High-sensitivity diamond magnetometer with nanoscale resolution

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Published online: 14 September 2008; doi:10.1038/nphys1075

The detection of weak magnetic fields with high spatial resolution is an important problem in diverse areas ranging from fundamental physics and material science to data storage and biomedical science. Here, we explore a novel approach to the detection of weak magnetic fields that takes advantage of recently developed techniques for the coherent control of solid-state electron spin quantum bits. Specifically, we investigate a magnetic sensor based on nitrogen-vacancy centres in room-temperature diamond. We discuss two important applications of this technique: a nanoscale magnetometer that could potentially detect precession of single nuclear spins and an optical magnetic-field imager combining spatial resolution ranging from micrometres to millimetres with a sensitivity approaching a few fT Hz$^{-1/2}$.

Over the past few decades, a wide variety of magnetic sensors have been developed using approaches including superconducting quantum interference devices (SQUIDs), the Hall effect in semiconductors, atomic vapour and Bose–Einstein condensate based magnetometry and magnetic resonance force microscopy. Here, we present a novel approach to high-spatial-resolution magnetic-field detection, using systems currently explored as quantum bits: isolated electronic spins in a solid. We focus on spins associated with nitrogen-vacancy colour centres in diamond, because they can be individually addressed, optically polarized and detected, and exhibit excellent coherence properties even at room temperature. Recently, coherent control of nitrogen-vacancy electronic spin qubits has been used to sense and manipulate nearby individual electronic and nuclear spins in a diamond lattice. In what follows, we describe how such a system can also be used for the precision sensing and imaging of external magnetic fields.

We discuss two types of potential implementation of such sensors. First, a single sensing spin confined in a nanoscale region can be brought in direct proximity to a magnetic-field source, such as an electron or nuclear spin. For example, a diamond nanocrystal containing a single nitrogen-vacancy centre can be attached to a tip of a scanning probe such as that of an atomic force microscope. Second, a bulk diamond sample with a high density of nitrogen-vacancy centres can be used to sense fields created by remote objects with ultrahigh sensitivity and submicrometre spatial resolution.

MAGNETOMETRY WITH SINGLE ELECTRONIC SPIN QUBITS

The operating principles of our approach are closely related to those of magnetometers based on spin precession in atomic vapours. In particular, detecting the relative energy shift induced by a magnetic field between two Zeeman sublevels enables precise determination of an applied d.c. or a.c. magnetic field. Ultimately, sensitivity is determined by the spin coherence time and by the spin projection noise. Although solid-state electronic spins have shorter coherence times than gaseous atoms, quantum control techniques can decouple them from the local environment and from each other, as we show below, leading to a substantial improvement in their sensitivity to external, time-varying magnetic fields, while retaining the desirable features of a robust solid sensor.

The canonical approach to detecting a Zeeman shift uses a Ramsey-type sequence as shown in Fig. 2a. A π/2-pulse creates a superposition of two Zeeman levels, which acquire a relative phase $\phi \propto (g\mu_B/h)br$ from the external field $b$ during the free evolution interval $r$, where $\mu_B$ is the Bohr magneton and $g \approx 2$ for nitrogen-vacancy centres. Another π/2-pulse transforms the relative phase into a population difference, which is measured optically and from which the Zeeman shift is inferred. For small $\phi$, the magnetometer signal $S$ (proportional to the induced population difference) depends linearly on the magnetic field: $S \approx (g\mu_B/h)br$. During the total averaging interval $T$, $T/r$ measurements can be made, yielding a shot-noise-limited sensitivity $\eta$ given by the minimum detectable field, $b_{\text{min}} \equiv \eta/\sqrt{T} \approx (h/g\mu_B)(1/\sqrt{T})$. 

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Increasing the interrogation time $\tau$ improves the sensitivity until random (environmental) perturbations lead to decay of the free-precession signal. In the case of solid-state spin systems, the coherence is limited by interactions with nearby lattice nuclei and paramagnetic impurities, resulting in an ensemble dephasing time $T_2^\ast$. Furthermore, there will be a finite number of fluorescence photons collected and detected, leading to extra photon shot noise, and a finite contrast to the Ramsey fringes. We describe these effects by a single parameter $C \lesssim 1$, which approaches unity for ideal, single-shot readout (see Methods section). The optimum sensitivity of a magnetometer based on a single electronic spin, achieved for $\tau \sim T_2^\ast$, is given by

$$\eta_{\text{ac}} \approx \frac{\hbar}{g\mu_B C \sqrt{T_2^\ast}}.$$  

For current experiments\textsuperscript{15}, with detection efficiency $\sim 10^{-3}$, $C \approx 0.05$ and $T_2^\ast \approx 1\mu$s. This yields an optimal sensitivity $\sim 1\mu$T Hz$^{-1/2}$. Improving the collection efficiency by using a tapered fibre or a plasmonic waveguide\textsuperscript{20} to $C \sim 5\%$ yields $C \approx 0.3$ and leads to a sensitivity $\sim 120$ nT Hz$^{-1/2}$.

Coherent control techniques can improve the sensitivity for a.c. fields. Owing to the long correlation times characteristic of dipolar interactions between spins in systems such as diamond—the principal source of dephasing—spin echo techniques can markedly extend the coherence time. Specifically, by adding an extra microwave $\pi$ pulse to the Ramsey sequence at time $\tau/2$, the Hahn echo sequence (Fig. 2a) removes the effect of environmental perturbations with a correlation time that is long compared with $\tau$. Thus, a signal field $b(t)$ oscillating in-phase with the pulse sequence produces an overall additive phase shift, leading to a total phase accumulation,

$$\delta \phi = \int_{t_0}^{t} b(t) \, dt - \int_{t_0}^{t} b^\prime(t) \, dt.$$  

For a signal field of frequency $\nu$ and initial phase $\phi_0$, $b(t) = b \sin (\nu t + \phi_0)$, this yields $\delta \phi = (g\mu_B/\hbar) b \nu t_0 \sin (\nu t + \phi_0)$, with $f(x, \phi_0) = (\sin^2(\nu t/4) \cos(x/2 + \phi_0)/x/4)$. In essence, the spin echo enables us to extend the interrogation time $\tau$ from the limit set by $T_2^\ast$ up to a value $T_2$, which is close to the intrinsic spin coherence time, at the cost of a reduced bandwidth and insensitivity to frequencies $\lesssim 1/T_2$. For maximal response to continuous-wave signals with known frequency and phase (assuming small $b$), we find $\tau = 2\pi/\nu$ and $\phi_0 = 0$ to be optimal. For signals with a time dependence that is $a$ priori unknown, it is useful to measure the signal variance, which provides equivalent performance (see Methods section). In either case, the sensitivity is improved by

$$\eta_{\text{ac}} \approx \frac{\pi \hbar}{2g\mu_B C \sqrt{T_2}}.$$  

The optimum sensitivity is achieved only for fields oscillating near $\nu \sim 1/T_2$. However, these results can be easily extended to higher frequency signals. In particular, for signal field oscillation periods shorter than the dephasing time, the interrogation time need not be restricted to the duration of one period, but can be multiples of it. Then, composite pulse sequences such as the Carr–Purcell–Meiboom–Gill\textsuperscript{21} (CPMG) sequence may perform
better at the expense of a reduced bandwidth. Furthermore, in ultrapure samples where nuclear spins' evolution leads to decay of the echo signal, the long correlation time of the nuclei leads to non-exponential decay of the echo signal. In this case, the CPMG sequence can increase the interrogation time, further reducing the minimum detectable field (see Fig. 2 and Methods section). Finally, another way to improve the magnetometer sensitivity is to use many sensing spins, where we can take advantage of the relatively high achievable density of spins in the solid-state (\(10^{12} \text{ cm}^{-3}\)) compared with atomic magnetometers (\(10^{11} \text{ cm}^{-3}\)) (ref. 24).

**IMPLEMENTATION WITH NITROGEN-VACANCY CENTRES**

We now discuss specific details of magnetometry using nitrogen-vacancy centres in diamond, developing an appropriate operating regime and determining the optimal sensitivities possible for current experimental technology. The fine structure of the electronic ground state of a nitrogen-vacancy centre, shown in Fig. 1a, is a spin triplet. The crystal field splits the \(m_s = \pm 1\) Zeeman sublevels from the \(m_s = 0\) sublevel by \(\Delta = 2\pi \times 2.87 \text{ GHz}\), enabling the use of electron-spin resonance (ESR) techniques even at vanishing external magnetic field. Furthermore, under application of green light, nitrogen-vacancy centres exhibit a transient, spin-dependent fluorescence, which enables optical detection of the spins. After the transient signal decays, the system optically pumps into the \(m_s = 0\) state, which prepares the system for the next measurement (see Methods section).

As a specific example, we focus on magnetometry in low external static magnetic fields (\(\lesssim 10 \text{ mT}\)). In this case, \(\Delta\) is the largest energy scale and sets the spin quantization axis parallel to the nitrogen-to-vacancy direction. The secular Hamiltonian, including a small external field \(\mathbf{B}(t) = (B_x, B_y, B_z)\), is

\[
\mathcal{H} = \hbar \Delta S_z^+ + g \mu_B B_x S_z,
\]

where \(B_x\) is the component of the magnetic field along the nitrogen-vacancy-centre’s axis and \(S_z\) takes the values \(m_s = 0, \pm 1\). Terms proportional to the perpendicular field are suppressed to order \(B_y, B_z / \Delta\) and do not depend on the field \(B_x\) being measured, and therefore may be neglected.

At low magnetic fields, the \(m_s = \pm 1\) manifold can be used to implement a vector magnetometer, sensitive only to components of the magnetic field along the centre’s axis. We focus on the \(m_s = \pm 1\) manifold as it has twice the energy splitting of the 0–1 manifold and is less affected by nuclear spin-induced decoherence at low fields, because internuclear interactions are suppressed by the large hyperfine field.

Coherent control of the nitrogen-vacancy-centre’s spin states is obtained using an ESR magnetic field oscillating at angular frequency \(\Delta\). ESR pulses linearly polarized along the z axis rotate the nitrogen-vacancy spin between the two-dimensional subspace of \(|0\rangle\) and \(|+\rangle = (|1\rangle + |−1\rangle) / \sqrt{2}\). To manipulate \(|\pm 1\rangle\) superpositions, extra control can be provided by a background oscillating reference field \([B_0 \sin(2\pi f/\tau)]\) along the z axis. Specifically, \(B_a = (\hbar / g \mu_B) \pi \tau / 8\) yields an optimal phase offset to achieve a magnetometer signal linear in the field strength (Fig. 2).

The sensitivity as a function of the signal frequency for both a.c. and d.c. detection is plotted in Fig. 2. For diamonds where natural abundance (1.1%) carbon-13 nuclei are the principal cause of dephasing, \(T_2^* \sim 1 \mu s\) and \(T_2 \sim 300 \mu s\) (ref. 18). Again using current experimental parameters, with \(C \approx 0.05\), and measurement and preparation time \(t_m \leq 2 \mu s\), we can optimize the sensitivity as a function of \(\tau\). Including corrections from decoherence with expected signal decay (see Methods section) \(\propto \exp(-\tau / T_1^*)\), we find: \(\eta_{\text{acc}} = (\pi \hbar / 2 g \mu_B) e^{(1/T_1^*)} \sqrt{\tau + t_m / \tau}\). We obtain optimal sensitivity of \(\eta_{\text{acc}} \approx 18 \text{ nT Hz}^{-1/2}\) for a single nitrogen-vacancy centre using current experimental collection efficiencies. Improved collection efficiencies (\(C = 0.3\)) would yield \(\eta_{\text{acc}} \approx 3 \text{ nT Hz}^{-1/2}\).

Note that spin \(T_1\) relaxation occurs on timescales much longer than milliseconds and may be safely neglected.

Finally, the observed dephasing times are independent of temperature from 4 to 300 K, owing in part to the vanishing polarization of the nuclear bath at small magnetic fields.

When more than one nitrogen vacancy centre exists in the sample, they can belong to four different crystallographic classes, each corresponding to the centres’ alignments along different (111) axes. To operate as a vector magnetometer along a controlled direction, a transverse (d.c.) magnetic field \(B_x \geq 0.3 \text{ mT}\) (see Methods section) detunes the other three classes’ levels such that the ESR field used for quantum control excites only spins with the desired crystallographic orientation, perpendicular to the external field. Thus, one in four spins contributes to the magnetometer signal.

**MAGNETOMETRY IN THE HIGH-DENSITY LIMIT**

A principle advantage of our approach over other spin precession magnetometers is the high achievable density \(n\) of sensing spins. This improves the sensitivity to fields that are homogeneous over the magnetometer volume, because the projection noise per unit volume decreases as \(1 / \sqrt{n}\). Nitrogen-vacancy centres can be created in controlled densities by implanting high-purity diamond with nitrogen ions and subsequently annealing the sample to recombine the nitrogen with vacancies. Assuming an initial nitrogen concentration \(\sim 10^{18} \text{ cm}^{-3}\) with a conversion \(f \sim 0.1\) to nitrogen-vacancy centres, we expect it will be feasible to create diamond crystals with a nitrogen-vacancy-centre density exceeding \(\sim 10^{17} \text{ cm}^{-3}\), with an average distance between centres of less than 10 nm. Even at these densities, effects such as superradiance do not have a role owing to the large spectral width of the nitrogen-vacancy fluorescence.

At high spin densities, nitrogen-vacancy paramagnetic impurities and nitrogen-vacancy/nitrogen-vacancy interactions may limit the sensitivity of the magnetometer. In particular, substitutional nitrogen impurities with one bound electron (P1 centres) become a sizable source of dephasing in high-density samples. The dipole–dipole interaction between these centres has a characteristic timescale \(T_2^* \approx (1 / \alpha n_{\text{tr}})^{1/3}\), where \(\alpha\) is of the order of the dipole coupling between electron spins, \((\mu_s / 4\pi)(g \mu_B)^2 / \hbar \approx 3 \times 10^{-13} \text{ s}^{-1} \text{ cm}^3\) and \(n_{\text{tr}}\) is the density of paramagnetic impurities. Qualitatively, this timescale corresponds to the rotation time of a single paramagnetic spin in the presence of the random field from the other paramagnetic centres. The timescale for interaction between this impurity bath and a given nitrogen-vacancy centre will be of the same order of magnitude. This suggests an exponential decay of spin echo coherence on a timescale \(T_2^*\) (see Methods section), in contrast to single nitrogen-vacancy-centre-based sensing, where nearby nuclear spins limit the coherence time.

To evaluate the effects of paramagnetic impurities, we assume a density \(n\) of nitrogen-vacancy centres and \(n_{\text{imp}} = n(1-f)/f\) of paramagnetic impurities, where \(f\) is the conversion factor described above. The relevant figure of merit is the sensitivity per root volume \(\eta_{\text{acc}}^V = \eta_{\text{acc}} \sqrt{V}\). We find

\[
\eta_{\text{acc}}^V = \frac{\hbar}{2 g \mu_B} \frac{\pi e^{(1/T_1^*)} C \sqrt{n \tau}}{C \sqrt{n \tau}} \times e^{(1/T_1^*)}.
\]

where we have taken into account that the sensing centres account for only one fourth of the nitrogen-vacancy centres.
in the sample. Here, we include both dephasing due to a bath of dipolar-coupled nuclear spins and the paramagnetic spin bath just discussed. In the high nitrogen-vacancy density and low-$f$ regime, $T_{2,\text{Carbon}} > T_{1} > T_{r}$, that is, carbon-$13$ is no longer the limit to echo lifetimes, but still limits inhomogeneous broadening. Then the optimum magnetometer sensitivity becomes: $\eta_{\text{opt}}^{V} = (\hbar \gamma_{C} / g_{\mu_{B}} C) \sqrt{2 \alpha \varepsilon (1 - f) / f}$. For $f = 0.1$ and $T_{2,\text{Carbon}} = 300 \mu s$, the optimum sensitivity is independent of the nitrogen-vacancy density over the range $n \approx 10^{12} - 10^{18} \text{ cm}^{-3}$, as is seen in Fig. 3a, and reaches a maximum sensitivity value $\eta_{\text{opt}}^{V} \approx 250 \mu T \text{ Hz}^{3/2} \text{ cm}^{-3/2}$ for $C = 0.3$. However, the optimum echo time depends on the nitrogen-vacancy density, $\tau = f / (1 - f) 2 n \alpha_{e}$, with higher density samples requiring higher detection frequencies. Finally, for $n \gg 10^{17} \text{ cm}^{-3}$, corrections due to finite preparation, control and measurement times can become important, and lead to the limitations in sensitivity at high nitrogen-vacancy density seen in Fig. 3a.

To push the sensitivity limits beyond the capping imposed by paramagnetic impurities, we can exploit more advanced forms of dynamical decoupling than spin echo. With appropriate external time-dependent controls, the system can be made to evolve under an effective, time-averaged Hamiltonian that is a suitable symmetrization of the undesired interactions. For example, driving the P1 centres through spin resonance at a rate much faster than the intrinsic decorrelation time, $T_{1}$, acts as a rapid spin echo for the nitrogen-vacancy centres without affecting the nitrogen-vacancy-centre’s magnetic-field-sensing capabilities. Furthermore, improving implantation and conversion techniques (by optimizing implant energies or by using cold implantation) could increase the ratio of nitrogen-vacancy centres to paramagnetic impurities. When the conversion efficiency exceeds 50%, interactions between nitrogen-vacancy centres become the primary source of noise, with a dephasing $\propto (\alpha \tau n)^{2}$. The coupling between the sensing nitrogen-vacancy centres is a $S_{z} S_{z}$ interaction that is not removed with spin echo. However, by using collective rotations driven by appropriate ESR pulses, the interaction can be successively rotated through the $x, y$ and $z$ axes for an equal time duration; so that on average the spins will experience an isotropic Hamiltonian, which commutes with the signal perturbation and thus allows the spin evolution necessary for magnetometry. Pulse sequences such as MREV (eight-pulse refocusing sequence introduced by Mansfield, Rhim, Ellis and Vaughan) can induce the desired evolution, and will be necessary in the high nitrogen-vacancy-centre density limit.

**SINGLE-SPIN DETECTION WITH A NANO-MAGNETOMETER**

Nitrogen-vacancy magnetometers can be applied to an outstanding challenge in magnetic sensing: the detection and real-space imaging of small ensembles of electronic and nuclear spins, with the long-term goal of resolving individual nuclear spins in a molecule. As the magnetic field from a single dipole decreases with distance as $1/r^{3}$, a magnetometer that can be brought into close proximity of the field source offers a clear advantage. A diamond nanocrystal or a single nitrogen-vacancy centre near the surface of a bulk crystal within a scanning set-up would enable a spatial resolution limited only by the distance between the nitrogen-vacancy centre and the object of study, not by the wavelength of the fluorescence signal. For example, consider a prototype system consisting of a crystal with a single nitrogen-vacancy centre at a distance $r_{0} \approx 10 \text{ nm}$ from the surface of the crystal. At this distance, the dipolar field from a single proton is $B_{p} \approx 3 \text{ nT}$, which is well within the projected limits for a single nitrogen-vacancy centre.

To examine a practical method to measure the magnetic field from a single spin, we consider a material with a varying nuclear spin density $n$, that is brought in close proximity (a distance $r_{0}$) to the nitrogen-vacancy-centre. At realistic temperatures, the thermal nuclear spin polarization of the material will be small. However, because only a few spins are involved, the distribution of spin configurations leads a large variance in the spin polarization, providing a substantial, albeit randomly oriented, magnetic field to be detected by the nitrogen-vacancy magnetometer. We find (see Methods section) that the field magnitude measured by our sensor will be characterized by a variance $B_{\text{rms}} \sim B_{0} \sqrt{N}$, where $N \approx 8 \pi n r_{0}^{3}$ is the effective number of spins contributing to the signal. This indicates that our prototype system has an effective spatial resolution determined only by the distance of the nitrogen-vacancy centre from the surface of the sample material, assuming we can position the sensor relative to the sample with stability much better than $r_{0}$.

At nuclear spin densities $\leq 10^{14} \text{ cm}^{-3}$, there is on average one or fewer nuclear spins in an effective sensing volume with $r_{0} \approx 10 \text{ nm}$. Hence, in this case, single spins could be measured. However, most organic molecules have substantially higher proton densities
∼ a magnet near the surface of a substrate can produce gradient fields from nitrogen-vacancy centres in a di
with a high density of nitrogen-vacancy centres. The signal with large field-of-view and optical wavelength-limited spatial resolution is comparable to micro-SQUID magnetometers but
is sensitive to magnetocrystalline anisotropy and to magnetic-field gradients.

The high magnetic-field sensitivity in a small volume offered by solid-state spin qubits such as nitrogen-vacancy centres in diamond can find a wide range of applications, from fundamental physics experiments to quantum computing applications to detection of NMR signals, surface physics and material science, and medical imaging and biomagnetism. Recently, proof-of-principle experimental demonstrations of such a sensor have been carried out by members of our collaboration and other groups. Further extensions could include the use of non-classical spin states, such as squeezed states induced by the spin–spin coupling. The sensitivity could also be improved by using synthesized, isotopically purified diamond containing a lower fraction of carbon-13, the main cause of dephasing at moderate nitrogen-vacancy densities, and by developing more efficient nitrogen-vacancy-centre creation techniques that do not result in high densities of paramagnetic impurities. On a more general level, these ideas could apply to a variety of paramagnetic systems or other types of solid-state qubit that are sensitive to different perturbations.

**METHODS**

**Spin-Hall effect magnetometry**

The spin triplets of the nitrogen-vacancy centre have a V-type level configuration. An external microwave field tuned to the Δ = 2.87 GHz resonance with its magnetic field linearly polarized along the X-axis drives transitions between |0⟩ and the superposition |±⟩ = (|+⟩ + |−⟩)/√2, whereas the state |−⟩ = (|+⟩ − |−⟩)/√2 is dark—it is decoupled from the field owing to destructive quantum interference. Application of a magnetic field aligned with the nitrogen-vacancy-centre Z-axis perturbs the interference, and enables complete quantum control of the spin triplet. In an echo sequence appropriate for magnetometry using the |±⟩ and |−⟩ states, the traditional π/2 − π − π/2 structure is replaced by π − 2π − π: the first pulse creates |±⟩, the second induces a relative π-phase shift between |±⟩ and |−⟩ and the third converts |−⟩ to |0⟩ while leaving |±⟩ population trapped in the |±⟩ manifold. We remark that for external fields in excess of a few milliteslas it may be more convenient to use the 0–1–1 manifold, as this different resonance frequencies would be necessary for using the ±1 manifold in this regime.

**A.C.-FIELD MEASUREMENT SCHEME AND BANDWIDTH**

a.c.-field detection requires synchronization of the pulse sequence with the external magnetic-field oscillations. When this is not practical or if the field phase ϕ varies randomly in time, successive measurements will give random readings distributed over the range of the function f(v, ϕ), where f is the main text) resulting in a zero average signal. In this situation, information about the field intensity is contained in the measured signal variance, provided the random phase correlation time τp satisfies: τp ≪ τc ≪ T (if τc > T, the total averaging time, the scheme presented in the main text could be used.)

More generally, our approach enables the detection of nanoscale variations in the chemical and physical environment.

We note that the present approach can surpass the sensitivity of SQUID's, Hall-bar and recently proposed optically-pumped semiconductor-based nano-magnetometers by more than an order of magnitude, with 10–2,000 times better spatial resolution. The ultimate limits to miniaturization of nitrogen-vacancy-centre nano-magnetometers, which are probably due to surface effects, are not yet well understood.

**IMAGING OF MACROSCOPIC MAGNETIC FIELDS**

In contrast to the nano-magnetometer approach outlined above, a macroscopic crystal of diamond containing many nitrogen-vacancy centres may be used as a high-sensitivity imaging magnetometer with large field-of-view and optical wavelength-limited spatial resolution. As an example system, we consider a crystal of diamond with a high density of nitrogen-vacancy centres. The signal from nitrogen-vacancy centres in a diffraction-limited setting, where a CCD (charge-coupled device) might be used to image the crystal, is divided into separate ‘pixels’, each pixel corresponding to a ∼(1 μm)3 volume element of the crystal. For nitrogen-vacancy centre densities of ∼1015–1017 cm−3 and C = 0.3, each pixel would have ∼100 pT Hz−1/2 a.c. sensitivity. This spatial resolution is comparable to micro-SQUID magnetometers but with four orders of magnitude higher magnetic-field sensitivity.

In such a scenario, diamond crystals could range from tens of micrometres to millimetres in size, and be physically integrated with fibre-based optics for a robust and practical magnetic-field imager.

Larger detector volumes further improve the sensitivity for whole-sample measurements. For example, a (3 mm)3 × 1 mm thick crystal can achieve an overall sensitivity of 3 fT Hz−1/2 with millimetre resolution. Reducing the ratio of paramagnetic impurities to nitrogen-vacancy centres could potentially lead to the detection of attotesla fields, opening the prospect of improved tests of fundamental symmetries and physical laws.

**ESR CONTROL TECHNIQUES**

The spin triplet of the nitrogen-vacancy centre has a V-type level configuration. An external microwave field tuned to the Δ = 2.87 GHz resonance with its magnetic field linearly polarized along the X-axis drives transitions between |0⟩ and the superposition |±⟩ = (|+⟩ + |−⟩)/√2, whereas the state |−⟩ = (|+⟩ − |−⟩)/√2 is dark—it is decoupled from the field owing to destructive quantum interference. Application of a magnetic field aligned with the nitrogen-vacancy-centre Z-axis perturbs the interference, and enables complete quantum control of the spin triplet. In an echo sequence appropriate for magnetometry using the |±⟩ and |−⟩ states, the traditional π/2 − π − π/2 structure is replaced by π − 2π − π: the first pulse creates |±⟩, the second induces a relative π-phase shift between |±⟩ and |−⟩ and the third converts |−⟩ to |0⟩ while leaving |±⟩ population trapped in the |±⟩ manifold. We remark that for external fields in excess of a few milliteslas it may be more convenient to use the 0–1–1 manifold, as this different resonance frequencies would be necessary for using the ±1 manifold in this regime.

The high magnetic-field sensitivity in a small volume offered by solid-state spin qubits such as nitrogen-vacancy centres in diamond can find a wide range of applications, from fundamental physics experiments to quantum computing applications to detection of NMR signals, surface physics and material science, and medical imaging and biomagnetism. Recently, proof-of-principle experimental demonstrations of such a sensor have been carried out by members of our collaboration and other groups. Further extensions could include the use of non-classical spin states, such as squeezed states induced by the spin–spin coupling. The sensitivity could also be improved by using synthesized, isotopically purified diamond containing a lower fraction of carbon-13, the main cause of dephasing at moderate nitrogen-vacancy densities, and by developing more efficient nitrogen-vacancy-centre creation techniques that do not result in high densities of paramagnetic impurities. On a more general level, these ideas could apply to a variety of paramagnetic systems or other types of solid-state qubit that are sensitive to different perturbations.
of the signal field and $W_0$ a windowing function. With a similar calculation, we obtain the windowing function for an $n_z$-cycle pulse sequence:

$$ W_{n_z}(\alpha, \tau) = \frac{1 - \sec(\tau \alpha/4)}{\tau \alpha/2} \sin(n_z \alpha/2). $$

This function has a band centre $\approx 2n_z/\tau$ and bandwidth (half-width at half-maximum) $\approx 4n_z/\tau$. We can evaluate the improvement in coherence times for the CPMG sequence in cases where a detailed understanding of the main source of decoherence is available. For the single-spin magnetometer, we can approximate the nuclear spin environment by separating contributions from distant nuclear spins, undergoing dipolar spin diffusion, and nearby nuclear spins, the evolution of which is frozen by the electron spin’s dipolar field. We can model the distant nuclear spins as an exponentially correlated Gaussian fluctuations field $\mathcal{B}$ with a correlation function $\langle \hat{B}(t) \hat{B}(t') \rangle \approx (\hbar/g_\mathcal{B})^2 \exp(-|t - t'|/T_C)$, where $T_C$ is the correlation time of the nuclear spins.

Within this model, the random phase accumulated during an echo sequence $(\mathcal{B}(t_0) = \mathcal{B}_0 \exp(-i\mathcal{B}_0(t_0)/T_C))$ is characterized by its variance, $(\mathcal{B}_0^2) \approx \tau^2/T_C^2$ for $T_C \gg \tau$, and the total appreciation time $n_s$ being less than the relaxation time of the electron spins. Recent experiments have shown that the relaxation time in ultrapure samples is $\approx 50$ ms (ref. 18), suggesting $n_s \approx 50$ cycles can result in an $(2n_z/\tau)^2 \approx 4$ overall improvement in sensitivity. Note that in practice this improvement will be limited by imperfections in the control pulses. For example, $\pi$-pulse errors of order $1\%$ will limit $n_s \approx 25$, resulting in the optimal sensitivity shown in Fig. 2.

**MEASUREMENT EFFICIENCY**

The state of the electronic spin is measured by spin-selective fluorescence. When illuminated by green light, nitrogen-vacancy centres in the $m_s = 0$ state undergo a cyclic transition $|\pm\rangle \rightarrow |\mp\rangle$ with a rate limited by radiative decay ($\gamma$ $\approx$ 13 MHz). At the same time, centres in the $m_s = \pm 1$ state are rapidly pumped into a dark singlet state, from which they decay to the $m_s = 0$ state after a time $\tau_r \approx 0.5$ ms. To enable good discrimination of the $m_s = 0, \pm 1$ states, the measurement time $\tau_m$ should be smaller than the optical pumping time $\tau_r$.

For a given photon collection efficiency $\eta_p$, an average of $\alpha_0 \approx (n_s \tau_p) \eta_p$ photons are detected from each spin in the $m_s = 0$ state and $\alpha_1 \approx (n_s \tau_p) \eta_p$ photons are detected from each spin in the $m_s = \pm 1$ manifold. We can estimate the combined effects of spin projection noise and photon shot noise for $N$ measurements as $N^{-1/2}/C$, recovering the formulae for sensitivity in the main text, with $C = \sqrt{1 + 2(\alpha_0 + \alpha_1)/(\alpha_0 - \alpha_1)}^2$. This includes the effects of photon shot noise and reduced contrast. For current experiments, a contrast $(\alpha_0 - \alpha_1)/(\alpha_0 + \alpha_1) \approx 0.3$ is observed. Efficiencies of $\eta_p \approx 0.001$ are achieved in current experiments and give $C \approx 0.05$. We find high collection efficiency $\eta_p \approx 0.05$ gives $C \approx 0.3$.

**EFFECTS OF DIFFERENT NITROGEN-VACANCY-CENTRE ORIENTATIONS**

To use an ensemble of nitrogen-vacancy centres as a vector magnetometer, the signal should originate from only one of the four different crystallographic axes. Under application of a d.c. transverse magnetic field $B_0, \hat{x}$, the other (spectator) centres (with crystalline axis $\hat{y}$) have their $|\pm\rangle$ levels split by $\hbar g_\mathcal{B} B_0 \cdot \hat{y}$. This detunes the spectator centres from the microwave field used for preparing and manipulating the $m_s = \pm 1$ subspace. For example, to use nitrogen-vacancy centres along the $(1, 1, 1)$ crystallographic axis, the ideal choice of $\hat{x}$ is to align it with the $(1, 1, 2)$ axis. We require the microwave Rabi frequency $\Omega \approx 3\pi/T_2^*$ for pulse errors to be smaller than our assumed measurement errors for the desired (111) axis. This translates to a requirement that $\hbar g_\mathcal{B} B_0 \approx 3\hbar \Omega /T_2^*$ for the other three axes. For $T_2^* \approx 1\mu$s, we require $B_0 \approx 0.3\mu$T. One intriguing development of nitrogen-vacancy-centre-based magnetometry would be to exploit the four crystallographic classes of nitrogen-vacancy centre to provide a full (three-dimensional) vector magnetometer, by changing the direction of the biasing transverse field $B_0$ in between measurements.

Errors due to inhomogeneities in the nitrogen-vacancy-centre property (for example, variations of the g-factor due to crystal strain) or to spatial inhomogeneities of the magnetic field can typically be neglected. Even for an average microscale field signal and a distribution of g-factors or field inhomogeneity of order 4%, the induced dephasing leads to a broadening of the signal that is smaller than the effects of $T_2^*$.

**COUPLING TO PARAMAGNETIC IMPURITIES**

The coupling of a nitrogen-vacancy electronic spin to other nitrogen-vacancy centres ($S_i$) and paramagnetic impurities such as nitrogen ($I_1, g_i \approx g$) is given by the magnetic dipolar interaction. To first order in $1/\Delta$, the secular dipolar Hamiltonian is given by $H_\text{sec} = \sum_i S_i \cdot \sum_j I_j$, and $H_\text{sec} = \sum_i S_i \cdot J_i$. The dipole interaction vector is $J_i = (\mu g_\mathcal{B} \mu_i^B_0/4\pi t_0 \delta(t_0 - t_0'))$, with the $z$ axis set by the nitrogen-vacancy crystal axis of the sensing spin centres.

We model the secular component of the dipole coupling between paramagnetic impurities and nitrogen-vacancy centres as $J_{i,j} \approx (\delta_{i,j} |\mu g_\mathcal{B} \mu_i^B_0/4\pi t_0 \delta(t_0 - t_0')|)$, and with a characteristic correlation time $\tau_e \approx (\hbar f/(\delta_{i,j} \omega_0 |\mu g_\mathcal{B} \mu_i^B_0/4\pi t_0 \delta(t_0 - t_0')|)^2$ exp($|t - t_0'|/\tau_e$). We can now calculate the expected spin–echo signal as a function of $D_T^2$, which scales as the square of the density of paramagnetic impurities. In this limit, when the correlation times and the interaction energy are at comparable scales, spin echoes decay exponentially as exp($-t/T_e$). We find $T_e \approx 4/\sqrt{\Delta^2}$, hence, for paramagnetic impurity densities of $10^{13}$ cm$^{-3}$, $T_e \approx 1$ ps.

At high densities, paramagnetic impurities and spectator nitrogen-vacancy centres may have sufficiently strong interactions to reduce the correlation time of the field-aligned component, $I_z$. Spectator nitrogen-vacancy centres may be optically pumped to their $m_s = 0$ state, reducing dynamical noise, reducing the effective temperature of the spectator system. However, spin echoes will not remove the effects of the paramagnetic impurities with short correlation times, and they may in fact limit the $T_e$ time and the corresponding bandwidth of the system. Experiments in systems with high nitrogen concentrations indicate exponential decay of echoes on a $\approx 5–20$ ms decay time. At high temperatures, the fluctuations of the potential values of the dipolar field reflect the $\sqrt{N}$ noise statistics from a set of $N$ spins. The mean-square of the $z$-component of the magnetic field is then:

$$ B_{z,\text{rms}}^2 = G^2 \left( \sum_{i,j} g_{ij} \langle r_i \cdot r_j \rangle \right)_{\text{pos}} = 2\pi G^2 \eta_p N. $$

Where we choose coordinates such that the half-plane begins at $z = -r_{31}$ and pos averages over the homogeneous distribution of spin positions, and we assume a density $n_i$ of dipolar spins, enabling us to replace the sum $\sum_i$ with an integral $\int_3 r_i d^3 r_i$. At high temperatures, the fluctuations of the potential values of the dipolar field reflect the $\sqrt{N}$ noise statistics from a set of $N$ spins. The mean-square of the $z$-component of the magnetic field is then:

$$ B_{z,\text{rms}}^2 = G^2 \left( \sum_{i,j} \langle r_i \cdot r_j \rangle \right)_{\text{pos}} = 2\pi G^2 \eta_p N. $$

Where the average over spin configurations at high temperature uses $\langle r_{31} \rangle_{\text{avg}} = \delta_{31} \delta_0 \langle I_{31} (I_{31} + 1)/3 \rangle$. We find in particular that the statistical fluctuations are consistent with $B_{z,\text{rms}} \approx B_0 \sqrt{N}$, where $N \approx n_i r_i^3$. More specifically, the effective number of spins detected $N$ can be estimated from the relation $B_{z,\text{rms}} \approx \langle |B_{z,\text{rms}}|/\sqrt{N} \rangle$. Thus, $N = \langle |B_{z,\text{rms}}|^2 \rangle \approx (1/1 + 1/3) (8\pi n_i r_i^3)$. For $I = 1/2$, this reduces to $N = 8\pi n_i r_i^3$, equivalent in effective detection volume to a half-sphere of radius $2.29r_i$. © 2008 Macmillan Publishers Limited. All rights reserved.
References


Acknowledgements

We gratefully acknowledge conversations with D. Awschalom, A. Cohen, J. Doyle, G. Dutt, J. Maze, E. Togan, P. Stamwex, J. Hodges, S. Hong and M. P. Ledbetter. This work was supported by the NSF, ONR, MURI (DARPA) and the David and Lucille Packard Foundation. J.M.T. is supported by the Pappalardo Fellowship. F.C. is supported by the RaMP Fellowship.

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In the version of this Article originally published, the x axis of Fig. 2b was labelled incorrectly, and should have appeared as shown below. This has now been corrected in the HTML and PDF versions.