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Near-unity nuclear polarization with an open-source \textsuperscript{129}Xe hyperpolarizer for NMR and MRI

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The exquisite NMR spectral sensitivity and negligible reactivity of hyperpolarized xenon-129 (HP\textsuperscript{129}Xe) make it attractive for a number of magnetic resonance applications; moreover, HP\textsuperscript{129}Xe embodies an alternative to rare and nonrenewable \textsuperscript{3}He. However, the ability to reliably and inexpensively produce large quantities of HP\textsuperscript{129}Xe with sufficiently high \textsuperscript{129}Xe nuclear spin polarization (\(P_{Xe}\)) remains a significant challenge—particularly at high Xe densities. We present results from our “open-source” large-scale (~1 L/h) \textsuperscript{129}Xe polarizer for clinical, preclinical, and materials NMR and MRI research. Automated and composed mostly of off-the-shelf components, this “hyperpolarizer” is designed to be readily implementable in other laboratories. The device runs with high resonant photon flux (up to 200 W at the RB D\textsubscript{2} line) in the xenon-rich regime (up to 1,800 torr Xe in 500 cc) in either single-batch or stopped-flow mode, negating in part the usual requirement of Xe cryocooling. Excellent agreement is observed among four independent methods used to measure spin polarization. In-cell \(P_{Xe}\) values of ~90%, ~57%, ~50%, and ~30% have been measured for Xe loadings of ~300, ~500, ~760, and ~1,570 torr, respectively. \(P_{Xe}\) values of ~41% and ~28% (with ~760 and ~1,545 torr Xe loadings) have been measured after transfer to Teflar bags and transport to a clinical 3 T scanner for MR imaging, including demonstration of lung MRI with a healthy human subject. Long “in-bag” \textsuperscript{129}Xe polarization decay times have been measured (\(T_1\) ~38 min and ~5.9 h at ~1.5 mT and 3 T, respectively)—more than sufficient for a variety of applications.

Putting the detection sensitivity provided by their high, nonequilibrium nuclear spin polarization, hyperpolarized (HP) noble gases (\textsuperscript{129}Xe and \textsuperscript{3}He) are used in a number of NMR/MRI applications (1). Human lung imaging (e.g., refs. 2-5) with HP\textsuperscript{129}Xe is of particular interest. Moreover, xenon is soluble in blood (6), other tissues (7, 8), and many biologically compatible liquids (9), and its proclivity for interacting with other substances and its wide chemical shift range make HP\textsuperscript{129}Xe a sensitive NMR probe of molecular and material surfaces (1, 10-12). In many applications, HP\textsuperscript{129}Xe can replace \textsuperscript{3}He, and the relative abundance of \textsuperscript{129}Xe can greatly reduce the impact of the worldwide \textsuperscript{3}He shortage (13) in these instances. Despite considerable progress (14-22), a major obstacle toward implementing HP\textsuperscript{129}Xe for clinical imaging has been the ability to reliably and inexpensively produce large quantities of HP\textsuperscript{129}Xe with high polarization (\(P_{Xe}\)). HP\textsuperscript{129}Xe is usually created via spin-exchange optical pumping (SEOP) (23), whereby the unpaired electronic spins of an alkali metal vapor (e.g., Rb) are polarized via optical pumping with circularly polarized light, and the polarization is transferred to noble gas nuclear spins during collisions. It is generally anticipated that high \(P_{Xe}\) is achievable only in the low xenon-density regime (18, 24), because (i) higher Xe densities increase the destruction of Rb polarization from nonspin-conserving collisions at a rate that is orders of magnitude worse than those of other gases like \textsuperscript{3}N\textsubscript{2} and \textsuperscript{3}He (25-27); and (ii) higher total pressures tend to quench the three-body van der Waals contribution to Rb-Xe spin exchange—leaving the less-efficient two-body term (18, 23). Most large-scale polarizers, in particular all that are available commercially, operate in this low-Xe density regime. Applied research with HP\textsuperscript{129}Xe is severely hampered owing to a lack of access to expensive proprietary hyperpolarizers, and to stimulate development of HP\textsuperscript{129}Xe applications, we present here a low-cost, “open-source” design.

In our recent work (22, 28, 29) exploring Rb/Xe SEOP under conditions of high resonant laser flux, an inverse relationship was found between the optimal temperature and the in-cell Xe density. This effect was exploited to achieve surprisingly high \(P_{Xe}\) values at high Xe densities (e.g., ~52%, ~22%, and ~11% at 50, 500, and 2,000 torr Xe in a 75-cc cell) with <30 W of laser power (22). Expanding upon these results by over an order of magnitude, our collaboration’s first-generation large-scale (~1 L/h) \textsuperscript{129}Xe polarizer should enable a variety of clinical, preclinical, and materials magnetic resonance spectroscopy/MRI applications. (Portions of this paper were presented at the 2012 Meeting of the Radiological Society of North America.)

Significance

Lung diseases comprise the third leading cause of death in the United States and could benefit from new imaging modalities. “Hyperpolarized” xenon-129 can overcome the ordinarily weak MRI signals from low-density species in lung space or dissolved in tissue; however, clinical progress has been slowed by the difficulty in preparing large amounts of hyperpolarized xenon with high magnetization, as well as the cost and limited availability of xenon hyperpolarization devices. We describe a unique low-cost “open-source” xenon “hyperpolarizer,” characterize its ability to produce xenon-129 with high magnetization, and demonstrate its utility for human lung imaging.


The authors declare no conflict of interest.

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\textsuperscript{3}For spin-1/2 particles, polarization is defined as follows: \(P = (N_+ - N_-)/(N_+ + N_-)\), where \(N_+\) and \(N_-\) refer to the numbers of spins in the \(m = +1/2\) and \(m = -1/2\) magnetic sublevels; at thermal equilibrium, \(P_{Xe}\) is only ~2.85 x 10\textsuperscript{-4} at 3 T and ~300 K.

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work were presented previously.\textsuperscript{3} Composed of mostly off-the-shelf components, our automated, modular polarizer is portable, easy to use, and employs an open-source design that is readily implementable in other laboratories. Unlike most clinical-scale Xe polarizers, which typically run in continuous-flow mode, our hyperpolarizer runs in either single-batch or stopped-flow mode in the Xe-rich gas regime (with up to 1,800 torr, or >90%, in a 500-cc cell). The design in part negates the usual requirement to cryo-cool the HP\textsuperscript{129}Xe—a process that otherwise increases the complexity of the device and can lead to losses of spin polarization during HP\textsuperscript{129}Xe accumulation, storage, phase transitions (30), and transfer. Four independent methods were used to measure spin polarization, including the following: in situ field-cycled near-IR spectroscopy (28) (to probe Rb electron spin polarization); in situ low-field \textsuperscript{129}Xe NMR (calibrated with thermal \textsuperscript{1}H NMR); gas transfer to 47.5 mT for \textsuperscript{129}Xe NMR (calibrated with thermal \textsuperscript{13}C NMR); and gas transfer and subsequent \textsuperscript{129}Xe NMR/MRI using a clinical 3 T scanner. Excellent agreement was observed among these different approaches; taken together, the \textsuperscript{129}Xe polarization values reported here represent, by a significant margin, the highest yet achieved at such high Xe densities—and establish the feasibility of attaining near-unity polarization in single batches with HP\textsuperscript{129}Xe quantities sufficient for clinical use (demonstrated here with HP\textsuperscript{129}Xe lung ventilation maps from a healthy human volunteer). Long \textsuperscript{129}Xe polarization lifetimes were obtained in Tedlar bags that are more than sufficient for gas-phase storage or accumulation necessary for some applications.

**Methods**

**XeNA Polarizer.** XeNA, self-contained in a mobile chassis, is composed of a laser system, an electromagnetically coated assembly, a vacuum/gas-handling manifold, on-board optical and NMR spectrometers (and other sensors) for quality assurance (QA) and feedback, and a computer/automation system (Fig. 1A). The optical path (Fig. 1B) begins with a 200-W laser diode array (QPC Lasers) that produces tunable, frequency-narrowed (29) laser output at the Rb $D_2$ line ($\sim$794.8 nm; nominal FWHM = 0.27 nm). The laser beam passes through an optical fiber and expands to 2-inch diameter before collimation and entrance into a polarizing beam-splitter (PBS). The s-polarized beam component (<10%) is reflected 90° and discarded; the main (p-polarized) beam is rendered circularly polarized with a quarter-wave (QWP) plate and directed into the oven, which contains a cylindrical glass optical pumping (OP) cell (2” o.d., 9.75”, 500 cc) previously coated with a silanizing agent (Surfasil, Pierce) to slow gas diffusion and loaded with <200 mg of Rb. The cell is loaded with a variable Xe/N$_2$ gas mixture; N$_2$ helps suppress reemission of unpolarized light from electronically excited Rb (14) and provides additional collision-broadening of the Rb absorption line (31). The laser beam enters/exits through 3-inch anti-reflection-coated sapphire windows. The beam is retro-reflected back through the cell via a 3-inch mirror and directed into the optics assembly, where the PBS reflects the beam into a beam block. The Teflon oven’s temperature is controlled via a heated/cooled gas line from a self-pressurized liquid N$_2$ dewar. The oven resides in a magnetic field provided by a four-coil electromagnet (Acutran, 23.6° i.d.; nominal fields of 5.26 or 5.04 mT, corresponding to 62.0-kHz resonance frequency for Xe$^2$ and Xe, respectively). The manifold directs gas flow from cylinders containing natural-abundance Xe (26.44% $^{129}$Xe), isotopically enriched Xe (–86% $^{129}$Xe), and N$_2$ gas, providing desired mixtures up to 2,000 torr. All experiments used naturally abundant $^{129}$Xe, unless otherwise stated. After passing through Q$_0$ getters, the gases are loaded into the OP cell. Following SEOP, the HP\textsuperscript{129}Xe mixture is flowed through perfluoroalkoxy (PFA) tubing and a Teflon filter (Millipore Watagard) to get any residual Rb before collection in a Tedlar bag. The cell’s glass valve has a mechanically actuated Teflon stopcock; the manifold valves are automated pneumatic or solenoid valves, with Teflon surfaces for those downstream of the cell. In situ QA is provided by Ocean Optics HR4000 near-IR and Magritek Kea2 low-field NMR spectrometers. The near-IR spectrometer’s fiber is positioned behind the 3-inch mirror; the NMR spectrometer uses a home-built NMR probe/oven.


$A = -\ln T = -\ln \left( \frac{I_{\text{hot}}}{I_{\text{cold}}} \right)$, \hspace{1cm} [1]

where $A$ and $T$ denote the absorbance and transmittance, respectively, and $I_{\text{hot}}$ and $I_{\text{cold}}$ are the integrated intensities of the transmitted laser spectra obtained when a gas-loaded cell is respectively “hot” and “cold” (i.e., with and without significant Rb vapor present). With some simple assumptions, absorbance values may then be compared under the same conditions—except with or without the SEOP magnetic field ($B_0$) on—to calculate an estimate of $\left< P_{\text{Rb}} \right>$, according to a simple relation derived from Beer’s Law (28):

\hspace{1cm} Fig. 1. The XeNA polarizer. (A) Schematic of the polarizer’s key components (liquid N$_2$ dewar not shown for clarity). The optical path (shown in B) is represented by (“/4”). For the gas cylinders, “H” and “E” designate Xe with naturally abundant $^{129}$Xe and isotopically enriched $^{129}$Xe, respectively.

\hspace{1cm} co. User operation of XeNA employs a GUI programmed in open-source software (processing.org), which drives a microcontroller box that houses the “brain” of the polarizer (Arduino Mega 2560 REV3 microcontroller board), solid-state relays, and a Bürkert manifold enabling pneumatic valve operation. Material costs for the hyperpolarizer were $<125,000.

Other Experimental Aspects. Xenon was purchased from Nova Gases. MR spectra and images were obtained at 47.5 mT (32) or 3-T field using a Magritek Kea2 with permanent magnet (559-kHz $^{129}$Xe frequency and 508-13-kHz $^{13}$C frequency) or a Siemens 3-T clinical MRI scanner (34.09-MHz $^{129}$Xe frequency), respectively. Elemental analysis of Tedlar bag contents (following SEOP runs) was performed by Element One to ensure that <50 ng of Rb was present in the bag after transfer.

### Results

An estimate of the spatial average of the Rb electron spin polarization, $\left< P_{\text{Rb}} \right>$, can be obtained from measurement of the transmitted light of the pump laser while the magnetic field is cycled (28) (Fig. 2). Spectra are recorded from the laser transmitted through the cell to calculate absorbance values:

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where $A_0$ is the absorbance when $B_0 = 0$. Examples of transmission spectra are shown in Fig. 2 for a cell containing 495 and 1,545 torr of Xe and $N_2$, respectively. Multiple near-IR spectra were obtained in rapid succession for each condition, and then averaged and integrated. One set of scans was obtained at 57 °C, ~10 min into the SEOP process; the corresponding values of $P_{RB}$ and $P_{Xe}$ measured with low-field NMR (see below) were 43 ± 9% and 48.3 ± 2.8%, respectively. $P_{RB}$ cannot be higher than $(P_{RB})$ (Eq. 3); however, the difference between these values is less than the uncertainty.

As part of the optimization process, the temperature of the cell was raised to 65 °C, and after several minutes the sequence was repeated. Improved values of $(P_{RB})$ and $P_{Xe}$ of 71 ± 6% and 57 ± 3%, respectively, were obtained.

Xe polarization was monitored directly via NMR under three different sets of conditions (Figs. 3 and 4). First, low-field $^{129}$Xe NMR can be used to measure $P_{Xe}$ within the OP cell. Fig. 3A shows an example of a high-$^{129}$Xe NMR spectrum obtained at 5.25 mT from a cell containing 1,545 torr of Xe following SEOP and cooldown. Low-field NMR is calibrated using a thermal $^1$H reference signal obtained using the same NMR circuit at the same field $^{15}$C reference signal obtained from 170 mmol of sodium $^{13}$C acetate in 99.8% $D_2O$ located in an identical phantom (Fig. 3D). Comparison with in situ 5.26-mT $^{129}$Xe NMR polarimetry taken from the OP cell during the same experiment before transferring the gas to 47.5-mT magnet (68.5 ± 3.9%) indicated no polarization loss within error, and thus a highly efficient $^{129}$Xe transport process. In future designs, we hope to implement the new approach by Saam and coworkers (33) to measure $P_{Xe}$ from corresponding shifts in the optically detected Rb ESR signal.

The hyperpolarizer was delivered to Brigham and Woman’s Hospital (Boston, MA) and installed adjacent to a clinical MRI suite in February 2012. Optimization of the cell cooldown and automated Xe-transfer processes allowed Xe transfer to Taldir bags via expansion and subsequent transport to a 3-T clinical MRI. Inset of Fig. 4A shows a high-field $^{129}$Xe NMR spectrum from a Taldir bag containing —800 cc of gas (38% Xe by volume, given a cell loading of 760 torr Xe/1,240 torr $N_2$). The $P_{Xe}$ value is calibrated using a thermal $^{129}$Xe NMR signal from a 3-L spherical reference sample containing 4 atm Xe with 86% $^{129}$Xe enrichment and 2 atm $O_2$ with a $T_1$ of 0.99 s. Again, little loss of polarization was suffered throughout the transfer/transport process, given values of 42.8 ± 2.4% and 41 ± 2% for $P_{Xe}$ measured in the cell before transfer and at 3 T in the Taldir bag, respectively. The latter $P_{Xe}$ corresponds to 1.46 mT from a cell containing 5 mM $CuSO_4$-doped water (Fig. 3B). As part of our initial tests of the Xe-transfer process, $P_{Xe}$ measurement was also performed at an intermediate field of 47.5 mT located ~2 m from the polarizer. Fig. 3C shows a high-field $^{129}$Xe NMR spectrum obtained at this field from a portion of HP$^{129}$Xe within a cell containing 300 torr Xe following automated transfer to a ~50-cc polypropylene spherical phantom connected to the polarizer via 0.25-inch o.d. PFA tubing and located within the field of the permanent magnet. $^{129}$Xe polarization (here, 68.7 ± 3.4%) was calibrated using a thermal $^{13}$C reference signal obtained from 170 mmol of sodium $^{13}$C acetate in 99.8% $D_2O$ located in an identical phantom (Fig. 3D). Comparison with in situ 5.26-mT $^{129}$Xe NMR polarimetry taken from the OP cell during the same experiment before transferring the gas to 47.5-mT magnet (68.5 ± 3.9%) indicated no polarization loss within error, and thus a highly efficient HP$^{129}$Xe transport process. In future designs, we hope to implement the new approach by Saam and coworkers (33) to measure $P_{Xe}$ from corresponding shifts in the optically detected Rb ESR signal.

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Assumptions include the following: ~100% circularly polarized light; negligible electronically excited Rb; spatially constant [Rb]; and $P_{RB} = 0$ when $B_0 = 0$ (which ignores effects of weak residual static fields). More precise results may be obtained by applying an orthogonal field during $B_0 = 0$ scans (or by orienting the polarizer orthogonally to the residual (e.g., Earth’s) field).
to a ~144,000-fold polarization enhancement over the thermal equilibrium value. When preparing multiple bags of HPXe, the total time to produce each bag is ~35-40 min, including ~15-20 min of SEOP/Xe-polarization time. The final 129Xe polarization and mole fraction depend on the cell-loading parameters. The strong NMR signals obtained from the HP129Xe gas allowed the polarization decay to be monitored under a variety of conditions. Fig. 4 shows 129Xe T1 relaxation data for HP129Xe in Tedlar bags recorded at 3 T, but stored at different field strengths: following exponential fits, T1 decay times of 38 ± 12 min and 5.9 ± 0.4 h were observed for 129Xe gas in Tedlar bags at 1.5 mT and 3 T, respectively. Such long T1 values bode well for a variety of applications where gas-phase storage and/or accumulation of HP129Xe is needed (24).

129Xe polarization values for various Xe densities, operating conditions, and measurement magnetic fields are summarized in Fig. 5A. The data exhibit good run-to-run and method-to-method agreement across the various measurements for the given Xe cell loadings (~300-1,600 torr), despite the fact that the Pxe values were measured at different fields, with different NMR acquisition methodologies, and different thermal reference samples. The data include values for in-cell Pxe values of 90.9 ± 5.2%, 57.1 ± 3.3%, 50.1 ± 2.9%, and 33.4 ± 1.9% measured for Xe loadings of 300, 495, 765, and 1,570 torr, respectively; Pxe values of 41 ± 1.6% and 28 ± 1.1% with ~760 and ~1,545 torr Xe loadings were obtained following transfer to Tedlar bags and subsequent transport to the 3-T scanner, permitting imaging with high signal-to-noise ratio (SNR) (~40) to be demonstrated (Fig. 5A, Inset). Moreover, these posttransfer values are ~2.7 times greater than previous bests in Pxe, achieved near such high Xe densities; when combined with the increase in cell volume, these results constitute a ~18-fold improvement in HP129Xe production over the previous best results obtained at such high in-cell Xe densities (22). Finally, recent Food and Drug Administration Investigational New Drug and Institutional Review Board regulatory approval has allowed initial experiments with human subjects to begin. For example, 3D HP129Xe MRI lung ventilation maps (Fig. 5B) and spectra have now been obtained from healthy human volunteers, demonstrating the utility of the polarizer for a variety of clinical applications.

**Discussion**

The performance of the polarizer is the result of a number of aspects of the SEOP process that are both fundamental and technical in nature (29). SEOP can be treated as a simple relaxation process (23), which, at steady state, simplifies to give the following (15):

\[
P_{\text{Xe}}(t = \infty) = \frac{\gamma_{\text{SE}}}{\gamma_{\text{SE}} + \gamma_{\text{Xe}}} \cdot P_{\text{Rb}},
\]

where \(\gamma_{\text{SE}}\) is the Rb/Xe spin exchange rate and \(\gamma_{\text{Xe}}\) is the 129Xe nuclear spin destruction rate (\(=1/T_1\)); thus, it is convenient to
categorize aspects of the hyperpolarizer and its operation as factors that help optimize \( P_{ RB} \), \( P_{ SE} \), and/or \( \Gamma_{ Xe} \), respectively. First, the most important factor in maximizing global \( P_{ RB} \) is the cell illumination by the laser. At a given position \( (r) \) within the cell, the local \( P_{ RB} \) is given by the following (15):

\[
P_{ RB}(r) = \frac{\gamma_{ OP}(r)}{\gamma_{ OP}(r) + \Gamma_{ SD}} \quad [4]
\]

where \( \gamma_{ OP}(r) \) is the local OP rate [given by the integrated product of the laser flux and the Rb absorption cross-section (31)], and \( \Gamma_{ SD} \) is the Rb electronic spin destruction rate, which is dominated by nonspin-conserving collisions with Xe [via the spin-rotation interaction (25, 27)] and is high under our conditions because of its proportionality to \( [Xe] \). Correspondingly, the laser power used here is also high—usually \( \sim 170 \) W, mostly resonant with the Rb D\(_1\) line. In fact, the resonant photon flux is roughly sixfold greater than in ref. 22, but this increase explains only part of the performance improvement. Care was also taken with the optical path to ensure complete illumination throughout the OP cell, with no dark regions, and with additional photon flux provided by retro-reflection. Control of the cell temperature is also key for SEOP optimization. Previously, it was shown that there can be an inverse relationship between Xe density and the optimal temperature for SEOP, where higher \( [Xe] \) favors lower cell temperatures (22, 29)—an effect that may be explained in part by the need to maintain uniform illumination throughout the cell. Although \( \gamma_{ SE} \) is proportional to the Rb density (34)—ostensibly favoring higher cell temperatures—maintaining a high “phonon-to-Rb” ratio (in part by limiting \( [Rb] \)) can be particularly important for preserving global \( P_{ RB} \) as Xe density is increased, and Xe-induced Rb spin-destruction becomes dominant. [An additional contribution may arise from poor energy dissipation caused by reduced thermal conductivity of Xe-rich mixtures—a possibility we are currently studying with in situ Raman spectroscopy (35).] For example, ongoing simulations predict a high \( \Gamma_{ SD} \) value of \( \sim 134,300 \) s\(^{-1}\) for the experimental conditions of the 65 °C data in Fig. 2, dominated by Xe collisions (25–27). However, the high resonant photon flux at the front of the cell should give nearly a ninefold greater optical pumping rate \( (\gamma_{ OP} \sim 1.16 \times 10^5 \) s\(^{-1}\)) using Eq. 4; such values for \( \Gamma_{ SD} \) and \( \gamma_{ OP} \) would correspond to \( P_{ RB} \sim 89.6\% \). Maintaining a relatively low Rb density allows the laser to penetrate efficiently to ensure high photon flux and Rb polarization throughout the cell, consistent with the experimental \( (P_{ RB} = 71 \pm 6\%) \); Thus, particularly in the limit where \( \Gamma_{ Xe} \) is negligible, maximizing \( P_{ RB} \) is more important than \( \gamma_{ SE} \), once \( \gamma_{ SE} \) is sufficiently high. Indeed, our high \( P_{ Xe} \) values were enabled in part by our ability to achieve long-in-cell \( ^{129}\)Xe \( T_1(=1/\Gamma_{ Xe}) \) decay times. As one example, following SEOP with a cell containing 495 torr Xe and cooldown to 33 °C (where Rb should be condensed), a cell \( T_1 \) of 1.9 ± 0.6 h was measured at 5.26 mT. Given a typical build-up time constant measured for one experiment of \( \gamma_{ OP} = (\gamma_{ SE} + \Gamma_{ Xe}) \) = 8.5 min, this \( ^{129}\)Xe \( T_1 \) translates to a spin-exchange rate of \( \sim 1.8 \times 10^{-9} \) s\(^{-1}\)—roughly an order of magnitude greater than \( \Gamma_{ Xe} \). Thus, according to Eq. 3, \( P_{ Xe} \) should closely approach \( P_{ RB} \) under these conditions, again in good agreement with the Fig. 2 experiments.

In addition to long HP\( ^{129}\)Xe relaxation times measured in OP cells, slow polarization decay was also observed following transfer to Tedlar bags. Generally, \( ^{129}\)Xe relaxation rates are determined by several potential contributions (adapted from ref. 24):

\[
\Gamma_{ Xe} = \Gamma_1 + \Gamma_2 + \Gamma_{ O2} + \Gamma_{ w}. \quad [5]
\]

where \( \Gamma_1 \) is an “intrinsic” contribution from spin-rotation interactions of transient and persistent Xe/Xe dimers \( \{\Gamma_1 = \Gamma_1 + \Gamma_P \} \) (36), \( \Gamma_2 \) results from diffusion through field gradients, \( \Gamma_{ O2} \) results from dipolar interactions with residual paramagnetic O\(_2\) in the gas mixture, and \( \Gamma_1 = \Gamma_{ w,c} + \Gamma_{ w,n} \) includes contributions from Xe/wall collisions involving interactions with surface spins of unpaired electrons and nuclei, respectively. \( \Gamma_{ w,c} \) can be neglected for Tedlar (polyvinyl fluoride); moreover, \( \Gamma_2 \) can be neglected for the 3-T measurements given the homogeneous field. Saam and coworkers recently provided a semiempirical formula for \( \Gamma_1 \) that when adapted for our conditions, is given by the following (24):

\[
\Gamma_1 = \frac{[Xe]}{56.1h} + \frac{1}{4.59h} \left[ 1 + (3.65 \times 10^{-3})B_0^2 \right] \left( 1 + 0.51 \frac{[N_2]}{[Xe]} \right)^{-1}, \quad [6]
\]

where the two additive terms respectively correspond to \( \Gamma_1 \) and \( \Gamma_{ O2} \). \( B_0 \) is in tesla, the gas densities are in amagat \[1 \text{ amagat} = 2.6873 \times 10^{15} \text{ cm}^{-3}\], the density of an ideal gas at 0 °C and 760 torr, and 0.51 is a factor that takes into account the differential breakup rate of persistent Xe dimers from collisions with N\(_2\). Using values for \( [Xe] \) and \( [N_2] \) of 0.35 and 0.58 amag, Eq. 6 predicts a limiting value of 1/\( \Gamma_{ Xe} \) = 7.8 h (with 10% uncertainty), in relatively good agreement with our experimental value of 5.9 ± 0.4 h at 3 T, and indicating effective suppression of other relaxation pathways. Given the expected absence of paramagnetic wall sites, if we assume that \( (\cdot) \) relaxation from wall collisions involving surface nuclear spins (37) (e.g., \( ^1\text{H}, ^{19}\text{F} \)) is essentially quenched at high field and \( (i) \) that the remaining contribution to \( \Gamma_{ Xe} \) is from residual O\(_2\) [with a relaxation of \( ~0.4 \text{ Hz/amag} \) (38)], then the differential relaxation rate would correspond to an upper limit for the O\(_2\) partial pressure of \( \sim 2 \times 10^{-2} \) torr—a reasonable amount. Although \( \Gamma_1 \) is the limiting factor for our high-field \( ^{129}\)Xe relaxation, the accelerated decay at 1.5 mT (\( T_1 \approx 38 \pm 12 \) min) is consistent with \( \Gamma_{ w,N} \) providing the dominant mechanism. Driehuys et al. (37) showed that, at low field (\( ~10 \) mT), dipolar relaxation with \( ^1\text{H} \) “wall” spins was the primary contributor to \( ^{129}\)Xe polarization decay in cells with organosilane coatings, with \( T_1 \sim 50 \) min at 1.5 mT and 297 K. Although Tedlar may lack the nanoscale Xe permeability of SurfaSil coatings (37), it does possess a high surface density of \( ^1\text{H} \) and \( ^{19}\text{F} \) spins and likely provides an attractive surface for transient Xe adsorption. More detailed studies of \( ^{129}\)Xe relaxation in Tedlar containers will be the subject of future efforts.

The polarizer’s performance can also be compared with the spin-transfer efficiency, \( \eta \), defined as follows (adapting from ref. 39):

\[
\eta = \frac{P_{ Xe} \cdot N_{ Xe}}{\Gamma_{ Xe} \cdot P_{ Xe} \cdot N_{ Xe}} \quad [7]
\]

Where \( N_{ Xe} \) is the number of \( ^{129}\)Xe spins, \( \Delta t \) is the photon absorption rate, and \( T_1 \) is the \( ^{129}\)Xe relaxation time in hot illuminated cells (taken here as \( \gamma_{ OP} \)). Treated effectively as a constant fundamental to Rb/Xe SEOP, \( \eta \) was found to be \( \sim 0.043 \) (39) (i.e., \( \sim 4.3\% \) of the light angular momentum should end up in \( ^{129}\)Xe spins). Unfortunately, polarizers generally do not approach such efficiency. For example, extrapolating from the prediction of \( \sim 25 \) cc-atm/h of 100%-polarized \( ^{129}\)Xe with 1 W of absorbed light, our \( \sim 90\)-W absorption would correspond to \( \sim 22.2 \) L-atm/h of 100%-polarized \( ^{129}\)Xe. We can calculate a “production efficiency”, \( \eta_p \), defined as the number of polarized \( ^{129}\)Xe spins produced versus photons absorbed during a SEOP run: \( \sim 0.94 \) L-atm/h of \( ^{129}\)Xe spins with \( P_{ SE} \) \( \sim 30\% \) (equivalent to \( \sim 0.28 \) L-atm/h of \( P_{ Xe} = 100\% \) \( ^{129}\)Xe spins [we assume 1.570 torr Xe, 295 K, SEOP time of 17.5 min, and 100% duty cycle, corresponding to \( \sim 3.5 \) L-atm/h Xe gas, with \( P_{ Xe} \sim 30\% \) for the \( ^{129}\)Xe fraction (26.44%)]), giving \( \eta_p \sim 0.0054 \). However, if we assume little \( P_{ Xe} \) loss with the use of 100%-enriched \( ^{129}\)Xe, these numbers would climb to \( \sim 1.1 \) L-atm/h and \( \eta_p \sim 0.021 \), respectively. Although more comparable to the theoretical.
limits, these values suggest room for improvement in future designs, and justify further study of fundamental aspects governing SEOP efficiency.

On a technical level, efficiency is aided by the presence of in situ monitoring of the laser absorption and $^{129}$Xe NMR signals, which allow real-time SEOP optimization. Afterward, $^{129}$Xe polarization losses are mitigated by (i) rapid cooldown of the cell while reducing the laser power (to further decrease the heat load, while still providing sufficient Rb illumination—a procedure that allows only a few percent loss of $P_{Xe}$; as well as (ii) suppressing exposure to O$_2$ and other paramagnetic materials throughout the gas lines, (iii) moving the HP $^{129}$Xe quickly into the sample or transport vessel, and (iv) avoiding Xe phase transitions and other relaxation-susceptible portions of the Xe phase diagram (30).

Indeed, Xe cryocollection/sublimation before transfer to the sample is optional for XeNA, but it is normally a requirement with other polarizer designs operating with standard “xenon-lean” gas mixtures. We should add that not cryocollecting the Xe does result in dilution of the HP $^{129}$Xe with N$_2$ gas and leaves behind significant fractions of the SEOP mixture in the cell following the expansion process; however, both of these issues can be mitigated, respectively, by using Xe-rich mixtures and by the addition of a large automated gas piston (15) where the cell contents can be expanded into a much larger volume before transfer.

**Conclusion**

We have presented results from a (~1 L/h) hyperpolarizer that produces batches of HP $^{129}$Xe sufficient for clinical use. The hyperpolarizer’s open-source design and automated operation should facilitate implementation of HP $^{129}$Xe technology into other laboratories and clinical settings. Good agreement was obtained among the four independent methods for characterizing the spin polarization. The ability to achieve high $P_{Xe}$ values at high Xe densities, combined with stopped-flow operation, negates the usual requirement of Xe cryoaccumulation and storage, and opens a door to greatly improved polarization efficiency for quadrupolar isotopes (40, 41) (e.g., $^{83}$Kr and $^{133}$Xe). Further improvements in designs and gas-handling efficiency will be manifested in our second-generation hyperpolarizer (now under development); other possible improvements in SEOP efficiency of the heavy noble gases may exploit Cs (42) or Cs/Rb hybrid SEOP schemes. These results, combined with ultra-long gas-phase polarization lifetimes and recent regulatory approval, bode well for a host of planned clinical applications with human subjects.

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