Phase-Encoded Hyperpolarized Nanodiamond for Magnetic Resonance Imaging

David E. J. Waddington1, Thomas Boele1, Ewa Rej1, Dane R. McCamey2, Nicholas J. C. King3, Torsten Gaebel1 & David J. Reilly1,4

Surface-functionalized nanomaterials are of interest as theranostic agents that detect disease and track biological processes using hyperpolarized magnetic resonance imaging (MRI). Candidate materials are sparse however, requiring spinful nuclei with long spin-lattice relaxation (T1) and spin-dephasing times (T2), together with a reservoir of electrons to impart hyperpolarization. Here, we demonstrate the versatility of the nanodiamond material system for hyperpolarized 13C MRI, making use of its intrinsic paramagnetic defect centers, hours-long nuclear T1 times, and T2 times suitable for spatially resolving millimeter-scale structures. Combining these properties, we enable a new imaging modality, unique to nanoparticles, that exploits the phase-contrast between spins encoded with a hyperpolarization that is aligned, or anti-aligned with the external magnetic field. The use of phase-encoded hyperpolarization allows nanodiamonds to be tagged and distinguished in an MRI based on their spin-orientation alone, and could permit the action of specific bio-functionalized complexes to be directly compared and imaged.

Diamond is a remarkable material with outstanding thermal, mechanical, optical, and electrical properties that have given rise to a diverse spectrum of technologies since it was found to be readily synthesizable in the 1950s. In the context of nanomedicine, synthetic nanodiamond can act as a bright, fluorescent marker1 of use in sub-cellular tracking of biological processes2, in vivo sequestering of migratory cell populations3 or as a non-toxic substrate for the delivery of chemotherapeutic payloads4–7. Further, the extremely weak nuclear magnetism in diamond makes the material interesting for hosting quantum information8 and for constructing quantum sensors that exploit its long-lived nuclear9 or electron10 spin-states.

The long lifetime of these spin-states has recently motivated the development of new nanodiamond-based theranostic agents that are detectable with magnetic resonance imaging (MRI) by acquiring signal from their 13C nuclei11,12. In natural abundance carbon however, spin-1/2 13C nuclei comprise only 1.1% of the diamond lattice, rendering their signal undetectable using conventional MRI. The presence of paramagnetic centers in diamond offers a means of boosting the weak signal from 13C using hyperpolarization to transfer the much larger thermal polarization of the centers to the 13C nuclei, by driving spin transitions optically13,14, or with microwaves at cryogenic temperatures11,15–18. Conversely, the presence of paramagnetic centers also shortens the nuclear spin relaxation time (T1), limiting the timescale over which hyperpolarized nuclei can be put to use19,20.

Although there has been significant work now showing that nanodiamond nuclei can be significantly hyperpolarized, a demonstration that this nanomaterial can be used in a true imaging context has been lacking, posing a barrier for biomedical applications. Overcoming the imaging barrier needs more than simply detecting the boosted signal from 13C15, since constructing spatially-resolved images also requires that the nuclear dephasing time (T2) be sufficiently-long, relative to the timescale set by the strength of practical magnetic field gradients. For this reason solid-state compounds with short T2 times have always presented a major challenge for imaging.

Here we address these challenges by demonstrating the use of nanodiamond for hyperpolarized 13C MR-imaging. Via electron paramagnetic resonance (EPR) measurements, we first show that synthetic nanodiamond

1ARC Centre of Excellence for Engineered Quantum Systems, School of Physics, University of Sydney, Sydney, NSW, 2006, Australia. 2ARC Centre of Excellence for Exciton Science, School of Physics, University of New South Wales, Sydney, NSW, 2052, Australia. 3The Discipline of Pathology, School of Medical Sciences, Bosch Institute, Sydney Medical School, University of Sydney, Sydney, New South Wales, 2006, Australia. 4Microsoft Corporation, Station Q Sydney, University of Sydney, Sydney, NSW, 2006, Australia. Correspondence and requests for materials should be addressed to D.J.R. (email: david.reilly@sydney.edu.au)
contains a concentration of impurities and dangling-bonds that is well suited to hyperpolarization (see Fig. 1(a), without significantly degrading the long nuclear T₁ time. In a major advance over previous work 11, we highlight how hyperpolarization via paramagnetic centers in nanodiamond opens up new avenues for preparing spin-populations with unique MRI contrast signatures.

Reasoning that spatially-resolved imaging requires suitably long spin coherence, data characterizing the dephasing times for ¹³C nuclei is presented, demonstrating the use of dynamical decoupling pulse sequences that preserve coherence and offer a means of improving image resolution. Testing the viability of hyperpolarized nanodiamond for MRI, we first acquire image data as a function of time, nanoparticle concentration, and size, before deploying the technique in a setting relevant to preclinical applications, acquiring a co-registered ¹³C-¹H image in a mouse post mortem. Together, these results help bridge the gap between our earlier basic demonstration of hyperpolarization11, and the requirements for imaging applications.

Finally, we speculate on the use of phase-encoded hyperpolarized MRI for addressing common challenges faced with targeted nanoparticle theranostics.

Figure 1. Hyperpolarization of Nanodiamond. (a) Nanodiamonds (blue) are host to spin-1/2 defects such as charged vacancy sites (1), dangling bonds (2) and substitutional nitrogen atoms (3). Resonant microwaves can be used to drive spin polarization from these defects to ¹³C nuclei in the lattice. (b) X-band EPR spectra of 2 µm HPHT (blue) and 2 µm NAT (orange) diamonds. The HPHT and NAT nanodiamonds have substitutional nitrogen impurities present at 95 ppm and 4 ppm respectively. Other spin-1/2 defects are present at concentrations of 115 ppm in the HPHT nanodiamonds and 28 ppm in the NAT nanodiamonds. A representative EPR absorption signal is also shown (inset, bottom right). Hyperfine splitting (A) of the EPR line results from nitrogen impurities. (c) Energy level schematic of Zeeman split electrons (e) and nuclei (n), showing possible flip-flop and flip-flop transitions in diamond. Transitions are numbered in order of increasing energy (1–6). (d) Hyperpolarization spectra of ¹³C as a function of microwave frequency for HPHT 210 nm (green) and HPHT 2 µm (blue) diamond particles. Signals are normalized to unity. Transitions shown in (c) are indicated by arrows ((1) 80.81 GHz; (2) 80.84 GHz; (3) 80.87 GHz; (4) 80.94 GHz; (5) 80.97 GHz; (6) 81.0 GHz). (e) ¹³C NMR spectra of HPHT (blue) and NAT (orange) 2 µm diamonds after 20 minutes of hyperpolarization, normalized to the NMR signal from thermal polarization (HPHT - black; NAT - brown). (f) ¹³C NMR spectra of HPHT (green) and NAT (purple) 210 nm nanodiamonds after 20 minutes of hyperpolarization, normalized to the NMR signal from thermal polarization (HPHT - black; NAT - brown).
Intrinsic Defects as a Source of Polarized Spins

Turning to the details of our experiments, we begin by comparing synthetic nanodiamond produced via the high-pressure high-temperature method (HPHT), to natural nanodiamond (NAT) samples, highlighting the suitability of each for hyperpolarized MRI. Both kinds of nanodiamond exhibit a series of intrinsic electronic defects, immediately apparent in their X-band EPR spectra, shown in Fig. 1(b). The generic features in the spectra can be attributed to a convolution of a hyperfine-split P1 center associated with substitutional nitrogen atoms, a narrow singlet component from vacancy sites, and a broad singlet that stems from dangling carbon bonds. We find that P1 center concentration in the HPHT diamonds (95 ppm) is more than an order of magnitude higher than the P1 center concentration in the NAT nanodiamonds (4 ppm) [see Supp. Fig. 1 and Supp. Table 1 for discussion of particle size dependent effects].

In the presence of an external magnetic field (B = 2.88 T), these paramagnetic defects provide a platform for nuclear hyperpolarization by microwave driving the manifold of hyperfine transitions shown in Fig. 1(c). Although room temperature hyperpolarization is possible, the most significant enhancements occur when microwaves are applied at cryogenic temperatures (4 K or below), transferring the large Boltzmann polarization of the electronic defects to closely coupled 13C nuclei. For spins in nanodiamond, hyperpolarization occurs via a mixture of the solid-effect and cross-effect and leads to a polarized nuclear population of order 8% at saturation [see Supp. Note 1 for discussion of polarization mechanisms]. Detuning the microwave frequency below or above the central EPR line drives hyperfine transitions that build a positive or negative enhancement in the 13C signal (see Fig. 1(d)). The sign of the signal indicates nuclear hyperpolarization that is aligned or anti-aligned with the external magnetic field. Hyperpolarizing HPHT and NAT nanodiamond yields the 13C signal enhancements shown in Fig. 1(e,f), for a mean particle diameter d = 2 μm and d = 210 nm respectively. We draw attention to the variation in signal enhancement with particle size and when using nanodiamonds with different concentrations of paramagnetic defects (HPHT versus NAT). These variations suggest further approaches to enhance hyperpolarization, dependent on specific applications.

Spin Relaxation and Spin Dephasing

The concentration and configuration of paramagnetic centers in nanodiamond carry-over into the characteristic times for (electron-mediated) nuclear spin relaxation, limiting the time for which hyperpolarization can be usefully deployed. Surprisingly, in their readily available synthetic form, HPHT nanodiamonds are already well-suited for hyperpolarization applications, retaining up to 11% of their polarization two hours after being transferred to an imaging platform. Decay rates for the smaller, 210 nm particles are faster but still sufficient for imaging applications, with 10% of hyperpolarization remaining after a period of 20 minutes [see Supp. Fig. 2 for data].

Comparing the relaxation data to the hyperpolarization spectra reveals that driving at select microwave frequencies leads to polarization that relaxes with a distinct nuclear T1 (colored arrows in Fig. 1(d)) correspond to colored curves in Fig. 2(a). This dependence, which has not previously been investigated, likely arises due to the microwave frequency selecting particular paramagnetic centers to act as hyperpolarization lattice sites for their surrounding nuclei. The same centers then dominate nuclear spin relaxation, such that different types of centers and their concentration will lead to different relaxation rates.

Examining the decay of the hyperpolarized state shows that the data is bi-exponential, containing short and long characteristic relaxation time components. Further, as a function of magnetic field, both components vary by nearly two orders of magnitude as the field is increased from B = 130 mT to 7 T, as shown in Fig. 2(b). We attribute this field dependence to the presence of electron and three-spin mediated processes, characteristic of nitrogen impurities. Rapid decay of hyperpolarization at low-field presents a practical challenge to the implementation of hyperpolarized nanoparticle imaging, requiring the use of magnetized sample shuttles to ferry compounds between hyperpolarization and imaging platforms [see Supp. Fig. 3 for details].

Turning to the key results of our paper, we note that imaging clinically-relevant concentrations of the nanodiamond brings new challenges beyond simply detecting hyperpolarization. In particular, the short T2 dephasing times of solids can limit the spatial resolution that is possible using typical scanner gradient magnetic field strengths to the extent that imaging is not possible, despite being able to significantly hyperpolarize the material. Fortuitously, we find the 13C nuclear spins in HPHT nanodiamond possess T2 times already suitable for demonstrating the technique, with substantial scope for improving spatial resolution using dynamical decoupling sequences. To see this, in Fig. 2(c) we compare the signal decay from a standard Hahn-echo sequence (dotted-line) to the signal produced by the use of a Carr-Purcell-Meiboom-Gill (CPMG) decoupling sequence, after the sample has been transferred from the polarizer to a 7 T imaging platform. Although the T2 seen with Hahn-echo is already suitable for sub-millimetre imaging (see Supp. Note 2 for discussion of resolution limits), we find that spin-coherence is preserved for a significantly longer time by decreasing the inter-pulse spacing in the CPMG sequence. This enhancement in coherence indicates that repeated π-pulses have the effect of refocusing the homonuclear dipolar coupling, significantly narrowing the spectral linewidth with promise to improve the imaging resolution of hyperpolarized nanodiamond.

MRI with Hyperpolarized Nanodiamond

Having established nanodiamond has a sufficiently long T2 to render imaging applications viable, we now present the main result of our work, the demonstration of nanodiamond imaging with hyperpolarized 13C MRI.

We restrict our imaging experiments to the HPHT diamonds due to their larger signal enhancement. We begin our study by preparing the “half-moon” phantom shown in Fig. 3(a), which consists of one chamber filled with water and one with an aqueous mixture of 2 μm diamond particles that have been hyperpolarized for 2 hours. The magnetic field that the diamond particles were exposed to was maximized at all times during the transfer and dissolution process to retain hyperpolarization [see Methods]. Once transfer of the phantom to the MRI system...
is complete, $^{13}$C imaging is performed immediately with a centrically-ordered, ultrafast gradient echo (GRE) sequence operating at $B_0 = 7$ T. Sequences such as GRE allow for very short echo times with small tip angles, making them well suited to imaging hyperpolarized dipolar systems such as diamond, where $T_2$ is short and polarization is nonrenewable between phase encoding steps.

Immediately following the acquisition of the $^{13}$C signal we perform conventional $^1$H MRI in order to generate a co-registered water-nanodiamond image, shown in Fig. 3(b). The $^1$H component (gray-scale) of the co-registered image clearly shows the location of water and structure of the phantom, with the $^{13}$C component (red-orange) indicating the location of the hyperpolarized nanodiamond solution. We repeat the experiment, now with a delay of 30 minutes between transfer of the phantom to the MRI system and subsequent imaging. Although the image taken after 30 min shows a significant reduction in $^{13}$C signal (down to ~25%, of the image that was acquired with no delay), we note that it still contains sufficient contrast to be able to identify the presence of nanodiamond. This demonstration constitutes a significant advance over hyperpolarized liquid compounds that are typically limited by relaxation times measured in seconds. Switching to the smaller 210 nm nanodiamonds yields images with a reduced signal-to-noise (SNR) (comparable to 30% of the 2 μm solution), with the images decaying more rapidly in keeping with the size dependence of $T_1$ [see Fig. 3(c) and Supp. Table 3].

Evaluating the sensitivity of $^{13}$C MRI in our current setup, we acquire images for aqueous solutions of hyperpolarized nanodiamonds in a range of concentrations from 20–200 mg mL$^{-1}$, as shown in Fig. 3(d). For the larger 2 μm particles, concentrations as low as 20 mg mL$^{-1}$ continue to yield an acceptable SNR ~ 11, corresponding to 57 μg of diamond per pixel. This particle mass sensitivity is similar to other particle imaging techniques based on hyperpolarization$^{19,23}$.

Figure 2. Decay of Hyperpolarization. (a) Signal decay of hyperpolarized 2 μm diamond particles at $T = 293$ K and $B_0 = 7$ T. Decay curves are shown for NAT 2 μm diamonds hyperpolarized at 80.87 GHz (purple) and HPHT 2 μm diamonds hyperpolarized at various frequencies (red - 80.81, yellow - 80.87 GHz, green - 80.94 GHz and blue - 81.0 GHz). Solid lines are double exponential fits to the data (see Supp. Table 2 for fit parameters). (b) $T_1$ relaxation time versus magnetic field for 2 μm diamond particles. Circular markers show the HPHT long (red), NAT long (black), HPHT short (blue) and NAT short (green) components of a double exponential fit to the hyperpolarized decay curve (raw data shown in Supp. Fig. 2). Dashed lines are a guide to the eye. (c) Signal magnitude of echoes in a hyperpolarized 2 μm diamond sample. CPMG traces are shown with various echo spacings (TE = 0.5 ms - blue; TE = 1 ms - red; TE = 2.1 ms - green). The fit to a Hahn echo experiment performed with thermally polarized 2 μm diamond is also shown (black dotted line; data shown in Supp. Fig. 5).
Animal Imaging

Aware that MRI phantoms can artificially enhance image quality, we demonstrate hyperpolarized nanodiamond imaging in a setting that is more realistic to the clinic by acquiring a co-registered $^{13}$C:$^1$H MRI of a post mortem three-week old laboratory mouse (Mus musculus). Using 2 $\mu$m HPHT nanodiamond, the sample was hyperpolarized for 2 hours before rapid dispersion into a syringe incorporating a 450 mT Halbach array to retain hyperpolarization. The mouse was positioned on a stage within an open-access permanent magnet and the hyperpolarized bolus then injected into the thoracic cavity. Transfer of the mouse to the 7 T MRI system again made use of a custom Halbach array. The total time between removal of the sample from the hyperpolarizer and insertion of the mouse in the MRI system was ~60 seconds.

Co-registered $^1$H:$^{13}$C images of nanodiamond in the mouse are shown in Fig. 3(e). The $^1$H image reveals anatomical features within the chest at sub-millimeter resolution. The $^{13}$C diamond image then clearly shows that the nanodiamond is confined to a region around the lungs. We would expect this localization to occur after injection into the thoracic cavity as the 2 $\mu$m particles are too large to cross membranes into other regions of the mouse\(^{19,32}\).

Phase-Encoded Hyperpolarization

The use of nanoparticles for future theranostic applications hinges on the effectiveness of their functionalized surfaces, which, for instance, can enable targeted take-up of particles by binding to specific cells and localizing at specific disease foci. Accompanying these active targeting mechanisms are passive biological responses to nanoparticles based on their size, morphology, or particle composition\(^{33–36}\). Having demonstrated hyperpolarized nanodiamond MRI, we now look to the future and present a new imaging modality enabled by the material properties of nanodiamond and which aims to directly address the challenge of resolving passive response mechanisms from the active targeted delivery of nanoparticles.

The goal of distinguishing actively targeted vectors from a passive uptake is illustrated in Fig. 4(a), where the yellow particles feature surfaces with receptor molecules in contrast to the bare nanoparticles shown in green. The challenge then is to establish an imaging modality that enables contrast between nominally identical nanoparticles, based on a labelling mechanism that does not require a difference in particle size, composition, morphology, or physical state and is sufficiently long-lasting to track biological distribution processes.

A novel means of tagging nanodiamond populations is evident in the data presented in Fig. 4(b) [reproduced from Fig. 1(d)], which shows the preparation of hyperpolarized states with equal magnitude, both aligned, or anti-aligned, with the external magnetic field by selecting the frequency of microwave driving. In each case the
populations of $^{13}$C spins are prepared predominantly in their ground or excited state, relaxing back to thermal equilibrium on a timescale $T_1$, as indicated in Fig. 4(b).

Imaging contrast between the ground and excited state populations is thus encoded in the opposing phases, rather than amplitudes of the nuclear magnetization. Demonstrating this modality, we hyperpolarize two identical samples of 2 $\mu$m nanodiamond, the first at frequency $f = \omega_e - \omega_n$ and the second at $f = \omega_e + \omega_n$, flipping the phase of the magnetization as indicated in the phantom shown in Fig. 4(c). Overlaid $^{13}$C magnitude (left) and phase (right) colormaps are shown with masks applied to regions with less than 8 times the root mean square value of the background signal. Small black spots in gray-scale $^1$H magnitude image are bubbles adhered to surfaces in the phantom. Scale bar is 3 mm in length.

Figure 4. Distinguishing Identical Nanoparticles with Phase-Encoded Hyperpolarization. (a) Schematic representation of functionalized nanoparticles targeting tumors both actively and passively. Active targeting in the vasculature relies on specific interactions between ligands on the nanoparticle surface and receptors expressed on the surface of, for instance, endothelial cells adjacent to tumors. Passive targeting is nonspecific to nanoparticle functionalization and can occur via the enhanced permeability and retention effect in regions of leaky vasculature at tumor sites. Potentially, phase-encoded hyperpolarization can distinguish between these two regimes. (b) Direction of hyperpolarized $^{13}$C magnetization versus the pump frequency. (c) Phantom schematic showing two 2 $\mu$m diamond samples (yellow and green) in a tube of water (blue). Teflon walls of the tubes containing diamond are also shown (pink). The sample shown in yellow was hyperpolarized at 80.87 GHz ($f = \omega_e - \omega_n$) for 20 minutes before transfer to the imager. The diamond sample shown in green was subsequently hyperpolarized at 80.94 GHz ($f = \omega_e + \omega_n$) for 4 minutes before transfer to the hyperpolarizer. (d) Imaging of the phantom described in (c). Overlaid $^{13}$C magnitude (left) and phase (right) colormaps are shown with masks applied to regions with less than 8 times the root mean square value of the background signal. Small black spots in gray-scale $^1$H magnitude image are bubbles adhered to surfaces in the phantom. Scale bar is 3 mm in length.

Discussion

Opportunities exist to significantly improve the concentration sensitivity of nanodiamond MRI reported here. At present, transfer of the sample from hyperpolarizer to imaging platform results in a substantial loss of polarization, stemming from the rapid relaxation that occurs in a low-field (<380 mT) environment. The use of high-field sample-shuttles during transfer can likely increase the signal by a factor of 10 or more [see Supp. Fig. 4]. Further, we draw attention to the use of dynamical decoupling pulse sequences in preserving coherence in dilute spin systems such as diamond, as shown in Fig. 2(c).

The $^{13}$C image resolution demonstrated here is sufficient for in vivo applications, being comparable to that shown recently for localization of silicon nanoparticles in an ovarian cancer model with MRI[10]. However, our
frequencies of 299.97 MHz and 75.381 MHz respectively. The microimaging gradient set produces gradient fields combining MRI and optical fluorescence to span sub-cellular to whole-body scales. Design will likely see the nanodiamond platform emerge as a valuable theranostic tool for multimodal imaging, given its promise in chemical bonding, functionalization, and the potential for combinations with other modalities. The combination of MR and fluorescence promises to enable multi-modality imaging and therapeutics in smaller scales.

In conclusion, we have demonstrated 13C magnetic resonance imaging using nanodiamonds, including the introduction of a new imaging modality based on phase-encoded hyperpolarization, exploiting the long relaxation times inherent to the diamond material system. Future improvements in materials, hardware, and sequence design will likely see the nanodiamond platform emerge as a valuable theranostic tool for multimodal imaging, combining MRI and optical fluorescence to span sub-cellular to whole-body scales.

Methods
Diamond Particles. Nanodiamonds used in this work were purchased from Microdiamant (Switzerland). Specific nanodiamond types were monocristalline, synthetic HPHT and NAT particles. Both types were used in sizes of 210 nm (0–500 nm, median diameter 210 nm) and 2 μm (1.5–2.5 μm, median diameter 2 μm). The 2 μm HPHT diamonds used here are well suited to injection as they have a zeta potential of −38 ± 7 mV in water. This zeta potential shows that they are aggregation-resistant, a characteristic important to benefiting fully from the surface of nanodiamond in biological applications45. The 2 μm particles display sedimentation on the timescale of hours, as it is to be expected for diamond particles larger than 500 nm, which is the point at which gravitational forces overcome Brownian motion. 210 nm HPHT nanodiamonds have a near identical zeta potential of −39 ± 8 mV with limited sedimentation observed over a period of weeks. For details of dynamic light scattering measurements and calculation of particle stability, see Supp. Note 4.

EPR Characterization. EPR spectra were measured using a Bruker EMX EPR Spectrometer operating at 9.735 GHz and room temperature. Resulting spectra were fit using Easyspin to a three spin component model11,46. Defect concentrations were calculated relative to the signal from an irradiated quartz EPR standard47.

Hyperpolarization of diamond. Hyperpolarization is performed in a 2.88 T superconducting NMR magnet with a homobuilf DNP probe48. 80–82 GHz microwaves are generated via a frequency multiplier (Virginia Diodes) and 2 W power amplifier (Quinstar) before being directed to the sample via a slotted waveguide antenna. A helium flow cryostat (Janis) was used to cool the sample to 4.5 K. NMR measurements in the polarizer were taken with a custom saddle coil tuned to the 13C Larmor frequency of 30.912 MHz. All hyperpolarization events begin with a 13C NMR saturation sequence (64 × π/2 pulses) to zero the initial polarization. Individual points in the frequency sweeps of Fig. 1(d) correspond to unique hyperpolarization events, showing the 13C NMR signal from a π/2 pulse after 60 s of microwave saturation at a constant frequency. NMR enhancement plots in Fig. 1(e,f) show the 13C signal magnitude after 20 minutes of microwave saturation compared with the signal seen after an identical period of thermal polarization. Absolute polarization values were calculated from the thermally polarized 13C signal at equilibrium and the expected Boltzmann polarization41. Diamond samples were hyperpolarized in teflon tubes, chosen for their transparency to microwaves and robustness to thermal cycling.

Transfer and dissolution. All post-transfer NMR signals were acquired in the same 7 T superconducting NMR magnet used for imaging. Adiabatic sample transfer between the DNP probe and the MRI scanner occurs via a series of “magnetic shields” based on permanent magnets and Halbach arrays (see Supp. Fig. 3 and Supp. Note 5 for further detail). Halbach arrays are especially useful for transferring samples between superconducting magnets as they experience no net translational force in external magnetic fields. Depolarization at 7 T was measured via small tip angle for 2 μm and 210 nm samples hyperpolarized for 2 hours and 1 hour respectively. Data were divided by (cos α)−1, where α is the tip angle and n the pulse number, to account for RF-induced signal decay.

T1 versus magnetic field strength was measured in the stray field of the imaging magnet with samples hyperpolarized for 20 minutes. Small tip angle measurements of the 13C signal magnitude were taken at 7 T before and after shuttling the sample to a lower field region for repeated periods of 30 s. The phantom in Fig. 3(a) was prepared in an approximately 200 mT permanent magnet by rapid thawing of a 13C hyperpolarized 2:1 mixture of nanodiamond and water. The hyperpolarized mixture was then mixed with additional water to give a 120 mg mL−1 concentration. Transfer to the MRI system occurred via a 110 mT Halbach array.

MRI Experiments. All MRI was performed in a 7 T widebore, microimaging system with 1H and 13C Larmor frequencies of 299.97 MHz and 75.381 MHz respectively. The microimaging gradient set produces gradient fields up to 250 mT m−1. Phantoms were imaged using a 10 mm, dual resonance 1H:13C NMR probe. 1H phantom images
were acquired with a GRE sequence with 60 μm × 60 μm pixel size and 6 mm slice thickness. 13C slice thickness was restricted to 20 mm by the active region of the detection coil. Concentration phantoms contained 140 μL of diamond and water mixture.

13C GRE image data in Figs 3(b–d) and 4(d) were acquired with a 64 × 32 matrix and pixel resolution of 0.7 mm × 0.6 mm. The refocusing time (TE) of 1.22 ms was minimized by ramping gradients to full strength and using short, 60 μs, excitation pulses (see Supp. Fig. 6 for complete timing parameters). Centrically-ordered phase encoding with a constant flip angle of 20 was used to increase SNR by 1.96 times at the expense of limited blurring in the phase encode direction 46.

All images displaying 13C data only were interpolated to 128 × 128 resolution by zero-filling and Gaussian filtering in k-space. Co-registered 13C images were interpolated to the resolution of the accompanying 1H image and values smaller than 3 times (Fig. 3(b) or 5 times (Figs 3e and 4(d)) the SNR made transparent to reveal the underlying structure in the 1H image.

Animal Handling and Ethical Approval. Female, C57BL6 mice, 3-week-old, were purchased from the Animal Resource Centre (ARC) (Western Australia, Australia). They were housed under specific pathogen-free conditions with free access to food and water. Animal experiments performed were ethically compliant with the pertinent University of Sydney Animal Ethics Committee, State and NHMRC regulations. The protocols used in our experiment were approved by the University of Sydney Animal Ethics Committee.

Animal Imaging. A 150 mg sample of 2 μm nanodiamonds was hyperpolarized for 2.5 hours in a 1:1 mixture with water with 0.2 mL of hot water added during dissolution. Small tip angle characterization showed that hyperpolarizing in water or ethanol gave the same 13C enhancement seen for dry samples. The resulting 0.5 mL bolus was injected into the thoracic cavity of a three-week old female mouse post mortem (11 g, mus musculus) with a 21 G needle and syringe before transfer to the MRI system. Hyperpolarized nanodiamonds were magnetically shielded with custom magnets when not in the polarizer or imager (further details in Supp. Fig. 7). Total time for dissolution, injection and transfer to the imager was approximately 60 s.

A 1H:13C double resonance microimaging probe with 18 mm bore size was used for imaging the mouse torso. 13C SNR of this probe was measured to be one third of the 10 mm probe used for phantom imaging. A 150 mg sample of 2 μm nanodiamonds was hyperpolarized for 2.5 hours in a 1:1 mixture with water with 0.2 mL of hot water added during dissolution. Small tip angle characterization showed that hyperpolarizing in water or ethanol gave the same 13C enhancement seen for dry samples. The resulting 0.5 mL bolus was injected into the thoracic cavity of a three-week old female mouse post mortem (11 g, mus musculus) with a 21 G needle and syringe before transfer to the MRI system. Hyperpolarized nanodiamonds were magnetically shielded with custom magnets when not in the polarizer or imager (further details in Supp. Fig. 7). Total time for dissolution, injection and transfer to the imager was approximately 60 s.

A 1H:13C double resonance microimaging probe with 18 mm bore size was used for imaging the mouse torso. 13C SNR of this probe was measured to be one third of the 10 mm probe used for phantom imaging. A conventional 1H spin-echo sequence was used for high-contrast anatomical imaging. 1H images were acquired with 1 mm slice thickness with a 256 × 256 matrix and pixel resolution of 230 μm × 176 μm. The same 13C GRE sequence was used as in Fig. 3(b–d) but with a gradient echo time of 0.90 ms and pixel size of 2.1 mm × 1.9 mm. Slice thickness was limited to less than 18 mm by the sensitive region of the probe.

Phase-contrast Imaging. To demonstrate phase-contrast imaging, a 2 μm HPHT sample was positively polarized with 80.87 GHz microwaves for 20 minutes and transferred to the MRI scanner. A second, nominally identical sample was then negatively polarized with 80.94 GHz microwaves for 4 minutes and transferred to the MRI scanner.

Data Availability The authors declare that the data supporting the findings of this study are available within the article and its Supplementary Information. Raw data is available from the corresponding author on request.

References
1. Fu, C.-C. et al. Characterization and application of single fluorescent nanodiamonds as cellular biomarkers. Proc. Natl. Acad. Sci. USA 104, 727–732 (2007).
2. Kucsko, G. et al. Nanometre-scale thermometry in a living cell. Nature 500, 54–58 (2013).
3. Getts, D. R. et al. Therapeutic inflammatory monocyte modulation using immune-modifying microparticles. Sci. Transl. Med. 6, 219ra7 (2014).
4. Weissleder, R., Nahrendorf, M. & Pittet, M. J. Imaging macrophages with nanoparticles. Nat. Mater. 13, 125–138 (2014).
5. Almeida, J. P. M., Figueroa, E. R. & Drezek, R. A. Gold nanoparticle mediated cancer immunotherapy. Nanomed-Nanotechnol. 10, 503–514 (2014).
6. Li, C. A targeted approach to cancer imaging and therapy. Nat. Mater. 13, 110–115 (2014).
7. Boles, M. A., Ling, D., Hyeon, T. & Talapin, D. V. The surface science of nanocrystals. Nat. Mater. 15, 141–153 (2016).
8. Maurer, P. C. et al. Room-Temperature Quantum Bit Memory Exceeding One Second. Science 336, 1283–1286 (2012).
9. Waldherr, G. et al. High-precision temperature mapping with a single electronic spin in diamond. Nat. Nanotechnol. 7, 105–108 (2012).
10. Grinolds, M. S. et al. Nanoscale magnetic imaging of a single electron spin under ambient conditions. Nat. Phys. 9, 215–219 (2013).
11. Rej, E., Gaebel, T., Boede, T., Waddington, D. E. J. & Reilly, D. J. Hyperpolarized nanodiamond with long spin-relaxation times. Nat. Commun. 6, 8459 (2015).
12. Parker, A. J. et al. Optically-pumped dynamic nuclear hyperpolarization in 13C enriched diamond. arXiv 1708.00561 (2017).
13. King, J. P. et al. Room-temperature spin hyperpolarization from optically pumped nitrogen vacancy centres in diamond. Nat. Commun. 6, 8965 (2015).
14. Ajoy, A. et al. Orientation-independent room temperature optical 13C hyperpolarization in powdered diamond. Sci. Adv. 4, eaar5492 (2018).
15. Dutta, P., Martinez, G. V. & Gillies, R. J. Nanodiamond as a New Hyperpolarizing Agent and Its 13C MRS. J. Phys. Chem. Lett. 5, 597–600 (2014).
16. Casabianca, L. B., Shames, A. I., Panich, A. M., Shenderova, O. & Frydman, L. Factors Affecting DNP NMR in Polycrystalline Diamond Samples. J. Phys. Chem. C 115, 19041–19048 (2011).
17. Bretschneider, C. O. et al. On The Potential of Dynamic Nuclear Polarization Enhanced Diamonds in Solid-State and Dissolution 13C NMR Spectroscopy. ChemPhysChem 17, 2691–2701 (2016).
18. Kozłowska, G. et al. Direct hyperpolarization of micro- and nanodiamonds for bioimaging applications – considerations on particle size, functionalization and polarization loss. J. Magn. Reson. 286, 42–51 (2018).
19. Cassidy, M. C., Chan, H. R., Ross, B. D., Bhattacharya, P. K. & Marcus, C. M. In vivo magnetic resonance imaging of hyperpolarized silicon particles. Nat. Nanotechnol. 8, 363–368 (2013).
20. Atkins, T. M. et al. Synthesis of Long T1 Silicon Nanoparticles for Hyperpolarized 29Si Magnetic Resonance Imaging. ACS Nano 7, 1609–1617 (2013).
21. Cunningham, C. H. et al. Hyperpolarized 13C Metabolic MRI of the Human Heart: Initial Experience. Circ. Res. 119, 1177–1182 (2016).
22. Rej, E., Gaebel, T., Waddington, D. E. J. & Reilly, D. J. Hyperpolarized Nanodiamond Surfaces. J. Am. Chem. Soc. 139, 193–199 (2017).
23. Waddington, D. E. J. et al. Nanodiamond-enhanced MRI via in situ hyperpolarization. Nat. Commun. 8, 15118 (2017).
24. Rammohan, N. et al. Nanodiamond-Gadolinium(III) Aggregates for Tracking Cancer Growth In Vivo at High Field. Nano Lett. 16, 7551–7564 (2016).
25. Barklie, R. C. & Guven, J. 13C hyperfine structure and relaxation times of the P1 centre in diamond. J. Phys. C Solid State Phys. 14, 3621–3631 (1981).
26. Loudner, J. H. N. & Wyk, J. A. Electron spin resonance in the study of diamond. Rep. Prog. Phys. 41, 1201–1248 (1978).
27. Taitsev, A. Optical Properties of Diamond. A Data Handbook (Springer, 2001).
28. Terblanche, C. J., Reynhardt, E. C., Rakitianski, S. A. & Van Wyk, J. A. 13C Spin-Lattice Relaxation in Natural Diamond: Zeeman Relaxation in Fields of 500 to 5000 G at 300 K Due to Fixed Paramagnetic Nitrogen Defects. Solid State Nucl Magen Reson 19, 107–129 (2001).
29. Milani, I. et al. A magnetic tunnel to shelter hyperpolarized fluids. Rev. Sci. Instrum 86, 024101 (2015).
30. Li, D., Dementyev, A. E., Dong, Y., Ramos, R. G. & Barrett, S. E. Generating unexpected spin echoes in dipolar solids with π pulses. Phys. Rev. Lett. 98, 190401 (2007).
31. Dong, Y., Ramos, R. G., Li, D. & Barrett, S. E. Controlling coherence using the internal structure of hard π pulses. Phys. Rev. Lett. 100, 247601 (2008).
32. Whiting, N. et al. Developing hyperpolarized silicon particles for in vivo MRI targeting of ovarian cancer. J. Med. Imag. 3, 036001 (2016).
33. Sykes, E. A., Chen, J. & Zheng, G. Investigating the Impact of Nanoparticle Size on Active and Passive Tumor Targeting Efficiency. ACS Nano 8, 5696–5706 (2014).
34. Kümajchan, S. et al. Passive versus Active Tumor Targeting Using RGD- and NGR-Modified Polymeric Nanomedicines. Nano Lett. 14, 972–981 (2014).
35. Colombo, M. et al. Tumour homing and therapeutic effect of colloidal nanoparticles depend on the number of attached antibodies. Nat. Commun. 7, 13818 (2016).
36. Neri, D. & Bicknell, R. Tumour Vascular Targeting. Nat. Rev. Cancer 5, 436–446 (2005).
37. Zheng, H. et al. Quantitative Rate Determination by Dynamic Nuclear Polarization Enhanced NMR of a Diels-Alder Reaction. Anal. Chem. 82, 8897–8902 (2010).
38. Salvati, A. et al. Transferin-functionalized nanoparticles lose their targeting capabilities when a biomolecule corona adsors on the surface. Nat. Nanotechnol. 8, 137–143 (2013).
39. Kreyling, W. G. et al. In vivo integrity of polymer-coated gold nanoparticles. Nat. Nanotechnol. 10, 619–623 (2015).
40. Prey, M. A. et al. Phosphorus-31 MRI of hard and soft solids using quadratic echo line-narrowing. Proc. Natl. Acad. Sci. USA 109, 5199–5205 (2012).
41. Balasubramanian, G. et al. Ultralong spin coherence time in isotopically engineered diamond. Nat. Mater. 8, 383–387 (2009).
42. Knowles, H. S., Kara, D. M. & Attariûe, M. Observing bulk diamond spin coherence in high-purity nanodiamonds. Nat. Mater. 13, 21–25 (2014).
43. Lee, M., Cassidy, M., Ramanathan, C. & Marcus, C. Decay of nuclear hyperpolarization in silicon microparticles. Phys. Rev. B 84, 035304 (2011).
44. Capozzi, A., Cheng, T., Boero, G., Rousset, C. & Comment, A. Thermal annihilation of photo-induced radicals following dynamic nuclear polarization to produce transportable frozen hyperpolarized 13C-substrates. Nat. Commun. 8, 15757 (2017).
45. Mochalin, V. N., Shenderova, O., Ho, D. & Gogotsi, Y. The properties and applications of nanodiamonds. Nat. Nanotechnol. 7, 11–23 (2012).
46. Stoll, S. & Schweiger, A. EasySpin, a comprehensive software package for pulse sequence simulation and analysis in EPR. J. Magn. Reson. 178, 42–55 (2006).
47. Eaton, G. R. et al. Signal-to-noise standard for pulsed EPR. J. Magn. Reson. 205, 109–113 (2010).
48. Cassidy, M. C. Hyperpolarized Silicon Particles as In-vivo Imaging Agents. Ph.D. thesis (Harvard Univ., 2012).
49. Zhao, L. et al. Gradient-echo imaging considerations for hyperpolarized 129Xe MR. J. Magn. Reson. B 113, 179–183 (1996).

Acknowledgements
The authors thank M.S. Rosen for useful discussions and M.C. Cassidy for constructing the hyperpolarization probe. We are also grateful to R. Marsh and A. Itkin for the supply of custom hardware for these experiments. For EPR measurements, the authors acknowledge use of facilities in the Mark Wainwright Analytical Centre at the University of New South Wales. This work was supported by the Australian Research Council Centre of Excellence Scheme (Grant No. EQuS CE110001013) and ARC DP1094439. We also acknowledge use of the nanoparticle analysis tools provided by the Bosch Institute Molecular Biology Facility at the University of Sydney. An earlier version of this manuscript was posted as a preprint on arXiv. This preprint is available at https://arxiv.org/abs/1709.01851 for download.

Author Contributions
D.E.J.W., T.B., T.G. and E.R. performed the hyperpolarized imaging experiments. D.E.J.W. and T.B. performed the EPR experiments and EPR analysis and D.R.M. provided advice on EPR. D.E.J.W., T.B., T.G., E.R. and N.J.C.K. performed the ex vivo experiment. T.B., D.E.J.W. and T.G. designed and built the magnetic arrays for sample transport. All authors interpreted, processed and analyzed the data. D.E.J.W., T.G. and D.J.R. wrote the manuscript.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-42373-w.

Competing Interests: The authors declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
