

Direct magnetic resonance detection of myelin and prospects for quantitative imaging of myelin density

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Edited by Alan P. Koretsky, National Institutes of Health, Bethesda, MD, and accepted by the Editorial Board April 27, 2012 (received for review September 13, 2011)

Magnetic resonance imaging has previously demonstrated its potential for indirectly mapping myelin density, either by relaxometric detection of myelin water or magnetization transfer. Here, we investigated whether myelin can be detected and possibly quantified directly. We identified the spectrum of myelin in the spinal cord *in situ* as well as in myelin lipids extracted via a sucrose gradient method, and investigated its spectral properties. High-resolution solution NMR spectroscopy showed the extract composition to be in agreement with myelin's known chemical make-up. The 400-MHz ¹H spectrum of the myelin extract, at 20 °C (room temperature) and 37 °C, consists of a narrow water resonance superimposed on a broad envelope shifted ~3.5 ppm upfield, suggestive of long-chain methylene protons. Superimposed on this signal are narrow components resulting from functional groups matching the chemical shifts of the constituents making up myelin lipids. The spectrum could be modeled as a sum of super-Lorentzians with a T₂* distribution covering a wide range of values (0.008–26 ms). Overall, there was a high degree of similarity between the spectral properties of extracted myelin lipids and those found in neural tissue. The normalized difference spectrum had the hallmarks of membrane proteins, not present in the myelin extract. Using 3D radially ramp-sampled proton MRI, with a combination of adiabatic inversion and echo subtraction, the feasibility of direct myelin imaging *in situ* is demonstrated. Last, the integrated signal from myelin suspensions is shown, both spectroscopically and by imaging, to scale with concentration, suggesting the potential for quantitative determination of myelin density.

myelin *in situ* | myelin NMR spectrum | super-Lorentzian fitting | ultrashort echo time

Myelin is a critical feature of nervous system white matter (WM) and accounts for 14% of the wet mass of WM (1). It is a lipid–protein bilayer that extends from the outer membrane of glial cells (i.e., oligodendrocytes in the CNS) and discretely winds around individual axonal fibers, leading to an increase in conduction velocity (1). By speeding conduction and reducing axonal energy requirements, myelin makes large and complex organisms possible. Myelin also contributes to the mechanical and functional structure of the axon. In addition, some oligodendrocytic cells and precursors can support action potentials themselves (2). Deficiencies of myelin lay at the core of numerous neurodegenerative disorders, such as multiple sclerosis and schizophrenia (1). These deficiencies may result from developmental or acquired abnormalities in oligodendrocyte function, which also leads to axonal degeneration. Assessment of myelin may reveal CNS abnormalities far beyond those associated with classic demyelinating diseases. MRI of myelin has the potential to characterize not only loss of this important component of the CNS but also to reveal axonal and supporting glial integrity and function.

A diverse assortment of experimental techniques has been applied toward the goal of observing and quantifying myelin. The common methods rely on optical microscopy of histologically

stained tissue samples (3). X-ray diffraction (4) and nonlinear optical techniques (5, 6) also provide insight into myelin ultrastructure. Unfortunately, all these techniques are destructive and thus applicable only to animal studies.

More recently, myelin-specific chemical contrast markers that selectively bind to myelin have emerged. Such agents are currently under development for both MRI (7) and positron-emission tomography (8). Although these techniques are potentially promising, concerns over toxicity may pose significant hurdles to their clinical implementation.

So far, MRI has had the greatest impact toward nondestructive myelin assessment in both laboratory animals and humans. Further, MRI has the added benefit that signal contrast originates from endogenous protons and hence is not reliant upon injectable chemical probes nor limited by contrast-related temporal delays.

To date, two indirect MR techniques applicable to studies *in vivo* have demonstrated histologically correlated sensitivity to myelin: magnetization transfer (MT) and T₂ relaxometry. In MT, cross-relaxation between myelin protons and tissue water is exploited (9). The signal attenuation resulting from off-resonance saturation (MT ratio) has been found to scale with myelin concentration (10). T₂ relaxometry yields T₂ spectra, typically by inversion of the Carr–Purcell echo decay using an inverse Laplace transformation (11). Spectral components with T₂ values ranging from 10 to 50 ms have been assigned to motionally restricted myelin water (12, 13) and have demonstrated strong correlation with myelin-specific histologic staining (12, 14).

Although MT and T₂ relaxometry have shown promise, they both rely on indirect detection of myelin through the interaction of water with myelin. This complex interaction is affected by nonmyelin loss-related changes, which can lead to ambiguities in data interpretation. For example, MT is sensitive not only to myelin content but also to axon density (15). Therefore, even though both techniques may distinguish normal from abnormal WM, they rely on the invariance of the myelin–water interaction.

Direct detection of myelin with MR would remove some complications in the analysis from the intermediate effects of the

Author contributions: M.J.W., H.H.O., and F.W.W. designed research; M.J.W., H.H.O., and S.L.W. performed research; M.J.W., H.H.O., S.L.W., C.L., P.-H.T., and F.W.W. analyzed data; and M.J.W., H.H.O., C.L., D.B.H., and F.W.W. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission. A.P.K. is a guest editor invited by the Editorial Board.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1115107109/-DCSupplemental.

interaction of water and myelin, and therefore may provide contrast specific to myelin concentration. However, direct detection is complicated by motional restriction of the lipid chains in the myelin bilayer, resulting in broad lines and, consequently, short lifetime of the MR signal.

Perhaps the first characterization of the NMR spectral properties of myelin was by Lecar et al. (16), who studied anhydrous preparations by wide-line proton spectroscopy, concluding that under these conditions the system is in a liquid-crystalline state. To the best of our knowledge, the first report of myelin proton transverse relaxation was by Ramani et al. (17). The authors performed a multiexponential fit of spin-echo decays on fixed human WM samples from normal and multiple sclerosis patients and reported a T_2 value of $\sim 50 \mu\text{s}$ for myelin protons. They were also able to detect lipid signals by magic-angle spinning proton NMR in slices of normal WM tissue but not in multiple sclerosis lesions. Recently, Horch et al. (18) investigated the T_2^* and T_2 relaxation characteristics of myelin and reported values of $\sim 70 \mu\text{s}$ as well as a broader distribution ranging from 50 to 1,000 μs .

The transverse relaxation properties of myelin suggest the need for ultrashort echo time (UTE) MRI methods, which entail collection of the free-induction decay immediately after excitation. Typical implementations include either 3D radial sampling with nonselective rf pulses (19) or 2D radial sampling with slice-selective half rf pulses and ramp sampling (20) as previously applied for the characterization of cortical bone matrix and bone water (21, 22).

UTE MRI has been used to image the short T_2^* (i.e., $< 1 \text{ ms}$) signal from human brain in vivo (23). Unlike applications to study bone, these implementations include long T_2^* suppression methods to attenuate the tissue water signal. Tissue water, because of its rotational mobility and high concentration, has an intense long T_2^* signal that, without suppression, overwhelms signal from short T_2^* components (Fig. S14). Although the images indicated the short T_2^* signal to be predominantly located in WM, no evidence was provided to link this signal to myelin.

In this work we examine the origin and nature of the short T_2^* signal of CNS tissue in freshly excised rat spinal cord (SC) in comparison with purified myelin lipid extract with multinuclear NMR. We further explore the potential for direct detection and quantification of myelin by UTE MRI and discuss the possibilities and technical hurdles associated with translating MRI-based quantification of myelin to the clinic.

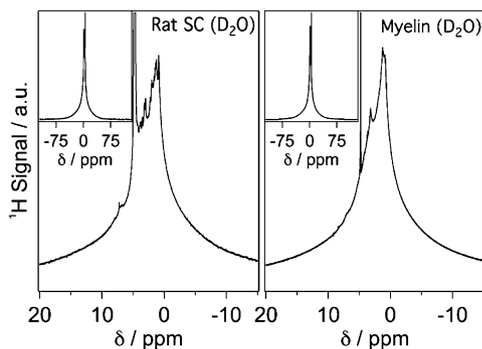


Fig. 1. The ^1H NMR spectra from rat thoracic SC after D_2O exchange of tissue water (Left) and myelin lipid extract suspended in D_2O (Right), showing remarkable similarity. Insets: Wide tails present in both spectra. Note that in addition to narrow resonances, likely stemming from proteins, the residual monodeuterium oxide (HDO) resonance is stronger in the tissue spectrum.

Results

High-Resolution ^1H NMR Spectra of Intact SC Tissue and Bovine Myelin Extract. Fig. 1 shows a comparison of the ^1H NMR spectra collected for a section of excised rat thoracic SC and a deuterium oxide (D_2O) suspension of myelin lipid extract. The SC was first immersed in 99.9% D_2O for 24 h to exchange tissue water with D_2O , and hence attenuate the bulk proton signal (Fig. S1B). The protein-free myelin lipid sample was chemically extracted from an intact isolated myelin sample, obtained via a sucrose gradient method (SI Materials and Methods) and quantitatively validated with proton-decoupled ^{13}C and ^{31}P NMR spectroscopy (Table 1 and Fig. S2 A and B).

Both ^1H spectra in Fig. 1 show a broad resonance with a superimposed narrow water resonance originating from residual HDO. Although the two broad resonances bear a high degree of similarity, a difference spectrum (i.e., tissue – extract) (Fig. S3) highlights the distinguishing features. First, the superimposed fine structure apparent in the SC spectrum is consistent with mobile proteins as they might occur in the cytoplasm, for example. Second, the difference spectrum contains a broad resonance as it might be expected from membrane proteins. The difference spectrum was generated such that the integrated signal area is 27.9% of the total spectrum, as expected according to the known protein fraction in myelin (details in SI Results). Small errors in the difference spectrum could arise because we are ignoring cytoplasmic proteins in this fraction.

UTE MRI of Intact SC. Fig. 2 shows a set of images of freshly excised rat SC, obtained with a 3D radial, ramp-sampled, dual-echo inversion recovery UTE (de-IR-UTE) pulse sequence (Fig. S4). Long T_2^* tissue water signal was attenuated via adiabatic inversion and complex echo subtraction. Adiabatic inversion was used to significantly reduce the signal intensity from tissue water, which would then be further attenuated with echo subtraction. We empirically selected TI to achieve the greatest WM intensity while minimizing gray matter (GM) intensity signal in the complex echo difference image in accordance with the expectation of low signal in GM considering its very low myelin content. Images were collected at both short (20 μs ; Fig. 2A) and long (1,200 μs ; Fig. 2B) TE. The magnitude of the complex difference image and signal profile (Fig. 2 C and D)

Table 1. Lipids of myelin with abbreviations used in the text

Myelin lipid	ID	Molar %		
		Norton*	NMR [†]	% labile $^1\text{H} \pm \sigma^\ddagger$
Cholesterol	CHOL	44.8	43.1	0.94 ± 0.04
Galactocerebroside	GC	17.5	19.6	2.20 ± 0.39
Galactosulfatide	GS	2.5	NA [§]	$0.28 \pm 0.05^\parallel$
Phosphatidylethanolamine	PE	3.4	3.9	0.24 ± 0.05
PE plasmalogen	PEpl	11.3	11.8	0.70 ± 0.13
Phosphatidylcholine	PC	8.0	7.8	0.00 ± 0.00
PC plasmalogen	PCpl	0.3	2.0	0.00 ± 0.00
Sphingomyelin	Sph	5.2	5.9	0.37 ± 0.08
Phosphatidylinositol	PI	0.7	2.0	0.19 ± 0.03
Phosphatidylserine	PS	0.2	3.9	0.11 ± 0.01
Total				5.05 ± 0.79

Comparison of average bovine myelin lipid molar ratios reported by Norton et al. (25) and quantitative multi-NMR methods. Also shown are average percentages of labile protons pertaining to each lipid component. PC, phosphatidylcholine; PE, phosphatidylethanolamine.

*From Norton et al. (23).

[†]Present study, ^{13}C and ^{31}P NMR.

[‡]Variability (σ , SD) due to lipid chain length [$\text{CH}_2(\text{CH}_2)_n$; $n = 10\text{--}25$].

[§]Not measured owing to a lack of an unambiguous resonance.

[¶]Assuming a GS molar percentage of 2.5%.

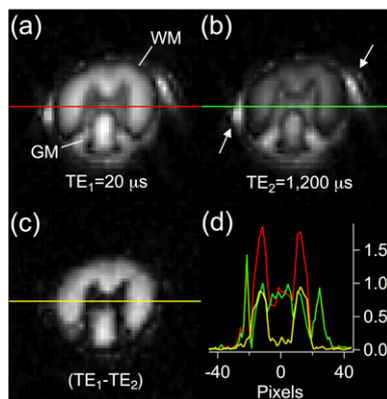


Fig. 2. The 3D de-IR-UTE images from rat thoracic SC averaged over five central slices. Magnitude images obtained for (A) TE = 20 μ s, (B) TE = 1,200 μ s, and (C) complex difference (A – B) (maximum-intensity range decreased by a factor of two to highlight myelin signal). (D) Intensity profiles across the three images (delineated as red, green, and yellow lines in A, B, and C, respectively) to show relative WM, GM, and background intensity. The most intense signal, present in the short- and long-echo profiles, originates from residual surface water. WM and GM are indicated in A, and arrows highlight residual surface water in B. The dark boundary observed at the GM/WM and WM/surface water interfaces in both echo images stems from partial voluming of adjacent regions with different T_1 resulting in destructive interference.

highlight the short T_2^* signal, which predominantly results from myelin protons.

MR Signal Dependence on Myelin Concentration. To separate the myelin and water peaks in the ^1H spectra of the myelin- D_2O suspensions, the spectra were modeled as a weighted sum of four super-Lorentzians (SL) for myelin resonances representing protons from general alkyl chain methylenes (noncholesterol, mostly from fatty acid residues), cholesterol alkyl chain methylenes, terminal methyls, and choline, and a Lorentzian for the HDO peak (details in *Materials and Methods*). Fig. 3A shows the results from fitting of the proton NMR spectrum of purified myelin suspended in D_2O . The fitting results were virtually identical for all myelin concentrations. Even though the signal envelope is very broad, relatively narrow resonances are also observed, likely due to proton pairs aligned with an average orientation at the magic angle relative to the static field (24).

Relative signal fractions, accounting for losses during excitation and acquisition, along with associated T_2^* distributions of the four SL components, were combined into a myelin T_2^* histogram (Fig. 3B). At 20 $^\circ\text{C}$, 26.4% of the total signal has an effective lifetime of <25 μ s, 51.8% of <100 μ s, and 91.6% of <1,000 μ s. At 37 $^\circ\text{C}$ these values are 16.9%, 44.8%, and 86.3%, respectively.

Fig. 4A shows a series of fitted myelin signals as a function of decreasing myelin concentration. The NMR signal areas for the total and separate spectral components (i.e., HDO and myelin) are plotted in Fig. 4B, indicating linear scaling with myelin concentration ($R^2 = 0.99$). We attribute the positive correlation of the water peak area with myelin concentration as resulting from labile protons from myelin constituents exchanging with D_2O to form HDO. The calculated average percentage of labile protons, for each of the 10 myelin lipid components, is listed in Table 1. The average signal contribution from the 0.1% impurity of the D_2O solvent, calculated as the y-intercept from the line of best fit (Fig. 4B), was subtracted from all of the HDO points, yielding an estimate of the labile myelin proton signal contribution. The predicted range of signal contributions from labile protons ($5.05\% \pm 0.79\%$) agreed well with the experimental HDO peak areas ($5.13\% \pm 2.00\%$). Given the excess D_2O used

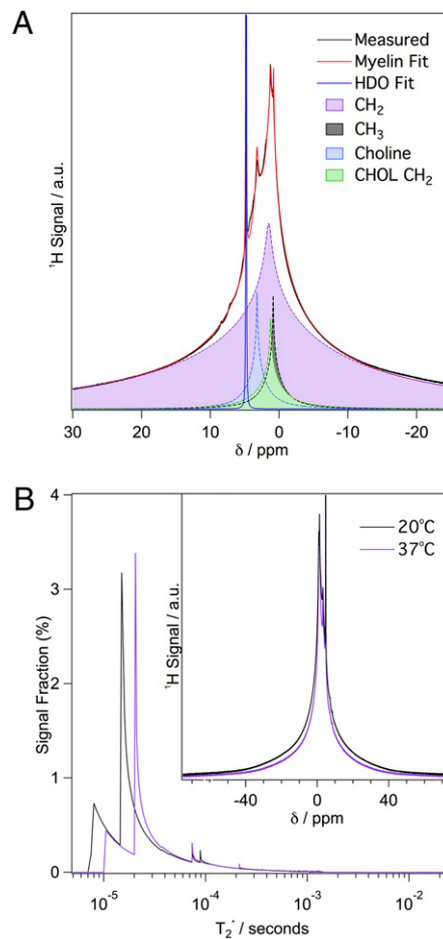


Fig. 3. ^1H NMR spectra and analysis of purified bovine myelin extract suspended in D_2O . (A) NMR spectrum (black) and SL fitting showing the resulting myelin (red) and HDO (blue) fits, as well as the four individual SL components of myelin (shaded). The four fitted SL components consisted of (i) SL containing 74.3% of the intensity, centered at 1.5 ppm, corresponding to the general alkyl chain methylene protons (CH_2), (ii) SL containing 12.4% of the intensity, centered at 0.9 ppm, corresponding to the terminal methyl protons (CH_3), (iii) SL containing 11.1% of the intensity, centered at 3.2 ppm, corresponding to the choline methyl protons (Choline), and (iv) SL containing 2.1% of the intensity, centered at 1.3 ppm, corresponding to the cholesterol alkyl chain methylene protons (CHOL CH_2). (B) T_2^* histogram of myelin components at 20 and 37 $^\circ\text{C}$ derived from the SL fitting. There are small T_2^* components that extend up to 26 ms, but these cannot be observed at the displayed scale. *Inset:* Myelin extract spectra collected at the two temperatures.

in the suspensions, it is reasonable to assume that all labile myelin protons had exchanged with deuterium.

Fig. 5 *Inset* shows the complex difference 2D projection de-IR-UTE image for a series of myelin suspensions of increasing concentration. Region of interest (ROI) average intensities from each of the myelin samples in the image are plotted in Fig. 5 and, analogous to the spectral data, are linearly correlated with myelin concentration ($R^2 = 0.98$).

On the basis of the relaxation characteristics of myelin (notably the lifetime of the various T_2^* components), Bloch equation simulations (details in *SI Materials and Methods*) accounting for losses sustained during the rf pulse sampling suggest that, under the spectral recording conditions, 81.7% of the total spectral signal is recovered, where the shorter T_2^* components account for the majority of the signal lost. The depletion of the imaging signal is more severe because it entails coherence losses during both

cannot be described with a basis set of exponential functions, the authors suggested that the resulting errors would be small for the case of myelin. Horch et al. analyzed free-induction decay signals of myelinated mammalian and amphibian nerves, as well as synthetic myelin at 4.7 T, yielding histograms of relaxation times. The authors detected significant components with T_2^* values of 20 and 70 μs in frog sciatic nerve, which they conjectured to arise from protein and methylene protons of myelin, respectively. In contrast, because our myelin extract was free of protein, the present data alternatively suggest the short T_2^* components (<25 μs) to arise from myelin lipids. This is not to imply that membrane proteins cannot contribute a short T_2^* component, as suggested by the broad resonance in the difference spectrum (Fig. S4) and in other reports (24, 33).

In bovine myelin suspended in D_2O , we found the integrated spectral areas to scale linearly with myelin concentration (Fig. 4), as did ROI intensities of the 2D projection de-IR-UTE images (Fig. 5), thus suggesting that quantitative myelin imaging may be feasible. Direct 3D de-IR-UTE imaging of a rat SC in situ at 400 MHz highlights the potential of such an approach, as demonstrated with images showing signal from the WM regions only. Absolute quantification would require a reference sample, ideally with relaxation and density properties matching myelin. The reference sample should also be chemically stable.

Previously, Waldman et al. (23) obtained images of the human brain using a slice-selective UTE along with soft-tissue suppression, essentially showing intense signal from WM regions of the brain, which they attributed to short T_2^* components. Under these conditions [i.e., selective half-*sinc* pulses of 400- to 600- μs duration (34)], all but the longest T_2^* components of the myelin protons would elude detection.

Our results indicate approximately 20% of the protons in myelin lipids to have an effective T_2^* less than 25 μs . Even under the more favorable conditions of our imaging experiments, the majority of these short T_2^* components remains undetectable. The very short lifetime of the signal has potentially adverse effects on the point-spread function (PSF) manifesting as blurring. A simple estimation based on the FWHM of the spectra (Fig. 3) predicts an intrinsic resolution (defined as the minimum achievable pixel size) on the order of 100 μm , which is approximately one pixel with our current imaging parameters (more details in *SI Materials and Methods*). It should be noted that although the blurring from the shortest T_2^* component would be greater, its impact on the PSF is negligible because this signal fraction remains virtually undetected.

A limitation that needs to be noted for this method is that it detects myelin solely on the basis of its T_2^* properties. Thus, errors in long T_2^* suppression may lead to signal misclassified as short T_2^* and hence falsely identified as myelin. Such errors could be accounted for and perhaps mitigated by tailoring a reference sample so as to contain water of comparable concentration and relaxation times to those of neural tissue. Last, there are other possible nonmyelin short T_2^* sources that could contribute to the UTE image intensity, including glial cell membranes, calcifications, tissue scars, vasculature, and hemorrhage (23), that would be indistinguishable from myelin.

Another potential problem could arise from saturation of the myelin signal via cross-relaxation (35). Even though adiabatic inversion of tissue water has minimal direct effect on the myelin lipid proton signal, transfer of magnetization from the water to the myelin proton pool could occur via chemical exchange or dipolar coupling. This mechanism would result in a reduction in the detected myelin signal, an effect that requires further scrutiny.

The potential for translation of the method to the clinic will be challenging. Nevertheless, it is encouraging to note that with dedicated hardware rf pulses of 20 μs or less have already been shown to be feasible on clinical equipment at 3 T field strength, as in recent work by Wu et al., who imaged the collagen matrix

of cortical bone (36). Further, at 37 $^\circ\text{C}$ the measured T_2^* values increase by *ca.* 30% (Fig. 3B), raising the minimum T_2^* value from 8 to 10.5 μs . At body temperature the scan parameters used in our current experiments at 9.4 T (except for a 20- μs hard pulse of 7.6 $^\circ$ flip angle to match the peak B_1 amplitudes of clinical head coils) predict 4.9% of the total myelin proton signal [i.e., 0.7% of the total proton signal given that myelin accounts for 14% of WM (1)] to be recoverable on a 3 T clinical MRI system (for calculation details see *SI Materials and Methods*).

Given that tissue proton concentration is ~ 100 M, the concentration of detectable myelin protons is approximately 700 mM. In comparison with proton MR spectroscopic imaging (MRSI) of brain metabolites, where the metabolite concentrations are on the order of 10 mM, detectable myelin proton concentration, and hence intrinsic SNR, should be a factor of approximately 70 greater than that of typical metabolites. However, this gain in SNR compared with MRSI is mitigated by the reduced sampling time imposed by the much shorter T_2^* of the protons in myelin compared with those in brain metabolites. We estimate reduced overall sampling time to result in a loss on the order of a factor of 10. Given a reported resolution for MRSI of 5–10 mm (37), we project the resolution achievable with our method to be roughly on the order of 2.5–5 mm with T_2^* -induced PSF blurring not exceeding 0.6 mm (*SI Results*).

Conclusions

We have characterized the spectral properties of the myelin proton signal in situ, as well as in reconstituted suspensions of myelin lipid extract. Our results show that the short T_2^* component of WM originates primarily from myelin lipid protons and further that direct imaging of these protons is possible even though the shortest components are not detectable. Last, our analysis suggests that translation from the laboratory to clinical MRI will be challenging.

Materials and Methods

All MR spectroscopy and imaging experiments were performed on a 9.4 T vertical-bore spectrometer/microimaging system (Bruker DMX 400) with Micro2.5 gradients (100 G/cm maximum strength) and BAFFA40 amplifiers.

Neural Tissue Preparation. SC samples were harvested from healthy adult Sprague-Dawley rats (Charles River Laboratories) and bovine spinal columns (Bierig Brothers Veal and Lamb Products). The rats were killed by carbon dioxide asphyxiation in accordance with an Institutional Animal Care and Use Committee-approved protocol. After killing, rat spinal columns were removed, and the SC was dissected out.

NMR Spectroscopy. High-resolution proton-decoupled ^{13}C NMR [Sweep width (SW) = 24 kHz, number of scans (NS) = 36,768, number of real and imaginary data points (TD) = 65,536, repetition time (TR) = 1.36 s, $\alpha = 30^\circ$] and proton-decoupled ^{31}P (SW = 3.23 kHz, NS = 8,536, TD = 8,192, TR = 1.27 s, $\alpha = 30^\circ$) spectra were collected for samples of purified bovine myelin extract, dissolved in a (5:4:2) ternary mixture of deuterated chloroform (99.8 atom % D; Acros Organics), methanol (99.8 atom % D; Acros Organics), and 0.2 M EDTA/water (99.9 atom % D; Sigma-Aldrich).

All ^1H NMR spectra were collected with the following parameters: SW = 100 kHz, NS = 256, TD = 262,144, TR = 3.6 s, $\alpha = 90^\circ$, pulse duration = 9.6 μs . Freshly excised SC sections (<2-h postmortem interval) were immersed in a perfluorinated oil (Fomblin-Y; Sigma-Aldrich) before experiments.

SL Fitting of Proton Spectrum. As described by Wennerström (30), only partial averaging of dipolar coupling via translational and rotational diffusion occurs, resulting in a dipolar-broadened liquid-crystalline lipid system with an SL lineshape that can be written as:

$$L(\omega) = \int_0^{\pi/2} \frac{\sin(\theta)}{|3\cos^2(\theta) - 1|} f \left[\frac{\omega - \omega_0}{|3\cos^2(\theta) - 1|} \right] d\theta \quad [1]$$

where ω_0 is the chemical shift, θ is the angle of the lipid bilayer surface normal with respect to B_0 , and $f(\omega - \omega_0)$ is any highly peaked lineshape such as

a Gaussian or Lorentzian. Assuming θ is uniformly sampled and setting $f(\omega - \omega_0)$ to be a Lorentzian, it can be seen from Eq. 1 that an SL is composed of a series of scaled Lorentzians. From the widths and intensities of these Lorentzians, the T_2^* distribution of a single SL can be calculated. Protons at different chemical shifts (e.g., alkyl chain methylenes, terminal methyls, and choline) are each expected to give rise to SL lineshapes (32).

Spectral fitting was performed in Matlab (Mathworks). Four SLs were used to represent general alkyl chain methylenes (noncholesterol, mostly from fatty acid residues), cholesterol alkyl chain methylenes, terminal methyls, and choline, whereas a single Lorentzian was used to model residual HDO. Because cholesterol alkyl chain methylenes sit deep within the lipid bilayer, it is reasonable to expect them to be more mobile than the general alkyl chain methylenes, therefore resulting in a narrower SL. The chemical shifts of each SL were set to the known isotropic shift of the various moieties, and the width and relative intensities were free parameters. The R^2 of the fit was greater than 0.99.

UTE MR Imaging. The 3D de-IR-UTE imaging (Fig. S4): SW = 200 kHz, TE = 20/1,200 μ s, TI = 500 ms, TR = 1 s, field of view = 15 mm, matrix size = 128 \times 128 \times 128, number of views = 5,342, pulse duration = 20 μ s. The sequence was based on that used by Anumula et al. (38). TI was determined empirically as the duration yielding optimal GM suppression (because GM is expected to have negligible myelin concentration) in a complex difference

image. A refocusing gradient was applied immediately after the first readout gradient, after which a second gradient echo was collected at TE = 1,200 μ s. A 3D image of the short T_2^* components was obtained as the complex difference of the two echo images (i.e., TE₁ – TE₂). A complex difference is necessary to distinguish the possible presence of both inverted and non-inverted voxel signals.

A 2D projection de-IR-UTE sequence was used to image the series of myelin/D₂O suspensions to avoid signal losses resulting from settling of myelin during scanning. The Mn doped water phantom served to identify the locations of the samples in the image. All experimental parameters were identical to those used in the 3D de-IR-UTE experiments.

All image reconstruction was done in Matlab (Mathworks) using a fast gridding algorithm (39) and incorporating k-space trajectory correction (40). All images were smoothed via bilinear interpolation with Image J (National Institutes of Health).

ACKNOWLEDGMENTS. We thank Joseph J. Sarver and Louis J. Soslowsky for providing a source of fresh rat CNS tissue and Jim Delikatny and Jeremy Magland for their help with the super-Lorentzian analysis. M.J.W. thanks Mary A. Selak for patient guidance and assistance perfecting the sucrose gradient-based isolation of myelin. This work was supported by National Institutes of Health Grant T32 EB00814 and US Department of Defense Award W81XWH-10-1-0714.

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