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Deciphering the complex paramagnetic NMR spectra of small laccase

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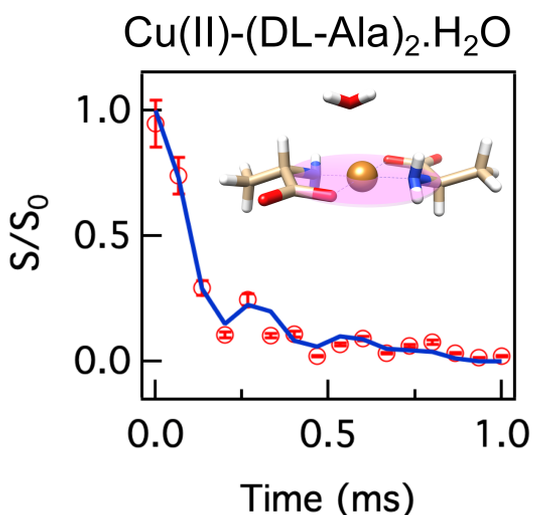
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Chapter 6

Dipolar dephasing for structure determination in a paramagnetic environment



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We demonstrate the efficacy of the REDOR-type sequences in determining dipolar coupling strength in a paramagnetic environment. Utilizing paramagnetic effects of enhanced relaxation rates and rapid fluctuations in $\text{Cu(II)-(DL-Ala)}_2\cdot\text{H}_2\text{O}$, the dipolar coupling strength for the methyl C-H that is 4.20 \AA away from the Cu^{2+} ion, was estimated to be $\sim 8.8 \pm 0.6 \text{ kHz}$. This coupling is scaled by a factor of ~ 0.3 in comparison to the rigid limit value of $\sim 32 \text{ kHz}$, in line with partial averaging of the dipolar interaction by rotational motion of the methyl group. Limited variation in the scaling factor of the dipolar coupling strength at different temperatures is observed. The C-H internuclear distance derived from dipolar coupling strength is similar to the crystal structure. The errors observed in the REDOR-type experiments are similar to those reported for diamagnetic systems. Increase in resolution due to the Fermi contact shifts, coupled with MAS frequencies of $30\text{--}35 \text{ kHz}$ allowed to estimate the hyperfine coupling strengths for protons and carbons from the temperature dependence of the chemical shift and obtain a high resolution $^1\text{H}\text{--}^1\text{H}$ spin diffusion spectrum. This study shows the utility of REDOR-type sequences in obtaining reliable structural and dynamical information from a paramagnetic complex. We believe that this can help in studying the active site of metalloproteins which are paramagnetic in nature at high resolution.

6.1 Introduction

Paramagnetic solid-state NMR has undergone a revolution in the past decade due to the development of high spinning frequency probes and tailored pulse sequences. Studying the ligands that directly coordinate the paramagnetic metal ions with NMR can provide insight into the structure and dynamics of the system.(1–4) Among the various interactions, dipolar interactions are attractive for characterizing fast cooperative dynamics that can be measured using rotational-echo double resonance (REDOR) type solid state NMR experiments including shifted-REDOR for probing anisotropic motions.(5–7)

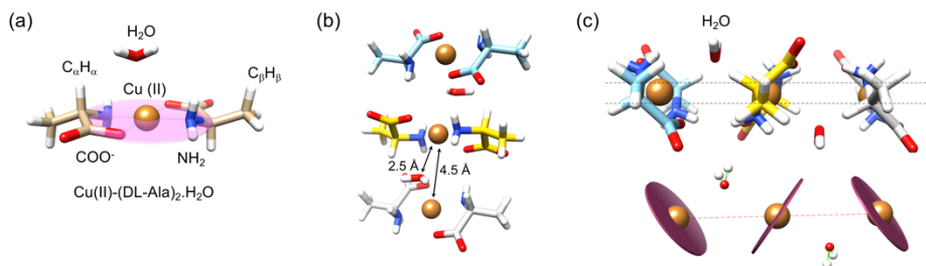


Figure 6.1. *Cu(II)-(DL-Ala)₂.H₂O* crystal structure (cif:1153761).(8) (a) Monomer depicting the plane of coordination and the respective proton and carbon orientations (b) Trimeric representation of chain with distance between copper atoms and between copper and oxygen of water and (c) Coordination plane of copper and position of water molecule.

In this study we show that rapid paramagnetic fluctuations favorably affect REDOR and shifted REDOR experiments therefore, those can be performed with high sensitivity and resolution. We find that internuclear dipolar couplings can be determined with high accuracy that is comparable to what has been achieved for diamagnetic systems. We have selected *Cu(II)-(DL-Ala)₂.H₂O* (Figure 6.1) as a model system to test the utility of the REDOR type sequences and to probe the C–H dipolar coupling of the methyl group in the alanine ligand, which is directly coordinating the paramagnetic metal center.(8–11) Recently, well-resolved heteronuclear dipolar correlation spectroscopy was reported for *Cu(II)-(DL-Ala)₂.H₂O*.(10, 12–14) The possibility to sustain internuclear dipolar transfer in the immediate vicinity of paramagnetic ions can be attributed to rapid paramagnetic fluctuations that average the anisotropic and traceless electron-nuclear dipolar interactions while leaving the isotropic contact shifts unaffected. Here, the J-coupling between the copper ions determines the fluctuation frequencies for averaging dipolar couplings, which are traceless without isotropic component and is intrinsic to a one-dimensional chain. This results in the motional narrowing from spin waves running along the chain.(15) This opens up a window of opportunity, since essential advantages of paramagnetism, such as rapid acquisition and increased spectral dispersion can be combined with the power of dipolar correlation spectroscopy for resolving static or dynamic structure quantitatively. To demonstrate this, we implement a REDOR method for *Cu(II)-(DL-Ala)₂.H₂O* by adapting the REDOR refocusing period to the fast rotation needed for signal detection in paramagnetic systems with their enhanced relaxation that limits the acquisition time. The *Cu(II)-(DL-Ala)₂.H₂O* form 1-dimensional chains in which the Cu^{2+} ions are weakly coupled.(11) The antiferromagnetic coupling $2J = -2.21 \text{ cm}^{-1}$ translates into a characteristic frequency of $\sim 70 \text{ GHz}$ for the electronic spin fluctuations.(11) The properties of enhanced relaxation and rapid averaging of electron-nuclear dipolar interactions by paramagnetic fluctuations that are much faster than the frequency scale of the heteronuclear dipolar coupling of $\sim 10 \text{ kHz}$ or less, allow to perform heteronuclear dipolar recoupling experiments with fast MAS in a very short time, much shorter than for diamagnetic species. In addition, we show how $^1\text{H} \square ^1\text{H}$ spin diffusion spectra and the temperature dependence of proton and carbon shifts from 1D Hahn-echo experiments can help to collect structural information from paramagnetic systems with ultra-fast MAS.

6.2 Results and discussions

NMR characterization of Cu(II)-(DL-Ala)₂.H₂O

There are several advantages when working with Cu(II) paramagnetic systems. Since Cu(II) has small susceptibility the Fermi-contact interaction is stronger than the paramagnetic relaxation enhancement (PRE).⁽¹⁶⁾ This results in a better resolution because of the increase in the spectral dispersion without adversely affecting the linewidth. Additionally, reduction of experimental time due to fast longitudinal relaxation compared to diamagnetic systems can be achieved. In contrast, with other metal ions such as Tb(III) the susceptibility anisotropy is large and may lead to resonances broadened beyond detection due to increase in PRE-effects.⁽¹⁶⁾ Figure 6.2 shows the 1D spectra of ¹H and ¹³C at different spinning frequencies and magnetic fields. The resolution is enhanced by the paramagnetic shifts at very fast MAS (> 20 kHz).^(12–14, 17–19)

The 1D proton rotor-synchronized Hahn-echo spectra (Figure 6.2a and b) show three prominent resonances attributed to CH_α at 8 ppm and CH₃1 at 32 ppm and CH₃2 at 21 ppm. We observed that the intensity of CH₃1 is 4 times that of H_α and CH₃2, while H_α and CH₃2 show similar intensities. The carbon spectrum shows three prominent resonances for C_α1 at -280 ppm, CO at -200 ppm and C_β1 at 180 ppm and two minor resonances for C_α2 at -258 ppm and C_β2 at 88 ppm, (Figure 6.2c, 6.2d and Table S6.1).^(10, 13, 19) The two components were attributed to two orientations of the methyl group as a consequence of crystal packing, where the major component (CH₃1/C_β1) is closer to the copper atom while the minor component (CH₃2/C_β2) is away.^(8, 20) The proton signals from the NH₂ group (data not shown) were reported to be at -132.7 and -159.5 ppm.⁽¹⁰⁾ The protons from the water molecule are generally broadened beyond detection due to fast exchange dynamics.^(10, 19) The methyl group of the major component was reported to be undergoing fast rotation which results in a single resonance for the three methyl protons (Figure 6.2a and b).⁽¹⁰⁾

A dipolar-INEPT experiment was performed to check the quality of the sample and the efficiency of polarization transfer via dipolar interactions, which was previously shown to be superior to cross polarization based transfer for paramagnetic systems.^(13, 21) It shows the similar spectrum (Figure 6.2d) as reported earlier albeit with better resolution and S/N allowing to identify the splitting of the methyl group response that arises due to crystal packing effects.^(8, 13, 20)

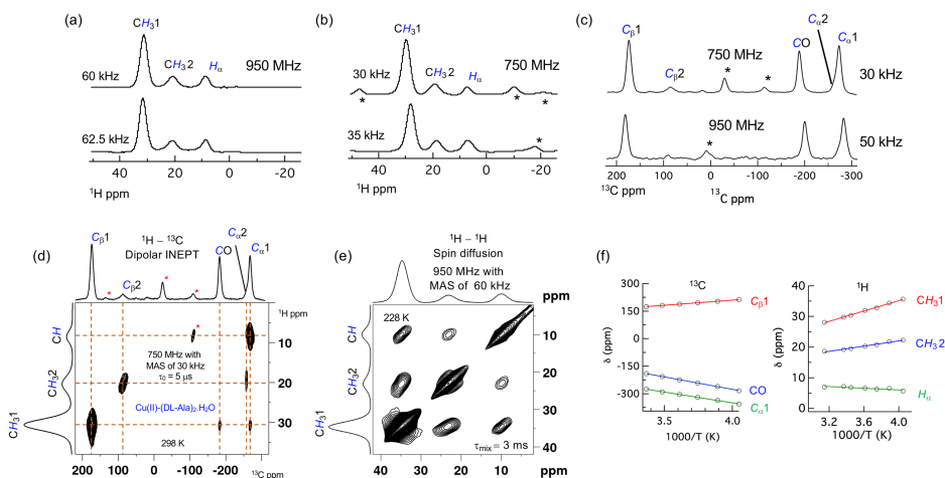


Figure 6.2. (a) 1D ^1H rotor synchronized Hahn-echo spectra at MAS frequencies of 60 and 62.5 kHz in an external magnetic field of 950 MHz; (b) 1D ^1H rotor synchronized Hahn-echo spectra at MAS frequencies of 30 and 35 kHz in an external magnetic field of 750 MHz; (c) 1D ^{13}C rotor synchronized Hahn-echo spectra at MAS frequencies of 30 and 50 kHz in an external magnetic field of 750 and 950 MHz respectively; (d) The 2D ^1H - ^{13}C dipolar-INEPT spectra at a MAS frequency of 30 kHz in an external magnetic field of 750 MHz with a transfer delay τ_0 of 5 μs at 298 K; (e) 2D ^1H - ^1H spin diffusion at a MAS frequency of 60 kHz in an external magnetic field of 950 MHz with a mixing time τ_{mix} of 5 ms at 228 K; (f) Temperature dependence of the chemical shift for carbon and protons where CH_31 and C_β are in red, CH_32 and carbonyl carbon CO are in blue and H_α and C_α are in green. Solid lines are the fit using equation S6.9. The spinning side bands in the upper panels are marked with asterisks. The assignments in panel (a) to (e) of ^1H and ^{13}C responses are shown in italics and blue color.

The 2D ^1H - ^1H spin diffusion spectra acquired at 950 MHz with a MAS frequency of 60 kHz (Figure 6.2e and S6.3), show that a high resolution homonuclear 2D spectra can be obtained under paramagnetic conditions with modification of the pulse sequence to consider the paramagnetic effects of increased relaxation and rapid fluctuations. Similar observations were reported for copper cyclam complexes.(21)

The temperature dependence of the chemical shift shows strong Curie behavior for CO, $\text{C}_\alpha1$, $\text{C}_\beta1$, CH_31 and CH_32 while weak anti-Curie behavior is observed for H_α (Figure 6.2f and Figure S6.2a). The hyperfine coupling strengths are summarized in table 1, obtained after fitting the data with equation 9 (Figure 6.2f).(22) It is interesting to observe that, although the H_α is close to the copper, the hyperfine coupling constant and hence the spin density on it is less than for H_β . This was also observed in DFT calculations for $\text{Cu(II)-(DL-Ala)}_2\cdot\text{H}_2\text{O}$ showing very little spin density localized on the H_α atom.(22, 23) The sign of the hyperfine coupling strength provides

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the spin polarization of the electron on the nucleus with respect to the external magnetic field. It was observed to be parallel for $C_{\beta}1$, CH_31 and CH_32 and anti-parallel for $C_{\alpha}1$, CO and H_{α} (Table 6.1 and Figure 6.2f). These observations are in line with the calculations performed on copper alanine complexes and previously reported data.(22) The estimated δ_{dia} from the fit of the temperature dependencies of the chemical shift (Figure 6.2f) is summarized in table 6.1. They differ from the expected diamagnetic chemical shift of DL-alanine (Table S6.3). This difference can be attributed to the differences in crystal packing of $Cu(II)-(DL-Ala)_2 \cdot H_2O$ and DL-alanine (Figure S6.6). The NMR chemical shift tensor for small molecules in solid state NMR differs significantly between different crystal forms due to changes in hydrogen bonding, stacking, ring current, $C-H \cdots \pi$ interactions etc.(24) DFT calculations of different crystal forms of purine derivatives show that the change in the diamagnetic chemical shift can be as high as 6.5 ppm, 20 ppm and 55 ppm for proton, carbon and nitrogen respectively.(24)

Table 6.1. Hyperfine coupling strength (A) and the estimated diamagnetic chemical shift from temperature dependencies of the resonances for carbon and protons.

| Nuclei | A (MHz) | δ_{dia} (ppm) |
|-----------------------------|-----------------|----------------------|
| $C_{\alpha}1$ | -3.7 ± 0.3 | 120 ± 4 |
| $C_{\beta}1$ | 1.8 ± 0.9 | -12 ± 1 |
| CO | -4.3 ± 0.4 | 265 ± 5 |
| H_{α} | -0.2 ± 0.04 | 11.0 ± 1.0 |
| CH_32^* | 0.5 ± 0.03 | 5.0 ± 0.9 |
| CH_31^* | 1.0 ± 0.03 | 1.5 ± 0.8 |

* The protons in question are indicated in bold.

The full width at half of the maximum (FWHM) of the resonances decreases with increasing temperature (Figure S6.2b and S6.2c). Most likely, an increase in the electronic relaxation rate with temperature reduces the paramagnetic relaxation enhancement.

Dipolar coupling strength estimation

Dipolar coupling measurements can provide insight into the dynamics of the molecule.(5, 25) Here, we have employed REDOR (Figure S6.4c) and shifted-REDOR (Figure S6.4d) to measure the dipolar coupling between 1H and ^{13}C of the methyl group in $Cu(II)-(DL-Ala)_2 \cdot H_2O$.(5, 6) The choice of these pulse sequences was based on their robustness for amplitude mis-setting, inhomogeneities in the radiofrequency field, carrier offset and CSA.(5) The methyl group was used as a probe because of the

presence of intrinsic methyl rotation that partially averages the dipolar coupling and its higher S/N ratio compared to that of other protons (Figure 6.2a and b).

Figure 6.3a shows the ^1H - ^{13}C REDOR curves of the C-H in CH_3I group at 277 K, 287 K and 297 K obtained at a MAS frequency of 30 kHz. The dipolar coupling strengths estimated after the fitting are 8.6 kHz, 10.0 kHz and 9.2 kHz respectively (Table 6.2 and Figure 6.3b). These values correspond with a scaling factor of ~ 0.3 (Table 6.2) relative to the rigid limit value of ~ 32 kHz, consistent with previously reported data for alanine in the diamagnetic crystalline state and similar to the theoretical scaling factor of 0.333.(26) The accuracy of the paramagnetic measurement is similar to that reported for the diamagnetic systems.(5, 6)

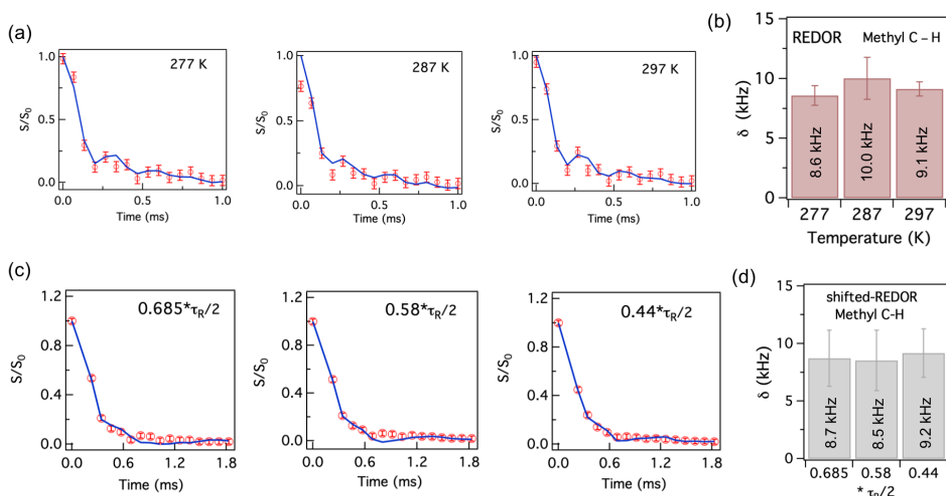


Figure 6.3. (a) REDOR curve for methyl C-H ($C_{\beta}1$ and CH_31) in $\text{Cu(II)-(DL-Ala)}_2\cdot\text{H}_2\text{O}$ at 277 K, 287 K and 297 K, the solid lines represent the fit; (b) The dipolar coupling strength estimated from fitting the REDOR curve for the methyl C-H ($C_{\beta}1$ and CH_31); (c) Shifted-REDOR curves at a MAS frequency of 30 kHz with different shift of the π pulse from ideal $\tau_R/2$ (16.67 μs). The solid blue lines are fit with asymmetric parameter η value of 0.1; (d) Dipolar coupling strength estimated from the shifted-REDOR curve fit of methyl C-H ($C_{\beta}1$ and CH_31).

The corresponding internuclear ^1H - ^{13}C ($C_{\beta}1$ and CH_31) distances derived after dividing the dipolar coupling strength with the scaling factor is ~ 1.0 \AA (Table 6.2) which is very similar to the crystal structure (1.0 \AA). (8, 23) This demonstrates the efficacy of the REDOR sequence in obtaining reliable distance information in a paramagnetic system. It is noteworthy that although the paramagnetic shift is temperature dependent the dipolar coupling strength is preserved. This property can be explored to characterize the dipolar coupling strength between the nuclei in the

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vicinity of the metal at the active site of the paramagnetic metalloproteins and consequently get insight into the dynamics.

Table 6.2. Dipolar coupling strength between C-H ($C_{\beta}1$ and CH_31) at different temperatures using REDOR with a MAS frequency of 30 kHz.

| Temp (K) | δ kHz | Scaling factor | Methyl C-H ($C_{\beta}1$ and CH_31) distance (\AA) |
|----------|----------------|-----------------|---|
| 277 | 8.6 ± 0.8 | 0.29 ± 0.03 | 1.1 ± 0.1 |
| 287 | 10.0 ± 1.7 | 0.34 ± 0.06 | 1.0 ± 0.2 |
| 297 | 9.1 ± 0.6 | 0.32 ± 0.02 | 1.0 ± 0.1 |

It was recently reported that using a variant of REDOR where the π pulses are shifted from its original position, known as shifted-REDOR (Figure S6.4d), is able to identify anisotropic motions present in the protein via the asymmetry parameter η easily by scaling down the oscillation.^(5, 6) We investigated the dephasing behavior in the presence of the paramagnetic metal center at a spinning frequency of 30 kHz and at 297 K with the π pulse shift (τ_S) of 0.685 or 0.58 or 0.44 times $\tau_R/2$ (Figure 6.3c). The dipolar coupling strength between the methyl C-H nuclei does not change with varying τ_S and it matches with the values obtained from REDOR (Table 6.3 and 6.4). The derived internuclear distance is $\sim 1.0 \text{ \AA}$ (Table 6.3), consistent with the crystal structure and similar to REDOR. Hence, shifted-REDOR appears also robust and reliable for obtaining structural information. The asymmetry parameter η of 0.1 gave the best fit, which is in line with the lack of anisotropic motion in the methyl group rotation. While there was no anomaly in the implementation of shifted-REDOR for Cu(II)-(DL-Ala)₂.H₂O, in shifted-REDOR (Figure 6.3c), the oscillatory behavior observed in the REDOR (Figure 6.3a) is diminished which can be due to the presence of additional protons (NH₂ and H α). This is consistent with the observations of Schanda *et al.* (2011), who showed that the presence of additional protons near the observed nuclei reduces the oscillatory nature of the shifted-REDOR for ubiquitin crystals.^(5, 6)

Table 6.3. Dipolar coupling strength of methyl C-H (CH_31) using shifted-REDOR and at different fraction of $\tau_R/2$ by which the π pulse is shifted at MAS frequency of 35 kHz.

| Fraction of $\tau_R/2$ shift | δ kHz | Scaling factor | Methyl C-H distance (\AA) |
|------------------------------|---------------|-----------------|--------------------------------------|
| 0.685 | 8.7 ± 2.4 | 0.30 ± 0.08 | 1.0 ± 0.3 |
| 0.580 | 8.5 ± 2.6 | 0.29 ± 0.09 | 1.1 ± 0.3 |
| 0.440 | 9.2 ± 2.1 | 0.32 ± 0.07 | 1.0 ± 0.2 |

In summary, both REDOR and shifted-REDOR sequences provide reliable estimates of the dipolar coupling strength and consequently information about the structure and methyl rotation.

6.3 Conclusions

This study shows that high resolution ^1H based solid-state NMR spectra can be obtained for a paramagnetic system at fast MAS (> 25 kHz). Using the high sensitivity of ^1H and the high resolution due to the paramagnetic nature of Cu(II), a temperature dependence study of the model compound Cu(II)-(DL-Ala) $_2$.H $_2$ O could be performed that shows that the shift correlates with the spin delocalization over the ligands. The fast MAS has enabled the implementation of ^1H - ^1H homonuclear spin diffusion, REDOR and shifted-REDOR pulse sequences. Our benchmark is the existing crystal structure. The error in the measured values were similar to that observed for diamagnetic systems and are largely systematic errors from e.g. thermal motions that effectively reduce the dipolar coupling from its rigid limit value. We present a technical step forward by showing that both REDOR and shifted-REDOR are quantitatively reliable in determining the dipolar coupling strength and internuclear distances near the paramagnetic metal center, which paves the way for future application in metalloproteins to probe fast dynamics. In addition, the results contribute to the converging and convincing evidence that paramagnetic solid-state NMR is increasingly accessible and can be interpreted in the same way as the NMR spectra of diamagnetic solids.

6.4 Supporting Information

Sample preparation and solid-state NMR experiments

Cu(II)-(DL-Ala)₂.H₂O complex was synthesized as reported earlier.⁽¹⁰⁾ All chemicals including the uniformly labelled ¹³C-DL-alanine were obtained from Sigma Aldrich (USA).

Solid-state NMR experiments were performed with AV-I (17.6 T) 750 MHz and AV-HDIII (22.31 T) 950 MHz spectrometers equipped with a 2.5 mm and 1.3 mm MAS probe, respectively, at spinning frequencies ranging from 30 to 62.5 kHz. 1D proton and carbon spectra were recorded with a rotor synchronized Hahn-echo sequence (Figure S6.4a). The high-power $\pi/2$ pulses were 2 μ s and 2.5 μ s for ¹H and ¹³C, respectively. A dead time of 4.5 μ s was subtracted from the refocusing period in the Hahn-echo pulse sequence. A comparison of spectra obtained with single 90° pulse (Figure S6.1) with rotor synchronized Hahn-echo spectra (Figure 6.2) confirms that there is little phase distortion in echo spectra obtained with a large spectral window (SW).⁽⁴⁾ The advantage of using rotor-synchronized Hahn-echo over single 90° pulse is elimination of probe artifacts as observed from Figure S6.1a and Figure 6.2a. This prompted us to use rotor-synchronized Hahn-echo pulse sequence to obtain 1D spectra for both carbon and proton for further study.

The 1D and 2D dipolar-INEPT sequence was adopted from Wickramasinghe *et al.*, (2008) with a $\pi/2$ pulse length similar to the rotor synchronized Hahn-echo experiment.⁽¹³⁾ Temperature dependent experiments were performed at temperatures between 247 K and 318 K. For the 2D ¹H–¹H spin diffusion (Figure S6.4b) experiment the mixing time was varied from 1 ms to 5 ms with a total acquisition time of 530 μ s and a delay of 30 ms between scans. 200 slices of data were acquired in the t1 dimension in a total experiment time of < 5 mins. REDOR and shifted-REDOR data were collected by direct excitation of ¹³C (Figure S6.4c and S6.4d).^(5, 6)^(27–29) The $\pi/2$ pulse lengths were 2 μ s and 2.2 μ s for ¹H and ¹³C respectively. Temperature calibration was done using samarium tin oxide and lead nitrate.^(30–32) Data analysis was performed with TOPSPIN 4.0.5 (Bruker) and Igor pro 6.37.

REDOR and shifted-REDOR data analysis

The REDOR and shifted-REDOR curves were fitted with SIMPSON 4.2.1.^(33, 34) The geometry of Cu(II)-(DL-Ala)₂.H₂O was adopted from Calvo *et al.*, and Zhang *et al.*, to generate the spin system parameters, dipolar coupling and Euler angles using SIMMOL-VMD.^(8, 23, 35, 36) The signal from the methyl C-H was used as a reporter to determine the dipolar coupling strength due to its intrinsic isotropic motion and

high S/N ratio (Figure 6.2a and b). The fitting algorithm as implemented in SIMPSON package was used to estimate the dipolar coupling.(33–35)

The rigid limit dipolar coupling strength of the methyl C-H is ~ 32.1 kHz which corresponds to an internuclear distance of 0.98 \AA .(23) It is known that methyl rotation partially averages the dipolar coupling between the C-H bond of the methyl group and therefore the rigid limit dipolar coupling is scaled according to(26, 37)

$$D_{avg} = S * D, \quad (S6.1)$$

with D being the dipolar coupling strength in the rigid limit. The scaling factor is

$$S = (3\cos^2\theta - 1)/2, \quad (S6.2)$$

where θ is the angle between the methyl C-H and the rotational axis, which in this case is the bond between $C_\alpha - C_\beta$. For a tetrahedral geometry of the methyl group $\theta = 109.5^\circ$. This gives a theoretical scaling factor of $S = 0.333$ (Figure S6.5) and $D_{avg} = 10.69$ kHz for the C-H bond in Cu(II)-(DL-Ala)₂.H₂O in the rotating methyl group. For the analysis of the data, the scaled $D_{avg} = 10.69$ kHz was changed to fit the experimental data. The average dipolar coupling strength was optimized to fit the REDOR curve using the fitting algorithm of SIMPSON. The homonuclear dipolar coupling between the three protons of the methyl group was also scaled by the theoretical value of 0.333 to give a $D_{avg} = 9.76$ kHz.(23) The codes for fitting and to obtain the spin-system are uploaded in https://github.com/rubindg/REDOR_fitting_simpson.

Theory of temperature dependence of Fermi contact shifted resonances

The chemical shift of the paramagnetic system is temperature dependent and the observed chemical shift is given by (16)

$$\delta_{obs} = \delta_{dia} + \delta_{FCS} + \delta_{PCS}, \quad (S6.3)$$

where δ_{dia} is the diamagnetic chemical shift and δ_{CS} , δ_{PCS} are the contributions from the Fermi contact shift (FCS) and pseudo-contact shift (PCS) respectively.(16, 38) For the nuclei of coordinating ligands $FCS \gg PCS$ and eq. 1 can be written as

$$\delta_{obs} \approx \delta_{dia} + \delta_{FCS}. \quad (S6.4)$$

The contribution from the FCS is given by(38)

$$\delta_{FCS} = \frac{A\langle S_z \rangle}{\hbar\gamma_I B_0}, \quad (S6.5)$$

with $A = \frac{2}{3} \hbar\gamma_I\mu_B g_e\mu_0\rho$ the isotropic hyperfine coupling that is proportional to the spin density ρ on a nucleus. The $\langle S_z \rangle$ is the expectation value of the z magnetization, \hbar is

the reduced Planck constant, γ_I is the gyromagnetic ratio of the nucleus in question, and B_0 is the external magnetic field.

Since the $\langle S_z \rangle$ is related to the magnetic susceptibility according to (38, 39)

$$\langle S_z \rangle = \frac{B_0 \chi}{\mu_B g_e \mu_0}, \quad (\text{S6.6})$$

where μ_B is the Bohr magneton, $g_e \approx 2.003$ is the free electron g -value, and μ_0 is the permeability of the vacuum, the FCS can also be expressed in terms of the magnetic susceptibility χ by

$$\delta_{FCS} = \frac{A \chi}{\hbar \gamma_I \mu_B g_e \mu_0}. \quad (\text{S6.7})$$

The system is a linear chain in the high temperature limit, since $2J = -2.2 \text{ cm}^{-1}$, which translates into $\sim 3 \text{ K}$. Therefore, we are in the high temperature limit, $|2J| \ll T$, and the experiments for $\text{Cu(II)-(DL-Ala)}_2 \cdot \text{H}_2\text{O}$ were performed at this limit. The magnetic susceptibility is given by (40)

$$\chi = \frac{S(S+1) \mu_B^2 g_e^2 \mu_0}{3 k_B T}, \quad (\text{S6.8})$$

where $S = 1/2$ is the spin of the Cu(II) ion, k_B is the Boltzmann constant and T is the temperature in Kelvin. Substituting eq. 6 in eq. 5, and in eq. 2 we obtain

$$\delta_{obs} \approx \delta_{dia} + A \frac{S(S+1) \mu_B g_e}{3 \hbar \gamma_I k_B T} \quad (\text{S6.9})$$

for the temperature dependent chemical shift, which was used to determine hyperfine couplings and spin densities by fitting to the data in Figure 6.3.

Supporting figures and tables

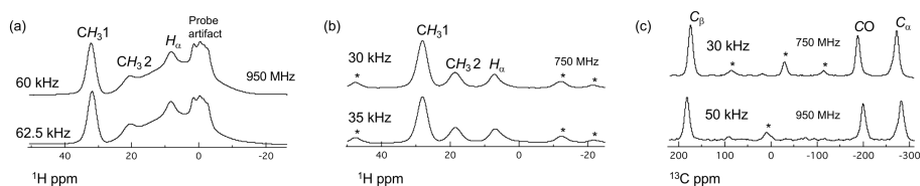


Figure S6.1. Single 90° pulse experiment for ^1H at (a) 950 MHz with MAS of 60 kHz and 62.5 kHz, (b) 750 MHz at MAS of 30 kHz and 35 kHz and for ^{13}C at (c) 750 MHz with MAS of 30 kHz and 950 MHz with MAS of 50 kHz. Resonances marked with * are the spinning side bands. ^1H and ^{13}C assignments are indicated with italics.

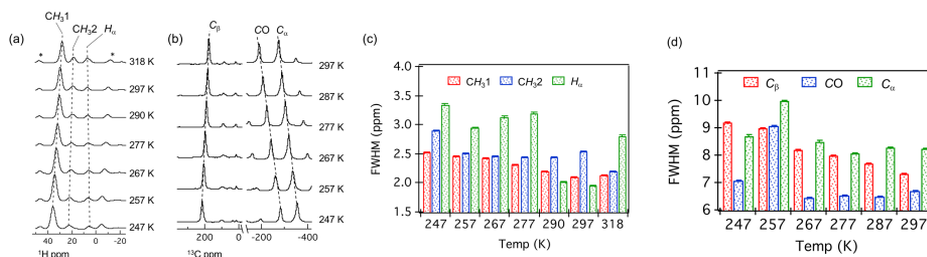


Figure S6.2. Temperature dependence of (a) Proton and (b) Carbon signals at 30 kHz MAS in a magnetic field of 14 T. The full width at half of the maximum FWHM of (c) Protons (red = $\text{CH}_3 1$, blue = $\text{CH}_3 2$ and green = H_α) and (d) Carbons (red = C_β , blue = CO and green = C_α) at different temperature. The peaks marked with stars are the spinning side bands while the smaller peaks in carbon spectra are from small impurities and spinning side bands. Assignments of ^1H and ^{13}C signals are indicated in italics.

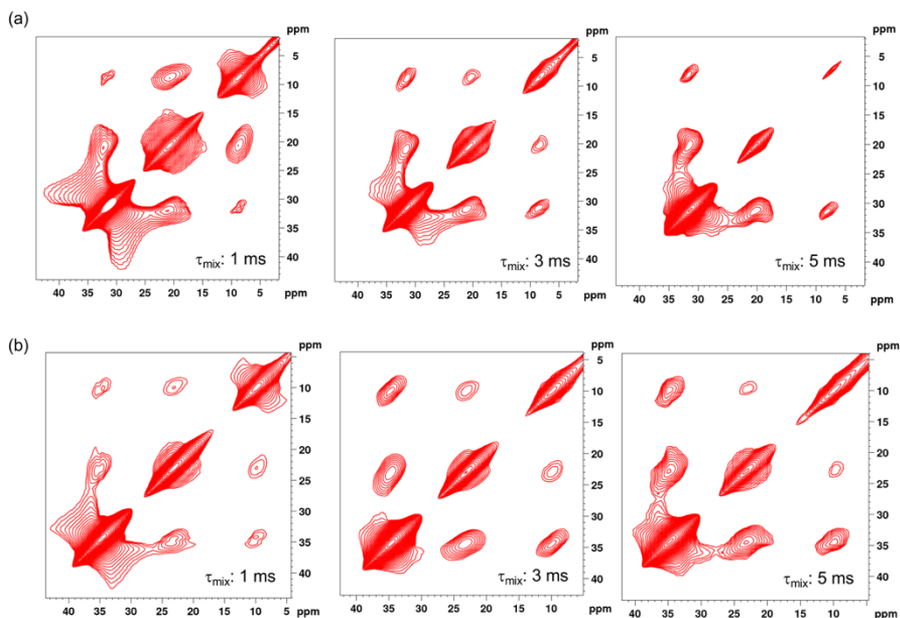


Figure S6.3. ^1H – ^1H spin diffusion spectra at (a) 300 K and (b) 228 K at 60 kHz MAS and 950 MHz ^1H frequency with mixing times of 1 ms, 3 ms and 5 ms.

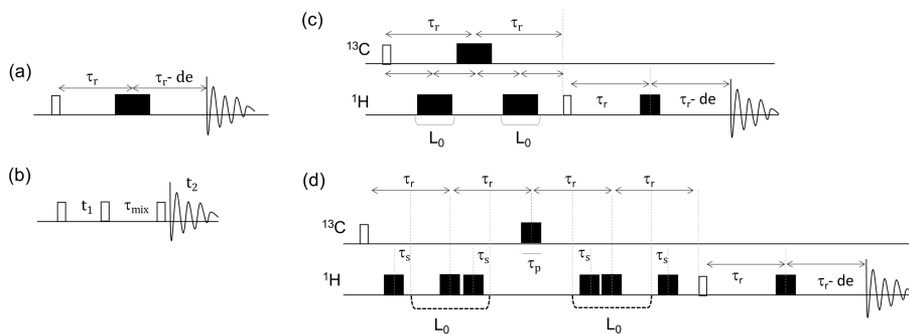


Figure S6.4. Pulse sequences used in this study. (a) Rotor synchronized Hahn-echo, (b) ^1H – ^1H spin diffusion, (c) Proton detected standard REDOR and (d) Proton detected shifted-REDOR. τ_r denotes the rotor period, t_1 and t_2 are the two-time domains for 2D spectra, τ_s is $(0.68 \text{ to } 0.44) \cdot \tau_r / 2$, τ_p is the π pulse length, de is the dead time of the instrument, τ_{mix} is the mixing time and L_0 is the loop counter for repeating the π pulses.

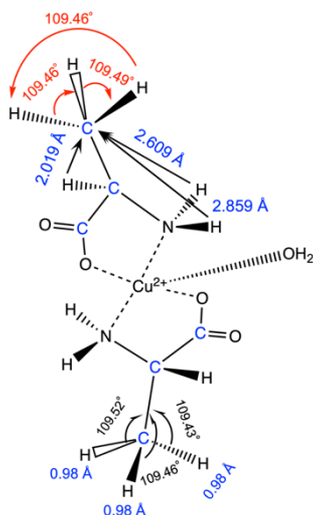


Figure S6.5. Geometry of $\text{Cu(II)-(DL-Ala)}_2 \cdot 2\text{H}_2\text{O}$ adopted from Zhang et al.,(23)

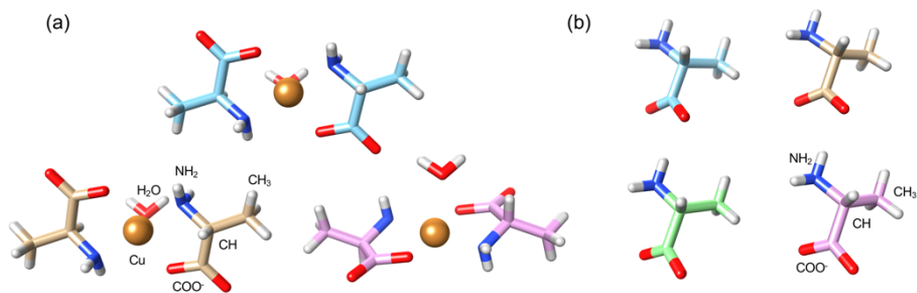


Figure S6.6. Crystal packing in a unit cell of (a) $\text{Cu(II)-(DL-Ala)}_2 \cdot 2\text{H}_2\text{O}$ (cif:1153761)(8) and (b) *DL*-alanine (cif:164668)(41).

Table S6.1. Assignment of ^1H spectra from the literature on $\text{Cu(II)-(DL-Ala)}_2\cdot\text{H}_2\text{O}$. (10, 19)
The magnetic field used was from 7.5 T to 11.75 T with spinning frequencies between 15 and 30 kHz.

| Protons of group | δ_{iso} (ppm) |
|----------------------|-----------------------------|
| CH_3 | 32 & 20.4 |
| CH | 8.4 |
| NH_2 | -132.7 & -159.5 |
| H_2O | 1 |

Table S6.2. Properties of $\text{Cu(II)-(DL-Ala)}_2\cdot\text{H}_2\text{O}$ experimentally determined.

| Properties | | Reference |
|--|----------------------------------|-----------|
| Intrachain AFM exchange coupling constant | $-2.27 \pm 0.08 \text{ cm}^{-1}$ | (11) |
| $2J_0$ at $T > 1.8 \text{ K}$ | | |
| g-factor | 2.091 ± 0.005 | (11) |
| Curie temperature from NMR | 6.5 K | (20) |
| Curie Temperature from $\chi_M(T)$ vs $\log T$ | 2.1 K | (8) |
| ^{13}C $T_{1\rho}$ @ 344 K | ms | |
| CO | 8.7 ± 0.1 | (17) |
| CH | 13.9 ± 0.3 | |
| CH_3 | 50.4 ± 1.9 | |
| Δ_{aniso} @ 297 K | ppm | |
| CO | -331 ± 4 | (17) |
| CH | -149 ± 10 | |
| CH_3 | 53 ± 2 | |

* The ^{13}C assignments are indicated in italics.

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