Measuring surface-area-to-volume ratios in soft porous materials using laser-polarized xenon interphase exchange NMR

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Abstract

We demonstrate a minimally invasive nuclear magnetic resonance (NMR) technique that enables determination of the surface-area-to-volume ratio ($S/V$) of soft porous materials from measurements of the diffusive exchange of laser-polarized $^{129}$Xe between gas in the pore space and $^{129}$Xe dissolved in the solid phase. We apply this NMR technique to porous polymer samples and find approximate agreement with destructive stereological measurements of $S/V$ obtained with optical confocal microscopy. Potential applications of laser-polarized xenon interphase exchange NMR include measurements of in vivo lung function in humans and characterization of gas chromatography columns.

81.05.Rm, 82.56.Lz, 68.47.Mn
Porous media are ubiquitous in nature, e.g., granular materials, foams, ceramics, oil- or water-bearing “reservoir” rocks, and animal and human lungs. Determining the structure of these materials is relevant to a wide range of scientific and technological problems, ranging from the coarsening of foams to the transport properties of subsurface fluids to cardiopulmonary physiology and medicine. In this paper we demonstrate a noninvasive technique to characterize the surface-area-to-volume ratio (S/V) of “soft” porous media: i.e., materials in which there is significant gas solubility in the solid phase. Such a description applies to many different materials, including porous polymer granulates used for filtering, porous polymer bead packs used for radio-immunoassay, resin columns used for chromatographic separation, and lung and sinus tissue in humans and animals.

S/V influences numerous interactions within porous media, including fluid diffusion and transport, electric charge distribution, and chemical exchange. For example, S/V is important in determining pulmonary function since the lung is the site of O$_2$ and CO$_2$ exchange between the body and the external environment. There are numerous techniques to measure S/V in porous media, including stereology with traditional and confocal microscopy, light scattering, and mercury intrusion porosimetry. However, only nuclear magnetic resonance (NMR) provides a noninvasive and nondestructive technique for determining 3D structure, including S/V.

In particular, NMR measurements of the time-dependent diffusion coefficient of a liquid filling the pore space has been used to determine S/V in a variety of porous media including glass beads, reservoir rocks, and biological samples. Recently, we extended this technique to NMR measurements of xenon gas diffusion within porous media, thereby probing longer length scales than with traditional liquid-infused diffusion measurements because of fast gas diffusion and long xenon spin polarization lifetimes. Gas diffusion NMR may also be used to study porous media that are not compatible with water-saturation, such as non-wetting polymers and in vivo lungs. However, for materials with pore size less than about 300 µm (such as lung alveoli), the high diffusivity of gases causes systematic errors in the determination of S/V because the rms diffusion distance
during the application of RF and magnetic field gradient pulses is comparable to the mean pore size \[12-14\]. Reducing this effect with faster gradient pulses, buffer gases, or higher gas pressures is impractical \[15\].

Thus we have devised an alternative NMR method (“xenon interphase exchange NMR”) to measure \(S/V\) in soft (i.e., xenon-soluble) porous media. In this technique we determine \(S/V\) from the rate of \(^{129}\text{Xe}\) polarization (not chemical) exchange between gas in the pore space and \(^{129}\text{Xe}\) dissolved in the solid. First, we infuse the sample with laser-polarized (i.e., highly spin-polarized or hyperpolarized) \(^{129}\text{Xe}\) gas. Next, we rapidly destroy (“quench”) all dissolved-phase \(^{129}\text{Xe}\) polarization using a suitable sequence of RF and magnetic field gradient pulses. The result is an initial condition of NMR-labeled (i.e., polarized) \(^{129}\text{Xe}\) in the gas phase with unpolarized \(^{129}\text{Xe}\) in the dissolved phase. We then measure the diffusion of \(^{129}\text{Xe}\) polarization into the dissolved phase by monitoring the recovery of the NMR signal at the chemically-shifted frequency of dissolved-phase \(^{129}\text{Xe}\). The rate of increase of the dissolved-phase \(^{129}\text{Xe}\) NMR signal is proportional to \(S/V\) (see analysis below).

Fast, high-resolution NMR measurements of the nuclear spin-\(\frac{1}{2}\) noble gases \(^3\text{He}\) and \(^{129}\text{Xe}\) are possible with optical pumping techniques using high-power lasers which increase the noble gas spin polarization to \(\sim 1 - 10\%\) \[13\]. Laser-polarized noble gas NMR is a powerful technique in the physical and biomedical sciences, e.g., in lung imaging \[17\]. For interphase exchange measurements, \(^{129}\text{Xe}\) is used rather than \(^3\text{He}\) because of two factors. First, \(^{129}\text{Xe}\) dissolves about ten times more readily than \(^3\text{He}\) in lipophilic compounds: e.g., \(^{129}\text{Xe}\) has a large dissolved/gas partition coefficient of \(\sim 0.25\) for lung tissue \[18\]. Second, the gas and dissolved-phase \(^{129}\text{Xe}\) NMR frequencies are typically separated (chemically-shifted) by \(\sim 200\) ppm, a relatively large value (the typical \(^3\text{He}\) chemical shift is less than 1 ppm) which makes it easy to manipulate or observe \(^{129}\text{Xe}\) spins selectively in either phase \[19\].

Xenon interphase exchange NMR may be particularly important for the human lung where the mean pore (alveolar) size is \(\sim 100\) \(\mu\text{m}\) \[3,4\]. A noninvasive method to measure and image \(S/V\) and gas exchange in the human lung could be an important tool for probing lung physiology and pathophysiology \[20\]; e.g., in the diagnosis and treatment of asthma.
and chronic obstructive pulmonary disease (COPD), the fourth leading cause of death in the U.S [21]. Traditional lung function tests of gas exchange parameters are seriously limited by the inability to provoke step changes in gas composition or properties deep in the lung. Deconvolving the effects of ventilatory dilution, tissue absorption, and blood perfusion is a major challenge. By contrast, xenon interphase exchange NMR provides an almost ideal step change in $^{129}$Xe polarization between tissue and gas, and thus offers a unique window into lung structure and function. For example, subsequent to our announcement of preliminary results using xenon interphase exchange NMR [22], Ruppert et. al. successfully applied a similar technique to studies of lung function in dogs [23]. Xenon interphase exchange NMR may also be a powerful tool in the characterization of non-living, soft porous media. For example, knowledge and control of $S/V$ in packed columns of polymeric beads - commonly used in molecular fractionation applications - is critical to achieving adequate molecular separation.

In the study reported here, we monitored the exchange of $^{129}$Xe polarization from the gas phase into the dissolved phase in a xenon-soluble porous polymer granulate (POREX Corp., Atlanta, GA), chosen as a model (with respect to mean pore size) for lung tissue. We investigated four different samples, which the manufacturer stated as having median pore sizes of 20, 70, 120, and 250 $\mu$m (obtained with mercury intrusion porosimetry on random production samples) and similar porosities of $\sim 30\%$. The interphase exchange experimental protocol was as follows. First, we filled a glass container completely with a porous polymer sample, excluding free gas outside the cylindrical polymer core. Next, we placed the sample in an NMR magnet, pumped out the air from the sample’s pore space, and allowed laser-polarized xenon gas to fill the space. (NMR imaging showed that the xenon gas uniformly filled the pore space within a few seconds.) We then used a frequency-selective RF pulse to quench the dissolved-phase $^{129}$Xe polarization, and observed the recovery of the dissolved-phase $^{129}$Xe NMR signal due to interphase exchange for times from 10 to 1000 ms. The NMR pulse sequence for the interphase exchange measurement is shown schematically in Fig. [4]. Note that we made an initial measure of the gas-phase $^{129}$Xe NMR signal using an RF pulse.
with a small flip angle $\alpha \sim 3 - 5^\circ$. Also, the selective RF pulse at the dissolved-phase $^{129}$Xe NMR frequency was not perfect and hence generated a small NMR signal from $^{129}$Xe gas in the pores. We exploited these gas-phase signals to correct the data for slowly decreasing $^{129}$Xe polarization due to both RF depletion and $T_1$ relaxation in the pores. An example of the observed interphase exchange spectra is shown in Fig. 2. We performed xenon interphase exchange experiments on two NMR spectrometers at different magnetic field strengths: a horizontal-bore GE Omega/CSI spectrometer operating at 4.7 T (55.3 MHz $^{129}$Xe NMR frequency) with field gradient strength up to 7 G/cm; and a modified SMIS spectrometer interfaced to a prototype IBM horizontal-bore magnet operating at 1.5 T (17.6 MHz $^{129}$Xe NMR frequency) with field gradient strength up to 2 G/cm. For each experimental run, we used the spin-exchange optical pumping technique \cite{16} in the fringe field of the NMR magnet and achieved $\sim 0.5\%$ $^{129}$Xe spin polarization for 2 bar-liters of xenon gas (natural isotopic abundance).

Calculation of absolute values for $S/V$ required prior characterization of the samples. We used thermally-polarized xenon NMR on both spectrometers to determine the porosity $\phi$ and $^{129}$Xe partition coefficient $b$ for the four porous polymer samples. We placed all 4 sample cells on a manifold along with an empty cell of equal volume, admitted $\sim 3$ bar of xenon gas (natural isotopic abundance) and $\sim 2$ bar of oxygen, and allowed the system to equilibrate over 2 - 3 days to achieve full penetration of xenon into the polymer samples. We then measured the $^{129}$Xe NMR gas signal in the empty cell ($s_{g0}$) as well as the equilibrium gas and dissolved-phase $^{129}$Xe NMR signals ($s_{gas}$, $s_{diss}$) in the four polymer samples. For each sample, $\phi = s_{gas}/s_{g0}$ and $b = (s_{diss}/s_{gas})\phi/(1 - \phi)$. We obtained similar results for $\phi$ and $b$ at 4.7 T and 1.5 T. Measurement of the dissolved-phase $^{129}$Xe diffusion coefficient $D_{diss}$ required large magnetic field gradients, which was only possible across small sample dimensions. Hence we cut additional pieces of each polymer sample to fit in standard 5 mm diameter NMR spectroscopy tubes and filled the tubes with $\sim 10$ bar xenon gas (isotopically-enriched to 90$\%^{129}$Xe). We then measured $D_{diss}$ using a 9.4 T Bruker AMX2 spectrometer with 50 G/cm field gradients, employing a modified pulsed-gradient technique previously
applied to gas-phase diffusion measurements on rocks and glass beads [12]. Table I lists the measured values of $\phi$, $b$, and $D_{\text{diss}}$, along with the $^{129}$Xe longitudinal spin relaxation times ($T_1$) for each porous polymer sample.

To determine $S/V$ from the xenon interphase exchange NMR measurements, we employed a one-dimensional diffusion model, asymptotically valid for short times. We assumed that the density of dissolved-phase $^{129}$Xe polarization, $\rho_{\text{diss}}(x, t)$, satisfies the diffusion equation on a half space, $x > 0$, where the non-trivial spatial coordinate $x$ is normal to the gas/solid interface. The quench of the dissolved-phase $^{129}$Xe polarization created the initial condition $\rho_{\text{diss}}(x, 0) = 0$, $x > 0$. Given the much greater gas-phase $^{129}$Xe density, we also assumed a fixed boundary condition $\rho_{\text{diss}}(0, t) = b\rho_{\text{gas}}$. Here, $\rho_{\text{gas}}$ is the density of gas-phase $^{129}$Xe polarization, which we found to be constant during an individual interphase exchange measurement except for the small effects of RF depletion and slow $T_1$ relaxation mentioned above. The solution of the diffusion equation for $\rho_{\text{diss}}(x, t)$ is given by the well known error function [24]. The number of dissolved-phase polarized $^{129}$Xe atoms $N_{\text{diss}}(t)$ can then be obtained by direct integration of the density or by solving the ordinary differential equation obtained by integrating the diffusion equation. The result is:

$$\frac{N_{\text{diss}}(t)}{V} = \rho_{\text{gas}} b \frac{S}{V} \sqrt{4D_{\text{diss}} t / \pi}$$  \hspace{1cm} (1)

where $S$ is the total pore-space surface area of the sample, and the Euclidean volume $V$ of the sample has been introduced as a common divisor on both sides of the equation. Noting that

$$\rho_{\text{gas}} = \frac{N_{\text{gas}}}{\phi V}$$  \hspace{1cm} (2)

where $N_{\text{gas}}$ is the number of gas-phase polarized $^{129}$Xe atoms (assumed constant except for RF depletion and $T_1$ relaxation), Eq. (1) can be rewritten as

$$\frac{N_{\text{diss}}(t)}{N_{\text{gas}}} = \frac{bS}{\phi V} \sqrt{4D_{\text{diss}} t / \pi}.$$  \hspace{1cm} (3)

$N_{\text{diss}}(t)$ and $N_{\text{gas}}$ are proportional to the frequency integrals over the observed $^{129}$Xe NMR
spectral peaks, $s_{\text{diss}}(t, \omega)$ and $s_{\text{gas}}(\omega)$, for the dissolved and gas phases respectively. Thus we have

$$I(t) = \frac{\int s_{\text{diss}}(t, \omega) d\omega}{\int s_{\text{gas}}(\omega) d\omega} = \frac{bS}{\phi V} \sqrt{4D_{\text{diss}}t/\pi}.$$ (4)

We determined $S/V$ by fitting the ratio of dissolved and gas-phase $^{129}$Xe NMR signals to $\sqrt{t}$ and using measured values for $b, \phi$ and $D_{\text{diss}}$

Concurrent with the NMR studies, we also characterized the four samples of polymer granulate using confocal microscopy in order to give independent measurements of $S/V$. We carefully sliced sections of each of the four polymer rods with different pore sizes, creating glass smooth surfaces of both the longitudinal and cross-sectional faces of the rods. We oriented these sectional samples on a Leica TCS NT confocal microscope configured for 488 nm excitation and reflected light imaging. The resulting microscope images showed the polymer granulates as solids, and the pore spaces as black voids. Computer-based image analysis (ImageSpace, Molecular Dynamics, Sunnyvale, CA) superimposed multiple, equally-spaced lines across the images, and interface boundaries on each line of the images were counted. Thus for each sample we determined the mean linear intercept, $L_m$, which is defined as the average distance between phase boundaries in an image. While $L_m$ is but one of several possible measures of pore size, it has the unique and powerful advantage of being stereologically related to the surface to volume ratio. In particular, $L_m$ is an unbiased estimator of $S/V$ in the image plane: $L_m = 2(S/V)^{-1}$. See Table II for the optically-determined values of $L_m$ and $S/V$.

We acquired three xenon interphase exchange data sets for each of the four different pore size polymer samples: two data sets at 4.7 T and one at 1.5 T. As an example, Fig. 3 shows an interphase exchange data set taken on the 4.7 T instrument. Each of the points plotted in Fig. 3 consists of the normalized average of three or four separate series of exchange spectra of the type shown in Fig. 2. The normalized xenon integral displayed on the ordinate of Fig. 3 is given by $\langle I(t) \rangle \phi \sin \alpha / b$ (see Eq. (4)), where $\alpha$ is the small flip angle of the RF pulse used in the initial gas-phase interrogation. All xenon interphase exchange NMR data
followed the expected $\sqrt{t}$ trend over the observed range of exchange times (out to $\sim 1$ s).

Fig. 4 compares $S/V$ values derived from the optical and NMR techniques for each of the four polymer samples: one optical and three NMR data sets. (See also Table I.) The solid line in Fig. 4 is the line of identity for the optical and NMR values of $S/V$; the dashed line is a line of regression constrained to go through the origin. The slope of the regression line is $1.27 \pm 0.18$, indicating approximate agreement between the optical and NMR measures of $S/V$. Sources of systematic error in the interphase exchange NMR measurement include: miscalibration of the flip angle $\alpha$ of the gas-phase interrogation RF pulse; inaccuracy in the dissolved-phase xenon diffusion coefficient measurements; and deviations from the simple, one-dimensional diffusion model used to derive $S/V$. For example, the very low dissolved-phase xenon NMR signal from the nominally 120 µm POREX sample prevented us from measuring $D_{diss}$ for this sample. Instead, we used the average of the measured $D_{diss}$ values for the nominally 70 µm and 250 µm samples to calculate $S/V$ from the interphase exchange measurements for the 120 µm sample.

We note that the manufacturer’s quoted pore sizes for the four porous polymer samples do not agree with our pooled optical and NMR measurements of $S/V$ (and hence $L_m$); even the rank order does not agree. This discrepancy may not be surprising, since the pore sizes given by the manufacturer were determined by mercury intrusion porosimetry [5,27], which is primarily a probe of minimum pore-throat size and thus not necessarily a good measure of $L_m$ or $S/V$ [28]. We emphasize that there are different choices of the definition of pore size; which choice is most appropriate will depend on the particular application.

In conclusion, we have demonstrated a new technique, xenon interphase exchange NMR, that provides a minimally invasive and nondestructive probe of the surface-area-to-volume ratio ($S/V$) of xenon-soluble porous materials. We have applied this technique to porous polymer samples and found approximate agreement with destructive stereological measurements of $S/V$ made with optical confocal microscopy.

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TABLES

TABLE I. For each porous polymer sample, measured values are listed for the porosity $\phi$, the $^{129}$Xe gas/polymer partition coefficient $b$, and the dissolved-phase $^{129}$Xe diffusion coefficient $D_{\text{diss}}$. Also listed are the $^{129}$Xe longitudinal spin relaxation time (polarization lifetime) in the dissolved phase ($T_{1\text{diss}}$) and the pore gas phase ($T_{1\text{gas}}$). The nominal pore size is that provided by the polymer manufacturer from mercury intrusion porimetry measurements on random production samples. Note: $T_{1\text{diss}}$ was the same at 4.7 T and 1.5 T; also, $D_{\text{diss}}$ was not measured in the nominally 120 $\mu$m sample because the dissolved-phase $^{129}$Xe NMR signal from this sample was too low to allow the experiment to be completed in a reasonable time. Listed measurement uncertainties do not include estimates of systematic errors.

| Nominal pore size ($\mu$m) | 20       | 70       | 120      | 250      |
|---------------------------|---------|---------|---------|---------|
| Porosity, $\phi$          | 0.45 ± 0.02 | 0.45 ± 0.03 | 0.46 ± 0.01 | 0.43 ± 0.02 |
| Xe partition coefficient, $b$ | 0.64 ± 0.02 | 0.52 ± 0.04 | 0.56 ± 0.03 | 0.63 ± 0.03 |
| Xe $D_{\text{diss}}$ ($\times 10^{-12}$ m$^2$s$^{-1}$) | 5.6 ± 0.5 | 7.0 ± 0.7 | n/a | 7.2 ± 0.7 |
| Xe $T_{1\text{diss}}$ (s) at 4.7 T | 8.1 ± 0.4 | 7.7 ± 0.3 | 7.6 ± 0.3 | 8.0 ± 0.3 |
| Xe $T_{1\text{gas}}$ (s) at 1.5 T | 21.8 ± 0.5 | 34.1 ± 0.5 | 35.4 ± 0.4 | 19.9 ± 0.2 |
| Xe $T_{1\text{gas}}$ (s) at 4.7 T | 41.6 ± 0.6 | 68.1 ± 1.4 | 70.0 ± 1.6 | 42.8 ± 0.6 |

TABLE II. The surface-area-to-volume ratios ($S/V$) derived from the optical and NMR measurements are listed for each of the four polymer samples. Also listed is the mean linear intercept, $L_m = 2(S/V)^{-1}$, determined from the optical microscopy measurements. Listed measurement uncertainties do not include estimates of systematic errors.

| Nominal pore size ($\mu$m) | 20       | 70       | 120      | 250      |
|---------------------------|---------|---------|---------|---------|
| $S/V$ from xenon NMR ($\mu$m$^{-1}$) | 0.030 ± 0.006 | 0.016 ± 0.005 | 0.016 ± 0.003 | 0.018 ± 0.003 |
| $S/V$ from microscopy ($\mu$m$^{-1}$) | 0.0229 ± 0.0004 | 0.0135 ± 0.0003 | 0.0122 ± 0.0003 | 0.0127 ± 0.0003 |
| $L_m$ from microscopy ($\mu$m) | 87.3 ± 1.7 | 147.8 ± 3.9 | 164.3 ± 4.5 | 158.0 ± 4.3 |
FIGURES

FIG. 1. Xenon interphase exchange NMR pulse sequence. The sequence can be divided into the following segments. A: interrogation of the gas-phase NMR signal \( s_{\text{gas}} \) with a small flip angle RF pulse; \( s_{\text{gas}} \) is proportional to the \(^{129}\text{Xe} \) polarization. B: semi-selective saturation train of RF pulses to destroy the dissolved-phase \(^{129}\text{Xe} \) polarization. C: interphase exchange time, \( t \). D: semi-selective 90° RF pulse to observe the recovery of the dissolved-phase \(^{129}\text{Xe} \) signal \( s_{\text{diss}}(t) \); this RF pulse also excites a small gas-phase signal, allowing correction for incremental loss of gas-phase magnetization. E: interphase exchange measurement, typically repeated ten times for each exchange time, \( t \).

FIG. 2. Dissolved-phase xenon NMR spectra as a function of interphase exchange time for the nominally 20 \( \mu \text{m} \) pore-size sample. The peaks at 0 ppm arise from the weak, far-off-resonance excitation of gas-phase polarized \(^{129}\text{Xe} \) when dissolved-phase \(^{129}\text{Xe} \) is pulsed semi-selectively. Following saturation, the dissolved-phase \(^{129}\text{Xe} \) peaks at 200 ppm show recovery as a function of exchange time. The observed spectra are scaled to normalize the integrals of each gas peak to the gas peak in the first exchange spectrum; this normalization accounts for the slow loss of gas-phase magnetization due to exchange, RF depletion, and \( T_1 \) relaxation.

FIG. 3. One set of xenon interphase exchange NMR data, acquired at 4.7 T for each of the four porous polymer samples. (The samples are indicated in the figure by the manufacturer’s claimed pore size.) Each data point represents the weighted mean of 3 to 4 normalized spectra of the type illustrated in Fig. 2, acquired in repeated runs of the experiment. The normalized xenon integral plotted on the ordinate is thus given by \( \langle I(t) \rangle \phi \sin \alpha / b \) (see Eq. (4)). Uncertainties for each interphase exchange spectrum were determined from the measured rms noise within the spectral bandwidth, normalized for variations in gas-phase magnetization; one-standard-deviation error bars were then calculated for the weighted means of the normalized xenon integrals. The error bars are larger for longer exchange times because these data were acquired later in each experimental run and hence had lower gas-phase magnetization, which correspondingly increased the normalized noise of the dissolved-phase xenon spectra.
FIG. 4. Comparison of polymer pore-space $S/V$ values obtained with optical microscopy and NMR. For each of the four polymer samples, three NMR experimental runs are plotted versus one optical measurement. The solid line is a line of unity for the $S/V$ values provided by the two measurement techniques. The dashed line indicates a regression between the NMR and optical $S/V$ values, constrained to pass through the origin; this regression line has a slope of $1.27 \pm 0.18$. 
S/V optical ($\mu m^{-1}$)

S/V NMR ($\mu m^{-1}$)

'120' $\mu m$

'250' $\mu m$

'70' $\mu m$

'20' $\mu m$