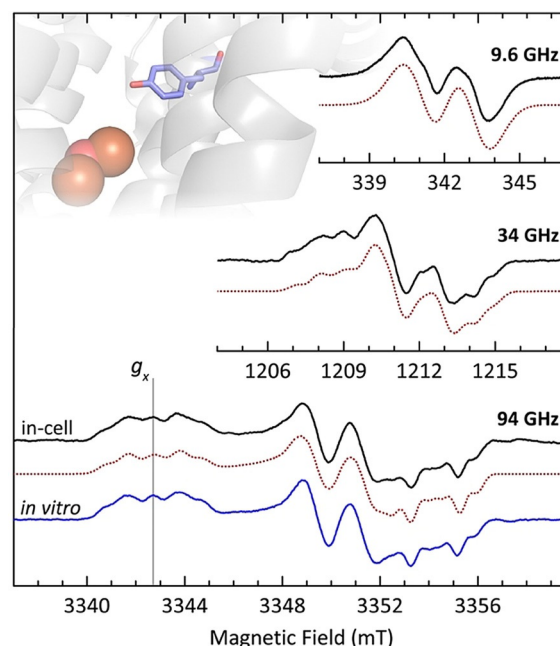


Figure 2. Continuous wave EPR (9.6 GHz, top) and first-derivative pulse EPR (34 GHz, middle and 94 GHz, bottom) spectra of Y_{122}^{\cdot} in whole *E. coli* cells are shown in solid black lines along with the corresponding EasySpin^[23] simulations in dotted red lines. See SI 2.6 for the background correction details. The 94 GHz EPR spectrum of in vitro Y_{122}^{\cdot} (solid blue line) is shown for comparison. Position of the g_x value is displayed with a grey vertical line. Inset: The isolated di-iron centre (orange and red spheres) and residue Y_{122} (blue sticks) are shown within the protein environment.

calculated.^[12d,20] The g_x value reports on the electrostatic environment of a tyrosyl radical and is affected by the changes in the radical's local environment, such as addition or loss of an H-bond.^[12d,20a,21] Furthermore, distributions in g_x value indicate multiple molecular orientations and/or radical environments. Another important structural parameter is the strength of the hyperfine couplings of C β -methylene protons. They are related to the dihedral angle between C β -H bonds and the ring plane, and therefore provide information on the structure of tyrosyl radicals.^[22]

Therefore, a substantial change in the structure of Y_{122}^{\bullet} would result in a different β -methylene proton hyperfine coupling, and thus distinct EPR line shape (SI 2.4), as it has been observed previously for residue Y_{731} .^[12b] Here, we detected a g_x value for in-cell Y_{122}^{\bullet} that is identical to that of in vitro one, resulting in nearly identical EPR line shapes for in-cell and in vitro Y_{122}^{\bullet} . The only marginal difference observed in the line shapes is due to the Mn^{2+} species that is inherent to the cells (see SI 2.5 and 2.6, as well as a comparison of two in-cell W-band spectra without background correction in Figure S4 B, SI 2.4). Overall, all of the EPR data and simulations clearly show that Y_{122}^{\bullet} generated within the cells is a single radical species with one set of magnetic parameters and a well-defined orientation. Additionally, our results demonstrated that structure and environment of Y_{122}^{\bullet} in the cells are highly similar to those of Y_{122}^{\bullet} in vitro. These results were not unexpected considering the isolated nature of Y_{122}^{\bullet} inside the protein, around 10 Å away from its surface.

H-Bonding Environment of Y_{122}^{\bullet} in Cells

Next, we employed ENDOR spectroscopy at 34 GHz to probe the H-bonding environment of Y_{122}^{\bullet} in the cells. Orientation-selective ENDOR reports on the number, separation distance (usually ≤ 15 Å) and orientation of the magnetic nuclei that are coupled to the observed radical, and thus provides structural information.^[24] We recorded ENDOR spectra at three field positions within the Y_{122}^{\bullet} EPR line corresponding to g_{xy} , g_y and g_{yz} molecular orientations shown in SI 2.6. Figure 3A illustrates the orientation-selective 1H Davies ENDOR spectra of Y_{122}^{\bullet} in the cells (black lines) and in vitro (blue lines) with the corresponding simulations (red dotted lines). Based on the previous 1H ENDOR studies on Y_{122}^{\bullet} conducted at 9 up to 263 GHz,^[19c,25] seven internal 1H hyperfine couplings were included in the simulations of the in vitro and in-cell spectra (see Table S3 and Scheme S1). As expected, the simulations were in excellent agreement with in vitro ENDOR and multi-frequency EPR data.

The 1H Davies ENDOR line shape of the in-cell Y_{122}^{\bullet} contained additional spectral features compared to the in vitro Y_{122}^{\bullet} . Therefore, the in vitro spectrum was subtracted from the in-cell one after normalizing the spectra to the C β -methylene proton ENDOR intensities (couplings around ± 28 MHz in Figure 3A). The resulting spectrum (orange trace in Figure 3B) is very similar to the 1H Davies ENDOR spectrum recorded for *E. coli* cells that contain overexpressed apo- $\beta 2$ without tyrosyl radicals (labeled *E. coli* cells, purple

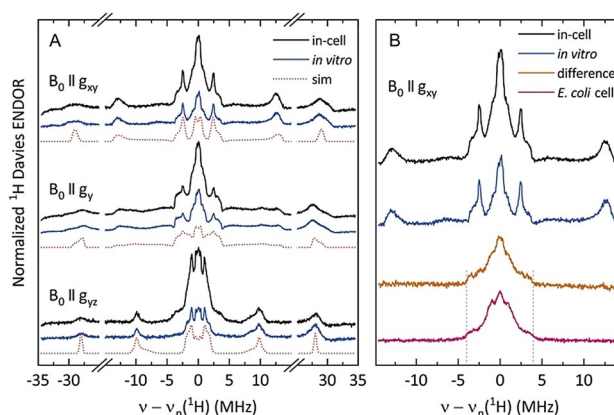


Figure 3. A) Orientation-selective 1H Davies ENDOR spectra of in-cell (black) and in vitro (blue) Y_{122}^{\bullet} recorded at 34 GHz at three field positions corresponding to $g = 2.0094$ ($B_0 \parallel g_{xy}$), 2.0059 ($B_0 \parallel g_y$), and 2.0005 ($B_0 \parallel g_{yz}$). Simulations (red dotted lines) were performed using the spin Hamiltonian parameters determined in Ref. 18c (Table S3). B) ENDOR spectra ($B_0 \parallel g_{xy}$) of the in-cell (black) and in vitro (blue) Y_{122}^{\bullet} ; the scaled subtraction result (orange), and Mn^{2+} -related 1H ENDOR features of *E. coli* cells not treated with iron (purple). Dashed vertical lines indicate spectral features associated with $[Mn(H_2O)_6]^{2+}$.

trace). This comparison clearly demonstrated that the additional proton hyperfine couplings observed for the in-cell sample are not related to Y_{122}^{\bullet} ; but rather to Mn^{2+} in *E. coli* cells as the only clearly detectable paramagnetic species in these cells was Mn^{2+} (SI 2.6). Indeed, the additional 1H ENDOR features detected in *E. coli* cells closely resemble the in-cell Mn^{2+} ENDOR line shape found in literature,^[26] with the broader shoulders reaching ± 3.8 MHz around the 1H Larmor frequency (dashed vertical lines) indicative of the water-ligated Mn^{2+} species.^[27] We note that the assignment of additional proton hyperfine couplings to the Mn^{2+} species is in line with identical g_x values detected for in-cell and in vitro Y_{122}^{\bullet} . An extra H-bond to Y_{122}^{\bullet} would make its local environment more polar and shift its g_x to lower values, as reported previously for tyrosyl radicals of distinct proteins coupled to hydrogens of similar strengths.^[12d,21] Furthermore, the 1H hyperfine coupling observed in the difference spectrum, which is similar to the A_y component of C $_3$ /C $_5$ protons, would result in additional splittings in the 94 GHz EPR spectrum as shown in Figure S4, which were not observed. Overall, the analysis of our orientation-selective 1H Davies ENDOR data combined with the g_x value and EPR line shape information demonstrated that the number of hydrogens coupled to Y_{122}^{\bullet} as well as the coupling strengths are the same in the cells and in vitro.

In-Cell Distance and Radical Distribution Information Obtained Using DEER Spectroscopy

In order to get insights into the wt- $\beta 2$ structure and in-cell radical distribution, we measured the distance between two Y_{122}^{\bullet} 's, each residing in a β monomer of *E. coli* RNR in whole cells (Figure 4). As orientation selectivity, which occurs when dipolar spectrum is dependent on the selected g -tensor

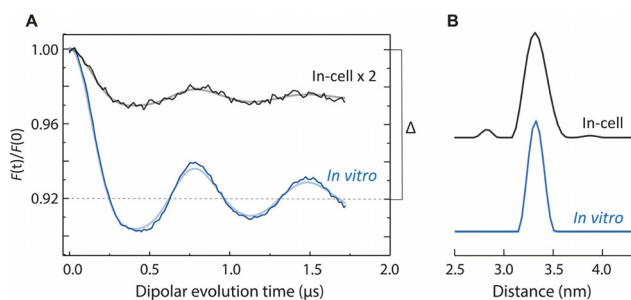


Figure 4. Background- and phase-corrected, normalized and orientation-averaged 34 GHz in-cell (black) and in vitro (blue) DEER traces of β_2 subunit of *E. coli* RNR (A) shown along with obtained $Y_{122^*}\text{-}Y_{122^*}$ distance distributions (B). DEER time traces were analysed with DeerAnalysis2019^[32] and the fits are overlaid in grey (in-cell) and light blue (in vitro). Details of the experiments and analyses are given in SI 2.7. The original in-cell trace was magnified by a factor of two for better visualization.

orientations,^[28] was reported previously for the in vitro $Y_{122^*}\text{-}Y_{122^*}$ pair,^[29] we employed orientation-selective DEER spectroscopy. Three DEER traces were recorded at distinct g -tensor orientations covering the whole EPR line, and the obtained dipolar traces were summed (see SI 2.7). Fourier-transform of in-cell and in vitro summed traces led to almost ideal Pake patterns (SI 2.7). The resulting orientation-averaged form factors and the extracted distance distributions are displayed in Figure 4. The analysis of the in-cell DEER data revealed a main mean distance at 3.32 nm with a distance distribution of $\sigma = 0.09$ nm (σ is the standard deviation of the distribution, which is assumed Gaussian). This distance belongs to the in-cell $Y_{122^*}\text{-}Y_{122^*}$ pairs, as it is identical to in vitro $Y_{122^*}\text{-}Y_{122^*}$ distances detected in this work (3.32 nm with $\sigma = 0.07$ nm) and previously (3.31 nm^[30]). An additional peak with an extremely small intensity emerged from the analysis of the in-cell data. It is most likely an artifact caused by the orientation-averaging procedure due to low signal-to-noise ratios (SNRs) of individual in-cell DEER traces, as well as the overlap between Y_{122^*} and Mn^{2+} EPR features (see SI 2.8 for details). The detected narrow in-cell distance distribution ($\sigma < 0.1$ nm) and the presence of orientation selectivity demonstrated the rigid nature of Y_{122^*} s, and more importantly, of the β_2 subunit within the cells. This conclusion is also supported by our EPR data that established Y_{122^*} generated within the cells as a single and structurally defined radical species.

Being able to detect a $Y_{122^*}\text{-}Y_{122^*}$ distance in the cells is not self-evident because in vivo radical distribution of β_2 is not known. In order to investigate the radical distribution within the cells, which was suggested to play a key role in RNR regulation and activity,^[2b,7] we analysed the modulation depth parameter Δ (shown in Figure 4A for the in vitro trace). Δ delivers information on the fraction of spin pairs in a sample, and is affected by the presence of monomers and other paramagnetic species.^[31] Comparison of DEER traces recorded at the same orientation and under identical experimental conditions showed that Δ values of in-cell samples were significantly lower than that of the in vitro one (1.3–2.1% vs. 5.3%, see Figure S10, as well as the red and black

traces in Figure 5C). A strong reduction in Δ for the in-cell samples was expected, given the large contribution from Mn^{2+} to the EPR signal at the observe position (SI 2.5 and Figure 5A). Note that the variations observed in the in-cell modulation depths were batch-dependent despite the identical sample preparation procedure (Figure S10). This outcome was not surprising given the number of various factors that could affect cell growth and were not controlled under our experimental conditions.

To decipher whether Mn^{2+} solely accounts for the reduction of Δ in the cells, we quantified its effect on Δ by adding various amounts of Mn^{2+} to in vitro samples (Figure 5, see Table S2 for details). As expected, the resulting calibration curve, which shows Δ as a function of relative Y_{122^*} contribution to the refocused echo, revealed a clear reduction in the Δ values with the increasing Mn^{2+} fraction (Figure 5C/D and SI 2.10 for details of the analysis). Δ of the in vitro wt- β_2 samples without Mn^{2+} depends neither on the protein nor on the spin concentration of the sample,^[29] as wt- β_2 dimer in vitro either contains a $Y_{122^*}\text{-}Y_{122^*}$ pair or no radical at all (“two or none” model^[6a]). Therefore, Mn^{2+} is the only reason for the reduced Δ of the in vitro samples. We note that Mn^{2+} ions are homogeneously distributed, and thus do not contribute to the detected distances and distributions throughout this work (Figure S9).

As seen in Figure 5D, the in-cell Δ values (red data points) fall consistently below the in vitro calibration curve (grey shaded area). This result suggests that the reduction of

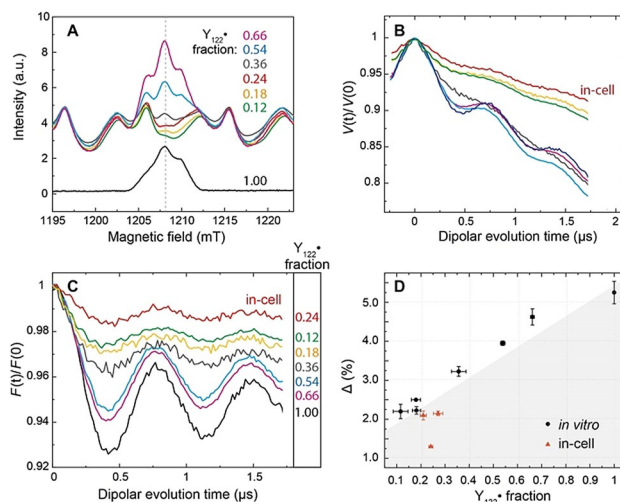


Figure 5. Modulation depth analysis of in vitro β_2 samples containing different equivalents of Mn^{2+} (see Table S2 for sample specifications). A) Q-band field-swept EPR spectra of RNR/ Mn^{2+} mixtures and an in-cell sample (red trace) recorded using refocused spin echo with the pump pulse applied at the primary echo position. The importance of the pump pulse in refocused echo experiments involving a high-spin species has been reported previously.^[33] The spectra were normalized to the Mn^{2+} intensity. B) Primary DEER traces and C) form factors of the corresponding samples (see SI 2.11 for details of the background correction). Y_{122^*} fractions were calculated as $I(Y_{122^*}) / [I(Mn^{2+}) + I(Y_{122^*})]$ at the observe position (dashed line in A). D) Calibration curve for the modulation depth as a function of relative Y_{122^*} contribution to the refocused echo. Area below the curve is shaded as a guide to the eye. For details, see SI 2.9 and 2.10.

Δ in the cells cannot be explained by the presence of Mn^{2+} alone. As Y_{122}^{\bullet} and Mn^{2+} were the only paramagnetic centres detected within the cells (Figures 2, S5, and S6), we conclude that some of the $\beta 2$ dimers present in the cells contain only one Y_{122}^{\bullet} , further reducing the in-cell Δ values. In vivo presence of 1 Y^{\bullet} per $\beta 2$ is consistent with the asymmetric $\alpha 2\beta 2$ complex structure.^[8] Although the in-cell fraction of $\beta 2$ subunits having only one tyrosyl radical could be calculated based on the fit resulted from the calibration curve, we refrain from such an analysis at this point. A detailed and controlled investigation of cell growth conditions such as growth time, added solutes, metal inhibition should be done to determine the exact in vivo monomeric 1 $\text{Y}^{\bullet}/\beta 2$ fraction, which is beyond the scope of this work and will be addressed in future studies. Overall, our data suggest a distinct radical distribution in *E. coli* cells compared to the in vitro “two or none” model, and to that of in vivo *S. cerevisiae*.

Additionally, we compared background slopes of the DEER dipolar traces, as the slope reports on the local spin concentrations in a sample.^[34] The background steepness of in-cell samples was highly similar to that of in vitro samples with the radical concentration of 22 μM and 5–6 equivalents of added Mn^{2+} (red vs. yellow and green traces in Figure 5B, see Table S2). In vitro DEER traces for samples with higher radical concentrations (70 and 150 μM) showed significantly steeper backgrounds (grey, blue, pink, and black traces in Figure 5B). This demonstrated that the local Y_{122}^{\bullet} concentration within the cells was substantially lower than 70 μM , and was similar to that of the in vitro sample, which was prepared to mimic the in-cell sample and contained 22 μM Y_{122}^{\bullet} (yellow trace in Figure 5B and SI 2.12). This result is in very good agreement with the intracellular Y_{122}^{\bullet} concentration of around 20–32 μM as determined by spin quantification at 9.6 GHz (SI 2.3). The detection of low intracellular Y_{122}^{\bullet} concentration is further supported by echo decay measurements. T_m of Y_{122}^{\bullet} within cells is either longer or the same as that of in vitro samples (see Figure S8).

Detection of $\text{F}_3\text{Y}_{122}^{\bullet}$ and $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\text{F}_3\text{Y}_{122}^{\bullet}$ Pairs in Cells

Incorporation of $\text{F}_3\text{Y}_{122}^{\bullet}$ was fundamental to being able to investigate in vitro radical transfer mechanism in RNR and to obtain the cryo-EM structure of the $\alpha 2\beta 2$ complex. Therefore, we explored the possibility of observing $\text{F}_3\text{Y}^{\bullet}$ in whole *E. coli* cells. Successful generation of $\text{F}_3\text{Y}_{122}^{\bullet}$ within the cells would be the first step towards gaining unique insights into the RNR catalysis under physiological conditions. In addition, it may serve as a new exciting probe to study in vivo radical transfer steps in other fundamental proteins containing amino acid radicals.

In order to generate $\text{F}_3\text{Y}_{122}^{\bullet}$, we followed the same procedure as that described for the in-cell generation of Y_{122}^{\bullet} . After over-expressing apo- $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\beta 2$ in the cells, we treated the harvested and washed cell cultures with Fe^{II} and O_2 to generate $\text{F}_3\text{Y}_{122}^{\bullet}$ (SI 1.2). Prior to EPR experiments, the viability of the cells and protein leakage from the cells were checked using cell counting and EPR experiments, respectively (SI 1.3 and 2.13). The resulting data showed that the

cells used for EPR measurements were intact and no detectable amounts of $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\beta 2$ protein pass through the *E. coli* cell wall. 9.6 and 34 GHz EPR measurements along with spectral simulations revealed the presence of the $\text{F}_3\text{Y}_{122}^{\bullet}$ species in the cells (SI 2.13). Spin quantification experiments yielded a $17 \pm 5 \mu\text{M}$ in-cell sample concentration of $\text{F}_3\text{Y}_{122}^{\bullet}$ (SI 2.13), in good agreement with that of the wt- $\beta 2$ in-cell samples.

Subsequently, in-cell and in vitro DEER measurements with samples containing $\text{F}_3\text{Y}_{122}^{\bullet}$ were performed (Figure 6). Four in vitro DEER traces recorded at different g -tensor orientations were summed to eliminate orientation-selection effects (see SI 2.14 for details). Such orientation-selective DEER measurements were not feasible with the in-cell sample because of poor SNR. We recorded an in-cell DEER trace at a molecular orientation at which the distance vector $\mathbf{r}_{\text{F}_3\text{Y}_{122}^{\bullet}\text{-F}_3\text{Y}_{122}^{\bullet}}$ is perpendicular to the magnetic field because the correct mean distances can be directly obtained from dipolar frequencies at this orientation.^[35] The distance analysis of this trace resulted in a mean distance of 3.37 nm (standard deviation of the in-cell distance distribution, σ , which is not totally reliable in the presence of orientation selectivity,^[32] is not given). The detected in-cell distance is assigned to a $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\text{F}_3\text{Y}_{122}^{\bullet}$ pair as it agrees extremely well with the in vitro distance detected here (3.36 nm with $\sigma = 0.11$ nm) and with the distance between oxygen atoms of $\text{F}_3\text{Y}_{122}^{\bullet}$ s in the recent crystal structure of $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\beta 2$ (SI 2.15).^[36]

Furthermore, we compared background steepness of in-cell and in vitro $\text{F}_3\text{Y}_{122}^{\bullet}$ DEER traces. Similar to the results obtained with Y_{122}^{\bullet} , this analysis showed that local $\text{F}_3\text{Y}_{122}^{\bullet}$ concentration within the cells is substantially lower than the detected in vitro concentration of $130 \pm 30 \mu\text{M}$ (SI 2.13 for spin quantification, and SI 2.14 for background comparison).

Conclusion

This study has revealed that the structure and electrostatic environment of the essential di-iron tyrosyl radical cofactor Y_{122}^{\bullet} in *E. coli* RNR are identical under in vivo and in vitro conditions. Here, we presented distance measurements be-

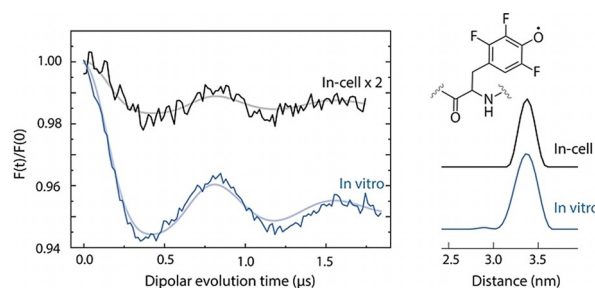


Figure 6. Background- and phase-corrected, normalized 34 GHz in-cell (black) and orientation-averaged in vitro (blue) DEER traces of $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\beta 2$ subunit of *E. coli* RNR (A) shown along with obtained $\text{F}_3\text{Y}_{122}^{\bullet}$ - $\text{F}_3\text{Y}_{122}^{\bullet}$ distance distributions (B). The structure of $\text{F}_3\text{Y}^{\bullet}$ is shown in the inset (B). DEER time traces were analysed by DeerAnalysis2019^[32] and the fits are overlaid in grey (in-cell) and light blue (in vitro). Details of the experiments and analyses are given in SI 2.14. The original in-cell trace was magnified by a factor of two for better visualization.

tween two native paramagnetic centres residing in an enzyme in intact and living cells where the protein under investigation is expressed. Distance measurements within the Y_{122}^{\bullet} - Y_{122}^{\bullet} pair provided insights into the in-cell structure and conformational rigidity of the β_2 subunit, and suggested a distinct in-cell radical distribution within this subunit. Detection of β_2 subunits having only one Y_{122}^{\bullet} strongly supports a model in which *E. coli* RNR activity is regulated by modulation of Y^{\bullet} concentrations in the cells.^[7] Our results serve as a basis for future experiments aimed at detecting and manipulating the factors influencing in-cell RNR activity. Additionally, we present spectroscopic detection of an unnatural amino acid radical in whole cells, and show that F_3Y incorporation does not affect the in vivo protein structure. Successful generation of $F_3Y_{122}^{\bullet}$ within the cells is the first step towards gaining unique insights into the RNR catalysis under physiological conditions. Furthermore, it showcases the possibility of unravelling the in vivo structure and role of tyrosyl radicals involved in other fundamental processes, such as photosynthesis, reduction of O_2 to water, and DNA repair by unnatural amino acid incorporation.

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Conflict of interest

The authors declare no conflict of interest.

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