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Measurement of ¹⁴N quadrupole couplings in biomolecular solids using indirect-detection ¹⁴N solid-state NMR with DNP†

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The quadrupolar interaction experienced by the spin-1 ¹⁴N nucleus is known to be extremely sensitive to local structure and dynamics. Furthermore, the ¹⁴N isotope is 99.6% naturally abundant, making it an attractive target for characterisation of nitrogen-rich biological molecules by solid-state NMR. In this study, dynamic nuclear polarization (DNP) is used in conjunction with indirect ¹⁴N detected solid-state NMR experiments to simultaneously characterise the quadrupolar interaction at multiple 14N sites in the backbone of the microcrystalline protein, GB3. Considerable variation in the quadrupolar interaction (>700 kHz) is observed throughout the protein backbone. The distribution in quadrupolar interactions observed reports on the variation in local backbone conformation and subtle differences in hydrogen-bonding; demonstrating a new route to the structural and dynamic analysis of biomolecules.

in biological and naturally occurring materials. However, solid-state NMR (ssNMR) studies of the 99.6% naturally abundant ¹⁴N isotope are challenging principally due to the integer spin number (I = 1)¹⁴N isotope difficult to manipulate and detect with ssNMR. Accordingly, ssNMR studies typically favour the spin-1/2 nucleus ¹⁵N.

The quadrupolar interaction is, however, extremely sensitive to the local structure and dynamics experienced at the ¹⁴N site, and its characterisation could provide a wealth of information not readily available from the spectra of the more commonly studied 15N isotope. 1-3 One area where 14N NMR may offer significant benefits is in the field of protein ssNMR. The quadrupolar interaction experienced by each of the amide nitrogens in the peptide backbone is potentially a sensitive reporter to the backbone conformation, secondary structure and dynamics. Indeed evidence from small model peptides indicates that differences in the nature of hydrogen bonding between α-helical and β-sheets conformers can result in significant differences in the magnitude of the 14N electric field gradient (EFG) at the amide nitrogen, regardless of the type of residue.^{1,4} Furthermore, the analysis of natural abundance ¹⁴N in proteins opens new avenues for the study of complex medical and environmental samples that cannot be labelled for NMR studies.3

A number of strategies have been developed in order to characterise the ¹⁴N quadrupolar interaction (QI) in a variety of organic and biological materials. These include direct detection of static wideline 14N ssNMR spectra with broadband excitation methods.⁵⁻⁹ Alternatively, the ¹⁴N overtone transition may also be directly detected since this can potentially provide improved resolution 10-14 and sensitivity 15,16 since the overtone transition is unaffected by the first order quadrupolar interaction. Several methods have also been proposed for indirectly detecting 14N that employ a spin-1/2 "spy" nucleus such as 13C or 1H to indirectly detect the fundamental 14N transition in a 2D experiment.2,17-24 The indirect detection strategy is perhaps the most promising for application to complex molecules with multiple ¹⁴N sites since it benefits from the resolution and sensitivity of the "spy" nucleus, and does not suffer the low excitation bandwidth of overtone methods. We have recently reported a technique for such indirect detection that exploits moderate rf fields close to the ¹⁴N Larmor frequency, as opposed to a period of free evolution as proposed by a number of other groups, 2,17-22 to generate coherence between the 14N and spy nuclei.24

Nitrogen is an important element in chemistry due to its prevalence and moderate electric quadrupole moment of the ¹⁴N nucleus. This results in ¹⁴N spectra that are anisotropically broadened by the first order quadrupolar interaction to several MHz, rendering the

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In this communication we demonstrate the feasibility of recording 13 C/ 14 N correlation spectra in a full-length protein using such an indirect detection scheme, allowing the determination of the distribution of 14 N quadrupolar couplings throughout the protein backbone of third IgG-binding domain of Protein G (GB3).

Dynamic nuclear polarization (DNP) offers significant improvements in sensitivity over conventional MAS-NMR, 25,26 aiding many biomolecular NMR studies.²⁷ Here we have applied high field MAS-DNP at a field of 18.8 T with sample temperatures of ca. 110 K in conjunction with 2D ¹³C/¹⁴N correlation spectra to enhance sensitivity and obtain spectra of a microcrystalline preparation of ¹³C-labelled GB3. Full experimental details are provided in the ESI.† For DNP measurements, two microcrystalline samples of GB3 were incubated with either 2 or 12.5 mM of the biradical AMUPol²⁸ in glycerol-d₈/D₂O (70:30 v/v). In the ¹H/¹³C cross polarization (CP) spectra of these two samples, shown in Fig. 1A and B, DNP enhancements ($\varepsilon_{\text{on/off}}$) of 3 and 18 were observed, respectively with relatively uniform enhancements throughout the spectrum, something mirrored in the corresponding proton-spin diffusion spectrum (ESI,† Fig. S2). Compared to spectra acquired at 273 K, a significant decrease in resolution is observed due to inhomogeneous broadening that arises upon freezing, although the resolution is favourable when compared to spectra of the homologous protein GB1 at 100 K²⁹ (Experimental details and a discussion on the resolution observed can be found in the ESI†).

The ¹⁴N filtered ¹³C spectra are shown in Fig. 1C and D. In contrast to the ¹H/¹³C CP-MAS spectra, the ¹⁴N filtered ¹³C signal shows only the ¹³C resonances of nuclei bound to a ¹⁴N spin. Furthermore, we notice that the ¹⁴N filtered ¹³C spectrum

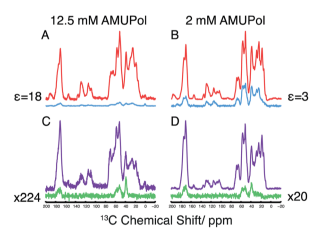


Fig. 1 DNP enhancements and 14 N transfer efficiencies recorded from ~ 30 mg of GB3 in AMUPol. All spectra recorded at 18.8 T, 100 K with 13.5 kHz MAS. CP-MAS 13 C NMR spectra in (A) and (B) were recorded on samples containing 12.5 mM AMUPol and 2 mM AMUPol, respectively. Spectra in red and blue show the intensity of the 13 C CP-MAS with and without microwave radiation irradiation respectively. The DNP signal enhancements (a) are indicated. Spin-echo (purple) and 14 N filtered spin-echo experiments (green) recorded with DNP on the same samples containing 12.5 mM (C) and 2 mM (D) AMUPol. 14 N filtered echo spectra scaling given with respect to corresponding spin-echo, with identical refocusing periods of 1.7 ms corresponding to the 14 N pulse length used in both experiments. The intensity of the spectra in each panel were normalised to reflect an equivalent number of acquisitions, and where necessary, scaled by the factor shown in the figure to aid visualisation.

is more intense at lower radical concentration, with a 10-fold improvement in signal intensity of the 2 mM AMUPol sample over the 12.5 mM AMUPol sample, when expressed as a fraction of the ¹³C spin echo signal under identical conditions. Typically, attenuation of the signal in the sample at high biradical concentrations is attributed to the enhanced T_2 relaxation arising from the presence of the biradicals in the sample. Here however ¹⁴N filtered intensities are compared to a ¹³C spin-echo signal whose refocusing periods match the duration of the ¹⁴N pulses, thereby compensating for any loss in signal due to increased ¹³C T_2 relaxation. This leads us to speculate that the attenuated signal observed at higher biradical concentrations arises through the enhanced relaxation of the multiple spin ¹⁴N-¹³C coherences during the rf driven recoupling This demonstrates the need to understand how DNP reagents influence the efficiency of different pulse sequences. 30,31 To assess the feasibility of conducting these experiments in the absence of DNP, measurement have made of both microcrystalline and lyophilised material where a greater amount of protein can be packed into the rotor. In the latter case a 1D ¹⁴N filter ¹³C spectrum could be acquired in ∼1 week, making 2D acquisition unrealistic.

The ¹⁴N/¹³C 2D correlation spectra of GB3 together with the corresponding 15N/13C correlation spectra processed to mimic the inhomogeneous broadening apparent in the samples measured with DNP are shown in Fig. 2. The most intense feature in the ¹⁴N/¹³C correlation spectra, with ¹⁴N shifts between 280 ppm and 320 ppm, is assigned to the primary amines in the lysine sidechain on the basis of the 14N and 13C chemical shifts. In the region corresponding to the Cα and CO chemical shift (50-55 ppm and 170-175 ppm ¹³C shift, respectively) a broad distribution of ¹⁴N resonances are observed between 310 ppm and 420 ppm. The ¹⁴N shifts cover a range of 110 ppm, a dispersion almost four times greater than observed in the amide region of the corresponding ¹⁵N/¹³C spectra. Notable in their absence, are resonances with ¹³C chemical shifts of 44-47 ppm arising from the glycine residues in GB3. ¹⁵N/¹³C 2D correlation spectra (data not shown) acquired under similar DNP conditions, show little perturbation in the Ca shifts of glycine residues, whilst studies of model peptides have indicated that the 14N QI is similar to that of other amino acids. We suggest that the absence of these resonances is due to the short T₂ of these residues arising from incomplete decoupling of the protons within the CH2 groups of the glycine residues with the proton decoupling fields available.

The increase in ¹⁴N shift dispersion in the ¹⁴N/¹³C spectrum of the protein, compared to the ¹⁵N spectra, is due to the contribution of the field dependent second order isotropic quadrupolar shift (SOIQS) to the ¹⁴N shift, in addition to the usual nitrogen isotropic chemical shift. The ¹⁴N SOIQS may be given by:³²

$$^{14}\text{N}\delta_{\text{Q}}^{\text{iso}} = \frac{3}{40} \left(\frac{\chi_{\text{Q}}}{\nu_0}\right)^2 \times 10^6$$
 (1)

where ν_0 is the Larmor frequency and χ_0 is the quadrupolar product:

$$\chi_{\mathcal{Q}} = C_{\mathcal{Q}} \sqrt{1 + \frac{\eta^2}{3}} \tag{2}$$

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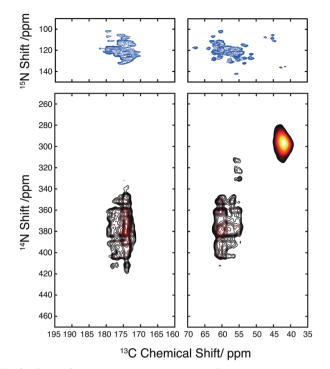


Fig. 2 Carbon/nitrogen correlation spectra of GB3. The top panels show the carbonyl and aliphatic regions of $^{13}\text{C}/^{15}\text{N}$ correlation spectra of U- $^{13}\text{C},^{15}\text{N}$ -GB3 recorded at 14.1 T, 293 K under 12.5 kHz MAS. Lower panels show corresponding regions of the $^{13}\text{C}/^{14}\text{N}$ correlation spectrum of U- ^{13}C GB3 containing 2 mM AMUPol recorded at 18.8 T, 100 K under 13.5 kHz MAS.

where $C_{\rm O}$ is the quadrupolar coupling constant (typically between 0 and 5 MHz for 14 N) and η the asymmetry parameter. While it has not been possible to assign any resonances in the ¹³C/¹⁴N spectrum of GB3 to specific amino acid residues, primarily due to the inhomogeneous broadening observed at 100 K, it is possible to characterise the distribution of quadrupolar couplings present. Subtracting from the centre of the amide region of the ¹⁴N spectrum of GB3, 365 ppm, the centre of the amide region of the ¹⁵N spectrum, 117.5 ppm, one can determine an "average" amide SOIQS in GB3 of 247.5 ppm at 18.8 T. From eqn (1) and (2), one can determine this to be consistent with $C_{\rm Q}$ values of 2.88–3.32 MHz, without any knowledge of the asymmetry parameter, η , of the EFG. Similarly, from the 110 ppm (310–420 ppm) distribution of ¹⁴N shifts measured at this field, using eqn (1) we can conclude that the ¹⁴N C_O magnitudes present in GB3 span 1.12 MHz, from 2.54 MHz to 3.66 MHz assuming that the asymmetry parameter is undefined. Studies of model peptides typically reveal asymmetry parameters ranging from 0.3 to 0.44 for amide bonds in the protein backbone reducing the span of possible to values of $C_{\rm O}$ to 724 kHz.

The observed dispersion of C_Q is dependent on both the local dynamics and electronic environment at each of the amide sites in the protein backbone, the latter reflecting the conformation and H-bond status of each site. The magnitude of C_Q measured agrees well with *ab initio* calculations⁴ and studies of model compounds including *N*-acetyl-valine, ¹ triglycine, ³³ and alanyl–glycyl–glycine¹⁷ suggesting the protein backbone is

immobile with little overall dynamic averaging of the quadrupolar interaction; as expected for this well-structured protein in a glass-frozen matrix at 100 K.

The dispersion in $C_{\rm Q}$ is however larger than previous studies of short model peptides, where changes from an α -helical to a β -strand conformation result in a variation of the ¹⁴N $C_{\rm Q}$ of \sim 200 kHz.⁴ The absence of any overall dynamic scaling of $C_{\rm Q}$ suggests that the larger distribution in $C_{\rm Q}$ reflects either smaller localised mobility leading to dynamic averaging of $C_{\rm Q}$ or the greater structural diversity present in the backbone of GB3 compared to earlier studies of model compounds.

To assess whether the dispersion of ^{14}N shifts reflects the distribution of backbone conformations present in GB3 captured here upon sample freezing at 100 K, we have modelled the $^{14}N/^{13}C$ correlation spectrum based on the SOIQS predicted from the backbone conformations in the crystal structure 34 and the backbone $^{15}N/^{13}C$ assignment. Qualitatively these modelled spectra mirror the experimental data (see ESI,† Fig. S3), with the intensity in regions where the SOIQS would correlate well with α -helical and β -strands structures. This highlights the potential for such studies to provide an oversight of the secondary structures in proteins.

In conclusion, we demonstrate that combining DNP at cryogenic temperatures with ¹⁴N indirect detection it is possible to characterise the quadrupolar interactions at ¹⁴N sites within a microcrystalline preparation of GB3. Despite the enhanced linewidths observed under DNP conditions which prohibit the identification of site specific resonances, we have demonstrated the feasibility of characterising the distribution of the quadrupolar interactions through the analysis of the unresolved envelope of 14N resonances in the indirect dimension. The distribution observed reveals that the 14N sites within the protein backbone exhibit a broad range of 14N EFGs that indicate that, in the case of GB3 at these temperatures, the observed shifts reflect the secondary structures adopted by each amino acid. The exquisite sensitivity demonstrated by the EFG to the backbone conformation and the possibility to characterise these sites, offers a novel route to the structural characterisation of biomolecules.

We envision that the utility of this method will be further enhanced through the use of additional dimensions that would alleviate some of the spectra crowding arising from the DNP conditions used. Furthermore, utilization of alternative spy nuclei such as protons would further boost sensitivity whilst facilitating the NMR analysis of biomolecules and other natural products that have previously proved intractable due to difficulties associated with labelling.

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Conflicts of interest

There are no conflicts to declare.

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