Enhanced measurement of residual chemical shift anisotropy for small molecule structure elucidation
Enhanced measurement of residual chemical shift anisotropy for small molecule structure elucidation†

Yizhou Liu, Ryan D. Cohen, Kirk R. Gustafson, Gary E. Martin and R. Thomas Williamson

A method is introduced to measure residual chemical shift anisotropies conveniently and accurately in the mesophase of poly-γ-(benzyl-L-glutamate). The alignment amplitude is substantially enhanced over common methods which greatly benefits measurements particularly on sp² carbons. The approach offers significant improvements in data accuracy and utility for small molecule structure determination.

Residual chemical shift anisotropy (RCSA) arises from incomplete rotational averaging of chemical shift anisotropy (CSA) as a molecule undergoes partial ordering in an alignment medium. RCSA provides structural information in the form of relative orientations between different CSA tensors in a molecule. Such information is particularly valuable for proton-deficient compounds in which one-bond ¹³C–¹H residual dipolar couplings (RDCs) are scarce. Historically, accurate RCSA measurements were plagued by the change in the isotropic component of the chemical shift tensor (ΔΔδiso) as a molecule is aligned by the addition of an alignment medium. RCSA measurements have occasionally been used as an alignment medium for acquiring RDC data in structure elucidation studies. In contrast, RCSA measurement only utilizes the simple ¹³C{¹H} experiment in which homonuclear dipolar couplings (RDCs) are scarce. While RCSAs from these methods, either alone or in conjunction with RDC data, have been successfully applied to the determination of constitution and configuration of a number of small molecules, a limitation exists in that RCSA values tend to be very small for sp³-hybridized carbons under these alignment conditions, and differentiation of subtle structural variations involving these carbons is expected to be challenging in the absence of RDC data.

Increasing the alignment amplitude is a logical step to address the problem of small RCSAs. In fact, certain liquid crystal (LC) media, such as that formed by poly-γ-(benzyl-L-glutamate) (PBLG), known to generate alignment amplitudes over an order of magnitude larger than constrained polymeric gels. For RDC measurement, a stronger alignment is not necessarily beneficial, as it simultaneously enlarges undesired interactions such as the homonuclear dipolar couplings, which severely broaden proton lines and degrade the performance of HSQC-type experiments typically used for RDC measurements. As a result, PBLG has only occasionally been used as an alignment medium for acquiring RDC data in structure elucidation studies. In contrast, RCSA measurement only utilizes the simple ¹³C{¹H} experiment in which homonuclear ¹³C–¹H dipolar interactions and homogenous interactions from proton spin diffusion are both eliminated by proton decoupling. Therefore, relatively sharp ¹³C lines can still be obtained even with strong alignment, making accurate RCSA measurement feasible. In fact, the RCSA phenomenon has previously been utilized in the chiral nematic (cholesteric) phase of PBLG for the qualitative differentiation of enantiomeric mixtures. Here we describe a simple but highly effective strategy for the accurate quantitative measurement of RCSA in PBLG mesophase for the purpose of small molecule structural elucidation. Data were collected for strychnine, retrorsine, and caula cardine A (Fig. 1a–c), for which RCSA data were also available for comparison from stretched poly-methyl-methacrylate (PMMA) or poly-2-hydroxyethyl-methacrylate (PHHEMA) gels.

The challenge for accurate RCSA measurement is to correct for ΔΔδiso associated with the PBLG phase transition at different concentrations. Our correction method is based on the premise

Fig. 1 Structures of strychnine (1a), retrorsine (1b), and caula cardine A (1c).
that the ΔΔδiso dependence on PBLG concentration is identical in both phases, such that the ΔΔδiso in the mesophase can be corrected for by extrapolating its value from its trend in the isotropic phase. The rationale and limitation of this approach are discussed in ESI†.

We successively added small amounts of PBLG to a strychnine solution and acquired a 13C[1H] spectrum after each addition (Fig. 2). Tetramethylsilane (TMS) at 4% (v/v) was added for 13C chemical shift referencing. The effect of the bulk susceptibility change at different PBLG concentrations is eliminated by TMS referencing. Although vibration-rotation coupling can generate anisotropic effects in a tetrahedral molecule,24–27 TMS aligns much more weakly than most compounds of interest and can therefore be used as a convenient reference in most cases.8

Alternatively, RCSA measurements can also be referenced by an arbitrary carbon in the compound itself, following a previously described procedure.7 As shown in Fig. 2a, the C21 sp2 resonance of strychnine gradually shifted upfield as PBLG was added up to 8.8% (w/v) due to ΔΔδiso and no CDCl3 residual quadrupolar coupling (RQC) was observed consistent with an isotropic solution. As the PBLG concentration reached 11.4%, a sudden large downfield resonance jump occurred due to RCSA (spectrum 5) concomitant with a CDCl3 RQC of 215.5 Hz, indicating the sample entered the mesophase. Additional PBLG increased both RCSA and RQC as the alignment strengthened. As shown in Fig. S3 (ESI†), ΔΔδiso follows a nearly linear relationship with PBLG concentration within the isotropic phase, although not strictly. The linearity is actually expected if the analyte and PBLG do not bind (i.e., the binding Kd is very large). This relationship is then used to remove the ΔΔδiso contribution in the mesophase. Importantly, because ΔΔδiso from PBLG addition is small in comparison to RCSA even for sp3 carbons (Fig. 2b), ignoring the surface interference effect, which is a higher order effect as described in the ESI† is not expected to cause large errors, at least when the PBLG concentration is not much higher than the mesophase critical concentration (Ccrit).

Based on the linear relationship, we propose a simplified three-measurement approach, using the terminology: I0–I1–A1. I0 is an isotropic measurement without PBLG, I1 is a second isotropic measurement at a PBLG concentration below but close to Ccrit, and A1 is an anisotropic measurement at a PBLG concentration above Ccrit. RCSA can be extracted from these measurements by the simple equation:

\[
\text{RCSA}_i = \left( \delta_{i,1} - \delta_{TMS,1} \right) - \left( \delta_{10} - \delta_{TMS,10} \right) - \frac{\text{PBLG}_{i1}}{\text{PBLG}_{10}} \left[ \left( \delta_{i,1} - \delta_{TMS,1} \right) - \left( \delta_{10} - \delta_{TMS,10} \right) \right]
\]

The first and second terms in parentheses on the right side of eqn (1) represent the TMS-referenced chemical shifts of carbon \( i \) in A1 and I0, respectively. Their difference defines the RCSA of carbon \( i \), as well as ΔΔδiso represented by the third term. Note that all chemical shift terms on the right are determined experimentally and the PBLG concentrations are also known. Therefore, the RCSA value of each carbon can be extracted.

As shown in Fig. 2c, the RCSA in PBLG is much larger than the corresponding RCSA in a stretched PMMA gel.8 The ~2-fold increase in line-width is amply compensated for by the over 34-fold enhancement of the measured RCSA value. Consequently, these large RCSAs greatly improve the quality of data and structural analysis, as demonstrated later. One noteworthy point is that the proton decoupling used for an isotropic sample may be insufficient to fully decouple the much larger couplings present in mesophase. For example, the alignment with 34.5% PBLG can generate 13C–1H RDCs of over 1 kHz so higher decoupling power and a decoupling sequence with wide broadband coverage, such as GARP or WURST, should be used for PBLG mesophase (see ESI† for experimental details).

The data quality is evaluated in Table 1. The generalized degree of order (GDO) was employed to measure the overall magnitude of molecular ordering.28 For a mostly rigid molecule, the GDO measures the alignment amplitude. Our previously reported RCSA data collected in PMMA gels were used for comparison. Clearly, even at relatively low PBLG concentrations, the GDOs are ~5–6 times larger than in PMMA gels; at the high end of this study, a GDO of 2% was obtained for strychnine, i.e., the anisotropic NMR interactions were measured in a liquid sample at 2% of the solid-state values. To the best of our knowledge, anisotropic interactions of this magnitude have not been previously demonstrated in high-resolution NMR for complex molecules.

RCSAs from these strong alignments yield significantly improved structural analysis, as reflected by the Q-factors. As shown in Table 1, Q-factors with all carbon atoms considered (Qall) are substantially better in PBLG mesophases than in PMMA gels. For example,
a $Q_{\text{all}}$ of 0.029 in 15.5% PBLG is a significant improvement over a $Q_{\text{all}}$ of 0.049 in the stretched PMMA gel for strychnine. Similarly, RCSAs from the rigid part of retorsine produced a vastly improved $Q_{\text{all}}$ of 0.033 in 15.1% PBLG, over a $Q_{\text{all}}$ of 0.104 in stretched PMMA.

This strong alignment is particularly beneficial for compounds constituted primarily of sp$^3$ carbons. We demonstrate this point by omitting all RCSA data from sp$^2$ carbons while using only those data from sp$^3$ carbons to calculate the Q-factors ($Q_{\text{sp3}}$) and alignment tensor parameters. As shown in Table 1, the improvement in $Q_{\text{sp3}}$ followed the trend of $Q_{\text{all}}$ for data in PBLG vs. PMMA gels, with $Q_{\text{sp3}}$ from the former being lower than 0.1 in most cases. To further evaluate the reliability of using only sp$^3$ carbon RCSAs for structural analysis, we compared the performance of the alignment tensor parameters determined with all of the data and those determined with only sp$^3$ data through calculations of their intertensor angles ($\theta$). As shown in Table 1, the intertensor angles are substantially smaller in PBLG solutions than in PMMA gels, with $\theta$ well below 10° in most cases, confirming that the reliability of using only sp$^3$ carbon data significantly improved. One exception was observed for retorsine in 12.6% PBLG, in which $\theta$ was slightly higher than in the PMMA gel, presumably due to inaccuracies of the alignment tensor determination with only a small number of sp$^2$ carbons. From Table 1, it is also apparent that the $Q_{\text{sp3}}$ factor under a larger alignment for strychnine at the high end of PBLG concentration did not improve, but instead slightly degraded, whereas the $Q_{\text{all}}$ factor barely changes. This divergence between $Q_{\text{all}}$ and $Q_{\text{sp3}}$ is possibly due to smaller sp$^3$ carbon RCSA values being more susceptible to imperfections in our $\Delta\Delta\phi$ elimination method at high PBLG concentrations where surface interference is expected to be more pronounced. Based on this observation, for a sp$^3$ carbon-rich compound, a PBLG concentration of about 15% is advisable based on available data.

In a recent study, we successfully differentiated 13 possible diastereomers of strychnine using only RCSA data collected in a PMMA gel. Here we conducted the same analysis with RCSAs collected in 15.5% PBLG as a representative example (Fig. 3a), and compared the performance of PBLG vs. PMMA gel data for this differentiation (Fig. 3b). The stereochemical definition follows that in our previous work. The correct diastereomer RSSRRS with a Q-factor of 0.029, shown on the left, is clearly distinguished from all other diastereomers that exhibited much larger Q-factors (Fig. 3a). The confidence level of differentiation can be quantitatively assessed by the ratio: $Q/Q_{\text{best}}$ where $Q$ is the Q-factor of an isomer being evaluated, and $Q_{\text{best}}$ represents the lowest Q-factor in the pool, which, in this example, is $Q_{\text{RSSRRS}}$. The larger this Q-factor ratio, the higher the level of confidence for the differentiation. Fig. 3b displays the Q-factor ratios of all 13 diastereomers over $Q_{\text{RSSRRS}}$ for both PBLG data (blue) and PMMA gel data (red). Clearly, the confidence level is about twice as good with PBLG data for those less discriminated isomers, such as RSRRRS and SSSRRS, and overall much better, in comparison to the results obtained with PMMA gels. We should mention that the data quality from the stretched PMMA gel, with a $Q_{\text{RSSRRS}}$ of 0.049, was actually quite good. However, that result is superseded by the enhanced quality afforded by the measurement in PBLG.

Next, as a more challenging example, we used RCSA data to differentiate caulaminde A (Fig. 1c) from its C26-inverted diastereomer. In a recent collaboration, we reported the structure of caulaminde A using NMR data including RDC and RCSA collected in a PHEMA gel. Although a combination of RDC and RCSA clearly favoured the proposed structure over other constitutional and stereoisomers, the weak alignment of caulaminde A in PHEMA clearly left room for further improvement. Indeed, alignment with 13.1% PBLG generated a large RCSA range of −40 to...
100 Hz, a vast enhancement over the −3.4 to 3.5 Hz range in PHEMA. The second best structural candidate in the previous study was the C26-inverted diastereomer. This chiral inversion swaps the position of the chlorine atom but minimally perturbs the overall conformation (Fig. 4). Despite the high level of structural similarity, RCSA data from PBLG strongly supports the proposed structure with a Q-factor of 0.077 over the C26 inverted diastereomer with a Q-factor of 0.328 (Fig. 4). Interestingly, the largest RCSA outlier from the correlation plot of experiment vs. theory, (circled in red in Fig. 4b), corresponds to the inverted sp³ carbon 26, which clearly reveals the inconsistency of this chiral inversion with experimental data. The RCSA data were collected with only 1.6 mg of material, and the actual measurements took 54, 69, and 131 minutes for I₀, I₁, and A₁, respectively, on a 600 MHz spectrometer with a 5 mm helium cryoprobe. This work further demonstrates the utility of the method for mass-limited samples.

Finally, anisotropic NMR work on scarce materials, such as many natural products, usually mandates sample recovery from the alignment medium after NMR measurements are completed. Here we also developed an efficient sample recovery procedure that achieves over 90% recovery for an array of test compounds soluble in ethyl acetate, as shown in Table S1 (ESI†). The recovery procedure is detailed in the ESI.†

In summary, we demonstrated with several examples that a simple first-order correction for PBLG concentration was sufficient to eliminate ∆Δνiso from RCSA measurement in the PBLG mesophase. The large RCSAs resulting from the strong alignment greatly improve structural analysis in general, especially that of sp³ rich compounds. The commercial availability of PBLG, the facile sample preparation and high analyte recovery comparable to polymer gels, and the easily implemented NMR experiment should facilitate utilization of RCSA data for small molecule structural analysis in a greater number of laboratories than has been heretofore possible with existing methods.

Conflicts of interest

There are no conflicts to declare.

Notes and references

1 G. Cornilescu, J. L. Marquardt, O. Marcel and A. Bax, J. Am. Chem. Soc., 1998, 120, 6836.
2 G. Cornilescu and A. Bax, J. Am. Chem. Soc., 2000, 122, 10143.
3 R. S. Lipsitz and N. Tjandra, J. Magn. Reson., 2003, 164, 171.
4 R. S. Lipsitz and N. Tjandra, J. Am. Chem. Soc., 2001, 123, 11065.
5 Y. Liu and J. H. Prestegard, J. Biomol. NMR, 2010, 47, 249.
6 G. Kummerlöwe, S. L. Grage, C. M. Thiele, I. Kuprov, A. S. Ulrich and B. Luy, J. Magn. Reson., 2011, 209, 19.
7 F. Hallwass, M. Schmidt, H. Sun, A. Mazur, G. Kummerlöwe, B. Luy, A. Navarro-Vázquez, C. Griesinger and U. M. Reinscheid, Angew. Chem., Int. Ed. Engl., 2011, 50, 9487.
8 N. Nath, M. Schmidt, R. R. Gil, R. T. Williamson, G. E. Martin, A. Navarro-Vázquez, C. Griesinger and Y. Liu, J. Am. Chem. Soc., 2016, 138, 9548.
9 E. Mevers, J. Sauri, Y. Liu, A. Moser, T. R. Ramadhar, M. Varlan, R. T. Williamson, G. E. Martin and J. Clardy, J. Am. Chem. Soc., 2016, 138, 12324.
10 Y. Liu, J. Sauri, E. Mevers, M. W. Peczuh, H. Hiemstra, J. Clardy, G. E. Martin and R. T. Williamson, Science, 2017, 356, 6333.
11 D. J. Milanowski, N. Oku, L. K. Cartner, H. R. Bokesch, R. T. Williamson, J. Sauri, Y. Liu, K. A. Blinov, Y. Ding, X. Li, D. Ferreira, I. A. Walker, S. Khan, M. T. Davies-Coleman, J. A. Kelley, J. B. McMahon, G. E. Martin and K. R. Gustafson, Chem. Sci., 2017, 9, 269.
12 P. Doty, A. M. Holzer, J. H. Bradbury and E. R. Blout, J. Am. Chem. Soc., 1954, 76, 4493.
13 R. R. Gil, C. Griesinger, A. Navarro-Vázquez and H. Sun, Structure Elucidation in Organic Chemistry: The Search for the Right Tools, Wiley-VCH, Weinheim, 1st edn, 2015, ch. 8, pp. 279–323.
14 C. M. Thiele and S. Berger, Org. Lett., 2003, 5, 705.
15 L. Verdier, P. Sakhaii, M. Zweckstetter and C. Griesinger, J. Magn. Reson., 2003, 163, 553.
16 C. Aroulanda, V. Boucard, F. Guibé, J. Courtieu and D. Merlet, Chem. – Eur. J., 2003, 9, 4536.
17 A. Marx and C. Thiele, Chem. – Eur. J., 2009, 15, 254.
18 L. Vrazil, M. Kratochvílovič-Simánková, T. Landovský, K. Polívková, J. Budka, H. Dvořáková and P. Lhotač, Org. Biomol. Chem., 2015, 13, 9610.
19 M. Reller, S. Wesp, M. R. M. Koos, M. Reggelin and B. Luy, Chem. – Eur. J., 2017, 23, 13351.
20 P. Lesot, D. Merlet, A. Meddour, J. Courtieu and A. Loevenstein, J. Chem. Soc. Faraday Trans., 1995, 91, 1371.
21 A. Meddour, P. Berdague, A. Hedli, J. Courtieu and P. Lesot, J. Am. Chem. Soc., 1997, 119, 4502.
22 M. Sfarati, P. Lesot, D. Merlet and J. Courtieu, Chem. Commun., 2000, 2069.
23 M. Rivard, F. Guillet, J. C. Fiaud, C. Aroulanda and P. Lesot, Tetrahedron: Asymmetry, 2003, 14, 1141.
24 N. J. D. Lucas, Mol. Phys., 1971, 22, 147.
25 J. W. Emsley and J. C. Lindon, Mol. Phys., 1974, 28, 1373.
26 E. E. Burnell and C. A. De Lange, J. Magn. Reson., 1980, 39, 461.
27 J. G. Snijders, C. A. De Lange and E. E. Burnell, J. Chem. Phys., 1982, 77, 5386.
28 J. H. Prestegard, H. M. Al-Hashimi and J. R. Tolman, Q. Rev. Biophys., 2000, 33, 371.
29 J. Sass, F. Cordier, A. Hoffmann, A. Cousin, J. G. Omichinski, H. Lowen, S. Grzesiek and M. Rogowski, J. Am. Chem. Soc., 1999, 121, 2047.
30 N. Marcó, R. R. Gil and T. Parella, Magn. Reson. Chem., 2017, 55, 540.