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Orientation selection in high-field RIDME and PELDOR experiments involving low-spin Co^{II} ions

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Orientation selective (OS) RIDME and PELDOR were conducted on a low-spin Co^{\parallel} complex coordinated by two nitroxide (NO) labelled 2,2':6',2"-terpyridine ligands. Co-NO RIDME at W- and Q-band gave insight into the relative orientation between the Co-NO interspin vector ($r_{\mathrm{Co-NO}}$) and the NO moiety. This was further supported by W-band Co-NO PELDOR that also allowed elucidating the relative orientation of the Co^{\parallel} and NO g-tensors. Differences to earlier predictions were confirmed by DFT calculations. Finally, NO-NO PELDOR allowed retrieving the mutual orientations between the NO-NO interspin vector ($r_{\mathrm{NO-NO}}$) and the NO moieties. The results demonstrate that OS-RIDME and -PELDOR can provide geometric structure information on a system containing a Co^{\parallel} ion and two nitroxides. Especially, the high sensitivity and ease of interpretation of RIDME at W-band opens avenues for new applications of Co^{\parallel} as orthogonal spin label.

Over the past two decades pulse dipolar electron paramagnetic resonance (PD-EPR) has become valuable in resolving biomolecular architectures^{1, 2} by allowing measurement of nanometre-range distances between intrinsic or chemically harboured unpaired electrons. Nitroxide (NO) labels and endogenous paramagnets are typically used in PD-EPR,³ while other approaches involve the substitution of naturally occurring diamagnetic metal ions by paramagnets.³ Pulsed electron-electron double resonance (PELDOR/DEER)⁴⁻⁶ has been a cornerstone for measuring spin-spin distances (1-10 nm and beyond)⁷ and relaxation induced dipolar modulation enhancement (RIDME)^{8, 9} has recently emerged as

$$\nu_{dip} = \frac{\mu_0 \mu_B^2 g_A g_B}{4\pi\hbar} \frac{1}{r_{AB}^3} (1 - 3\cos^2\theta_{AB}) \tag{1}$$

 \hbar is the reduced Planck's constant, μ_0 the permeability of vacuum, μ_0 is the Bohr Magneton, $g_{\rm i}$ is the g-factor of spin i, r_{AB} is the scalar distance between the two spins and $\theta_{\rm AB}$ is the angle between the distance vector and the external magnetic field. In most cases, a random distribution of θ_{AB} can be assumed leading to the well-known frequency distribution, called Pake pattern. This distribution features "horns" and "edges" which correspond to the distance vector perpendicular to the magnetic field vector (v_{\perp} at $\theta_{\rm AB}$ = 90°) and to a less probable distance vector parallel to the field (v_{\parallel} at $\theta_{\rm AB}$ = 0°), respectively.

Interest in RIDME has been growing over the past three years 10-21 especially for measurements involving spectrally broad paramagnetic metal ions. Here, all spins can act as B spins and contribute to a larger modulation depth often leading to increased sensitivity. 11, 15, 22, 23 In PELDOR the limited excitation of broad EPR spectra by rectangular pulses can lead to selection of certain g-tensor components, a phenomenon referred as 'orientation selection' (OS).^{4, 22-31} If the spin centres are connected with flexible linkers as is often the case in site directed spin labelling³² OS will not affect the distance measurements. However, pronounced OS in combination with rigidly attached spin centres can lead to breakdown of the approximation of a Pake pattern as frequency response. In other words, selection of certain q-tensor values can lead to selection of certain molecular orientations which in turn conditions the values of θ_{AB} making v_{dip} in eq. 1 dependent on the spectral components excited by the mw pulses. With the

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an attractive alternative. PELDOR and RIDME employ microwave (mw) pulses to observe one electron spin (A) whilst inverting a coupled electron spin (B). Inversion of the B spin in PELDOR is induced by a coherent mw pulse, whilst in RIDME it is by longitudinal relaxation.^{9, 10} The systematic variation of the timing of the B-spin inversion leads to a modulation of the A spin signal with the dipolar coupling frequency $(v_{\rm dip})$.

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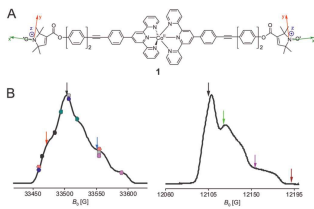


Fig. 1. Structure of complex 1 indicating the molecular axes of NO as investigated by our measurements (A); Field swept spectra of 1 optimized for NO at W- and Q-band (B, left and right, respectively). Circles indicate the detection/inversion positions of NO-NO PELDOR and arrows the detection and inversion position of Co-NO RIDME and PELDOR

special exception of Gd^{III 33} and Mn^{II 34} spins, OS of paramagnetic metals severely increases with high-fields, due to g-anisotropy, which can require measurements at multiple fields to reliably determine the distance(s). On the other hand, OS measurements can also yield atomic-level geometric and electronic structure information, as highlighted in OS-PELDOR of tyrosyl radicals in ribonuclease reductase.³⁰ Strikingly, the accuracy and precision of structural insight provided by OS-PELDOR allowed benchmarking DFT-based modelling parameters for nucleic acids.35

RIDME data also contains OS information with respect to the A spin. OS-RIDME effects were observed on a Cu" system measured at Q-band frequencies; however, they were not analysed to extract geometric information. 19

Cobalt spins have rarely been exploited in PD-EPR, except for one broadband PELDOR study on a Co^{II} porphyrin system at Xband.³⁶ In this work, a novel Co^{II} complex **1** having NO spins as the terminal parts of the terpyridine ligands³⁷ was synthesized

1 is expected to be rigid with negligible exchange coupling between the spins²³ and Co^{II} is known to form low-spin (S = 1/2) complexes with axial g-tensor with strong field ligands such as terpyridines.³⁸ Dependent on substitution, solvation and counter-ions spin-crossover is known to occur at temperatures above those commonly used for EPR distance measurements.39,40

Here, we employed Co-NO OS-RIDME at W- and Q-band frequencies and Co-NO OS-PELDOR at W-band to derive structural information for 1. The results were additionally supported by NO-NO OS-PELDOR at W-band.

Structural model and predicted results

From W-band measurements on similar but shorter and covalently linked bis-nitroxide chemical models^{31, 41} we can predict the components of the Pake pattern that will dominate in OS-PELDOR and -RIDME measurements in 1. To first order, one can approximate the NO bond to be along the backbone, but to fully rationalise⁴² the OS- PELDOR data the NO moiety

had to be modelled rotating freely on a cone with an opening angle of 25° about the molecular backbone. 41 In the Co-NO RIDME experiment the longitudinal relaxation of cobalt is assumed to be isotropic and $v_{\rm dip}$ will solely be determined by the orientations of the NO selected by mw pulses observing the A spin. This will be highly complementary to Gd^{III}-NO PELDOR at W-band where $v_{\rm dip}$ has been shown to be entirely determined by the NO orientations excited.⁴³ Using the common definition of the NO principal axes and the axial gtensor of $Co^{\parallel 38}$ we approximate NO g_x to be parallel to the Co-NO and NO-NO inter-spin vectors (r_{Co-NO} and r_{NO-NO} , respectively) and therefore to the molecular backbone, while NO g_v and g_z to be perpendicular (Fig. 1, A). Accordingly, when exciting NO g_x spins we expect to select molecules with the NO bonds parallel to the magnetic field. Thus, $\emph{r}_{\text{Co-NO}}$ will be confined to be parallel to the field and modulation frequency is expected to correspond to "the parallel component" (v_{\parallel}). Substituting $\theta_{AB} = 0^{\circ}$ in eq. 1 will yield in absolute values the double frequency of the v_{\perp} case ("the perpendicular component" when θ_{AB} = 90°). Detecting NO g_y and g_z is expected to select $\emph{r}_{\text{Co-NO}}$ perpendicular to the magnetic field vector. Furthermore, the geometry is assumed to fix the NO g_x components of both ligands collinearly, while g_{v} and g_{z} components can become interchangeable due to ester bond rotation. Comparison of NO-NO PELDOR experiments with previous results on a similar but shorter covalent bis-nitroxide allows to assess the overall formation and rigidity of the expected NO-Co-NO "yardstick".

The Co-NO PELDOR experiment is expected to reveal information complementary to RIDME further correlating the component of Co g-tensor selected by the observer pulses with the different NO orientations as shown for a similar Cu^{II}nitroxide model compound at X-band.⁴² Due to instrumental bandwidth restrictions, only high fields (equivalently low qvalues) of Co^{II} spectrally close to NO spins can be excited (see Supporting Information (SI), Fig. S1). According to literature³⁸ this component should correspond to a Co q-tensor orientation having the vector between Co^{II} and the central pyridine nitrogen parallel to the field. Thus, according to the approximations made above $\emph{r}_{\text{Co-NO}}$ should be parallel to the magnetic field vector ($\theta_{AB} = 0^{\circ}$) and ν_{\parallel} should be observed when NO g_x spins are inverted, while no modulation is expected when NO g_v or NO g_z spins are inverted.

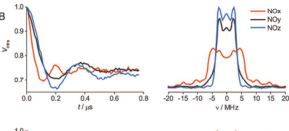
Nitroxide-nitroxide PELDOR

Measurements correlating the xx (denoting the detection and inversion positions, respectively, throughout the text; Fig. 2 A black), yy (dark cyan), zz (pink), zx (salmon), zy (light gray) and xy (blue) NO orientations (and details in SI) are in excellent agreement with previous experiments on shorter covalent bisnitroxides, $^{31,\ 41}$ demonstrating that the expected complex is indeed formed and is equally rigid to a covalently bonded system.

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0.9 0.8 0.7

-2.50 -1.25 0.00 v / MHz 1.25



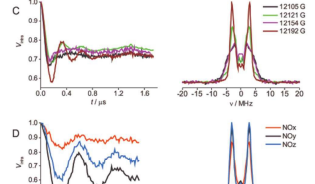


Fig. 2. Background corrected PELDOR and RIDME experiments (left) and corresponding dipolar spectra (right). NO-NO PELDOR at W-band (A); Co-NO RIDME at W- (B) and Q-band (C, here the field values are given due to admixture of NO g_x , g_y and g_z); Co-NO PELDOR at W-band (D). The colours correspond to the colours used in Fig. 1, B.

0.8

-20 -15 -10 o v/MHz 5 10 15

Co-nitroxide RIDME

0.5

Co-NO OS-RIDME was performed at W- and Q-band frequencies (Fig. 2, B and C, respectively) detecting NO spins at different positions of the spectrum (arrows on Fig. 1, B, left and right) with average measurement time 20 min. The Wband RIDME spectra exhibit significant double frequency contributions, v_{\parallel} , when observing on the NO $g_{\rm x}$ component (red arrow) indicating that this orientation must be parallel to $r_{\text{Co-NO}}$. Performing RIDME at y- and z-orientations (black and blue arrows, respectively) gave a modulation frequency, v_{\perp} , corresponding to molecules with $\emph{r}_{\text{Co-NO}}$ perpendicular to the field. At Q-band the dependence of the RIDME signal on the position within the NO spectrum is less pronounced. To better resolve the parallel component of the Pake pattern at 12154 G more selective pulses were applied (SI, Fig. S21-S23). The effects observed in Q-band RIDME are similar to those

observed on the analogous $\operatorname{Cu}^{\text{II}}$ complex 19 and results are in agreement with the NO-NO OS-PELDOR.

Beyond the ease of resolving NO orientation with respect to $r_{\text{Co-NO}}$ providing structural information, performing Co-NO RIDME, itself, opens new routes to metal-based labelling strategies. Firstly, Co^{II} RIDME, in contrast to Co^{II} PELDOR, is orientation-free with respect to cobalt. Secondly, employing high-field ligands commonly used for Mn^{II} and Gd^{III} labelling strategies³³ are expected to yield low-spin Co^{II} tags that in contrast to Mn^{II} and Gd^{III} will not suffer from overtones^{13, 17, 44} during the RIDME experiment since the free electron will undergo only one $(-1/2 \leftrightarrow +1/2)$ transition. Therefore, lowspin Co^{II} tags seem promising for fast, high-sensitive RIDME applications at low-frequency bands.

Co-nitroxide PELDOR

Co-NO OS-PELDOR (Fig. 2, D) was conducted at W-band inverting NO (arrows in Fig. 1, B, left) while observing the low q-component of Co^{II} . The data show that exciting any orientation of NO selects molecules with $\emph{r}_{\text{Co-NO}}$ perpendicular to the field (v_{\perp}) . It follows that if the low g component is assigned to the backbone parallel to the field as predicted above, Co-NO PELDOR would yield the parallel component, v_{\parallel} , while little or no modulation depth would be observed detecting NO g_{v} , g_{z} . However, for all orientations v_{\perp} was observed with the largest modulation depths for Co-NO $g_{\rm v}$ and Co-NO g_z measurements (SI, Table S6) and only very little for Co-NO g_x . These data can only be rationalized if the low Co gcomponent corresponds to an orientation of the backbone perpendicular to the magnetic field. Now, inverting NO q_v or q_z will select complexes with the molecular backbone perpendicular to the magnetic field yielding v_{\perp} and large modulation depth as observed experimentally. To validate this finding, we performed DFT calculations on the bis-terpyridine dication of Co^{II} (details in SI). The calculations show significant deviation from axiality. The eigenvalues were estimated to be 2.005, 2.087, 2.141, with the smallest *q*-value (highest field) corresponding to an eigenvector perpendicular to the Co^{II}-N bond of the central pyridine (corresponding to the backbone of our model system). Thus, the backbone is perpendicular to the magnetic field in excellent agreement with our Co-NO PELDOR data.

Conclusions

Overall, we have demonstrated the feasibility of performing Co-NO OS-RIDME allowing reconstruction of the relative orientation between $r_{\text{Co-NO}}$ and the NO q-tensor. Co-NO OS-PELDOR also gave insight into the Co g-tensor orientation, and was further supported by DFT calculations. NO-NO OS-PELDOR supported our structural model and comparison with earlier data on shorter covalent analogues confirmed the assumed rigidity of 1. Through a combination of W-band OS-RIDME and -PELDOR we have demonstrated that low-spin Co^{II} can be straightforwardly employed for exploring the molecular structure of systems of unknown geometry. The high sensitivity of high power W- and Q-band RIDME make it a

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highly appealing choice among PD-EPR methods for measuring distances involving low-spin Co^{II} ions.

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Conflicts of interest

There are no conflicts to declare.

Notes and references

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