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ARTICLE

An optimal acquisition scheme for Q-band EPR distance measurements using Cu²⁺-based protein labels

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Recent advances in site-directed Cu²⁺ labeling of proteins and nucleic acids have added an attractive new methodology to measure the structure-function relationship in biomolecules. Despite the promise, accessing the higher sensitivity of Q-band Double Electron Electron Resonance (DEER) has been challenging for Cu²⁺ labels designed for proteins. Q-band DEER experiments on this label typically require many measurements at different magnetic fields, since the pulses can excite only a few orientations at a given magnetic field. Herein, we analyze such orientational effects through simulations and show that three DEER measurements, at strategically selected magnetic fields, are generally sufficient to acquire an orientational-averaged DEER time trace for this spin label at Q-band. The modeling results are experimentally verified on Cu²⁺ labeled human glutathione S-transferase (hGSTA1-1). The DEER distance distribution measured at Q-band shows good agreement with the distance distribution sampled by molecular dynamics (MD) simulations and X-band experiments. The concordance of MD sampled distances and experimentally sampled distances adds growing evidence that MD simulations can accurately predict distances for the Cu²⁺ labels, which remains a key bottleneck for the commonly used nitroxide label. In all, this minimal collection scheme reduces data collection time by as much as six-fold and is generally applicable to many octahedrally coordinated Cu²⁺ systems. Furthermore, the concepts presented here may be applied to other metals and pulsed EPR experiments.

Introduction

Pulsed electron paramagnetic resonance (EPR) distance measurements^{1–7} have had a major impact in a variety of biophysical contexts, including the measurement of conformational changes,^{8–15} determination of relative packing in protein-protein^{16–18} and protein-DNA complexes,^{19,20} and the elucidation of ligand and metal binding sites in proteins.^{21–23} Additionally, these experiments have been introduced both *in vitro* as well as *in-cell*.^{24–27} Such measurements are enabled by site-directed spin labeling methodology that has predominantly used nitroxide spin labels.²⁸ Recently, several attractive schemes for labeling proteins and nucleic acids with metal ions have been developed to enhance the reach of spin labeling methodology in biophysics.^{29,30} In particular, site-directed Cu²⁺ labeling of proteins can provide distance distributions that are up to five-times narrower than similar measurements using nitroxide-based labels.³¹ The Cu²⁺ labeling method for protein relies on the strategic placement of two histidines which can bind to the Cu²⁺ ion. The labeling of the double-histidine (dHis) site is achieved by using either a Cu²⁺-iminodiacetic acid³² or a Cu²⁺-nitrilotriacetic acid (NTA)³³ chelate to prevent non-specific

binding of Cu²⁺ elsewhere in the protein. The labeling approach is facile, can be implemented in a wide variety of buffers,³⁴ and a range of pH.³⁵ In addition, distance measurements using dHis labeling and Relaxation Induced Dipolar Modulation Enhancement Spectroscopy (RIDME) can be performed at sub-micromolar concentrations.³⁶ The dHis-Cu²⁺ labeling method has been shown to provide enhanced resolution to the multi-lateration of native metal binding sites,²² measurement of relative orientations in proteins,³⁷ induced conformational changes,^{38,39} and the measurement of site-specific dynamics even on β -sheets.⁴⁰ For DNA, a nucleotide-independent labeling approach that can directly report on backbone distances^{41,42} and DNA conformational changes⁴³ is also available. There is also an increasing development of using Cu²⁺ to label DNA by creating a quadruplex structure.^{44–46} On the other hand, due to the large spectral bandwidth of Cu²⁺, compared to nitroxides, the sensitivity of the label is an area for improvement.

The advent of high-field instrumentation, especially at Q-band (ca. 35 GHz) has been particularly impactful by providing more than an order of magnitude enhancement of sensitivity compared to X-band measurements (ca. 9.5 GHz).⁴⁷ Overall, the improved sensitivity has been especially valuable for nitroxide based Double Electron-Electron Resonance (DEER) distance measurements. Despite this potential, distance measurements involving Cu²⁺ at Q-band have been limited, due to difficulties in properly sampling all the orientations of the inter-spin vector in DEER.^{37,44} At Q-band the spectral bandwidth of the Cu²⁺-spectrum is ca. 5 GHz due to the large anisotropy of the g-tensor. On the other hand, the pulses that are typically used

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excite only a bandwidth of ca. 100-300 MHz⁴⁸ due to resonator and pulse amplifier limitations. Consequently, DEER measurements at a given magnetic field sample only some orientations of the inter-spin vector. To ameliorate these orientational effects, a DEER experiment at Q-band require the collection of data at different magnetic fields. For example, earlier Q-band DEER work on a dHis-Cu²⁺ labeled protein utilized seventeen different magnetic fields to obtain the same distance distribution observed using X-band DEER measurement at a single magnetic field.³⁷ This constraint at Q-band can often make X-band measurements more practical, despite their decreased sensitivity. On the other hand, the initial work at Q-band likely suffered from oversampling, and therefore, there is a critical need to establish the *minimum* number of fields and the associated averaging scheme for practical distance measurements.

In this work, we establish an optimal approach for collecting dHis-Cu²⁺-based distance measurements at Q-band frequency by focusing on the human glutathione S-transferase (hGSTA1-1) enzyme. We first used MD simulations to identify the distance and orientational information of the Cu²⁺-labeled sites. This information was then used as initial values to determine the number of angles excited as a function of magnetic field using a Monte-Carlo scheme. The information on angles was then used to establish an excitation scheme that is expected to appropriately sample molecular orientations. We then demonstrate through simulations that the minimal acquisition scheme is applicable for any possible orientation. Thus, the acquisition scheme can be used without the need for prior structural information. Finally, this scheme was validated by experiments on the protein at both Q- and X-bands, and by MD simulations.

Experimental

Protein expression, purification, and sample preparation

All experiments were performed on the S-hexylglutathione (GSHex) bound form of hGSTA1-1. The K211H/E215H mutant was expressed and purified based on the previously published protocol,⁴⁰ except the cell growth was done in Luria Broth instead of Terrific Broth media. hGSTA1-1 is a homodimer such that a single dHis mutant provides two Cu²⁺-NTA binding sites. The purified protein was concentrated and aliquoted to ca. 200 μ M in buffer (pH = 6.5) with 150 mM NaCl and 50 mM sodium phosphate then stored at -80 °C.

In order to prepare the EPR samples, a 10 mM Cu²⁺-NTA stock was prepared as described previously^{33,34,49} and the GSHex ligand was purchased from Sigma Aldrich. All samples in this work were prepared with 3-N-morpholinopropanesulfonic acid (MOPS) buffer to facilitate efficient Cu²⁺-NTA binding to dHis.³⁴ Each EPR sample was prepared with 800 μ M GSHex, 800 μ M protein and 800 μ M Cu²⁺-NTA (a 1:1:1 ratio to ensure that the concentration of both protein and ligand are 10⁴ times over K_D of GSHex to hGSTA1-1)⁵⁰ in 50 mM MOPS buffer in D₂O (pH = 7.4) with 100 mM NaCl. All samples were incubated at 4 °C for

35 min to achieve maximum loading efficiency, and subsequently flash frozen in liquid MAP-Pro Propylene/propane gas with 50 % D₆-glycerol added as cryoprotectant. A step-by-step protocol for spin labelling and freezing has been published recently.⁴⁹

MD simulation of dHis-Cu²⁺-NTA labeled protein

The MD simulation was performed starting from the crystal structure of GSHex-bound hGSTA1-1 (PDB: 1K3L).⁵¹ To generate parameters for the ligand GSHex, the PDB of GSHex was extracted from the crystal structure of liganded hGSTA1-1. Using the Antechamber package in Amber18, charges were derived using the AM1-BCC method.^{52,53} Force field angles and dihedral terms were assigned with the General Amber Force Field (GAFF).⁵⁴ Sites 211 and 215 of each monomer were mutated to histidine for the incorporation of Cu²⁺-NTA. The force fields for Cu²⁺-NTA have been developed in previous work.⁵⁵ The hGSTA1-1 was simulated with the ff14SB AMBER force field.⁵⁶ Solvent waters were treated with the TIP3P water model.⁵⁷ The labeled protein was solvated in a cubic box with the box size of ca. 60 Å³. Sodium and chloride ions were added to neutralize the system. The MD simulation was performed with the pmemd program as part of the AMBER18 software package. The solvated system was first energy minimized with a harmonic force constant applied to the protein and ligand, which was gradually released from 20 to 0 kcal/mol/Å² over 12000 steps. The energy minimized system was then equilibrated using a Berendsen barostat at 298 K for 3.2 ns with a decreasing harmonic force constant from 20 to 0 kcal/mol/Å².⁵⁸ The system was then gradually heated from 0 to 298 K. Equilibration was reached after 2 ns. The simulation at the production phase was maintained at 298 K by Langevin thermostat using a 5.0 ps⁻¹ collision frequency and collected for 200 ns. A 2 ps time step for motion integration was used for equilibration and production simulations. The system pressure was kept at 1 atm with a pressure relaxation time of 1.0 ps. The SHAKE algorithm was used to restrain the bonding to hydrogens. Periodic boundary condition and particle mesh Ewald were applied to accurately treat the long-range electrostatic interactions.

Monte-Carlo simulation

To build an *in-silico* model of our sample, we implemented a home-written Python⁵⁹ code, available at the Saxena Lab GitHub

(github.com/SaxenaLab/MonteCarloSimulationCopper/). The process starts with the generation of 10,000 vectors, representing g_{\parallel} of Cu²⁺ spins (Spin A). For each Spin A, we generated another spin (Spin B) separated by a vector of length r . The g_{\parallel} tensor of Spin B is rotated by an angle γ with respect to the g_{\parallel} tensor of Spin A. Similarly, the g_{\perp} of Spin B is rotated by an angle η with respect to the g_{\perp} of Spin A. For each pair, another vector, representing the inter-spin vector, is generated with an orientation of χ with respect to the g_{\parallel} tensor of Spin A. The angles γ , χ , and η , depicted pictorially in Fig. 1, were

sampled from three different Gaussian distributions defined by the user. These vectors represent 10,000 spin-pairs of Cu²⁺-labeled hGSTA-1 sample. After generating the vectors, the spin-pairs were then randomly rotated to simulate spin-pairs with random orientations.

From each spin, we calculated the effective g and hyperfine interaction term, A , as a function of angle, ϕ , between the applied magnetic field and g_{\parallel} with the following equations:⁶⁰

$$g(\phi) = \sqrt{g_{\perp}^2 \sin^2 \phi + g_{\parallel}^2 \cos^2 \phi} \quad (1)$$

$$A(\phi) = \frac{\sqrt{A_{\perp}^2 g_{\perp}^4 \sin^2 \phi + A_{\parallel}^2 g_{\parallel}^4 \cos^2 \phi}}{g_{\perp}^2 \sin^2 \phi + g_{\parallel}^2 \cos^2 \phi} \quad (2)$$

Both the g and A tensors were experimentally determined, as shown in Fig. S1. The effective g and A values of each spin were then used to calculate the resonant field, B_{res,m_i} , of the spin using the following equation:

$$B_{res,m_i} = \frac{h\nu - Am_i}{g\beta_e} \quad (3)$$

where h is Planck's constant, ν is the microwave frequency, β_e is Bohr Magneton, and m_i is the nuclear quantum number. For our analysis, we set ν as 34.15 GHz to approximate a Q-band frequency. Then, at each resonant field, we built a Lorentzian line-shape with an arbitrary maximum intensity of 1 and a broadening parameter, β , of 40 G. The Lorentzian curve is built as follows:

$$I(B) = \sum \frac{\beta^2}{m_i(B - B_{res,m_i})^2 + \beta^2} \quad (4)$$

where β is the broadening parameter. By summing all the Lorentzian functions from all spins, we were able to generate a field-swept spectrum. The parameter β was chosen as 40 G to best fit with experimental field sweep (cf. Fig. S2).

In addition to the field-swept spectra, we used $I(B)$ to determine whether a spin with a given ϕ can be excited at a given field. Specifically, we consider a spin to be excited only when the intensity of $I(B)$ at a specific field is above a defined threshold parameter, α . The α was set to 0.4 in our analysis to ensure optimal sampling of the number of spins (cf. Fig. S3). The sampling of number of spins is also dependent on the choice of β . As $I(B)$ becomes broader due to an increase in β , α must be adjusted to maintain optimal sampling of the spins. Therefore, both α and β are critical in our simulations. More details for the choices of α and β are provided in the results section and the ESI.

Excitation profile of θ

To understand the origin of orientational selectivity, the excitation of θ angles can be studied. θ is defined as the angle

between the inter-spin vector, \mathbf{r} , and the applied magnetic field B_0 (Fig. 2A). To include orientational effects in DEER experiments, DEER signal for a two-Cu²⁺ system can be expressed as:⁶¹

$$V(t)_{intra} = 1 - \iint \lambda (1 - \cos [\frac{k}{r^3} (1 - 3\cos^2 \theta)]) P(r) \xi(\theta) d\theta dr \quad (5)$$

where λ is the modulation depth, k is the constant containing the g values of the two spins, r is the distance between the two spins, $P(r)$ is the distribution function of the distance, and $\xi(\theta)$ is the geometrical factor describing the probability of exciting θ , which is defined as:⁶²

$$\xi(\theta) = \frac{1}{2} \sum_{m_{i_1}, m_{i_2}} \langle k_{x_a}^3 k_{x_b}^2 \sin \varphi_{1a} (1 - \cos \varphi_{2a}) (1 - \cos \varphi_{3b}) + k_{x_b}^3 k_{x_a}^2 \sin \varphi_{1b} (1 - \cos \varphi_{2b}) (1 - \cos \varphi_{3a}) \rangle_{\phi, \delta\omega_1, \delta\omega_2} \quad (6)$$

where m_{i_i} is the nuclear quantum number of the i th spin, k_{x_a} and k_{x_b} are defined as the ratio of the resonance frequency of the excited spins versus the frequency at observer and pump frequency, respectively, φ_{i_a} is the flip angle of the first spin by the i th pulse, φ_{i_b} is the flip angle of the second spin by the i th pulse, and $\delta\omega_i$ is the inhomogeneous broadening of the observer or the pump pulses. If all orientations of the spin-labeled molecules are properly sampled, the geometrical factor $\xi(\theta) = \sin(\theta)$.

EPR measurements

To determine g and A -tensors and the coordination environment for Cu²⁺-NTA labeled liganded hGSTA1-1, continuous wave (CW) EPR experiment and three-pulse electron-spin echo envelope modulation (ESEEM) experiments^{63,64} were performed with a Bruker ElexSys E680 X-band FT/CW spectrometer with a Bruker EN4118X-MD4 resonator. The CW-EPR experiment was performed at 80 K and the ESEEM experiment was performed at 20 K. The CW was collected at microwave frequency ca. 9.70 GHz, with a modulation frequency of 100 kHz, modulation amplitude of 4 G, and a sweep width of 2000 G centered at 3100 G. A total of 1024 data points were collected with an attenuation of 30 dB, conversion time of 20.48 ms, and data was averaged over 50 scans. The X-band sample contains 100 μ L of 400 μ M hGSTA1-1 dimer, 800 μ M GSHex and 800 μ M Cu²⁺-NTA in 50 mM MOPS buffer prepared with D₂O (pH = 7.4) with 100 mM NaCl, 50% D₆-glycerol, placed in quartz tube with I.D. = 3 mm and O.D. = 4 mm. The CW spectrum was simulated with EasySpin.⁶⁵ More details of the ESEEM measurements are provided in ESI.

To acquire distance distributions between labels within liganded hGSTA1-1, four-pulse DEER experiments were performed at 18 K at both X-band and Q-band frequencies.¹ For the X-band DEER, the measurement was performed on a Bruker ElexSys E680 X-band FT/CW spectrometer equipped with a Bruker EN4118X-MD4 resonator and a 1 kW amplifier. For the Q-band DEER, the measurements were performed with a Bruker ElexSys E580 X-band FT/CW spectrometer with a Bruker

ER5106-QT2 resonator and a 300 W amplifier. The pulse sequence used was $(\pi/2)v_A - \tau - (\pi)v_A - \tau + t - (\pi)v_B - T - t - (\pi)v_A - T$ -echo.⁶⁶ For X-band DEER, rectangular pulses were used. The lengths of the observer $(\pi/2)v_A$ and $(\pi)v_A$ were 8 and 16 ns, respectively. The pump pulse had a length of 16 ns. The interval, t , was incremented by a step size of 30 ns over 202 points. For Q-band DEER, the $(\pi/2)v_A$ and $(\pi)v_A$ pulses were rectangular pulses with lengths of 12 ns and 24 ns, respectively. To increase the modulation depth, an 80 ns chirp pulse was used for the pump pulse. The pump pulse was set with a frequency from -200 MHz to -100 MHz relative to the observer frequency. A 16-step phase cycling was used. The duration, t , was incremented with the same step size as X-band DEER over a total of 237 points. We performed DEER measurements with pump pulses placed 116 G, 566 G, and 746 G, lower than the magnetic field with the highest intensity of the Field Swept Electron Spin Echo (FS-ESE) Spectrum. To further test the sufficiency of the collection scheme, seven additional DEER were collected with pump pulse placed at fields that are ca. 641 G, 516 G, 466 G, 416 G, 334 G, 250 G, and 165 G lower than the magnetic fields with the highest intensity of the FS-ESE spectrum. DEERAnalysis2021⁶⁷ was used to analyze the data.

Results and discussion

In this work, we focused on the conformation of the $\alpha 9$ helix of hGSTA1-1 in the presence of ligand GSHex. The conformation and dynamics of this helix is intimately related to the function of this protein.^{68,69} For this purpose, we generated a K211H/E215H hGSTA1-1 mutant, as described in previous work,³⁸ which was subsequently labelled with Cu^{2+} -NTA. Due to the homodimeric nature of hGSTA1-1, a single dHis mutant provides two labelled sites, one on each subunit, for distance measurements.

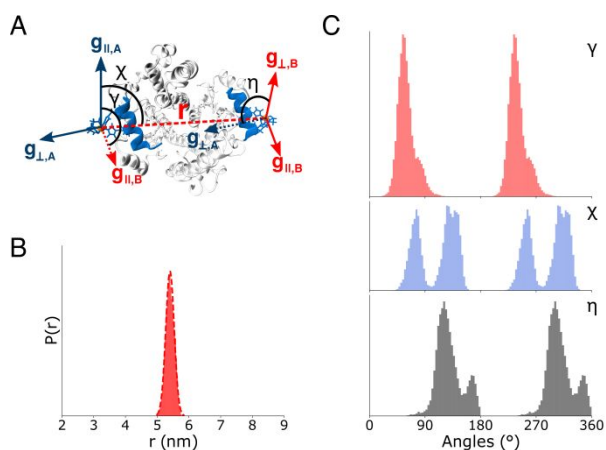


Fig. 1 (A) Definition of g -tensors and relative angles within dHis- Cu^{2+} -NTA labeled hGSTA1-1. γ is the angle between $g_{\parallel,A}$ and $g_{\parallel,B}$; χ is the angle between $g_{\parallel,A}$ and the interspin vector r ; and η is the angle between $g_{\perp,A}$ and $g_{\perp,B}$. (B) Cu^{2+} - Cu^{2+} distance distribution from a 200 ns MD run. The most probable distance is 5.3 nm, and the standard deviation is ca. 0.2 nm. The distance distribution was reasonably reproduced by a Gaussian shown by the red dashed line. (C) The distribution of each orientational angle.

MD simulation provides initial angle and distance distributions

We first, established the anticipated distance distribution and orientational distribution of the g -tensors of the two Cu^{2+} sites, using MD simulations. The orientation distribution is characterized by the three angles γ , χ , and η , shown in Fig. 1A. The angle, γ , is defined as the angle between $g_{\parallel,A}$ and $g_{\parallel,B}$; χ is the angle between $g_{\parallel,A}$ and inter-spin vector r ; and η is the angle between $g_{\perp,A}$ and $g_{\perp,B}$. For Cu^{2+} distances, orientational selectivity is strong for narrower orientational distributions.^{37,55,61,70} Therefore, a reference model providing Cu^{2+} - Cu^{2+} distances with relative orientations is pertinent to our analysis. We performed a 200 ns MD simulation of GSHex-bound hGSTA1-1 (PDB: 1K3L)⁵¹ with Cu^{2+} -NTA labels introduced into $\alpha 9$ at site K211H/E215H. In the MD simulation, we used the force fields parameters for dHis- Cu^{2+} -NTA that were recently developed.⁵⁵

From the MD simulation, we obtained the distance, r , between the two Cu^{2+} centers, plotted in Fig. 1B. The Cu^{2+} - Cu^{2+} distance distribution is centered around 53 Å, with a standard deviation σ_r of ca. 2 Å. We then extracted the relative orientations of the Cu^{2+} spin labels from the MD trajectory. The imidazole nitrogen atoms bind to Cu^{2+} equatorially, leading to the g_{\parallel} perpendicular to the equatorial plane. The directions of g_{\parallel} and g_{\perp} in the dHis- Cu^{2+} -NTA complex are defined in ESI (Fig. S4). The distributions of γ , χ , and η are shown in Fig. 1C. For each γ , χ , and η distribution under 180° , there is an identical distribution at $\gamma + 180^\circ$, $\chi + 180^\circ$ and $\eta + 180^\circ$, respectively. This n° and $n^\circ + 180^\circ$ distribution pattern is expected given that g_{\parallel} can be oriented “up” or “down” with respect to the equatorial plane due to the symmetry of the d_{z^2} orbital that the electron spin resides in. On the other hand, χ shows a bimodal distribution below 180° . A more careful examination of the MD traces, shown in Fig. 2, suggests that the Cu^{2+} adopts two preferred orientations, likely due to conformations of the $\alpha 9$ helix, as has been observed previously.^{38,40}

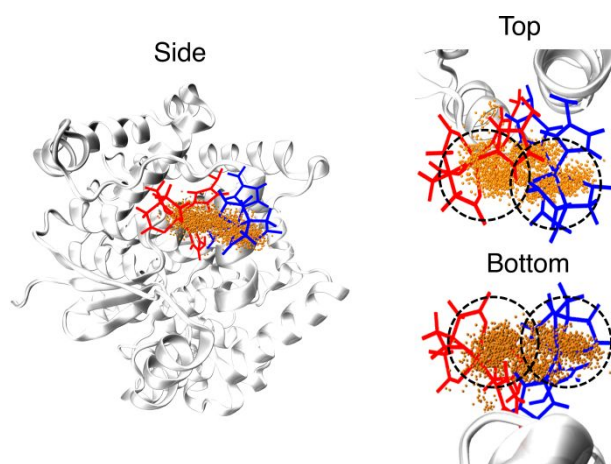


Fig. 2 Cu^{2+} spatial distribution. The helix $\alpha 9$ shows two conformations (indicated by Cu^{2+} -NTA conformations in red and blue sticks) in MD that result from a slight turn of the helix around the loop. Thus, the Cu^{2+} atom (yellow dots) shows a bimodal spatial distribution, indicated by the two black dashed circle, leading to a bimodal distribution in χ below 180° .

Finally, the standard deviation of γ , χ , and η are 14° , 10° and 21° , respectively. This standard deviation for these angles is consistent with earlier estimates on Cu^{2+} coordination to dHis sites in proteins and polynucleic acids that were obtained using a combination of MD and density functional calculations.^{29,55,71} In these systems, the Cu^{2+} coordination is elastic leading to a fluctuation in the bond angles and bond lengths of the Cu^{2+} coordinating atoms. Such fluctuations create a distribution in the directions of g_{\parallel} which generates a distribution in the γ , χ , and η angles.

The MD simulation, thus provide reasonable estimates for the standard deviations for the three angles, χ , γ and η . The greater the value of these standard deviations the lower the effects of orientational selectivity.^{61,70}

The optimal DEER collection scheme

Next, we used the information on expected orientation widths for dHis- Cu^{2+} labelled proteins obtained from MD simulations to devise an optimal collection scheme. First, we used Monte-Carlo methods⁷² to generate an *in-silico* sample containing 10,000 Cu^{2+} -labeled proteins, as shown in Fig. 3. Details are in the Experimental section. Each blue and green dot in Fig. 3A represents Spin A and Spin B, respectively, in a doubly Cu^{2+} labeled protein. The g -tensors and inter-spin vector, r , orientations are shown in Fig. 3B. In the *in-silico* sample, each set of Spin A, Spin B, and r is arranged with a given set of γ , χ , and η and r sampled from Gaussian distributions shown in Fig. 3D. For χ , γ and η , we used the most probable angles of 75° , 60° and 129° , respectively, as the mean angles for their respective Gaussian distributions. Note, that we ignored the

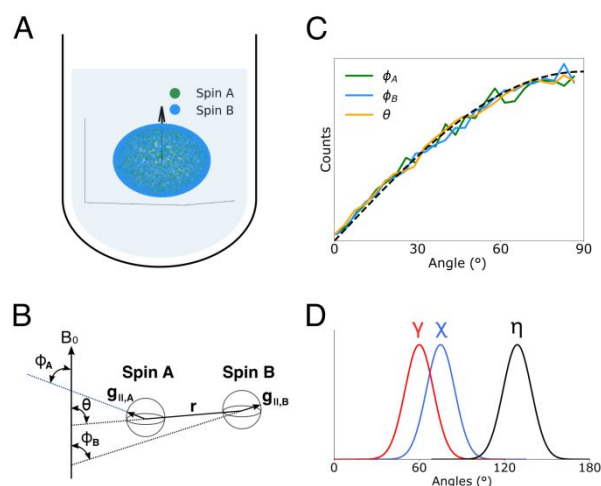


Fig. 3 *In-silico* sample consisting of randomly oriented vectors with two spins. (A) When all doubly labeled molecules are randomly oriented, the orientation of inter-spin vector, r , will also be random, as depicted by the spherical figure. The black arrow represents the applied magnetic field, while blue and green dots on the sphere represent orientations of the g_{\parallel} of Spin A and Spin B generated by the Monte-Carlo simulation. (B) A vector representation of two spins separated by inter-spin vector, r . The orientations of each spin can be described with ϕ while the orientation of r can be described with θ . (C) Plot of the distribution of θ , ϕ_A and ϕ_B in the Monte-Carlo simulation. The plot follows a sinusoidal curve depicted by the black dashed line, consistent with a random distribution. (D) Gaussian distributions of γ , χ and η used for the Monte-Carlo simulations.

bimodality for χ and the inherent symmetry for each angle (cf. Figure 1) We also used a standard deviation of 10° for each angle, which is a conservative estimate for the dHis- Cu^{2+} labeled proteins.^{29,37,55} Together, these choices make the *in-silico* sample more prone to orientational selectivity. Fig. 3C shows a count of the number of inter-spin vectors as a function of the angle, θ , between the inter-spin vector and the applied magnetic field. This probability distribution is sinusoidal, as expected for a random distribution.

To obtain a robust Q-band DEER, the data must be acquired in a manner that samples the θ distribution shown in Fig. 3C.⁶² On the other hand, the pulses in DEER are finite and can excite only some θ values at a given magnetic field. The excited θ values are dependent on the Cu^{2+} orientations, ϕ_A and ϕ_B , that are excited by the pump and excite pulses in DEER. Therefore, we first identified fields that can efficiently excite the largest number of ϕ angles.

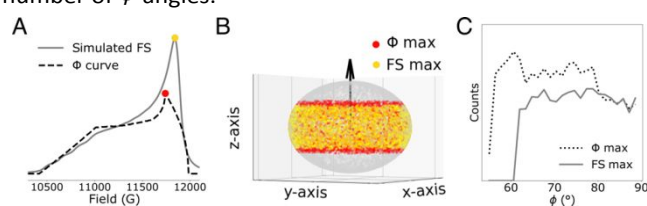


Fig. 4 (A) The spectrum in solid grey shows the simulated field-swept spectrum while the dotted line is the Φ curve described by equation (7). The Φ curve represents the number of ϕ excited at a given field. The max of both the field sweep and the Φ curve is marked by a yellow circle and a red circle, respectively. (B) Each dot on the sphere represents ϕ of a spin in the Monte-Carlo simulation. The yellow dots represent the spin-pairs excited when DEER is performed at the maximum of the field sweep. In contrast, the red dots represent the spin-pairs excited when DEER is performed at the maximum of the Φ curve. (C) Number of excited spins in DEER versus angle at the two magnetic fields.

We used the *in-silico* sample to calculate the resonant fields, B_{res,m_i} , of each Cu^{2+} -spin (A or B) as a function of ϕ_A or ϕ_B using equations (1-3). From this information a DEER signal can also be easily calculated. At each resonant field, we built a Lorentzian line-shape as a function of the magnetic field, $I(B)$, described by equation (4). Then, we summed each Lorentzian to build a field-swept spectrum, shown in Fig. 4A. Since each Lorentzian can be identified with the angle, ϕ , the number of ϕ excited at a given field, $\gamma(B)$, can be calculated as:

$$\Phi(B) = \int_0^{90^\circ} \left(\begin{cases} 1, & I_\phi(B) \geq \alpha \\ 0, & I_\phi(B) < \alpha \end{cases} \right) \sin\phi \, d\phi \quad (7)$$

where α is an intensity-threshold parameter. The expression of $\Phi(B)$ contains two terms. The bracketed first term represents a counter recognizing whether a ϕ value can be excited at a given field. The ϕ value is excited only when $I_\phi(B)$ is greater than or equal to α . The second term describes the relative probability of spins for a given ϕ . Overall, $\Phi(B)$ quantifies the number of ϕ excitable at a given magnetic field. The curve of $\Phi(B)$ is shown in Fig. 4A. We will refer to this curve as the Φ curve for the rest of this document. The Φ curve depends on the choice of α and breadth of the Lorentzian, β . The breadth of each Lorentzian was chosen by fitting the experimental FS-ESE spectrum of hGSTA1-1 (cf. Fig. S2). The optimal value of α , was

determined by examining the number of excited spins as a function of α for several different fields across the simulated FS-ESE spectrum (cf. Fig. S3). For a β of 40 G, an α of 0.4 leads to a stable count of spins at different fields. The optimal value of α for several values of β are shown in Fig. S3.

Fig. 4A shows the Φ curve generated for the *in-silico* sample shown in Fig. 3A. The value of $\Phi(B)$ (dotted lines) is overlaid on the simulated FS-ESE spectrum. The maximum number ϕ angles are excited at a field of 11738 G (shown by red circle on Fig. 4A), about 100 G lower than the maximum of the FS-ESE spectrum. To exemplify this point, we determined the spin-pairs excited by DEER at either the max of the Φ curve, defined as $B_{\phi 0}$, or the max of the FS-ESE spectrum. For simplicity, we assumed a square excitation profile from hyperbolic secant CHIRP pulses.⁷³ Specifically, we first identified the spins that can be excited by a pump pulse with a bandwidth of 100 MHz, which is set either at the maximum of the Φ curve or at the maximum of the field sweep. Next, we identified the spins that can be excited by a ca. 38 MHz observer pulse at the field 54 G lower than the pump field. These pulses are chosen to replicate a reasonable DEER on a commercial resonator that has ca. 200 MHz bandwidth. Finally, we identified the spins from the pump excitation that are paired with the spins from the observer excitation. These spin-pairs in our *in-silico* sample contribute to the intra-molecular DEER signal at a given field.

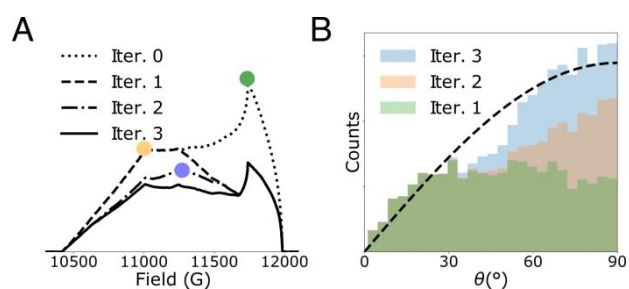


Fig. 5 (A) The Φ curve showing the number of excitable ϕ angles versus magnetic field. After excitation at one field, the spin-pairs that were excited were removed to generate the curve for subsequent iterations. The maximum ϕ angle for each iteration is labeled with a dot. We iterated this process until residual ϕ angles cannot be further reduced, and this curve is shown by the black solid line. (B) Excitation profile of θ angles. The black dashed sinusoidal curve depicts the ideal excitation of θ , and the histogram shows the cumulative θ excitation after each DEER at the three identified fields.

The ϕ angles excited at $B_{\phi 0}$ (red dot) versus the maximum of FS-ESE spectrum (yellow dot) are shown in Fig. 4B. At the maximum field sweep intensity, the excited spin-pairs have ϕ range of ca. 62° to 90° . In contrast, a more extensive range of ϕ from ca. 56° to 90° can be excited at the maximum of the Φ curve. The increase in the number of ϕ is due to the large parallel component of the hyperfine tensor, A_{\parallel} . More importantly, this observation indicates that DEER at 100 G lower than the maximum of ESE-FS spectrum is the most optimal for probing the largest number of ϕ for Cu^{2+} -labeled systems.

Fig. 5B shows the distribution of θ that is sampled at $B_{\phi 0}$ (green histogram) by DEER. Data at only this field is clearly insufficient to achieve ideal excitation, which is shown by the dashed line. Therefore, we identified additional fields that can excite the rest of the spin-pairs. Fig. 5A, shows the recalculated Φ curve,

labeled as Iteration 1, that excludes spins that are excited by the DEER at $B_{\phi 0}$. The maximum of the resultant Φ curve, $B_{\phi 1}$, is shown by the orange dot on the curve. The field, $B_{\phi 1}$, which is ca. 827 G lower than the maximum of the FS-ESE spectrum, represents a field that can excite the largest number of ϕ angles from the unexcited spins. We repeated the identification of the θ angles that are sampled at the additional field by DEER, shown as the orange histogram in Fig. 5B. Measurement of DEER at the additional fields improve θ sampling, especially for angles between 60° to 90° . We reiterated this process and identified a new maximum at a field, $B_{\phi 2}$, which is ca. 580 G lower than the maximum of the FS-ESE spectrum. From these three fields, we see that the overall distribution of sampled θ is reasonably close to ideal, indicating that the three identified fields are the most promising for Cu^{2+} DEER.

The three identified fields are based on the Φ curve. Note, however, that a change in the linewidth parameter, β , can potentially affect the Φ curve [cf. equation (4)]. To test the robustness of our approach, we calculated the Φ curve for different values of β . Details are provided in ESI. For each β , we determined the value of α that ensures optimal sampling of excited spins as shown in ESI (cf. Fig. S3). As shown in Fig. S3, the shape of Φ curve and the resulting three identified fields remain consistent with different β . These results provide further credence to the Φ curve method. However, it is evident from Fig. 5A that there are still residual ϕ angles that are not excited even after four iterations.

To gain further insight, we visualized the orientations of

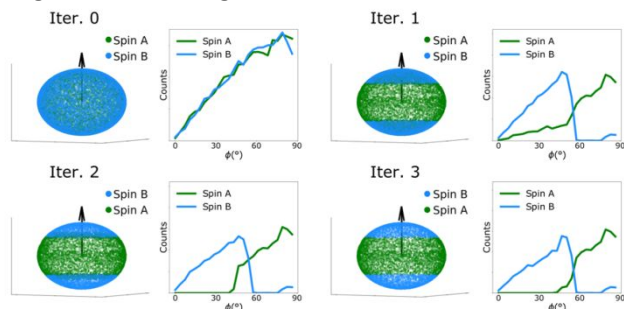


Fig. 6 Each spherical figure shows the ϕ angles of unexcited Spin A and Spin B at each iteration. The number of unexcited spins versus angle ϕ for Spin A and B is also shown. Initially, the orientations of unexcited Spin A and Spin B follows the same distributions as Figure 3C. However, the first DEER at $B_{\phi 0}$ excited spin pairs that have both Spin A and B in the region of ϕ between ca. 56° and 90° . As a result, the leftover spins in that ϕ region only consist of Spin A (cf. Iter. 1) that are paired to Spin B outside of that ϕ region. In Iteration 2, the unexcited spins in the region of ϕ between 0° and 46° only consist of Spin B which are paired to Spin A in a region of ϕ above 46° . After three DEERs in Iteration 3, Spin A and Spin B are largely isolated into two different regions of ϕ .

unexcited spin-pairs, Spin A and Spin B, throughout the sample in each iteration, shown in Fig. 6. Initially both Spin A and Spin B have diverse orientations. After the first iteration, there are no spin-pairs that have both spins at ϕ of ca. 56° to 90° . After the second iteration, there are no spin-pairs where both spins are in the region of ϕ between 0° to ca. 42° . Finally, after the third iteration, the leftover spin-pairs only consist of A spins with ϕ between ca. 56° to 90° that are paired with B spins with ϕ between 0° and ca. 56° . In other words, the final unexcited

spin-pairs primarily consisted of A spins oriented to the perpendicular region while paired with B spins oriented to the parallel region. Consequently, the resonant fields of most unexcited spin-pairs are too far apart to be excited by conventional DEER with a resonator bandwidth of 200 to 300 MHz.

Quantitatively, we found that DEER at the three fields cannot excite ca. 4,500 spin-pairs out of 10,000 spin-pairs. However, we observed ca. 800 leftover spin-pairs that have ϕ around 56° for both Spin A and Spin B. These 800 spin-pairs can be excited by a fourth DEER at a field between B_{ϕ_1} and B_{ϕ_3} (ca. 338 G lower than the maximum of the FS-ESE spectrum). In this case, the fourth DEER only improves the sampling of θ slightly (cf. Fig. S5). Next, we simulated the Q-band DEER signal by summing up the contribution from each spin pair that was excited at the three fields. The simulated DEER is shown by the solid blue line in Fig. 7. Superposed on the curve is the ideal DEER signal (black dashed line) if all possible spin pairs were excited. Comparison of the two traces indicates that DEER experiments performed at the three identified fields are sufficient to obtain the appropriate modulations in the time-domain signal. Furthermore, an additional DEER does not significantly improve the DEER signal (cf. Fig. S5).

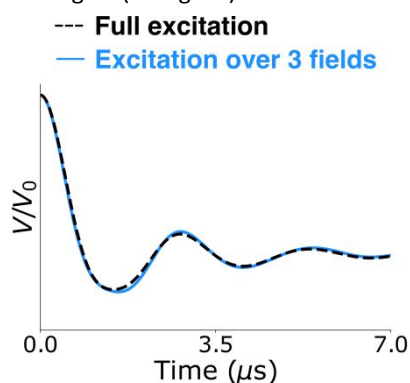


Fig. 7 Simulated averaged DEER time trace using the optimal collection method in solid blue line is compared to an ideal DEER time trace with all spins excited (black dashed line). The simulation assumes a Gaussian distance distribution with mean distance = 5.3 nm and $\sigma_r = 0.2$ nm.

Note, the case shown in Fig. 7 has been calculated for the orientational distribution shown in Fig. 3D. In order to ensure that the results are general, we calculated DEER traces for 125 combinations of angle χ , γ and η . In each case, the standard deviation of each angle was 10° . Fig. 8 shows simulated DEER signal with $\eta = 0^\circ \pm 10^\circ$, and for several different values of χ and γ that range from 0° to 90° . Simulations for $\eta > 0^\circ$ are shown in ESI (cf. Fig. S6 to S9). For all orientations the summed DEER signal averaged over the three identified fields is reasonably identical to the ideal DEER time trace. However, we noticed deviations between the simulated DEER time trace and the expected time trace in a small number of cases (4 out of 125 cases) when $\eta = 0^\circ$, $\chi = 20^\circ$, $\gamma = 40^\circ$ and 60° ; $\eta = 0^\circ$, $\chi = 40^\circ$, $\gamma = 60^\circ$ and 80° . To evaluate the significance of the discrepancy in distance distributions, we analyzed the DEER time traces with DeerAnalysis21.⁶⁷ In the distance analysis, random noise was added to the DEER signals to represent real-life measurements.

We show, in Fig. S10, that the resulting most probable distance as well as the distribution shape generally agree with the expected distance distribution. However, in these cases, a small peak at with an intensity roughly 10% of the main peaks is observed, due to residual orientational effects. This feature originates from the 800 spin-pairs that remain unexcited. In these cases, that contain a small feature and addition an additional DEER with pump pulse place 338 G lower than the maximum of FS-ESE spectrum is sufficient to suppress the minor peak (cf. Figure S10). Overall, DEER measurements at three magnetic fields are valid in most cases, but an additional DEER measurement may be useful to allow confident interpretation of any minor peaks in the distance distribution. The scheme is not dependent on distances of the system (eq.6). Therefore, no prior knowledge of distances or structures are required.

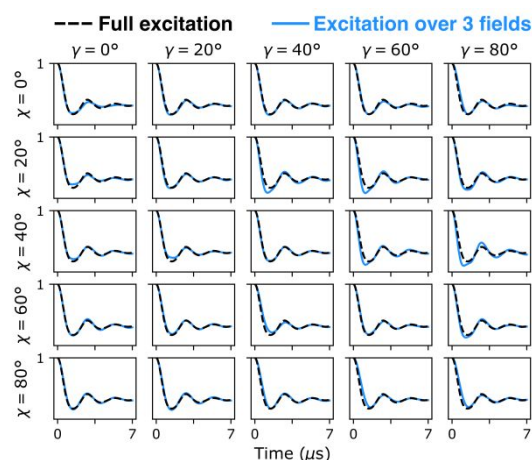


Fig. 8 Averaged DEER time traces (blue) simulated for the optimal DEER collection method compared to full excitation (black dashed). The simulation set $\eta = 0^\circ$ and $\sigma_\gamma = \sigma_\chi = \sigma_\eta = 10^\circ$, and iterates through γ and χ from 0° to 90° .

Assumptions and generality of the acquisition scheme

The modeling is encouraging and suggests that three magnetic fields might be sufficient to acquire robust DEER data for Cu^{2+} under the following conditions. First, the parallel component of the hyperfine tensor should be ca. 161 G. Such large values of hyperfine, which are typical for many octahedrally coordinated Cu^{2+} centers,^{74–76} ensure that many orientations are excited at a given magnetic field. Second, the standard deviation of the orientational distribution is ca. 10° or larger. The value of the standard deviation is consistent with expectations for sites with moderate binding affinity such as dHis.^{37,55,77,78} However, for substantially larger orientational distributions, as seen for the Cu^{2+} -2,2'-dipicolylamine DNA label,^{29,79} a DEER at a single field is sufficient. On the other hand, more fields may be needed for systems that exhibit rigid coordination to Cu^{2+} ,^{61,80–83} which will lead to lower distribution widths for angles χ , γ and η . Finally, the bandwidth of the pump pulse should be ca. 100 MHz or larger. A value of 100 MHz is conservative and can be readily achieved by pump pulses lengths of around 10 ns or using arbitrary waveform generators and resonators on commercial instrumentation. Furthermore, increasing the pump excitation bandwidth to 300 MHz can potentially reduce the DEER

acquisition scheme to only 2 fields (cf. Fig. S11). The acquisition scheme can be reduced further to one field with the recent development of pent loop-gap resonators⁸⁴ and ultra-wideband arbitrary waveform generators (cf. Fig. S12).^{85,86}

DEER on liganded hGSTA1-1

We next verified the acquisition scheme experimentally, by collecting DEER data on K211H/E215H hGSTA1-1 mutant which was labelled with Cu²⁺-NTA. Due to the homodimeric nature of hGSTA1-1, a single dHis mutant provides two labelled sites, one on each subunit, for distance measurements. Fig. 9A shows the FS-ESE spectrum of K211H/E215H hGSTA1-1 and the three pump fields at which DEER was acquired is indicated by solid vertical lines. The values of specific fields are in the Experimental section. We were able to obtain a dipolar evolution time of ca. 7 μ s due to the long phase memory relaxation time achieved (over 9 μ s, cf. Fig. S13) by deuteration of the solvent and the glycerol as has been recently reported.⁸⁷ Such a dipolar evolution time is important to resolve a ca. 50 Å distance, and is a dramatic improvement over previous efforts where short phase memory times limited the dipolar evolution time to ca. 4 μ s.^{38,87} The primary DEER traces are shown in ESI (Fig. S14).

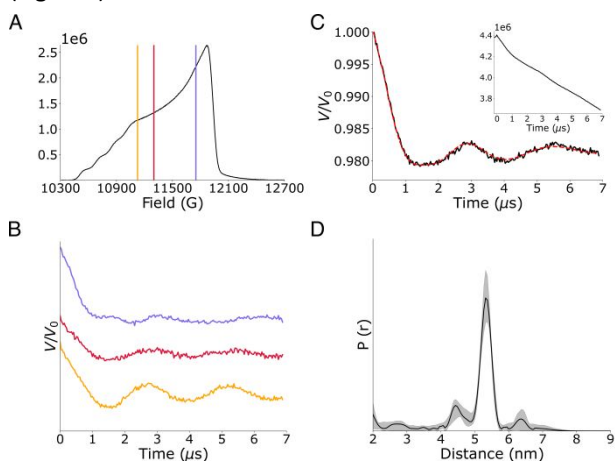


Fig. 9 (A) Field-swept-electron spin echo spectrum of dHis-Cu²⁺-NTA labeled hGSTA1-1. The magnetic fields for the pump pulses are labeled with color coded vertical lines. (B) The background subtracted DEER time domain at each magnetic field. The data was offset on the y-axis for better visualization. (C) The background subtracted average DEER time trace is plotted in black solid line, with Tikhonov fit shown as a red dashed line. The inset is the summed primary DEER time trace. (D) Distance distribution is analyzed by comparative DEER analysis (CDA). The uncertainty in the distribution is shown by grey shading.

The background subtracted DEER data using DEERAnalysis⁶⁷ at these fields are shown in Fig. 9B. The time traces are not identical at the three fields, which is a clear indicator of orientational selectivity.

Each time-domain DEER signal was normalized to the intensity of the FS-ESE spectrum (cf. Fig. S14) and summed to obtain the field average DEER signal (cf. inset of Fig. 9C). The background subtracted DEER signal and the fit from Tikhonov regularization is shown in Fig. 9C. Fig. 9D shows the resulting distance distribution using ComparativeDEERAnalyzer (CDA).^{88,89} CDA generates a consensus distance distribution and uncertainty

estimate using DEERNet and DEERLab analysis programs. DEERNet utilizes a deep neural network and automatically analyzes the contribution to the baseline from intermolecular dipolar interactions, which reduces potential user bias from the analysis.⁸⁸ DEERLab is a single step automated fitting program that utilizes Tikhonov regularization, removing all user bias.⁸⁹ The uncertainty of the distributions is plotted in grey shading. The individual results from the different analysis programs are provided in ESI (Fig. S15). A single distance centered around 5.3 nm, labeled by the red dashed vertical line in Fig. S15, was consistently shown by each analysis tool.

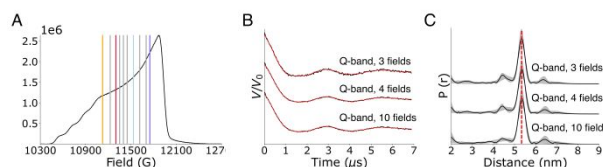


Fig. 10 Comparison of Q-band DEER collected at 3, 4 and 10 magnetic fields. (A) FS-ESE spectrum with DEER time traces collected at one additional magnetic field labeled with blue, and at six more magnetic fields with grey vertical lines. The fields are in addition to the 3 optimal fields that are shown by colored lines in purple, red and yellow. (B) DEER time trace averaged over 3, 4 and 10 magnetic field. The time traces are offset at y-axis for visualization. (C) Distance distribution by CDA. The uncertainty is indicated by the grey shading. The vertical line marks the most probable distance at 5.3 nm. The distance distributions for 3 fields, 4 fields, and 10 fields agree within the uncertainty.

To further confirm that three fields are sufficient, we collected 7 more traces at different magnetic fields (cf. details in the Experimental section) marked by a blue and six grey vertical lines in Fig. 10A. Fig. 10B shows that collecting one or even seven additional DEER measurements does not change the time trace. The resulting distance distributions, shown in Fig. 10C, are identical within error. This experimental observation is consistent with our DEER simulations where additional DEER does not significantly improve the sampling of θ and the DEER signal (cf. Fig. S5).

The proposed acquisition scheme is a dramatic improvement of the DEER protocol on commercial instrumentation. Using the optimal scheme, we were able to obtain the expected distance with only three measurements at different magnetic fields. The data collection time at each magnetic fields was between 1 h to 3.5 h, so the total DEER data collection time was ca. 7 h. In contrast, seventeen magnetic fields were used to measure the distance on a dHis-Cu²⁺ labeled protein at Q-band.³⁷ The optimal scheme therefore reduces the cryogenics cost and data collection time by at least 5 to 6 fold. Note, however, that the acquisition of data at different fields is useful if orientational information is needed.³⁷

X-band DEER and MD results further proves the validity of the summed Q-band DEER

Next, we acquired DEER data at X-band to further validate the Q-band DEER results. For dHis-Cu²⁺ labeling orientational selectivity is minimal at X-band and acquisition at only one field is needed.^{31–33,37,55} The DEER time trace at both X-band and Q-band are shown in Fig. 11A. The X-band time trace was collected for ca. 6 μ s to ensure sufficient signal-to-noise ratio while

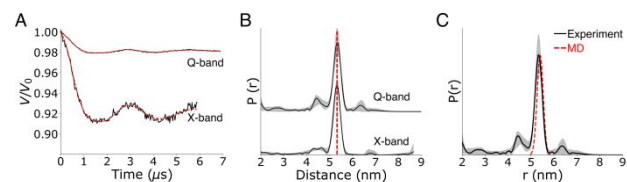


Fig. 11 (A) Summed Q-band DEER signals compared to X-band DEER. The modulation depth for X-band DEER is 7.7%, whereas the Q-band DEER shows a modulation depth of 2.0%. (B) Distance distribution by Consensus DEER analysis of the summed Q-band data compared to the X-band data. The uncertainty is indicated by the grey shading. The most probable distance, shown by the red-dashed line, was the same for X-band and Q-band data. (C) Cu²⁺-Cu²⁺ distance distribution from MD simulation is compared to the EPR distance distribution. The most probable Cu²⁺-Cu²⁺ distance by DEER measurement and MD simulation which agree within the uncertainty level of the experiment.

obtaining enough acquisition time for two modulations of a ca. 60 Å distance. The X-band time trace shows similar modulation frequency seen in the averaged Q-band signal, and the resulting distance distributions from X-band and Q-band are similar, as shown in Fig. 11B.

Finally, in Fig. 11C we compare the experimental distance distribution with the distribution obtained from MD simulations. The agreement between the two distributions is remarkable. More importantly, this result adds to previous evidence⁵⁵ that MD simulations can accurately predict the EPR distance measurements for Cu²⁺-NTA labeled proteins. The accurate prediction of distances by modeling remains a critical bottleneck for nitroxide labels.^{90,91} This body of work therefore foreshadows future work that combine MD simulations with EPR distance constraints to incisively probe protein structure and function.

Conclusions

In this work, we have demonstrated an efficient procedure to perform dHis-Cu²⁺-based Q-band DEER distance measurements. The optimal acquisition scheme can generate a high-resolution distance distribution with five to six times less collection time. Using a novel Monte-Carlo approach, we established that collecting data with the pump pulses at three magnetic fields (ca. 100 G, 580 G and 827 G lower than the maximum of the FS-ESE spectrum) is generally optimal. The summed data from these fields leads to orientational-independent DEER measurements. An additional DEER measurement (ca. at the magnetic field ca. 338 G lower than the maximum of the FS-ESE spectrum) is suggested to allow confident interpretation of the minor features in the distance distributions. Using these three magnetic fields, we collected a high-quality distance distribution for hGSTA1-1 and verified the obtained distribution with data at X-band and MD simulation. Importantly, we demonstrated that performing Q-band DEER measurement at

seven additional fields did not show further orientational averaging. This protocol will benefit greatly from recently developed ultra-wideband arbitrary waveform generators⁸⁵ and pent loop-gap resonators.⁸⁴

While this work focuses on the DEER using Cu²⁺-based labels, future application of our Monte-Carlo approach can be applied to ESEEM experiments for Cu²⁺-based systems that also shows orientational selectivity effects.^{92,93} We can also apply this approach to Cu²⁺ distance measurements by alternative pulsed-dipolar spectroscopy techniques.^{94,95} Additionally, this approach can be adapted for other Cu²⁺ systems with higher binding affinity^{61,80–83} and other paramagnetic spins with large spectral bandwidth such as Co²⁺ and Fe³⁺.^{96,97} Further understanding of the orientational selectivity effect will expand the power of pulsed-EPR techniques and make EPR measurements more resource-conscious and widely accessible to the scientific community.

Author Contributions

X.B. performed the experiments, molecular dynamics simulations, and analysed the data. Z.H. wrote the Python program for carrying out the Monte-Carlo simulations and Z.H. and H.R.H. analysed the theoretical results. S.S. conceived the project and supervised all aspects of the work. X.B., Z.H., H.R.H., and S.S. prepared the manuscript. All authors reviewed the manuscript and agreed to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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