Magnetic resonance measurements of cellular and sub-cellular membrane structures in live and fixed neural tissue

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We develop magnetic resonance (MR) methods for measuring real-time changes of tissue microstructure and membrane permeability of live and fixed neural tissue. Diffusion and exchange MR measurements are performed using the large static gradient produced by a single-sided permanent magnet. Using tissue delipidation methods, we show that water diffusion is restricted solely by lipid membranes. Most of the diffusion signal can be assigned to water in tissue which is far from membranes. The remaining 25% can be assigned to water restricted within membrane structures in the range of 200–1400 nm at the cellular, organelle, and vesicle levels. Diffusion exchange spectroscopy measures water exchanging between membrane structures and free environments at 100 s⁻¹.

tissue microstructure|diffusion exchange spectroscopy | porous media | membrane permeability| NMR-MOUSE Correspondence: basserp@mail.nih.gov

Introduction

The first diffusion tensor images of brain tissue showed diffusion anisotropy in the white matter (1). It was postulated that this anisotropy is due to myelin membranes and other cellular components impeding water mobility more in the direction perpendicular to the oriented fibers than parallel to them. By process of elimination, Beaulieu concluded that the origin of diffusion anisotropy in white matter is due to axonal cell membranes (2, 3). New methods to clear lipid membranes while leaving other tissue components intact (4, 5) have confirmed directly that diffusion becomes isotropic and diffusivity approaches the value of free water after complete delipidation (6).

A characteristic of magnetic resonance (MR) is that spin magnetization retains the history of motions encoded during the experimental pulse sequence (7). Diffusion MR measures the spin echo (8) signal attenuation of nuclear spins which displace randomly in the presence of a magnetic field gradient during a defined diffusion encoding time (9–11). Signal attenuation is achieved by increasing the gradient strength, the gradient duration, and/or the diffusion encoding time, and the effect can be summarized in a single variable, b (12). The diffusion MR signal from a single species such as freely diffusing water with self-diffusion coefficient D_0 attenuates as $\exp(-bD)(13)$, while water diffusing within restricted environments attenuates more slowly (14–18). The diffusion MR signal of water in heterogeneous materials such as biological tissue would be expected to contain a multitude of components arising from water in different microenvironments, which restrict water diffusion to varying degrees (19).

Investigation of neural tissue using diffusion MR has led to numerous reports of two dominant diffusivity regimes. As a result, a biexponential model, which assumes that the signal is a sum of two components with distinct diffusion coefficients (20-22), has been used to fit the water diffusion signal attenuation. Albeit simple, a two-component model cannot capture the complexity and heterogeneity of the underlying microstructure of biological tissue. With the advent of model-independent empirical data inversion techniques, the diffusion signal attenuation could be described by a distribution of diffusion coefficients (23), a phenomenological representation which makes no assumptions about tissue microstructure (24). Pfeuffer, et al. (23), along with Refs. (25, 26), suggest that the diffusion coefficient distribution can be used to investigate the microstructural properties of neural tissue.

Nuclear spins may also exchange along the diffusion coefficient distribution by moving between microenvironments, causing diffusion coefficients of components to shift and appear closer together on the distribution. Exchange can be measured from the change in apparent diffusion coefficients with encoding time (27-30). Alternatively, MR can store the spin history from one encoding b_1 during a mixing time t_m and recall it for a second encoding b_2 (31) to measure exchange along the distribution (32). The standard diffusion measurement is one-dimensional (1-D) in that there is one encoding variable b and one measured parameter D. By encoding the spins twice (b_1, b_2) , the diffusion exchange spectroscopy (DEXSY) sequence becomes 2-D (32). 2-D DEXSY measures the relationship between spins' diffusion coefficients at two separate instances (D_1, D_2) to show exchanging and non-exchanging components (33, 34).

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The full DEXSY sampling of the 2-D $b_1 - b_2$ space (32) is too time-consuming for scanning live specimen. Recent research shows that there is some redundancy in the data (34–37) and alternative DEXSY-based approaches may measure exchange with fewer data points (34, 38, 39). Cai, et al. (39), developed a rapid measurement of the exchange fraction, f, from just four points in the $b_1 - b_2$ space.

Larger gradient strengths and gradient durations probe smaller structures (10). Hardware constraints cap the maximum strength of gradient coils of MRI systems at a few Tesla/m. With long gradient pulse durations and encoding times, diffusion MR microstructural resolution is predicted to be limited to structures larger than a micron (40). However, when water pools communicate during encoding, signal components become blurred, further limiting resolution (41). Gradient pulses last at least a millisecond, which sets a lower limit for the encoding time (42). Surprisingly, an experiment dating back to the origins of MR (8, 9), performed in a large static gradient field, can break this microstructural resolution limit (43). Low-cost, portable, bench-top, singlesided NMR devices with greater than 10 T/m static gradients (44) can probe sub-micron structures (45) that ordinarily cannot be resolved from larger microscale structures using state-of-the-art pulsed gradient MR systems with lower maximum gradient strengths (46). Displacement encoding within a static gradient field occurs by using radiofrequency (RF) pulses (8, 9) to switch the "effective gradient" (11), allowing for diffusion encoding times as short as 100 μ s (47). This permits resolution of sub-micron structures that can contain rapidly exchanging water pools (45). The static gradient 1-D diffusion (48) and 2-D DEXSY (49) experiments can then be used to probe cellular and sub-cellular components and water exchange between them.

In this paper we adapt 1-D diffusion and 2-D diffusion exchange methods to perform measurements with a singlesided MR system having static gradients, to investigate the cellular and sub-cellular structures in isolated neonatal mouse spinal cord. We develop a system to support both live and fixed spinal cords during NMR measurements such that data could be compared directly. Diffusion coefficient distributions show signal from water in various restricted environments. Diffusion signal attenuation can separate restrictions in the window of 200-1400 nm, indicating that sub-cellular structural resolution is achieved. DEXSY reveals the water pools that exchange between restricted and free environments on a timescale of 10 milliseconds, indicating that subcellular membrane structure can only be measured with encoding times less than 10 ms. Replacing the protons of water with deuterium (D₂O) decreased the signal from all components of the distribution equally, indicating that the majority of the signal is coming from water. Delipidation of membranes by the surfactant Triton X indicated that restriction was caused by lipid membranes and not proteins.

Results

1-D diffusion measures 25% of water to be restricted. Signal was acquired over an entire region including the spinal cord sample and artificial cerebro-spinal fluid (aCSF) bathing the sample. The contribution from the aCSF surrounding the sample needed to be quantified. Fig. 1 shows distributions of diffusion coefficients for a fixed spinal cord bathed in aCSF (a) and after removing the aCSF (b). These can be compared to the distribution from only aCSF filling the RF coil (c). Differences in the fraction of the most mobile diffusion coefficient component (integral of $P(D/D_0)$ for $D/D_0 > 0.17$) indicates that aCSF accounts for only 5% of the signal in (a). The solenoid coil itself does an excellent job isolating signal from the tissue. Comparing the distributions in (a) and (b) demonstrate that the solenoid RF coil gets signal from the spinal cord specimen filling its interior (see Methods)).

Diffusion coefficients are nondimensionalized by the diffusion coefficient of aCSF $D_0 = 2.15 \times 10^{-9} \text{ m}^2/\text{s}$ found by monoexponential fits (see supporting information (SI) Fig. S1). aCSF is well described by a single diffusion coefficient and thus its distribution should be a delta function at $D/D_0 = 1$. However, data inversion yields a smooth, broad distribution because of regularization, which is needed to stabilize distribution estimates (50).

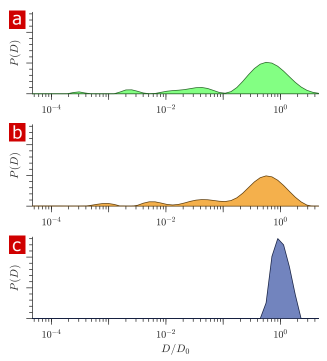


Fig. 1. Diffusion coefficient distributions. a) Distribution of diffusion coefficients from a fixed spinal cord bathed in aCSF. b) Distribution from the same spinal cord after removing the aCSF bath. c) Distribution from only aCSF filling the RF coil.

Signal attenuation and diffusion coefficient distributions from 1-D diffusion measurements performed on a fixed spinal cord specimen are presented in Fig. 2. Signal attenuation from measurements of pure aCSF is also shown for comparison. Attenuation models are used to define signal and distribution components as free (13), localized near surfaces but not completely restricted (17, 18) and restricted (14–16) (described in the SI Appendix). Signal is plotted as a function of the non-dimensionalized diffusion encoding variable $b \times D_0$. Exponential attenuation is expected for fluids diffusing freely, as shown by Eq. S1. N.B. The largest *b* corresponds to 3,000,000 s/mm², two to three orders of magnitude larger than what is typically reached in conventional pulsed gradient diffusion MRI studies. aCSF signal is mono-exponential and quickly attenuates to the background noise level. Spinal cord signal attenuation is multiexponential over the entire range of *b* and does not fully attenuate, implying the presence of multiple highly restricted pools.

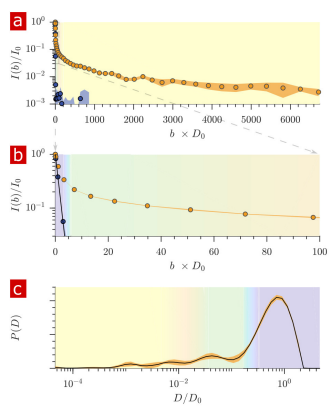


Fig. 2. Diffusion measurement of a fixed spinal cord. a) Mean (circles) and standard deviation (shaded bands) of the signal intensity from five diffusion measurements, spaced 54 minutes apart, performed on a fixed spinal cord (orange) and three measurements performed on aCSF (purple) at 25° C. The signal intensity is plotted as a function of $b \times D_0$ where $D_0 = 2.15 \times 10^{-9} \text{ m}^2/\text{s}$ is the diffusion coefficient obtained from monoexponential fits of the aCSF. b) Signal intensity from the zoomed in area shown in (a). c) The distribution of diffusion coefficients resulting from inversion of the data. The purple, green, and yellow shading across plots signifies water which is free, localized, and restricted respectively. The color gradient is meant to signify a continuous change between diffusion regimes rather than sharp boundaries.

The distribution of apparent diffusion coefficients is shown in Fig. 2 (c). The majority of the distribution has $D/D_0 > 0.3$, and this is labeled the free water component. Humps extend to lower values of D/D_0 and these represent signal which is more and more restricted and on smaller length scales, as indicated by the color gradient.

Full DEXSY measures water exchanging between free and restricted environments 100 times per second. 2-D DEXSY labels spins based on their local mobility at two instances which are separated by the mixing time variable t_m (32). This permits the direct measurement of water movement from one environment (e.g. A) to another (B) as well as water moving in reverse (B) to (A) to fulfill mass conservation. In the case that water exchanges between environments on intermediate timescales (greater than the diffusion encoding time and less than the longitudinal relaxation time T_1 which causes spins to forget their encoding), the exchange increases and saturates as a function of t_m (34, 39, 51, 52). The classic analysis of DEXSY is as a joint 2-D probability distribution showing relationships between the apparent diffusion coefficients of water populations during the first encoding period, D_1 and the second encoding period D_2 (34). Integrated probability density at a point or region (D_1, D_2) indicates the probability of a spin being at D_1 during encoding 1 and D_2 during encoding 2. Non-exchanging water populations have $D_1 = D_2$, defining a diagonal line across the distribution whereas exchanging water populations are located off the diagonal. A representative 2-D DEXSY distribution of spinal cord is shown in Fig. 3 (a). The distribution is divided into a 3x3 grid for the possible exchange pathways between components A $(3.2 \times 10^{-4} - 1 \times 10^{-2})$, B $(1 \times 10^{-2} - 3.2 \times 10^{-1})$, and C $(5.6 \times 10^{-1} - 1 \times 10^{1} D/D_{0})$, shown by the color coding and labels. The integrated probability density from each region represents an exchange (off-diagonal) or nonexchange (on-diagonal) fraction. The distribution shows exchange between free water, labeled as C, and restricted water, labeled as B and A. Additionally, there appears to be exchange between restricted components. Stacked plots at $t_m = 0.2, 4, 20$, and 160 ms (b) show the increase in probability for components exchanging with free water, regions AC + CA and regions BC + CB and a decrease in probability for the non-exchanging components, e.g., region AA appears to decrease to near zero at the longest t_m .

Fig. 4 shows exchanging (a) and non-exchanging (b) fractions, f from integrating regions of the 2-D probability distributions. The build-up (a) and decay (b) of f over t_m are fit with a first order rate equation to obtain apparent exchange rates (AXRs), presented in the figure caption. Measurements on six different spinal cords show consistent exchange behavior. Restricted components exchange with free water with AXR $\approx 100 \text{ s}^{-1}$. f_{AB+BA} does not increase with t_m , indicating that the DEXSY measurement is primarily sensitive to water exchanging between restricted and free environments and not between and among different restricted environments.

Rapid exchange measurement agrees with full DEXSY.

Full DEXSY measurements at four mixing times took eight hours—too long to measure exchange in living tissue. Therefore, we strove to develop a method to rapidly measure exchange (39) (see SI). The rapid measurement provides an apparent exchange fraction f, a diffusion-weighted average of exchange between all water pools. The full DEXSY can resolve multiple exchanging water pools and the exchange pathways between them (53, 54). Although the rapid measurement lacks the full DEXSY's resolution of multicomponent exchange, it provides enhanced temporal resolution, both with respect to t_m and experimental time, by sidestepping the need for 2-D data inversion (55) and by acquiring the data much more rapidly. The protocol used here acquired the data at one exchange fraction (one mixing time) per minute.

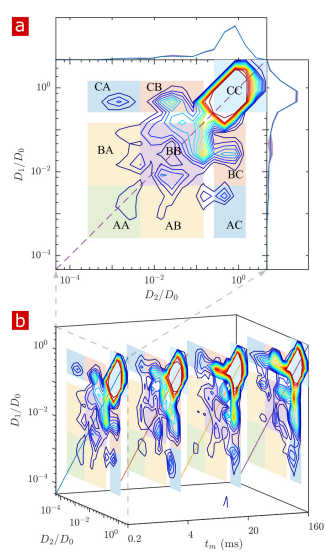


Fig. 3. Full 2-D DEXSY diffusion exchange distribution for a fixed spinal cord. a) Exchange distribution measured with mixing time $t_m = 0.2$ ms. Distributions show exchanging (off-diagonal) and non-exchanging (on-diagonal) components. These components are lumped into regions A, B, and C, shaded and labeled in a 3×3 grid for each exchange combination. The range of $P(D_1, D_2)$ is set to add detail to components A and B, but cuts off the top of the most mobile region CC. The marginal distributions $P(D_1)$ and $P(D_2)$ are presented on the sides, with means (solid blue lines) and standard deviations (shaded bands around lines) from 3 measurements. b) A stacked view of distributions measured with mixing time $t_m = [0.2, 4, 20, 160]$ ms. With increasing t_m , the probability density builds up in regions of free and restricted water exchange AC + CA and BC + CB and decays for non-exchanging restricted water regions AA and BB. The build-up and decay is quantified in Fig. 4.

Exchange fractions from the rapid measurement are presented in Fig. 5 for the same specimen as used for the full DEXSYs (Figs. 3 and 4). The AXR from three repeat measurements was $110 \pm 30 \text{ s}^{-1}$. This value is not different statistically from the results of the full DEXSY measurement, validating the rapid measurement method. Additionally, the value from repeated measurements on five fixed samples was $110 \pm 6 \text{ s}^{-1}$ at 25° C, indicating high reproducibility between specimen.

Restricted diffusion measures sub-micron structures.

After free water has fully attenuated, restricted water signal attenuation is exponential with the diffusion encoding time τ

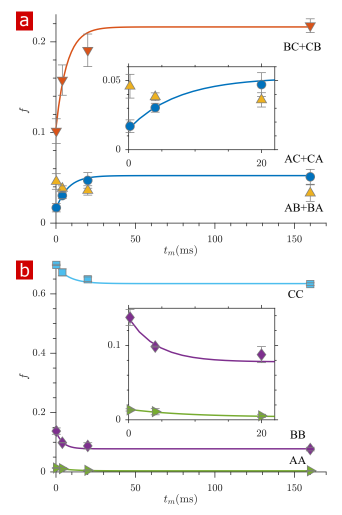


Fig. 4. Fractions of exchanging and non-exchanging water. a) Fractions of water which exchange between regions A, B, and C, (defined in Fig. 3) as a function of mixing time, t_m . Exchange fractions $f_{\rm AC+CA}$ and $f_{\rm BC+CB}$ are fit with a first-order rate model with estimated apparent exchange rates AXRs (mean±standard deviation from three measurements) 140 ± 80 and 140 ± 140 s⁻¹ respectively. $f_{\rm AB+BA}$ does not follow the expected first order rate model. b) Fractions of non-exchanging water. Non-exchange fractions $f_{\rm AA}$, $f_{\rm BB}$, and $f_{\rm CC}$ are fit with a first order rate model with estimated AXRs $10\pm100, 230\pm170$, and 130 ± 100 s⁻¹ respectively. Insets zoom in on the initial rise in exchange. Error bars are the standard deviations from three repeat measurements.

and the size of the restriction (14-18), as presented in Eqs. S2–S5. Attenuation models for water restricted in spheres of radius R indicate that with q = 15.3 T/m the diffusion experiment provides a 200-1400 nm window on restriction radii (Fig. 6 (a)). Signals from water in restrictions smaller than R = 200 nm do not attenuate significantly enough to differentiate. In restrictions larger than R=1400 nm, signal from water far from the surfaces attenuates as free water and signal from water near the surface attenuates as localized water (restricted on one side but free on the other). The long time behavior of the diffusion signals are analyzed to estimate a radius of restriction in Fig. 6 (b). Exchange also causes attenuation which is exponential with τ (45). The estimate accounts for attenuation due to exchange, utilizing the measured AXR. The estimated radius is R = 900 nm. This can be viewed as a volume-averaged restriction length, filtering out water in structures with R > 1400 nm.

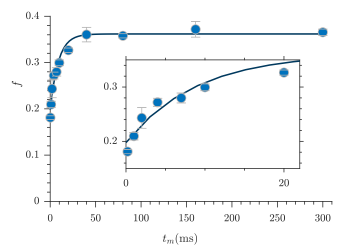


Fig. 5. Rapid measurement of exchange fractions. A fit of the first order rate model estimated an apparent exchange rate, AXR = $110 \pm 30 \ {\rm s}^{-1}$. Inset shows a zoomed-in region of the initial rise in exchange.

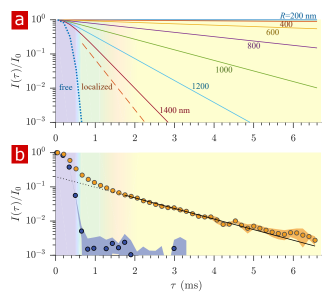


Fig. 6. Estimating membrane structure sizes from the diffusion signal attenuation. a) Signal intensity is simulated for water restricted in spherical compartments of varying radius between R = 200 - 1400 nm (Eq. S3, solid lines) (16), for water localized near a surface in larger restrictions (Eq. S2, dashed line) (17, 18), as well as water diffusing freely (Eq. S1, dotted line) (13). Signal is plotted as a function of the variable τ (rather than $b \sim \tau^3$). b) Signal is re-plotted from Fig. 2. Signal at $\tau \geq 1.8$ ms is with the model for water restricted in spherical compartments in the limit of long τ (Eq. S5, solid line) (16) incorporating the AXR=110 s⁻¹ (45), estimating a radius R = 900 nm. The dotted line extrapolates back to $I/I_0 = 0.2$.

NMR recordings do not affect viability of spinal cord. The signal attenuation from diffusion measurements performed on live spinal cords (n=9) is compared to that of fixed spinal cords (n=6) and only aCSF in Fig. 7 (a). The signal attenuates slightly faster although not significantly as seen by the standard deviations. The mobility of water on the timescale of milliseconds is very similar in live and fixed specimen.

After two hours of NMR measurements and four to seven hours post-dissection, spinal cords (dissected on postnatal (P) day P2, P3, and P4) were assessed for viability by recording motoneuronal electrical activity after stimulation of a dorsal root. Mono- and polysynaptic reflexes were recorded in all preparations (n=4), Fig. 7 (b), indicating that neither the experimental setup nor the protocol compromised the neuronal excitability of the spinal cord.

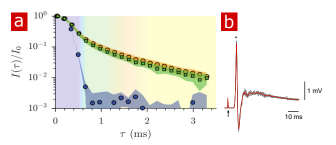


Fig. 7. Diffusion in fixed vs. live. a) Signal intensity from diffusion measurements performed at $25^{\circ}C$ on live samples (n=9) (green squares), fixed samples (n=6) (orange circles) and aCSF (blue circles) plotted as a function of the variable τ . (b) Mono- and poly- synaptic reflexes were recorded from the L6 ventral root of live samples (n=4) after NMR measurements. Stimulation was done on the homory-mous dorsal root. The grey lines are 5 successive stimuli (30 s interval) and the superimposed red line is the average signal. The arrow indicates the stimulus artifact and the star the monosynaptic reflex.

NMR measurements are primarily sensitive to water. To determine whether biomacromolecules were contributing to the signal observed in the spinal cords, rapid exchange and 1-D diffusion measurements were recorded in real-time as a specimen was washed with aCSF made with 99.8% deuterium water (D₂O aCSF). (Results are presented in Fig. S2). After two successive washes, proton signal decreased to values similar to D₂O aCSF alone. All diffusion coefficient distribution components decreased similarly after D₂O washing. Components of the distribution which are not from water would still remain after removing H₂O. Therefore, all distribution components are primarily made up of water. This points to water rather than biomacromolecules accounting for the vast majority of the measured signal.

Delipidation shows membranes to be the sole source of restriction. Triton X surfactant was used to remove lipid membranes from spinal cords in order to determine the effect of membranes on water restriction. The aCSF bathing the spinal cord was replaced with aCSF with Triton X while rapid exchange and 1-D diffusion measurements were repeatedly performed (n=2). Fig. 8 shows exchange fractions (a) and 1-D distributions from select time points (b).

After the addition of 1% Triton X at 1 hr, the exchange fractions decreased slowly and reached a plateau. Washing to 5% Triton X at 50 hrs decreased the exchange fractions further until they again reached a plateau.

The diffusion coefficient distributions Fig. 8(b) (and SUP. MOVIE S1) show that the delipidation process removed the barriers for restriction. Therefore, lipid membranes, not the remaining macromolecules, are the source of restriction of water diffusion. The fraction of restricted water (Fig. S3) decreases and plateaus similarly to the exchange fractions.

About 6% of the signal appears restricted after 120 hrs of delipidation. This is signal from Triton X, which as a 5% solution imparts a 6% signal at $D/D_0 = 0.01$ (see Fig. S4). Samples were also studied after full delipidation and wash-

Samples were also studied after full delipidation and washing away Triton X (Fig. 9). The diffusion signal attenuation,

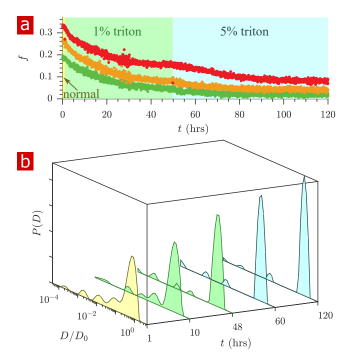


Fig. 8. Timecourse study of Triton X delipidation. a) Exchange fractions from rapid exchange measurements with $t_m = 0.2$ (green), 4 (orange), and 20 ms (red) measured throughout the timecourse, as the sample was washed to aCSF with 1% Triton X, and then 5% Triton X. b) Representative diffusion coefficient distributions from 1-D diffusion measurements performed at different times before and after addition of Triton.

(a) and (b), shows that 95% of the signal is monoexponential with $D/D_0 = 0.90$. The diffusion coefficient distribution (c) shows one major free diffusion peak which is not significantly different from the diffusion coefficient distribution of pure aCSF. Some small peaks which are not seen unless P(D) is magnified lead to a 1% restricted component. Without membranes there is essentially only one diffusive environment throughout the whole sample.

Discussion

We present NMR methods which use the diffusion of water to probe cellular and sub-cellular membrane structures on submillisecond and millisecond timescales. Much of the advance was possible because the large static gradient overcomes (43)many hardware (42) (e.g. slew rate, eddy currents, maximum gradient strength) and biological (e.g. peripheral nerve stimulation (56)) limitations of pulsed gradients. In 1996 Köpf, et al., realized this capability on biological tissue, performing diffusion measurements using a 50 T/m static gradient in the stray field of a superconducting 9.4 T magnet (57). Another stray field study by Carlton, et al., used an 18 T/m static gradient to measure bacteria concentrations (45). Bacteria cell membranes are roughly a micron in diameter, similar to subcellular structures in tissue. They noted the static gradient experiment provided more intracellular signal compared to pulsed gradient experiments on bacterial systems (46) due to less exchange during the shorter diffusion encoding time. We re-purposed a low field single-sided permanent magnet (44) which, due to its profiling capabilities (58), has found a num-

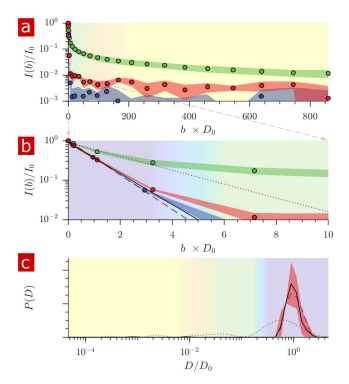


Fig. 9. Diffusion measurement after delipidation. a) Diffusion signal intensity from measurements on spinal cords performed at 25° C after delipidation (n=2) with 10% Triton X and after washing the Triton away. The mean (circles) and standard deviation (shaded bands) of the attenuation are plotted for the delipidated samples (red) alongside pure aCSF (purple) and fixed undelipidated spinal cords (n=6) (green). b) The initial attenuation of signal. Monoexponential fits of the attenuation for moints 2–4 yielded 2.15 ± 0.01, 1.94 ± 0.02 and 0.87 ± 0.14 × 10⁻⁹ m²/s for the aCSF, delipidated, and undelipidated spinal cords and are shown as the dashed, solid, and dotted lines respectively. c) Diffusion coefficient distribution of the delipidated spinal cords, for which the mean (solid line) and standard deviation (shaded band around line) are not significantly different from the pure aCSF (dashed line). The distribution from a fixed, undelipidated spinal cord (dotted line) is also shown for comparison. The purple, green, and yellow shading across the plots signifies water which is free, localized, and restricted.

ber of niche applications in materials science and engineering, biology and medicine, and cultural heritage (59–61). The large 15.3 T/m gradient allowed for the attenuation of tissue water signal below $I/I_0 = 0.01$ in a diffusion encoding time of 6.6 ms.

A large signal-to-noise ratio (SNR) and Gaussian zero mean noise was necessary to resolve slowly attenuating signals above the noise and to not confuse the signal with a baseline noise floor. In general, SNR > 100 is needed for diffusion coefficient distribution analysis (62) and as a rule-of-thumb this allows for resolution of populations comprising as little as 1% of the signal. Performance tuning led to very stable measurements and SNR > 600. These modifications included a 2000 echo CPMG readout, 25 μ s echo time, a sample-specific solenoid RF coil, a wet/dry chamber without circulation, and noise reduction/isolation.

The solenoid RF coil permitted low-power, 2 μ s RF pulses, high filling factor, and maximized signal from the spinal cord filling its interior relative to aCSF. Previous studies on live ex-vivo neural tissue utilized MR imaging (30, 63–67). However, their analysis was on regions of interest (ROIs) which encompassed the entire sample, indicating that no additional specificity was obtained from the imaging. Because 95% of the signal came from spinal cord tissue, imaging was not necessary. This let us achieve high SNR and sufficiently rapid measurements.

The wet/dry chamber kept the liquid environment still while the gas environment provided oxygenation. Previous diffusion MRI studies on live ex-vivo neural tissue provided oxygen to the sample through perfused aCSF (30, 63–68). Media perfusion can cause convection artifacts in the diffusion measurement (69). Researchers typically implemented start-stop diffusion MR protocols, with aCSF perfusion between MR measurements (65, 66). However, a steady concentration of oxygen is preferable and better represents the in-vivo environment. The wet/dry chamber provided a constant supply of oxygen to the tissue while avoiding convection artifacts, creating ideal conditions for diffusion measurements on live tissue. We also directly demonstrate that spinal cords are alive after hours of NMR measurements by recording electrical responses.

Our novel experimental setup reveals signal from highly restricted pools. We determined that these pools are exclusively membrane-restricted water. Previous nerve tissue studies reported two (1-D encoding (23, 25, 26)) and three (2-D, integrated diffusion-relaxation (19)) resolvable water mobility components, spanning 2 orders of magnitude. Here, distributions of diffusion coefficients showed components that are 3 orders of magnitude lower than free water. Similar values have been reported for other, larger, proton bearing molecules naturally occurring in biological tissue (70). However, by replacing the water in the tissue with deuterated water, we determined our methods to be sensitive primarily to water and not macromolecules.

After full delipidation, the free water peak of the diffusion coefficient distribution accounts for 99% of the water (Fig. 9), vs. roughly 75% of the water before delipidation. Signal attenuation is monoexponential with $\langle D \rangle / D_0 = 0.90$, vs. multiexponential with $\langle D \rangle / D_0 = 0.40$ before delipidation. Deviation from monoexponential, Gaussian signal attenuation is due to restriction by lipid membranes. From micron (2, 3, 6) to sub-micron length scales, water restriction in tissue is due solely to membranes.

After removing restrictions, hindrances to water diffusion can be thought of as arising from volume obstruction by macromolecules such as proteins within the tissue. Obstruction models of water (solvent) self-diffusion incorporate only the volume fraction of macromolecules, θ , as a free parameter and are generally adequate models in the limit of low (< 0.1) volume fraction (71). Such models predict $0.03 < \theta < 0.07$ for $\langle D \rangle / D_0 = 0.90$. Neural tissue consists of 8% proteins (6) and thus can account for the obstruction effects. Leuze, et al. used another lipid clearing method to determine that lipids are the dominant source of MRI contrast. We can now additionally say that proteins act as simple obstructions, reducing water diffusion only slightly from D_0 . This finding can be compared to previous reports of water diffusion in cytoplasm isolated from red blood cells, showing $\langle D \rangle / D_0 = 0.70$ (72). Although we used shorter diffusion encoding times, delipidated samples showed monoexponenital, Gaussian diffusion signal attenuation with b, indicating that microstructural information is averaged out during the encoding time and should show no additional encoding time dependence (73). The decreased diffusivity from cytoplasm (72) could arise from the presence of organelles and membrane particles still present in the supernatant after lysing and centrifuging the red blood cells.

The static gradient diffusion measurement provided a window on membrane structures in the range of 200–1400 nm. The neonatal mouse spinal cord contains mostly gray matter (74, 75). Structures within 200–1400 nm include cells and portions of cells with small radii such as axons, dendrites and dendritic spines, myelin, and a number of membranous organelles such as nuclei, mitochondria, endoplasmic reticula, and vessicles (76), in addition to extracellular sheets and tunnels (77). Static gradient methods are complementary to pulsed gradients, which provide enhanced resolution to micron-sized cell membrane restrictions, but are unable to attenuate signal from sub-micron cells and organelles (78, 79).

Full 2-D DEXSY distributions showed restricted water exchanges with free water, but could not resolve exchange between restricted water pools. The rapid measurement was designed to hone in on the exchange between restricted and free water pools and AXRs were consistent with results from the full DEXSYs. Results indicate that we have developed a non-invasive, sufficiently rapid method of measuring exchange across membranes in live tissue. The AXR $\approx 100 \text{ s}^{-1}$ is significantly faster than intracellular-extracellular water exchange rates measured in neural tissue ($\approx 0.5 - 5 \text{ s}^{-1}$) (41, 80-83). Such fast turnovers are not unheard of, e.g. red blood cells show similar ($\approx 100 \text{ s}^{-1}$) rates (27, 28, 30) due to their high expression of aquaporin (84). In addition to membrane permeability, the other factor affecting exchange is the ratio of membrane surface to volume. This ratio increases with smaller structure sizes. Therefore, fast AXRs can be explained by the large surface to volume ratios of the structures which the measurement is sensitive to.

The methods were used to follow penetration of deuterated water into the tissue as well as the delipidation of the tissue via Triton X. The timescale of water penetration was ≈ 1 min, consistent with mass transport theory, but the timescale of Triton X penetration and delipidation was ≈ 1 day, longer than predicted (≈ 3 hrs) (see SI). An increased time is expected due to the reaction front which develops as Triton X delipidates, slowing its overall penetration.

Previous studies addressing the microstructural resolution limit of diffusion MR treated the problem as solely limited by gradient strength (40). Higher exchange causes the size of structures to be overestimated. The fast exchange rates measured in the current work indicate that resolution is limited by the diffusion encoding time in addition to the maximum gradient strength; exchange cannot be neglected.

Methods

The experimental test chamber (Fig. 10(a)) was designed to support live spinal cord for hours without requiring oxygenated artificial cerebro-spinal fluid (aCSF) flow, thus avoiding flow-related measurement artifacts (69). The gastight wet/dry chamber was fabricated at the NIH/NINDS mechanical workshop. The assembled chamber had two environments—a static liquid environment with aCSF and above it a gas environment with a slow flow of humidified 95% O₂ and 5% CO₂ gas. The sample temperature can be controlled in the range of $7 - 37^{\circ}$ C. Sample temperature was monitored by a PicoM fiber optic sensor (Opsens Solutions Inc., Québec, Canada) and regulated by a shallow water bath surrounding the chamber. The bottom portion of the chamber was made of aluminum to provide good heat conduction to the media. (See SI for additional information.)

In order to compare data between live and fixed tissue, oxygenated aCSF was used as the buffer solution for all experiments. A spinal cord was placed inside the solenoid RF coil within the chamber half-filled with aCSF previously bubbled with 95% O_2 and 5% CO_2 . The chamber was sealed and connected to gas flow with humid 95% O_2 and 5% CO_2 .

Mouse spinal cord dissection, fixation, and delipida-

tion. All experiments were performed on Swiss Webster wild type (Taconic Biosciences, Rensselaer, NY, USA) between one day after birth to postnatal day 4 and were carried out in compliance with the National Institute of Neurological Disorders and Stroke Animal Care and Use Committee (Animal Protocol Number 1267-18). The mouse spinal cords were isolated and placed in a dissecting chamber perfused with cold Low-Calcium High Magnesium aCSF (concentrations in mM: 128.35 NaCl, 4 KCl, 0.5 $CaCl_2 \cdot H_2O$, 6 $MgSO_4 \cdot 7H_2O$, 0.58 $NaH_2PO_4 \cdot H_2O$, 21 $NaHCO_3$, 30 Dglucose) bubbled with 95% O_2 and 5% CO_2 . To expose the spinal cords, a ventral laminectomy was performed, and they were subsequently isolated together with the ventral roots and ganglia. Spinal cords were roughly (anterior-posterior length \times lateral width \times ventral-dorsal height) $15 \times 1 \times 1.5$ mm, increasing with days postnatal.

Prior to live spinal cord transportation, the cord was placed in a sealed 50 ml tube with 10 ml aCSF previously bubbled with 95% O_2 and 5% CO_2 . The air in the tube was flushed with 95% O_2 and 5% CO_2 .

For fixed experiments, at the end of dissection the cords were fixed in 4% paraformaldehyde overnight at 4°C. Fixative was then replaced with aCSF three times over the course of two days to remove any residual paraformaldehyde.

Triton X-100 (Sigma-Aldrich) nonionic surfactant was used to delipidate spinal cords. Samples (n=2) were studied during delpidation by replacing the aCSF media with aCSF media containing a specified % of Triton X during NMR recording. Samples (n=2) were also studied after delipidation with 10% Triton X in phosphate buffered saline (PBS) for two days, removal of Triton X by periodically replacing the PBS media for two more days, and equilibration in aCSF for a final day. **NMR hardware.** NMR measurements were performed at 13.79 MHz proton frequency with a Kea2 spectrometer (Magritek, Wellington, New Zealand). A PM-10 NMR MOUSE (Magritek, Aachen, Germany) permanent magnet (44) provided a B₀ magnetic field, specially designed to be constant along an x-y (20 mm \times 20 mm) plane parallel to the magnet's surface, but decreases rapidly and linearly in the perpendicular (y-direction) from the surface, providing a strong static magnetic field gradient (See Fig. 10(c)) (58). The NMR MOUSE was raised or lowered with a step per motor with a step size of 50 μ m in order to move B₀ = 0.3239 T, 13.79 MHz, to the precise depth within the sample (17 mm from the surface of the magnet). At this depth the magnetic field gradient g = 15.3 T/m, or 650 KHz/mm.

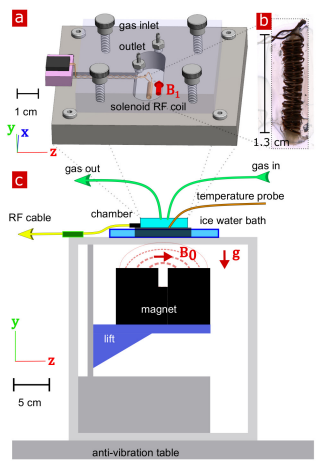


Fig. 10. Experimental setup. a) 3-D technical drawing of the test chamber. b) Image of the solenoid RF coil containing a fixed, delipidated specimen. c) Technical drawing of the experimental setup. The magnet is drawn in the "service" position to show the field lines extending from one magnetic pole to the other. To perform measurements, the magnet would be raised such that the B_0 was correctly positioned relative to the sample. Vectors B_1 , g and B_0 point in the x, y, and z directions respectively

Double-wrapped (length \times inner diameter) 13×2 mm solenoid radiofrequency (RF) coils (Fig. 10 (b)) and an RF circuit were built in-house. The solenoid connected to the circuit board with detachable pin connectors. Tune and match used two trimmer capacitors with range 1–23 pF(NMAJ25HV, Knowles Voltronics). RF pulses were driven by a 100 W RF pulse amplifier (Tomco, Adelaide, Australia). (See SI and Fig. S5 for additional information

and circuit design.)

NMR experimental methods. NMR measurements were performed in Prospa 3.22 (Magritek). All measurements set repetition time (TR) = 2 s, 90°/180° pulse times = 2 μ s and amplitudes = -22/-16 dB, and acquired 2000 CPMG echoes with 25 μ s echo time. The acquisition time and dwell time were 4 and 0.5 μ s respectively, leading to roughly a 400 μ m slice thickness. The lift was positioned such that the signal was at a maximum, thus providing a slice through the center of the solenoid. Signal was phased such that the component from the real channel was maximum and the mean of the imaginary channel component was zero. Measurements were performed without regulating temperature (22 – 26°C) or else at a controlled temperature 25°C when specified in figure captions.

Diffusion measurements were performed using the spin echo sequence (48) (Fig. S6(a)). τ was incremented linearly from 0.05 to 6.55 ms in 43 data points (corresponding to *b* values from 1.4 to 3,130,000 s/mm²) or, for live and some fixed specimen, from 0.05 to 3.3 ms in 22 points. (See SI for additional information.)

The DEXSY sequence (Fig. S6(b)) was written in-house and used eight phase cycle steps. For full 2-D DEXSY measurements (32), data points were acquired on a $21 \times 21 b_1, b_2$ grid by incrementing τ_1 linearly from 0.200 to 3.3 ms in an inner loop and τ_2 from 0.213 to 3.313 ms in an outer loop. For the rapid exchange measurement, points were acquired as a function of b_s and b_d by varying τ_1 and τ_2 accordingly. For rapid exchange measurements (39), the standard 4-point acquisition used one point at $b_s = 200$, $b_d = 20 \text{ s/mm}^2$, and three points along $b_s = 4500 \text{ s/mm}^2$ with $b_d = -4300, -150$, and $b_d = 4300$. Unless otherwise specified, the t_m list was [0.2, 4, 20, 160] ms for full DEXSYs and [0.2, 1, 2, 4, 7, 10, 20, 40, 80, 160, 300] ms for the rapid exchange measurement. (See SI and table S1 for sequence details and phase cycles.) Standard CPMG T_2 (10 s TR, 8000 echoes) and saturation recovery T_1 (1 s TR, 21 recovery points logarithmically spaced to 10 s) measurements were performed, with all other parameters consistent with diffusion and exchange measurements. In all cases, data was monoexponential. (See Figs. S7 and S8.) The T_2/T_1 values were $275 \pm 5/1870 \pm 10$ ms for aCSF (3 measurements), $155 \pm 13/972 \pm 53$ ms for fixed spinal cords (n=10/4), and $176 \pm 35/1030$ ms for fixed spinal cords after delipidation (n=3/1).

NMR data analysis. 1-D distributions were fit using ℓ_2 regularization (50) and singular value decomposition (85, 86), with 50 grid points logarithmically spaced from 10^{-13} to 10^{-8} m²/s, and the regularization parameter chosen using the generalized cross validation (GCV) method (87). 2-D distributions were fit with an algorithm which uses ℓ_2 regularization and singular value decomposition (85, 86), with 21x21 grid points logarithmically spaced from 10^{-13} to 10^{-8} m²/s and the regularization parameter chosen by the L-curve method and held constant for all experiments. Exchange fractions were calculated from the rapid exchange measurement using Eqs. S8–S10 with $D_e = 10^{-9}$ and $D_i =$

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 10^{-11} m²/s. AXRs from both full DEXSYs and the rapid exchange measurement were estimated from fits of a first order rate model, Eqs. S11–S12 (34, 39, 51), incorporating a non-zero initial condition to account for exchange during encoding (52). All analysis was performed using Matlab. (See SI for additional information.)

System characteristics led to high SNR diffusion measurements. Although SNR is highly dependent on the magnetic field strength, the decrease in SNR at low field is boosted by refocussing the signal 2000 times in a CPMG train for each data point (48). Moreover, the solenoid RF coil maximized the sample filling factor, increasing SNR roughly 10-fold from previous flat RF designs (88). RF pulses used little power, permitting short 2 μ s RF pulse durations and producing negligible heat. The coil design allowed for short echo times which reduced relaxation during acquisition in the CPMG train, again boosting SNR. Significant attention was given to shielding and grounding the equipment to minimize noise pickup. All together, 1-D diffusion measurements obtained SNR > 600 (defined as the I_0/σ where σ is the standard deviation of the real channel measured at the noise floor).

Electrophysiological recording. Electrical activity from motoneurons was recorded with suction electrodes into which individual ventral roots (L6 or T10) were drawn after NMR measurements (n=4). The recorded signals were filtered (between 0.1 and 3 kHz) and amplified (gain: 1000), digitized at 10 kHz (Digidata 1500 B) and stored digitally on a computer. Episodes of data were analyzed off-line using Matlab. To elicit monosynaptic responses in motoneurons, the homonymous dorsal roots were stimulated with a single electrical pulse (250 μ s duration) repeated 5 times at 30 s intervals. The threshold for a given spinal root was defined as the lowest current intensity at which that root had to be stimulated to elicit a monosynaptic response in 5/5 attempts. Recordings were obtained at 5 × threshold.

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Theory

Diffusion signal attenuation models. Attenuation of the MR signal in a spin echo diffusion experiment under a static magnetic field gradient can exhibit three diffusion regimes corresponding to three different characteristic length scales: the restriction length l_s , the diffusion length, $l_d = \sqrt{D_0 \tau}$, and the dephasing length, $l_g = (D_0/\gamma g)^{1/3}$ where $D_0 = 2.15 \text{ m}^2/\text{s}$ is the diffusion coefficient of water in artificial cerebrospinal fluid (aCSF) at 25°C, τ is the time between the first 90° and the 180° radio frequency refocussing pulses of the spin echo sequence or 1/2 the echo time TE, and γ is the gyromagnetic ratio (1). The diffusion length is the average distance that water diffuses during the time τ . In the spin echo diffusion measurements, τ was linearly increased from 0.05 to 3.3 or to 6.6 ms, corresponding to $l_D = 0.33$, 2.7, and 3.7 μ m respectively. The dephasing length is the distance that spins must displace in order to dephase by 2π radians. For the 15.3 T/m gradient used here, $l_g = 800 \text{ nm}$. The shortest of these three length scales determines the regime that applies to the diffusing spins, and thus dictates the asymptotic behavior of the spin echo decay. What follows is a basic interpretation of the attenuation regimes, as taken from Hürlimann, et al., (1). A complete discussion of the regimes, higher order terms to the attenuation, and a historical account of the field was provided by Grebenkov (2).

Free diffusion regime. The free diffusion regime occurs when l_D is the shortest characteristic length scale. This regime was first described by Hahn (3) and then the diffusion coefficient of water was measured by Carr and Purcell (4), both using static magnetic field gradients. In this regime, the signal decays by (5)

$$I(\tau)/I_0 = \exp\left(-\frac{2}{3}D_0\gamma^2 g^2 \tau^3\right)$$

= $\exp\left(-\frac{2}{3}\left(\frac{l_D}{l_g}\right)^6\right)$
= $\exp\left(-bD_0\right).$ (S1)

Water diffusion is often modeled as Gaussian with an effective or apparent diffusion coefficient D, rather than D_0 , and in the limit of low attenuation $D = \langle D \rangle$. "Apparent" implies that the measured diffusion coefficient will depend on the experimental parameters (6). This is particularly true when using Eq. S1 to model signal which includes water in other regimes. The use of *b* coefficient or factor comes from diffusion MRI literature (7).

Localization regime. The localization regime occurs when l_s is the shortest characteristic length scale. In this regime, signal near the restrictive surfaces will dephase more slowly than signal farther away. While the entire decay curve can be quite complicated (2), in the asymptotic long-time (τ) limit the signal was experimentally characterized by Hürlimann et al. (1) and theoretically modeled by Stoller, et al. (8), and de Swiet et al (9), and shown to attenuate as

$$I(\tau)/I_0 = c \frac{D_0^{1/3}}{\gamma^{1/3} g^{1/3} l_s} \exp\left(-a_1 D_0^{1/3} \gamma^{2/3} g^{2/3} \tau\right)$$

= $c \frac{l_g}{l_s} \exp\left(-a_1 \left(\frac{l_D}{l_g}\right)^2\right),$ (82)

where $a_1 = 1.0188$ and is, importantly, independent of the confining geometry. The prefactor c varies depending on the geometry and equals 5.8841 for water restricted between parallel plates (9). Note that l_s affects the fraction of signal present in the asymptotic limit but does not affect the decay. By varying τ under a static gradient, we see that the signal attenuates exponentially with τ , $(l_D/l_g)^2$ or $(bD_0)^{1/3}$.

Motional averaging regime. The motional averaging regime occurs when l_g is the shortest characteristic length scale. Signal attenuates very slowly and water can diffuse across the restricted volume many times before dephasing appreciably. Signal decay in the motional averaging regime was first experimentally measured by Wayne and Cotts (10) and subsequently modeled by Robertson (11). Neuman derived the signal attenuation models for water restricted between parallel plates and within cylinders oriented perpendicular to g, and within spheres (12). We focus on the model for spheres of radius R for which the signal attenuates by

$$I(\tau)/I_0 = \exp\left(-\frac{2\gamma^2 g^2}{D_0} \sum_{m=1}^{\infty} \frac{\alpha_m^{-4}}{\alpha_m^2 R^2 - 2} \left(2\tau - \frac{3 - 4\exp(-\alpha_m^2 D_0 \tau) + \exp(-\alpha_m^2 D_0 2\tau)}{\alpha_m^2 D_0}\right)\right)$$
(83)

where α_m is the *m*th root of

$$\alpha_m R J'_{3/2}(\alpha_m R) - \frac{1}{2} J_{3/2}(\alpha_m R) = 0$$
(S4)

for which the first 5 roots are $\alpha_m R = 2.0815$, 5.940, 9.206, 12.405, and 15.579 (13). In the limit of long τ relative to the timescale to diffuse across the restriction, Eq. S3 becomes

$$I(\tau)/I_{0} = \exp\left(-\frac{8}{175} \frac{R^{4} \gamma^{2} g^{2}}{D_{0}} \left(2\tau - \frac{581}{840} \frac{R^{2}}{D_{0}}\right)\right)$$
$$\approx \exp\left(-\frac{4}{175} \left(\frac{l_{D}}{l_{g}}\right)^{2} \left(\frac{l_{s}}{l_{g}}\right)^{4}\right)$$
(S5)

where the final approximation drops the $581/840R^2/D_0$ as insignificant. In the long-time limit, decay models for other geometries vary from Eq. S5 by a scaling within the exponential, e.g., rather than 8/175 for spheres, the scaling factor is 1/120 for parallel plates, and 7/296 for cylinders (12).

As in the long-time limit of the localization regime, decay of signal in the motional averaging regime is exponential with τ , $(l_D/l_g)^2$ or $(bD_0)^{1/3}$. Exchange also occurs on the timescale of τ and can be incorporated into the decay model by multiplying by $\exp(-2\tau AXR)$ where AXR is the apparent exchange rate (13).

The effect of motional averaging can be reached in the extreme case of when δ approaches Δ in pulsed gradient measurements, which researchers have commented leads to restrictions appearing smaller than they actually are (14–16). Given the gradient strength limitations on human MRI scanners, the clinical translation of advanced diffusion MRI methods requires the use of gradient pulses with maximum amplitude for efficient diffusion encoding (17). Consequently, clinical implementations of many advanced diffusion MRI preparations can be adjusted to effectively resemble/be equivalent to experiments in a static gradient field.

Decay models for the various regimes are compared in Fig. 6 of the main manuscript.

Supplementary results

Penetration timescales. The methods were used to follow penetration of deuterated water D_2O into the tissue as well as the delipidation of the tissue via Triton X. For the D_2O wash (Fig. S2), I_0 decreased to 0.36 two minutes after washing, indicating that water in the tissue communicates with the aCSF on timescales of minutes. Exchange and restricted fractions decreased during delipidation on the timescale of roughly one day (Fig. S3). Mass transport theory estimates the timescale to equilibrate a concentration gradient across the tissue as ≈ 1 min for water and ≈ 3 hrs for Triton X based on $t = r^2/4D$ (18) with specimen radius r = 0.7 mm and measured $D = 2.15 \times 10^{-9}$ m²/s for water and $D = 1.3 \times 10^{-11}$ m²/s for Triton X (see supplementary Fig. S4). The timescale of water penetration was consistent with mass transport theory (18), but the timescale of Triton X benetration and delipidation was longer than predicted. An increased time is expected due to the reaction front which develops as Triton X delipidates, slowing its overall penetration.

Testing the rapid exchange measurement on fixed spinal cord tissue. Given that this is the first time the rapid exchange measurement has been used with a static gradient as well as on anything other than an ideal phantom, a full characterization of the signal seemed necessary. Details of the rapid exchange measurement method can be found in Cai, et al., 2018 (19), and the supplementary methods section (below). In this section we test the behavior of the signal acquired as a function of b_d , b_s , and t_m on fixed spinal cord and compare results to the predicted exchange behavior (19).

The curvature along slices of $b_s = 4500 \text{ s/mm}^2$ as a function of b_d at different mixing times is shown in Fig. S9. The signal is concave up with maximum at $b_d = \pm 4500$, minimum at $b_d = 0$, and roughly symmetric about $b_d = 0$, as expected. Exchange increases with mixing time, also as expected. From this, we can conclude that the 4-point method (discussed below) can capture the exchange with maximal sensitivity.

The optimal b_s maximizes the finite difference (Eq. S8) and provides optimal sensitivity to exchange in the presence of noise. However, different b_s values may diffusion-weight the measurement towards exchange between different pools. Fig. S10 shows that the finite difference reaches a maximum near $b_s = 6000 \text{ s/mm}^2$. The value $b_d = 4500 \text{ s/mm}^2$ used in this study is thus near the optimum. Additionally, from this data it was found that the exchange rate is not significantly different between b_s values, indicating that AXR is not very sensitive to b_s .

The four-point method allowed for high temporal resolution of exchange. Fig. S11 shows a dense sampling of f as a function of t_m , acquired overnight on a fixed spinal cord specimen. The data shows the $t_m = 0.1$ ms point to be not well behaved, with f decreasing from $t_m = 0.1$ to 0.2 ms, but increases from $t_m = 0.2$ ms onwards, indicating that $t_m = 0.2$ is a good point for the minimum t_m . The exchange plateaus near $t_m = 100$ ms and stays roughly constant to $t_m = 300$ ms, indicating that $t_m = 300$ ms is a good point for the maximum t_m because it captures the maximum, steady state exchange, and it does not show does not show T_1 relaxation effects (in particular due to differences in T_1 between exchanging pools (19)). The data was fit with the first-order rate model, Eq. S12, and the value (presented in the figure caption) was not statistically different from the less-dense sampling used throughout the paper. The dense sampling shows that the data is not fully explained by the first order rate model; it rises up quicker and plateaus slower. However, rather than fitting a model with more parameters, e.g. a

model with two AXRs, we choose to stick to the first order rate model with one AXR. The extimated AXR = $113.7 \pm 5.2 \ s^{-1}$ is not statistically different from the values measured from the AXR measured from the standard $11 \ t_m$ point protocol used throughout the text, indicating that the 11 point protocol does not bias the measured AXR. The dense sampling also iterates $f(t_m = 0) \neq 0$, thus calling for a non-zero initial condition (discussed below).

Supplementary Materials and Methods

Test chamber. The bottom portion of the experimental test chamber was made of aluminum to provide good heat conduction to the media. A bored-out rectangular hole with a glass cover slide glued to the bottom held the media, solenoid, and spinal cord. Aluminum parts that contacted aCSF were coated with a thin layer of RTV silicone to avoid corrosion. The top of the chamber was made from poly(methyl methacrylate) (PMMA) with two inlets for inflow and outflow of the gas and a hole for the PicoM fiber optic temperature sensor (Opsens Solutions Inc., Québec, Canada). Temperature measurements were not affected by RF and did not induce noise in the RF system. The temperature of the bath was monitored and recorded continuously. The top slide was secured and mounted to the bottom part with four screws. The NMR solenoid coil was glued to the glass cover slip bottom with a hot glue gun. Two separate chambers and solenoid coils were built for live and fixed spinal cord specimen.

NMR hardware. The solenoid radiofrequency (RF) coils and the circuit were built in-house. Solenoids were made from wrapping two concentric layers of AWG 30 copper wire around a #2-56 plastic screw totaling 39 turns resulting in 2 mm inner diameter 4 mm outer diameter, 1.3 cm length. The resulting coils had an inductance $L \sim 600$ nH and impedance $X \sim 52 \Omega$ at 13.79 MHz.

The solenoid connected to a circuit board with detachable pin connectors. (The circuit design and the circuit resonance spectrum when immersed in PBS is shown in Fig. S5.) The circuit used two trimmer capacitors (NMAJ25HV, Knowles Voltronics) with tunable range 1–23 pF for tune and match. The circuit board was connected to the Kea2 spectrometer by a 50 Ω coax cable. The coil matched to -34 dB at 13.79 MHz when immersed in aCSF. RF pulses were driven by a 100 W RF pulse amplifier (Tomco, Adelaide, Australia).

1-D spin echo diffusion. A standard pulse sequence (SEdec in Prospa) was used for measuring diffusion with a static gradient (20) built off of a spin echo followed by a Carr–Purcell–Meiboom–Gill (CPMG) echo train (3, 4, 21), as shown in Fig. S6a. The phase cycle list was four scans long (22).

Signal from the CPMG train is summed up as one data point. This summation provides a significant boost in the signal-to-noise ratio (SNR). Only signal from the real channel (rather than signal magnitude) is taken. This leads to zero-mean Gaussian rather than Rician noise, a significant benefit for multiexponential signal analysis. The echo time of the spin echo τ is incremented in successive loops of the experiment to encode for diffusion. For water and with large gradient the attenuation due to T_2 relaxation during τ is insignificant. Signal can be modeled using Eq. S1 for pure liquids such as water. In heterogeneous materials such as biological tissue, water in different parts of the material experience different hindrances and restrictions. Each sub-ensemble of water molecules has its own effective self-diffusion coefficients. The signal can be modeled as arising from the distribution of effective or apparent self-diffusion coefficients of the water in the different environments using

$$I(b)/I_0 = \int_0^\infty P(D)e^{-bD}dD.$$
 (S6)

Static gradient DEXSY sequence and phase cycles. Attention was paid to phase cycles for the static gradient DEXSY sequence (Figure S6) due to each RF pulse being imperfect and exciting multiple coherence pathways when the inhomogeneity of the magnetic field is greater than the bandwidth of the RF pulses (23). The phase cycle list was eight scans long, shown in Table S1.

Although the phase cycle list is not exhaustive, we found the signal to be well-behaved on a non-exchanging two-pool system comprised of a capillary filled with polydymethylsyloxane bathed in water. Signal as a function of b_1 or b_2 were symmetric and decayed the same as signal as a function of b from the 1-D SEdec sequence. Additionally, signal was flat along slices of constant b_s and the 2-D DEXSY map showed two diffusion coefficients along the $D_1 = D_2$ diagonal equal to D_{water} and D_{PDMS} . One exception was that the phase cycles let through signals which do not form a gradient echo when they see the storage pulse, but do form a gradient echo upon acquisition, thus seeing the sequence as a 1-D stimulated echo diffusion. This became apparent due to additional refocussing when $b_1 = b_2$. This was found to be an issue with a previous miniature flat RF coil design (24) but went away when switching to the solenoid RF coil, thus it is likely an issue of B₁ inhomogeneity of the miniature flat RF coil (25). To avoid refocusing this signal, points were never acquired exactly on $b_1 = b_2$. Also note that the static gradient acts as a crusher during the storage interval. The sequence selects both compensated and uncompensated signals (26). The sequence can be compared and contrasted to another DEXSY sequence developed for static gradients but using stimulated echoes for diffusion encoding (27)

Fig. S6b shows the static gradient spin echo DEXSY sequence. In this sequence, molecules are encoded for their diffusion coefficient in their local environment during the first interval τ_1 . Magnetization is then stored for a mixing time, t_m during

which time molecules move freely and may exchange between diffusive environments. (Note that this definition of t_m is like the definition used for T_2-T_2 (28), and is different from the original definition presented by Callaghan and Furó which included the gradient pulse duration (29). We choose not to use that definition since τ is changing in a DEXSY measurement whereas the definition of t_m should require that it be constant throughout the whole experiment.) Molecules are again encoded for diffusion during τ_2 and then signal is acquired in a CPMG train.

Full DEXSY. The classic way to analyze the full 2-D DEXSY is as a diffusion exchange distribution, related to the signal through

$$I(b_1, b_2) = \int_0^\infty \int_0^\infty P(D_1, D_2) e^{-b_1 D_1 - b_2 D_2} dD_1 dD_2,$$
(87)

a 2-D version of Eq. S6. The diffusion encoding variables, b_1 and b_2 are varied by independently varying τ_1 and τ_2 . Molecules which do not exchange between environments will have the same diffusion coefficient during τ_1 and τ_2 contributing to populations on the diagonal of the 2-D distribution (see, e.g., Fig. 2a and b in (19)). Molecules which do exchange and thus are encoded with different diffusion coefficients between the two τ will contribute to off-diagonal exchange peaks in the 2-D distribution.

Rapid exchange measurement. Alternatively to the full DEXSY, we recently introduced a rapid method for measuring exchanging fractions which relied on curvature of the raw signal after a variable transformation (30). In particular, this method shows that exchange between diffusion environments results in the raw data being curved up along a slice of constant $b_s = b_1 + b_2$ (see Fig. 2c and d in (19)). The exchanging fraction scales with $(\partial^2 I/\partial b_d^2)$ where $b_d = b_2 - b_1$. The second derivative can be approximated with the 2nd order finite difference method,

$$\frac{\partial^2 I}{\partial b_d^2}\Big|_{b_d=b} \approx \frac{I\Big|_{b_d=b-\Delta b_d} - 2I\Big|_{b_d=b} + I\Big|_{b_d=b+\Delta b_d}}{\Delta b_d^2},$$
(S8)

omitting higher order terms. The greatest sensitivity to exchange is when the central point is acquired at $b_d = 0$ and the edges are acquired at $b_d = \pm b_s$. Normalizing by a datapoint acquired with no diffusion weighting $b_s = 0$ removes relaxation effects. This is a relative measure of exchange and is enough to provide image contrast in MRI, to look at time-varying processes, or to measure exchange rates (discussed below).

In our previous publication we developed the theory for obtaining the exchanging fraction, f, from $(\partial^2 I/\partial b_d^2)$, which for a two-site exchange model results in (19)

$$f = \left(\frac{\partial^2 I}{\partial b_d^2}\Big|_{b_d=b}\right) \frac{e^{b_s D_s}}{\cosh(b_d D_d) D_d^2}$$

= $\left(\frac{\partial^2 I}{\partial b_d^2}\Big|_{b_d=0}\right) \frac{e^{b_s D_s}}{D_d^2}$ (S9)

where

$$D_s = \frac{(D_e + D_i)}{2}; \ D_d = \frac{(D_e - D_i)}{2}.$$
 (S10)

Fitting exchange parameters. Exchange rates can be estimated from the full DEXSY or the rapid measurement by repeating the measurement with multiple mixing times which span the exchange process. Exchange parameters can be determined assuming exchange between diffusion environments is governed by a first-order rate law of the form $df_{i,e}/dt = k_{i,e}f_{i,i} - k_{e,i}f_{e,i}$ with rate constants $k_{i,e}$ and $k_{e,i}$ (28). The data we present calls for a nonzero initial condition; $f_{i,e}(t_m = 0) = f_{i,e,0}$ (discussed below). The resulting two-site exchange model is:

$$f_{i,e}(t) = f_{e,i}(t) = \frac{f(t)}{2} = \left(\frac{f_e k_{e,i}}{k_{e,i} + k_{i,e}} - f_{e,i0}\right) \left(1 - e^{-(k_{i,e} + k_{e,i})t}\right) + f_{e,i0}$$

$$= \left(\frac{f_i k_{i,e}}{k_{i,e} + k_{e,i}} - f_{i,e0}\right) \left(1 - e^{-(k_{e,i} + k_{i,e})t}\right) + f_{i,e0},$$
(S11)

with equilibrium fractions f_e and f_i . With either the full DEXSY or the rapid measurement, exchanging fractions as a function of mixing time can be fit with a 3-parameter model of the form

$$f(t_m) = (f_{SS} - f_0) \left[1 - e^{-kt_m}\right] + f_0$$
(S12)

to estimate the initial exchange fraction $f_0 = 2f_{i,e0}$, the steady-state exchange fraction $f_{SS} = \frac{2f_i k_{i,e}}{k_{i,e}+k_{e,i}}$, and the characteristic exchange rate $k = k_{i,e} + k_{e,i}$ (called the apparent exchange rate, AXR, in the text).

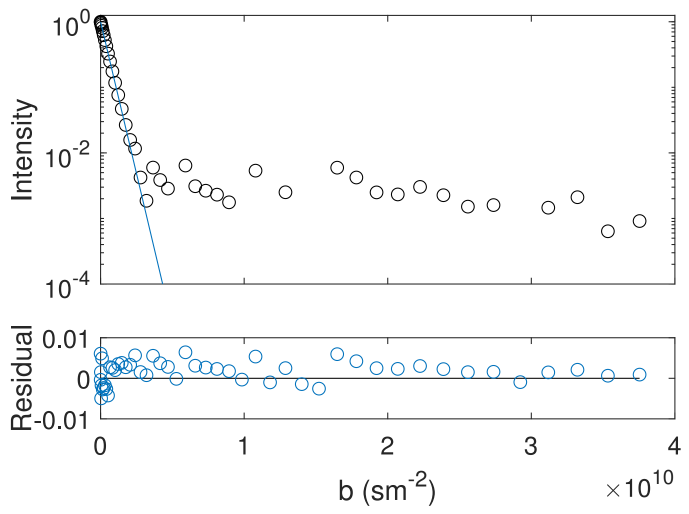


Fig. S1. 1-D diffusion data, single exponential fit and residuals for aCSF at 25C, 50 points. A fit of the first 15 points, attenuating the signal to $I/I_0 = 0.1$, provides $D_0 = 2.153 \pm 0.014$ Table S1. $D_0 = 2.153$

$\overline{\phi_1}$	ϕ_2	ϕ_3	ϕ_4	ϕ_5	ϕ_6	ϕ_{rec}
0	$+\pi/2$	0	0	$+\pi/2$	$\pi/2$	π
π	$-\pi/2$	0	0	$+\pi/2$	$\pi/2$	0
0	$+\pi/2$	π	0	$+\pi/2$	$\pi/2$	0
π	$-\pi/2$	π	0	$+\pi/2$	$\pi/2$	π
0	$+\pi/2$	0	π	$-\pi/2$	$\pi/2$	0
π	$-\pi/2$	0	π	$-\pi/2$	$\pi/2$	π
0	$+\pi/2$	π	π	$-\pi/2$	$\pi/2$	π
π	$-\pi/2$	π	π	$-\pi/2$	$\pi/2$	0

The data we present calls for a nonzero initial condition. A previous study of T_2-T_2 exchange in a polymer–solvent system near the glass transition also observed finite exchange when $t_m \approx 0$ (31). Models have shown significant exchange during the encoding periods τ can lead to exchange peaks at $t_m = 0$ (32).

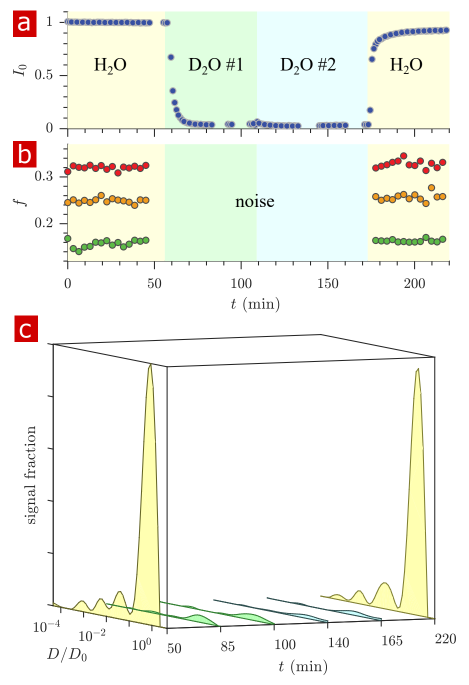


Fig. S2. Timecourse study of D_2O wash, in which the sample was washed in two steps from aCSF to aCSF made with deuterated water, showing a) the proton signal intensity from the rapid exchange measurement data normalized to remove T_1 effects at different mixing times and b) exchanging fractions from rapid measurements with $t_m = 0.2$ (blue), 4 (red), and 20 ms (orange). 1-D diffusion measurements were performed at points throughout the time-course (seen as breaks in the data in (a) and (b)) and distributions are presented as signal fractions ($P(D) \times I_0$) in (c). All distribution peaks decreased after washing with D_2O indicating that the distribution is indicative of water mobility, with no single component being solely from non-water molecules.

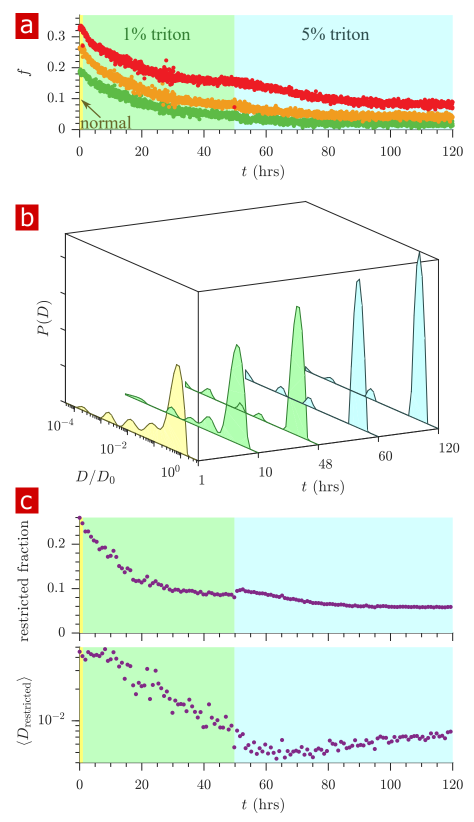


Fig. S3. Timecourse study of Triton X delipidation, in which the sample was washed to aCSF with 1% Triton X, and then 5% Triton X showing a) exchanging fractions from rapid measurements with $t_m = 0.2$ (green), 4 (orange), and 20 ms (red), b) representative distributions throughout the timecourse, and c) the fraction and mean diffusion coefficient of signal arising from restricted water ($\langle D_{\text{restricted}} \rangle$ with $D/D_0 < 0.17$). A bump in the restricted fraction seen upon addition of the 5% Triton is due to the Triton X itself which forms 5 nm micelles (33). The loss of the restricted diffusion peaks and the decrease in the exchanging fractions shows that membranes are the cause of water restriction.

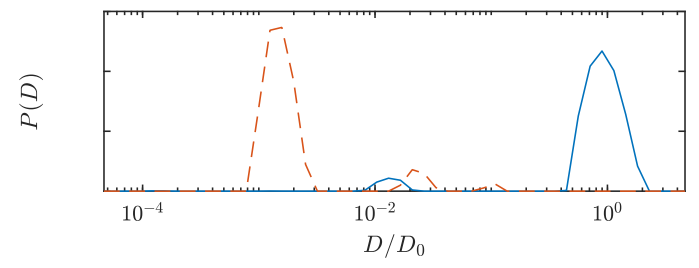


Fig. S4. Diffusion coefficient distributions of 5% Triton X in aCSF (solid blue line) and pure triton (dashed red line). The standard diffusion measurement protocol (43 pts. $\tau = 0.05 \rightarrow 6.6$) was used. The 5% Triton X shows an additional peak at $D/D_0 = 0.013$ due to formation of 5 nm micelles (33). Pure Triton X shows a major diffusion coefficient component at $D/D_0 = 0.001$.

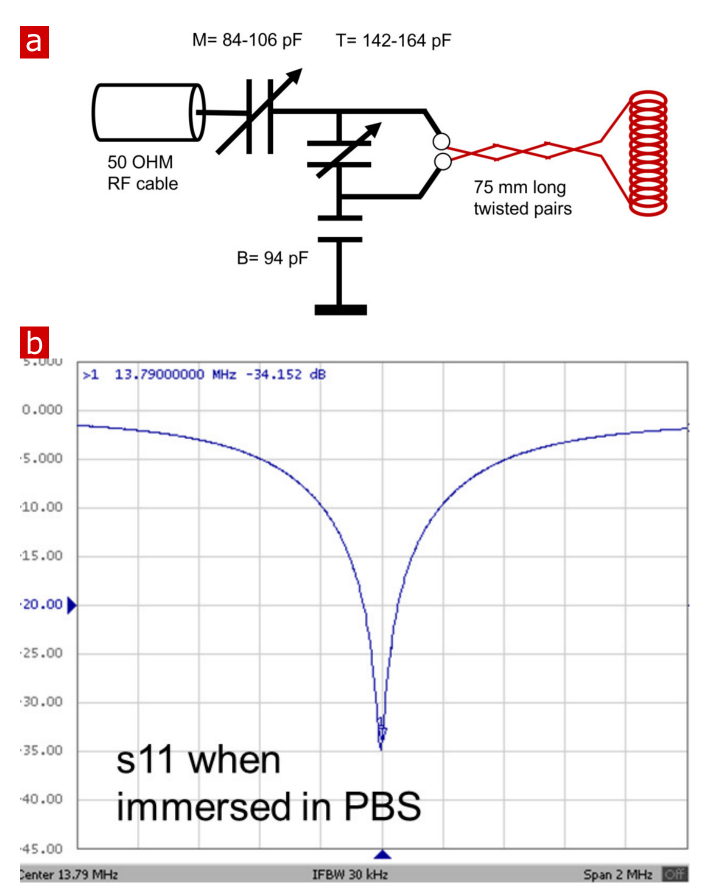


Fig. S5. a) Drawing of the circuit for the RF showing the capacitance of the tune (T), match (M), and balance (B). Note a single wrapping of the solenoid was drawn rather than the actual double-wrap for visual simplicity. b) Circuit resonance spectrum when immersed in PBS and after tuning and matching to 13.79 MHz.

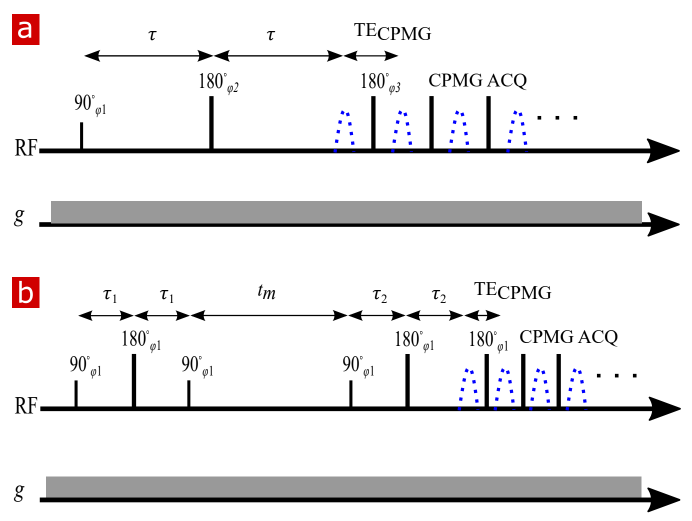


Fig. S6. (a) Spin echo pulse sequence for measuring diffusion with a static gradient. τ is varied to control *b*. (b) DEXSY pulse sequence for measuring exchange with a static gradient. The two SE encoding blocks with τ_1 and τ_2 , varied independently to control b_1 and b_2 , are separated by t_m . Signal is acquired in a CPMG train for both (a) and (b).

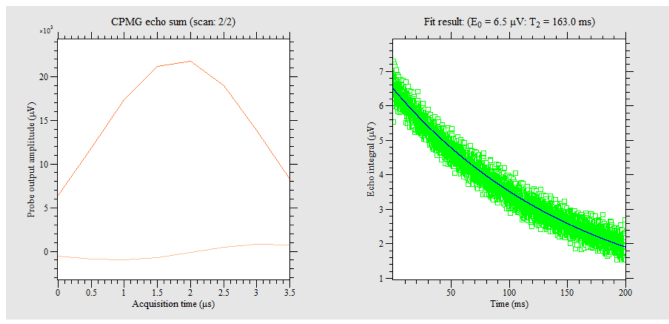


Fig. S7. representative Carr-Purcell-Meiboom-Gill (CPMG) T_2 measurement (10 s repetition time (TR), 8000 echoes) on fixed spinal cord showing the echo shape (real signal phased maximum and imaginary signal phased to zero) on the left and real signal decay and exponential fit on the right. Figures are output from the Prospa (Magritek, New Zealand) software.

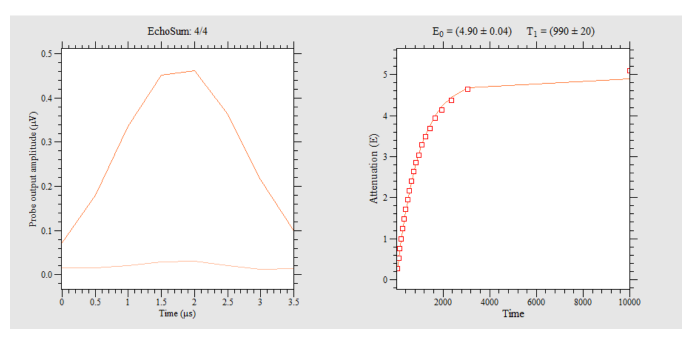


Fig. S8. representative saturation recovery T_1 measurement (1s TR, 21 recovery time pts. logarithmically spaced from 50 ms to 10s) on fixed spinal cord showing the echo shape (real signal phased maximum and imaginary signal phased to zero) on the left and real signal recovery and exponential fit on the right. Figures are output from the Prospa (Magritek, New Zealand) software.

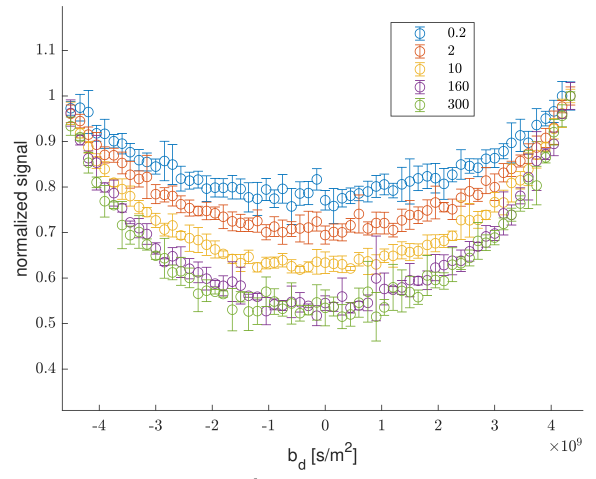


Fig. S9. The curvature along slices of $b_s = 4500 \text{ s/mm}^2$ as a function of b_d highlighting the increase in the depth of the curvature with increasing t_m , shown in figure legend. The increased depth is due to increased exchange (19, 30)

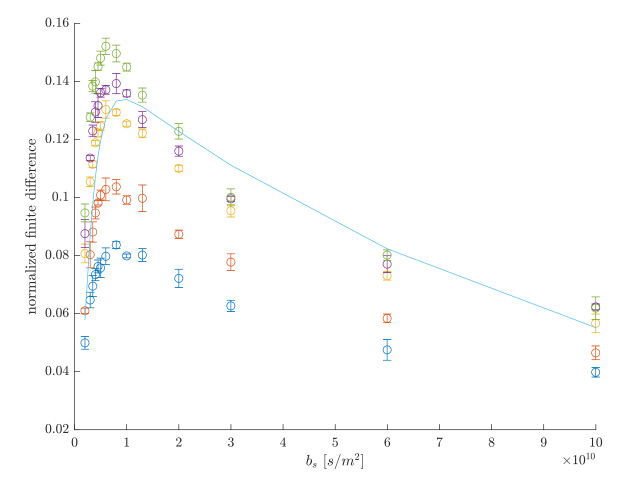


Fig. S10. Difference between $2I_{bd=0}$ and $I_{bd=+bs} + I_{bd=-bs}$, normalized by $I_{0,0}$, a function of b_s , showing the optimal b_s for measuring the largest curvature response as occurring $b_s = 6000 \text{ s/mm}^2$. The line is a prediction of the finite difference for a two-site system, from Eq. 8 in Ref. (19) using f = 0.15, $D_e = 10^{-9}$, and $D_i = 10^{-11} \text{ s/mm}^2$.

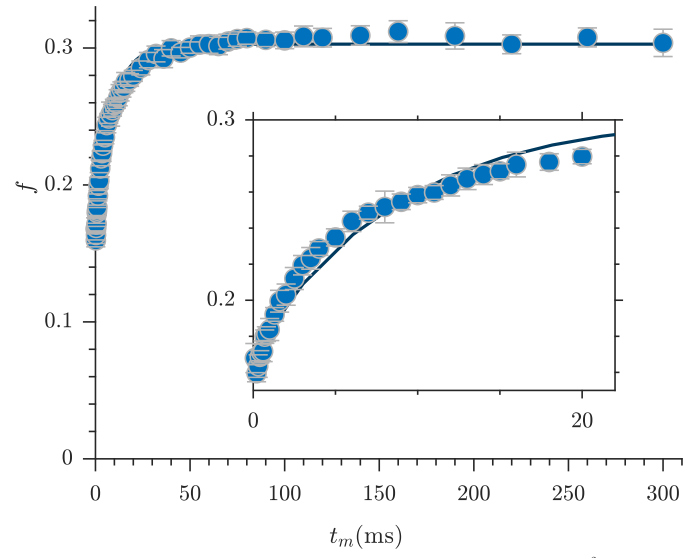


Fig. S11. High temporal resolution exchange rate measurement from the rapid measurement with $b_s = 4500 \text{ s/mm}^2$ and 53 t_m points between 0.1 and 300 ms. First order rate model fits estimate AXR=113.7 ± 5.2 s⁻¹ (solid line).

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