

Nanoscale Magnetic Imaging with a Single Nitrogen-Vacancy Center in Diamond

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(Article begins on next page)

HARVARD UNIVERSITY Graduate School of Arts and Sciences

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DISSERTATION ACCEPTANCE CERTIFICATE

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"Nanoscale magnetic imaging with a single nitrogen-vacancy center in diamond"

presented by Sungkun Hong

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Date October 11, 2012

Nanoscale magnetic imaging with a single nitrogen-vacancy center in diamond

A DISSERTATION PRESENTED BY Sungkun Hong to The School of Engineering and Applied Sciences

IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF Doctor of Philosophy In the subject of Applied Physics

> Harvard University Cambridge, Massachusetts October 2012

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Nanoscale magnetic imaging with a single nitrogen-vacancy center in diamond

Abstract

Magnetic imaging has been playing central roles not only in fundamental sciences but also in engineering and industry. Their numerous applications can be found in various areas, ranging from chemical analysis and biomedical imaging to magnetic data storage technology. An outstanding problem is to develope new magnetic imaging techniques with improved spatial resolutions down to nanoscale, while maintaining their magnetic sensitivities. For instance, if detecting individual electron or nuclear spins with nanomter spatial resolution is possible, it would allow for direct imaging of chemical structures of complex molecules, which then could bring termendous impacts on biological sciences. While realization of such nanoscale magnetic imaging still remains challenging, nitrogen-vacancy (NV) defects in diamond have recently considered as promising magnetic field sensors, as their electron spins show exceptionally long coherence even at room temperature.

This thesis presents experimental progress in realizing a nanoscale magnetic imaging apparatus with a single nitrogen-vacancy (NV) color center diamond. We first fabricated diamond nanopillar devices hosting single NV centers at their ends, and incorporated them to a custom-built atomic force microscope (AFM). Our devices showed unprecedented combination of magnetic field sensitivity and spatial resolution for scanning NV systems. We then used these devices to magnetically image a single isolated electronic spin with nanometer resolution, for the first time under ambient condition.

We also extended our study to improve and generalize the application of the scanning NV magnetometer we developed. We first introduced magnetic field gradients from a strongly magnetized tip, and demonstrated that the spatial resolution can be further improved by spectrally distinguishing identical spins at different locations. In addition, we developed a method to synchronize the periodic motion of an AFM tip and pulsed microwave sequences controlling an NV spin. This scheme enabled employment of 'AC magnetic field sensing scheme' in imaging samples with static and spatially varying magnetizations.

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Author List

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Introduction

Advancements in magnetic detection and imaging have contributed immensely to a wide range of scientific communities from fundamental physics and chemistry to practical applications such as the data storage industry and medical sciences. One classic example is nuclear magnetic resonance [7, 77] which has led to powerful applications, such as magnetic resonance imaging [54] (MRI). Over the past few decades, many advanced magnetic imaging schemes have been developed with improved sensitivities and spatial resolutions: among those are magnetic force microscopy [55], scanning Hall probe microscopy [12], superconducting interference devices [41], and magnetic resonance force microscopy [99]. However, they are often times limited by operating conditions such as cryogenic temperatures and/or high vacuum, hindering their use towards imaging systems which require ambient conditions.

Recently, negatively-charged nitrogen-vacancy (NV) color centers in diamond



Figure 1.1.1: Nitrogen-vacancy color centers in diamond. (a) Level diagram of a single nitrogen-vacancy center in diamond. (b) Optically detected electron spin resonance as a function of magnetic fields along NV axis.

have been proposed as a promising system for nanoscale magnetic field sensing [4, 18, 58, 88]. It has been shown, both experimentally and theoretically, that NV centers offer excellent magnetic field sensitivities [58, 88]. Moreover, since NV centers are point defects tightly confined in a crystal lattice, they can be brought in proximity to magnetic samples within a few nanometers, allowing for nanometric spatial resolution. These sensing capabilities are maintained under ambient conditions (room temperature and atmospheric pressure), and can in principle work in liquid, which is crucial for biological imaging. Over past few years, these exciting properties have led to rapid progress in developing NV-based magnetometers.

1.1 MAGNETIC FIELD SENSING WITH NV CENTERS

The NV center is a point defects in diamond consisting of a substitutional nitrogen with a vacancy in its nearest neighbor lattice site (Fig. 1.1.1a). The

negatively charged state forms a spin triplet in the orbital ground state. The crystal field splits these spin sublevels, resulting in the $m_s = 0$ state in the lowest energy state, and the $m_s = \pm 1$ sublevels lifted by 2.87 GHz. An additionally applied external magnetic field splits the $m_s = \pm 1$ sublevels, which isolates an effective spin-1/2 system. The population of this spin-1/2 system can be read out and initialized optically via spin-dependent fluorescence and optical pumping, respectively. These optical readout and initialization schemes, along with the coherent microwave manipulation, allow for full control of the NV center's spin states.

Magnetic field sensing with NV centers is based on Zeeman splitting of the NV's spin states [4, 58, 88], where the $m_s = \pm 1$ states split in proportion to the magnetic field along the N-V axis. This Zeeman energy shift can be read out optically. Optically detected electron spin resonance spectra [30] (ESR), for instance, can be used for magnetic field sensing, as spin resonance frequencies shift proportional to external magnetic fields (Fig. 1.1.1b). A well-established methodology to precisely determine Zeeman shifts is to monitor changes in spin precession, as widely used in spin-based magnetometers such as atomic vapor cells [11]. This type of approach uses a Ramsey-type measurement sequence [88], where additionally accumulated phase $\varphi = \gamma B \tau (\gamma = g e \mu_B / \hbar i s)$ the gyromagnetic ratio of an electron spin) during a fixed evolution time τ reflects the local magnetic field B (Fig. 1.1.2a). This accumulated phase is converted to a population difference, which can be read out via NV's spin-dependent fluorescence. In the limit of small magnetic field, the corresponding shot-noise-limited sensitivity, or minimally detectable field 8Bopt within a second, can be expressed by

$$\delta B_{opt} \sim rac{2}{\gamma C} rac{1}{\sqrt{F_{mean} N au_{opt}}}$$

where F_{mean} is the mean number of photons collected per one measurement shot, C is the fluorescence contrast between the spin states $m_s = 0$ and $m_s = 1$, N is the



Figure 1.1.2: Magnetic field sensing with NV centers. (a) Magnetic field detection scheme with ESR pulse sequences. Measurement of static magnetic field is based on a Ramsey pulse sequence (top). The measurement starts with preparing an NV spin's state to an equal superposition of $m_s=0$ and $m_s=1$ (or -1) by applying $\pi/2$ -pulse. During a free evolution time, τ , phase is accumulated in proportion to the field. The second $\pi/2$ -pulse converts the phase to the population of the NV's spin state, which then can be measured through spin-dependent fluorescence. Sensitivity of this measurement is limited by the coherence time of the Ramsey sequence. An echo-based pulse sequence can improve the magnetic field sensitivity (bottom), provided that it measures AC magnetic field with its period matched to the total free evolution time. It allows for a longer free evolution time by prolonging the coherence time. (b) Electron spin coherence times of an NV center in high-purity diamond. The figures are adapted with permission from Reference [14]. $\bigcirc 2006$. American Association for the Advanced of Science. The coherence time in Hahn echo signal (242 μ s) is two orders of magnitude longer than the coherence time of Ramsey signal ($2 \mu s$).

number of NV centers participating in the measurement, and τ_{opt} is the optimal evolution time, which would normally be determined by NV's phase coherence time. The coherence time, otherwise limited by environmental fluctuations, can be extended by orders of magnitude, using decoupling sequences such as a Hahn-echo squence [17, 88] (Fig. 1.1.2b). Using these techniques, often referred as 'AC sensing schemes', an NV center becomes sensitive only to alternating magnetic fields [88], requiring modulation of targeted magnetic sources. For paramagnetic spins, this can be achieved by conventional resonant driving methods. Using this AC sensing scheme, it has been shown that a single NV center in high purity diamond is already capable of 30 nT/\sqrt{Hz} sensitivity [58], and can achieve 4 nT/\sqrt{Hz} by isotope engineering [5].

A critical advantage of magnetic sensing with NV centers is that they are atomic-sized point defects in solids. Nanoscale magnetic imaging generally requires the sensor's close proximity to the sample of interest in addition to small detection volume of the sensor, since the magnetic fields from individual spins, fundamental building blocks of magnetism, falls off as $1/r^3$. Fulfilling these requirements is a challenging problem in other spin precession based magnetometers, as the spatial extents of sensing atomic clouds are limited by sizes of their traps. However for a single NV center, crystal lattices naturally provide a tight and stable trapping of NV centers within only a few angstroms. Furthermore, it has been recently reported that NV centers can be formed within 5 nm from the diamond surface while maintainng coherence times longer than $100 \ \mu s$ [69]. This combination of tight confinement and nanometer-scale proximity to the diamond surface suggests that it is feasible to bring NV centers in a few nanometers from the sample. In addition, these solid-state aspects of NV centers are also advantageous for artificially engineering NV centers. For instance, well-established techniques for defect engineering, such as ion-implantation [76] and delta-doping [69], can be readily applicable to create NV centers at few nanometer position accuracy.



Figure 1.2.1: Scanning magnetometry with a single NV center. (a) Schematic of scanning NV magnetometry. (b) Schematic of an AFM probe with diamond nanocrystal containing a single NV center. (c) Spatial image of a magnetic field profile from a magnetic structure acquired by a diamond nanocrystal probe. The images are adapted with permission from Reference [4]. ©2008. Nature Publishing Group.

1.2 Scanning magnetometry with a single NV center

A natural approach to implement nanoscale magnetometry with a single NV center is utilizing scanning probe technique with the NV center residing at the end of the tip [13, 18] (Fig. 1.2.1a). In this configuration, a well-established atomic force microscopy (AFM) technique allows for bringing the sensing NV to a sample of interest in nanometer scale. In addition, to collect spin-dependent fluorescence from an NV center, conventional far-field optics can be used. Finally, a microwave source is installed to manipulate the spin state of an NV center.

As already discussed in the previous section, maintaining a short stand-off distance between the sensing NV centers and the sample is crucial for nanoscale imaging. The first scanning NV magnetometer was implemented by using commercially available diamond nanocrystals hosting a single NV center grafted at the apex of an AFM tip [4] (Fig. 1.2.1b). The size of nanocrystals, nominally 40 nm in diameter, ensures proximity of an NV center to the sample and corresponding magnetic spatial resolution within a few tens of nanometers. Magnetic images could then be acquired by scanning the NV center with respect to the sample, while recording NV's spin-dependent fluorescence in response to the sample's local magnetic field.

This first scanning NV magnetometer successfully demonstrated feasibility of magnetic imaging using NV centers as a local probe. However, the magnetic field sensitivity was limited by the NV sensor's short coherence time due to high concentration of paramagnetic defects in commercial diamond nanocrystals.

1.3 DESCRIPTION OF THIS THESIS

This thesis describes experimental progress in advancing a scanning NV magnetometer with nanoscale resolution, while achieving high magnetic sensitivity at the same time. First, Chapter 2 presents a new approach to realizing such scanning NV magnetometer. This scheme is based on monolithic diamond nano-pillars with single NV centers artificially implanted within a few tens of nanometers from the tip surface [52]. As these devices are fabricated from high-purity diamond, longer coherence times, compared to commercial diamond nanocrystals, are acquired. Moreover, the nano-pillars are designed to support efficient optical wave-guiding of the NV's fluorescence by more than a factor of 5 compared to conventional far-field collection, while requiring an order of magnitude less optical excitation power [3]. This combination of long coherence time and increased photon collection efficiency results in excellent magnetic field sensitivity. As a result, unprecedented combination of high magnetic field sensitivity (56 nT/\sqrt{Hz}) and nanoscale spatial resolution ($\approx 25 nm$) were demonstrated. Chapter 3 presents nanoscale imaging of the magnetic field from a single electron spin using our scanning NV devices, under ambient condition. This work constitutes the first room-temperature magnetic imaging of a single electron spin.

In addition to maximing the magnetic field sensitivity and therefore pushing the imaging capability to a single spin level, an orthogonal approach to improve spatial resolution using magnetic field gradients [29] is discussed in Chapter 4. In this experiment, we demonstrated that identical proximal NV spins become spectrally distinguishable under strong magnetic field gradients, allowing for selective manipulation of the spins at distant locations. This method was applied to spatially image optically irresolvable proximal NV spins.

Finally, in Chapter 5, experimental study of coupled system of an NV spin and a mechanical oscillator is presented [36]. By synchronizing the motion of the mechanical oscillator to the pulsed spin-addressing sequences, coherent control of the hybrid mechanical/spin system was achieved. This scheme was then used to extend 'AC magnetic sensing scheme' to image samples with static and spatially varying magnetizations.

2 A robust scanning diamond sensor for nanoscale imaging with single nitrogen-vacancy centers

2.1 INTRODUCTION

The nitrogen-vacancy (NV) defect center in diamond has potential applications in nanoscale electric and magnetic field sensing [4, 18, 19, 58, 88], single-photon microscopy [15, 84], quantum information processing [64], and bioimaging [59]. These applications rely on the ability to position a single NV center within a few nanometers of a sample, and then scan it across the sample surface, while preserving the center's spin coherence time and readout fidelity. However, existing scanning techniques, which use a single diamond nanocrystal grafted onto the tip of a scanning probe microscope [4, 15, 44, 82], suffer from short spin coherence times due to poor crystal quality, and from inefficient far-field collection of the fluorescence from the nitrogen-vacancy center, Here, we demonstrate a robust method for scanning a single NV center within tens of nanometers from a sample surface that addresses both of these concerns. This is achieved by positioning a single NV center at the end of a high-purity diamond nanopillar, which we use as the tip of an atomic force microscope. Our approach ensures long NV spin coherence times (75 μ s), enhanced collection efficiencies of NV fluorescence due to waveguiding, and mechanical robustness of the device (several weeks of scanning time). We are able to image magnetic domains with width of 25 nm, and demonstrate a magnetic field sensitivity of 56 nT/\sqrt{Hz} at a frequency of 33 kHz, which is unprecedented for scanning NV centers.

NV-based nanoscale sensing is possible because the NV center forms a bright and stable single-photon source [45] for optical imaging, and has a spin-triplet ground state that offers excellent magnetic [58] and electric [19] field sensing capabilities. The remarkable performance of the NV center in such spin-based sensing schemes, is the result of the long NV spin coherence time [5], combined with efficient optical spin preparation and readout [30]. These properties persist from cryogenic temperatures to ambient conditions, which distinguishes the NV center from other systems proposed as quantum sensors such as single molecules [60], or quantum dots [13].

Reducing the distance between the NV center and the sample of interest is crucial for improving spatial resolution. Past experiments aimed at implementing scanning NV microscopes were focused on grafting diamond nanocrystals onto scanning probe tips [4, 15]. Although used successfully in the past, this approach suffers from the poor sensing performance of nanocrystal-based NV centers, for which the spin coherence times are typically orders of magnitude shorter than for NVs in bulk diamond [58]. Here, we present a novel approach that overcomes these drawbacks and thereby realizes the full potential of bulk NV-based sensing schemes in the scanning geometry relevant for nanoscale imaging. In particular, we have developed a monolithic 'scanning NV sensor' (Fig. 2.2.1 a), which uses a diamond nanopillar as the scanning probe, with an individual NV center artificially created within 10 nm of the pillar tip through ion implantation [38]. Long NV spin coherence times are achieved as our devices are fabricated from high-purity, single-crystalline bulk diamond, which brings the additional advantage of high mechanical robustness. Furthermore, diamond nanopillars are efficient waveguides for the NV fluorescence band [3], which for a scanning NV device yields record-high NV signal collection efficiencies.

2.2 Experimental set-up and device fabrication for the scanning NV sensor

Figure 2.2.1b shows a representative scanning electron microscope (SEM) image of a single-crystalline diamond scanning probe containing a single NV center within 10 nm of its tip. To prepare such devices, a series of fabrication steps are performed sequentially, including low-energy ion implantation for NV creation, several successively aligned electron-beam lithography steps and reactive ion etching [34]. An essential element to this sequence is the fabrication of micrometer-thin, single-crystalline diamond slabs that form the basis of the scanning probe device shown in Fig. 2.2.1b. A detailed description of the fabrication procedure of these slabs and the resulting devices can be found in the Supplementary section 2.7.1. Our scanning diamond nanopillars have typical diameters of 200 nm and lengths of 1 µm and are fabricated on few-micrometer-sized diamond platforms that are individually attached to atomic force microscope (AFM) tips for scanning (see Fig. 2.2.1b and Supp. 2.7.1 for details of the attaching process). Our fabrication procedure (Fig. 2.2.1c) allows for highly parallel processing, as shown in the array of diamond devices depicted in the SEM image in Fig. 2.2.1d. Close to 30% of the diamond nanopillars in our samples contain single, negatively charged NV centers. Other devices contain more than one NV, or one NV in a charge-neutral state, which is unsuitable for magnetometry. From these 30%, we select the NV centers that exhibit the highest



Figure 2.2.1: Experimental set-up and probe fabrication for the scan-

ning NV sensor. (a) Schematic of the set-up, consisting of a combined optical and atomic force microscope (AFM). We use a 532 nm laser (green arrows) to address the scanning NV center through its red fluorescence (red arrows). The scanning NV center resides in a diamond nanopillar (inset) and its proximity to the sample is maintained by means of AFM feedback. (b) SEM image of a single-crystalline diamond nanopillar probe (false color coded in red) with a single NV center in its tip (see Fig. 2.3.1). (c) Brief depiction of the fabrication process for scanning single-crystalline diamond NV sensors. Electron-beam lithography is used to define nanopillars and platforms from the top and bottom sides of a few-micrometer thin diamond membrane. Patterns are then transferred to the diamond by reactive ion etching. (d) SEM image of a finalized array of diamond platforms with nanopillars. In all panels, dotted rectangles highlight diamond nanopillars.

photon count rates and longest spin coherence times and mount these single-NV nanopillars onto AFM tips to yield the finalized scanning probe shown in the SEM picture in Fig. 2.2.1b. We note that these scanning devices were fabricated from a [001]-oriented diamond crystal, resulting in NV orientation and magnetic-field sensing along an axis tilted by 54.7° from the nanopillar direction.

To use the scanning NV sensor and characterize its basic spin and optical properties, we used a combined confocal and atomic force microscope as sketched in Fig. 2.2.1a. The set-up was equipped with piezo positioners for the sample and an AFM head to allow for independent scanning with respect to the optical axis. Optical addressing and readout of the NV center was performed through a long-working-distance microscope objective (numerical aperture, NA = 0.7). Microwave radiation for coherent NV spin manipulation was applied using a gold bonding wire attached in proximity to the NV center (see Supp. 2.7.2).

2.3 Characteristics of a single NV center in a scanning diamond nanopillar

Figure 2.3.1a shows a confocal scan under green laser illumination (excitation wavelength, 532 nm) of a typical single scanning NV device. The bright photon emission emerging from the nanopillar (white circle) originates from a single NV center, as indicated by the pronounced dip in the photon-autocorrelation measurement (Fig. 2.3.1b) and the characteristic signature of optically detected NV electron-spin resonance (ESR) [30] (Fig. 2.3.1c), all obtained on the same device. Importantly, we confirm that photon waveguiding through the nanopillar [3] persists despite the close proximity of the NV to the tip of our fabricated nanopillar devices. For example, the data in Fig. 2.3.1c were obtained at 100 μ W excitation power and demonstrate single NV counts approaching 2.2 \cdot 10⁵ counts per second (c.p.s.)—an approximately fivefold increase in detected fluorescence intensity compared to an NV observed under similar



Figure 2.3.1: A single NV center in a scanning diamond nanopillar. (a) Confocal image of red fluorescence from a single-crystalline diamond probe (see side view SEM image in Fig. 2.2.1b). Fluorescence counts are normalized to $I_0 = 1.5 \cdot 10^5$ c.p.s. The encircled bright feature stems from fluorescence of a single NV center in the nanopillar. b, Photon autocorrelation measurement $(g_2(\tau))$ for NV fluorescence in the scanning nanopillar device. Data with $g_2 < 0.5$ (grey-shaded region) demonstrate the presence of a single photon emitter in the nanopillar. (c) Optically detected ESR identifies the single emitter in the nanopillar as an NV center. The two possible NV spin transitions [30] are split by the NV electron Zeeman splitting $2\gamma_{NV}B_{NV}$, where γ_{NV} = 2.8 MHz/ G is the NV gyromagnetic ratio and B_{NV} is the magnetic field along the NV axis (here, B_{NV} 103 G). (d) Spin-echo measurement for the NV center in the diamond nanopillar device. The envelope fitted to the characteristic NV spin-echo decay (see Supp. 2.7.3) yields the NV spin coherence time of $T_2 = 74.8 \ \mu s$. Data in panels b–d were all taken on the same device. Green excitation power: 100 μ W in b–d, and 400 μ W in a.

conditions in an unpatterned diamond sample. We thus significantly increase the fluorescence signal strength from the scanning NV and at the same time minimize exposure of the samples to green excitation light, which is especially relevant for possible biological or low-temperature applications of the scanning NV sensor. Using well-established techniques for coherent NV-spin-manipulation [37], we characterized the spin-coherence time, T_2 , of the same NV center studied so far. Spin-coherence sets the NV sensitivity to magnetic fields and limits the number of coherent operations that can be performed on an NV spin; it is therefore an essential figure of merit for applications in magnetic field imaging [88] and quantum information processing [64]. Using a Hahn-echo pulse sequence, we measured the characteristic single NV coherence decay [91] shown in Fig. 2.3.1d; from the decay envelope we deduce a spin coherence time of $T_2 = 74.8 \,\mu$ s. We note that this T_2 -time is consistent [88] with the density of implanted nitrogen ions $(3 \cdot 10^{11} \text{ cm}^{-2})$ and conclude that our device fabrication procedure fully preserves NV spin coherence. Combining measurements of the T₂-time with the fluorescence count rate and NV spin readout contrast as obtained in Fig. 2.3.1, we obtain a maximal 'AC' magnetic field sensitivity [88] of 56 nT/ $\sqrt{\text{Hz}}$ at a frequency of 33 kHz and (based on data in Fig. 2.3.1c) and a 'DC' sensitivity of 6.0 μ T/ \sqrt{Hz} . We note that both 'AC' and 'DC' magnetic field sensitivities could be further improved by using spin-decoupling sequences $\begin{bmatrix} 17 \end{bmatrix}$ and optimized parameters for spin-readout [20], respectively.

2.4 NANOSCALE MAGNETIC FIELD IMAGING WITH THE SCANNING NV SENSOR

To characterize the resolving power of the scanning NV sensor, we imaged a nanoscale magnetic memory medium consisting of bit-tracks of alternating (out-of-plane) magnetization with various bit-sizes. Figure 2.4.1 illustrates our method and results. The scanning NV sensor operated in a mode that imaged contours of constant magnetic field strength $(B_{\rm NV})$ along the NV axis through the continuous monitoring of red NV fluorescence, in the presence of an ESR

driving field of fixed frequency ω_{MW} and typical magnitude $B_{MW} \approx 2$ G (as determined from NV Rabi oscillations (not shown)). We detuned ω_{MW} by δ_{MW} from the bare NV spin transition frequency, $\omega_{\rm NV}$, but local magnetic fields due to the sample changed this detuning during image acquisition. In particular, when local fields brought the spin transition of the NV into resonance with ω_{MW} , we observed a drop in NV fluorescence rate, which in the image yielded a contour of constant $B_{\rm NV} = \delta_{\rm MW} / \gamma_{\rm NV}$, with $\gamma_{\rm NV} = 2.8$ MHz/G being the NV gyromagnetic ratio. We simultaneously acquired two such images by applying sidebands to $\omega_{\rm NV}$ with $\delta_{MW} = \pm 10$ MHz (dark and bright arrows in Fig. 2.4.1c). Normalization of the pixel values in the two data sets then directly provided a map of magnetic field contours with positive and negative values of B_{NV} (here, with $B_{NV} = \pm_3 G$) and at the same time helped to reject low-frequency noise. Fig. 2.4.1a shows a resulting scanning NV magnetometry image of two stripes of magnetic bits (indicated by the white dashed lines) with bit-spacings of 170 nm and 65 nm. The shape of the observed domains is well reproduced by calculating the response of the NV magnetometer to an idealized sample with rectangular magnetic domains of dimensions corresponding to the written tracks (Fig. 2.4.1e and Supp. 2.7.4).

The spatial resolution of an NV magnetometer is affected by the distance of the NV center to the sample. Therefore, approaching the NV sensor more closely to the magnetic sample revealed magnetic bits with average sizes of \approx 38 nm, as shown in Fig. 2.4.1b. We note that in this image, due to the large field gradients generated at the boundaries between domains, we could observe transitions between magnetic field lines on length-scales of \approx 3 nm (full width at half maximum of the line-cut in Fig. 2.4.1d). An even further decrease of NV-sample distance allowed us to image yet smaller domains, \approx 25 nm in width (Fig. 2.4.1f), but with a reduced imaging contrast caused by strong magnetic fields transverse to the NV axis, which occur in close vicinity to the sample's surface [46] (see Supp. 2.7.5). One of the disadvantages of using a hard drive to characterize our tip is that the local magnetic fields are very large and exceed the

Figure 2.4.1: Nanoscale magnetic field imaging with the scanning NV sensor. (a) NV magnetic field image of bit-tracks on a magnetic memory, highlighted by dashed white lines. We plotted normalized data, I_{norm} = $I_{MW,1}/I_{MW,2}$ (see text and (c)), to reveal magnetic field lines corresponding to $B_{\rm NV} = \pm_3$ G (see inset in a). Additionally, a bias magnetic field of $B_{\rm MW} \approx 52$ G was applied to determine the sign of the measured magnetic fields. (b) Magnetic image obtained as in a, but with the NV-sample distance decreased by an estimated 50 nm. Bringing the NV closer to the sample increases the magnetic field magnitude at the NV sensor, and improves the imaging spatial resolution, allowing us to image magnetic bits, \approx 38 nm in width. Approximate NV-sample distances are noted in the schematics illustrating the experimental configuration, with the sensing NV center fixed on the optical axis and the magnetic sample scanned below the pillar. Total image acquisition times were 11.2 minutes (50 ms/pixel) for a, and 12.5 minutes (75 ms/pixel) for data in b, with laser powers of 130 μ W. The color bar applies to a and b. (c) Optically detected ESR of the sensing NV center. For magnetic field imaging, we modulate an applied microwave field between two frequencies ($\omega_{MW,1} = 2.766$ GHz and $\omega_{MW,2} = 2.786$ GHz) and collect NV fluorescence counts ($I_{MW,1}$ and $I_{MW,2}$, respectively) in synchrony with the microwave modulation. (d) Line-cut along the white line indicated in b (averaged across six adjacent pixels) indicating the sensor's ability to spatially distinguish magnetic field lines separated by \approx 3 nm (limited by the local magnetic field gradient). (e) Calculated NV response for the experimental situation in a, assuming a simplified magnetic sample (see Supp. 2.7.4) (f) Magnetic image as in a and b for a different experimental realization. Here, due to a further decrease in the NV-sample distance, the smallest observed domains have average sizes of 25 nm as determined for the six domains found in the 150 nm interval marked by the blue arrows. The reduced imaging contrast in f results from the close proximity of the NV to the sample (see text).

Figure 2.4.1 (Continued)





Figure 2.5.1: Nanoscale fluorescence quenching imaging of the scanning NV sensor. (a) Scanning the diamond pillar over a sharp metallic tip leads to a bright, circular feature due to sample-topography (see Supp. 2.7.8). Positioning the metallic tip exactly at the location of the NV center (red square), however, yields a sharp dip in NV fluorescence. The illustration shows the experimental configuration used in this experiment. (b) Zoomed-in image of the red square region in a. The observed fluorescence quenching dip has a spatial resolution ≈ 20 nm. (c) AFM topography image obtained simultaneously with the data in b. Blue scale bars represent 100 nm displacement in all directions. Image acquisition times were 30 minutes (320 ms/pixel) and 2.7 minutes (250 ms/pixel) in a and b, respectively at laser power of 35 μ W.

typical dynamic range of our technique. However, such experiments provide valuable information regarding NV-sample distance, and consequently the spatial resolution achieved in imaging. In particular, we estimate the distance between the scanning NV and the sample to be comparable to 25 nm, based on the smallest magnetic domain-sizes we observed.

2.5 Determining the location of a single NV center in a diamond nanopillar

To independently verify the NV's proximity to the diamond surface, we have conducted a measurement where we scanned a sharp metallic tip (< 20 nm in

diameter, see schematic in Fig. 2.5.1a and Supp. 2.7.7) over the NV-containing pillar to image the NV's location. The imaging contrast consisted of the detected NV fluorescence in the far-field changing when the NV was in close proximity to the metallic tip [10]. Owing to the strong dependence of NV fluorescence rate on the distance between the NV and the metallic sample [10] (here, due to partial fluorescence quenching and local modifications of excitation light intensity), we could use this technique to precisely locate the position of the NV center within the diamond nanopillar. The resulting data (Fig. 2.5.1a) showed signatures of the topography of the scanning diamond nanopillar (bright ring in the NV fluorescence signal, see Supp. 2.7.8 for details). More importantly, however, while the front-end of the diamond probe scanned over sharp metallic tip, we observed a dip in NV fluorescence (red square in Fig. 2.5.1a and zoomed image in Fig. 2.5.1b) when the metallic tip was positioned at the location of the NV center. This feature is not accompanied by any topographic features and is thus attributed to partial quenching of NV fluorescence due to the sharp metallic tip (see Fig. 2.5.1c and Supp. 2.7.8). The Gaussian width (double standard deviation) of 25.8 nm of this fluorescence quenching spot was likely still limited by the size of the metallic tip and therefore marks an upper bound to our ability to localize the NV center within the pillar. Importantly, such data allow us to find the position of the single NV center with respect to the topography of our device (Fig. 2.5.1c), which will greatly facilitate precise alignment of our sensing NV center with respect to (magnetic) targets in future sensing and imaging applications.

2.6 DISCUSSION AND OUTLOOK

The biggest remaining uncertainty to the distance between the scanning NV center and the sample is vertical straggle in the NV implantation process, which is still poorly understood [90]. Naturally occurring, stable NV centers have been observed as close as 3 nm from diamond surfaces [9] and thus future advances in the controlled creation of NV centers [73] should allow us to further improve NV-sample distance and therefore spatial resolution in scanning NV imaging by
about one order of magnitude. Additionally, the coherence properties of artificially created NV centers close to the diamond surface could be further improved by appropriate annealing techniques [63] or dynamical decoupling [17] which would both significantly improve the magnetic sensing capabilities of the scanning NVs. We note that for magnetic field imaging, our current ability to resolve individual magnetic domains already equals the typical performance of alternative methods [42, 94], with the added advantages of being non-invasive and quantitative.

The magnetic field sensitivity we demonstrated here with the scanning NV sensor compares well to the performance realized previously with single NV centers in ultrapure, bulk diamond samples [58]. Additionally, the mechanical robustness and durability of our diamond probes (up to several weeks of scanning with the same tip) illustrate the advantage of our method over alternative approaches to scanning NV magnetometry [4, 15]. Compared to other physical systems used for nanoscale magnetic imaging, NV centers in monolithic diamond scanning probes stand out due to the excellent photostability of the NVs, the possibility of room temperature operation and the chemical inertness of diamond, which allow for magnetometry operation even under harsh environmental conditions. To conclude, we note that the scope of applications of our scanning NV probe extends far beyond magnetic imaging. For example, our devices are ideally suited to use as an optical sensors [60, 84] and form an interesting platform to coherently couple the scanning NV spin to other spin systems such as phosphorus in silicon [39], other NV centers, or carbon-based spin qubits [80]. Quantum information could thereby be transferred between a stationary qubit and our scanning NV center, and from there to single photons [89] or other qubit systems such as long-lived nuclear spin qubits in the diamond matrix.

2.7 SUPPLEMENTARY SECTION

2.7.1 DIAMOND TIP FABRICATION

Devices were fabricated from a sample of high purity, single crystalline diamond (Element Six, electronic grade diamond, < 5 ppb nitrogen; thickness 50 μ m). We implanted the sample with atomic nitrogen at an energy and density of 6 keV and $3 \cdot 10^{11}$ cm⁻² (leading to a nominal mean NV depth of ≈ 10 nm). Subsequent annealing at 800°C for two hours yielded a shallow layer of NV centers with a density of ≈ 25 NVs/ μ m² and a depth of \approx 10 nm. We then etched the sample from the non-implanted side to a thickness of \approx 3 μ m using reactive ion etching (RIE, Unaxis shuttleline). We employed a cyclic etching recipe consisting of a 10 min Ar/Cl₂ [48] etch, followed by 30 min of O_2 [34] etching and a cooling step of 15 min. This sequence was essential to maintain the integrity of the diamond surface during the few-hour etching time. On the resulting thin diamond membrane, we fabricated an array of diamond nanopillars on the top side by using electron-beam lithography and RIE as described previously [34]. Next, we performed a second lithography step on the back-side of the diamond slab, which defined platforms to hold the diamond nanopillars. A final RIE process transferred the resist pattern to the sample, and fully cut through the diamond membrane to yield in the structure shown in Fig. 2.2.1d.

To mount a pre-selected diamond platform on an AFM tip, we used a focused ion beam (FIB) system (Zeiss NVision 40) equipped with a nanomanipulator (Omniprobe AutoProbe 300) and ion-assisted metal deposition. We used tungsten deposition or SEM-compatible glue (SEMGLU, Kleindiek) to attach a diamond platform to a quartz AFM tip and then used FIB cutting to release the diamond platform from the bulk. With a properly aligned FIB, this process does not contaminate the scanning diamond nanopillar, and yields a NV/AFM probe as shown in Fig. 2.2.1b.

2.7.2 Combined confocal and atomic-force microscope

We used a homebuilt microscope combining optical (confocal) imaging and AFM. The optical microscope was based on a long working-distance microscope objective (Mitutoyo ULWD HR NIR 100x, 0.7NA). The AFM was tuning-fork based, controlled using commercial electronics (Attocube ASC500) and mounted using a home-built AFM head. Both the sample and the AFM head were fixed on three-axis coarse and fine positioning units (Attocube ANPxyz101 and ANSxyz100, respectively) to allow positioning of the diamond tip with respect to the fixed optical axis and subsequent scanning of the sample with respect to the diamond probe.

Optical excitation of the NV center was performed using a diode-pumped solid-state laser (LaserGlow LRS-0532-PFM-00100-01) at a wavelength of 532 nm. Pulsed excitation for coherent NV spin manipulation used a double-pass acousto-optical modulator (AOM) setup (Isomet, AOM 1250C-848). ESR was driven with a microwave generator (Rhode Schwartz, SMB100A) and amplifier (MiniCircuits, ZHL-42W). The microwave field was delivered to the NV center through a gold bonding-wire (25 μ m diameter) which was mounted as a short-cut termination to a semi-rigid coaxial cable. The wire was brought in close proximity (\approx 50 μ m) to the scanning NV to minimize the required microwave power. Both the microwave source and the AOM were timed using a computer-controlled trigger-card (Spincore, PulseBlasterESR-PRO-400).

2.7.3 Fit to spin-echo data.

To obtain the NV T_2 -time form the spin-echo measurement presented in Fig. 2.3.1d, we fitted the data to a sum of gaussian peaks, modulated by a decay envelope $\propto \exp[-(\tau/T_2)^n]$, i.e., we employed the fit-function [14]

$$\exp[-(\tau/T_{2})^{n}]\sum_{j}\exp[-((\tau-j\tau_{\rm rev})/T_{\rm dec})^{2}].$$
 (2.1)

Taking T_2 , n, τ_{rev} and T_{dec} as free fitting parameters, we found $T_2 = 74.8 \pm 4.3 \,\mu s$, $n = 1.7 \pm 0.2$, $\tau_{rev} = 16.7 \pm 0.1 \,\mu s$ and $T_{dec} = 7.6 \pm 0.2 \,\mu s$ for the data shown in Fig. 2.3.1d (errors indicate 95% confidence intervals for the nonlinear least squares parameter estimates).

2.7.4 SIMULATION OF MAGNETIC IMAGES

In order to reproduce the magnetic images obtained with the scanning NV sensor, we performed a model-calculation of the local magnetic fields in proximity to the hard-disc sample we imaged in our experiment. The magnetic domains were approximated by an array of current-loops in the sample-plane as illustrated in Fig. 2.7.1a. We chose the sizes of the loops to match the nominal size of the magnetic bits on the sample (bit-with 200 nm and bit-length 125 nm and 50 nm for the tracks in the figure) and set the current to 1 mA (corresponding to a density of \approx 1 Bohr magneton per (0.1 nm)²), which we found to yield the best qualitative match to the magnetic field strengths observed in the experiment. We then applied Biot-Savart's law to this current-distribution to obtain the magnetic field distribution in the half-plane above the sample.

Fig. 2.7.1b shows the resulting magnetic field projection onto the NV center at a scan height of 50 nm above the current loops. The NV direction was experimentally determined to be along the ($[0\bar{1}1]$) crystalline direction of the diamond nanopillar (in a coordinate-system where x-, y- and z- correspond to the horizontal-, vertical and out-of plane directions in Fig. 2.7.1b), by monitoring the NV-ESR response to an externally applied magnetic field (using 3-axis Helmholtz-coils). We then allowed for slight variations of the NV orientation due to alignment errors between the diamond crystallographic axes and the scan directions to find the NV orientation that reproduced our experimental data best. With this procedure, we found an NV orientation ($\sqrt{2}sin(\varphi)$, $\sqrt{2}cos(\varphi)$, 1)/ $\sqrt{5}$, with $\varphi = \pi 162/180$.

Finally, we used this magnetic-field distribution to calculate the response of the NV center to a magnetometry scan as described in the main text. For this, we



Figure 2.7.1: Simulation of NV response to bits of a magnetic memory. (a) Current distribution used to simulate the magnetic bits imaged in this work. Red (blue) loops indicate currents of 1 mA in the (counter-)clockwise direction. (b) Magnetic field generated by the current-distribution in (a), projected on the NV axis at a height of 50 nm above the current loops. The NV axis was tilted by 37° out of the scan-plane ([$i\overline{11}$] crystalline direction) with an in-plane component as illustrated by the blue arrow. (c) NV magnetometry response obtained from the magnetic field distribution in (b), assuming a Lorentzian NV-ESR response and microwave detunings as in the original experiment (see text). Note that this experimental situation only resolves magnetic field lines corresponding to the microwave detuning (here: $B_{NV} = \pm_3$ G).

assumed a Lorentzian ESR response with a full-width at half maximum of 9.7 MHz, a visibility of 20 % and two external microwave sources with detunings \pm 10 MHz from the bare ESR frequency, all in accordance with our original experimental parameters.

2.7.5 NV magnetometry in close proximity to a strongly magnetized sample

The presence of a strong magnetic field B_{\perp} , transverse to the NV axis leads to a reduction of contrast in optically detected ESR and moreover reduces the overall fluorescence intensity of the NV center [23]. These effects result from a mixing of the NV spin-levels in the optical ground and excited states of the NV center in the presence of B_{\perp} . Such mixing on one hand allows for spin non-conserving



Figure 2.7.2: Quenching of NV fluorescence and ESR contrast in harddisc imaging. (a) Total NV fluorescence I_{norm} as a function of sample position for an NV in close proximity to the hard-disc sample. I was normalized to the average fluorescence intensity of $I_{o} \approx 15000$ cps in the scan. Dark regions in the scan correspond to individual magnetic domains and are caused by strong magnetic fields transverse to the NV axis which occur in close proximity to the domains. (b) NV magnetic image recorded simultaneously with (a). Data acquisition and integration time per pixel was analogous to the magnetic imaging described in the main text. However here, due to strong transverse magnetic fields, NV ESR contrast almost completely disappeared and prevented NV magnetic imaging using optically detected ESR. The color-bar applies to (a) and (b). (c) Line-cut along the white line in (a), averaged over 7 adjacent pixels. $I_{\rm norm}$ shows a periodicity of ≈ 64 nm, indicating a bit-width of 32 nm. (d) Fluorescence approach curve on the magnetic memory medium. NV fluorescence I was normalized to the fluorescence rate $I_{\infty} = 27'$ 000 cps when the NV center was far from the sample. In contact with the magnetic sample (last data-point to the right), NV fluorescence was reduced by almost a factor of two compared to the NV counts far from the sample. (e) Magnetic imaging with the same NV sensor: Even in close contact to the sample, NV magnetic imaging using ESR is still possible, albeit with a strongly reduced ESR contrast and signal to noise ratio compared to the data shown in the main text. Data in (e) was acquired over 180 minutes, for the smallest resolvable magnetic domains (top third of image) we measure a mean width of 16.5 nm. The laser power was set to 100 μ W in a-d and 60 μ W in e.

optical transitions and on the other hand suppresses the spin-dependence in shelving from the NV excited state (triplet) to the metastable NV singlet state. Both, spin-conservation under optical excitation and spin-dependent shelving are responsible for the non-zero contrast in optically detected ESR of NV centers [26] and consequently, their suppression with transverse magnetic fields explains the disappearance of NV magnetometry features when closely approaching a strongly magnetized sample.

Fig. 2.7.2a shows the raw NV fluorescence counts observed when scanning an NV in a diamond nanopillar in close proximity (estimated 10 – 20 nm distance between NV and sample surface) to the sample. Dark features appear when the NV is scanned over magnetic bits that enhance B_{\perp} , while the inverse happens when B_{\perp} is reduced (or the longitudinal field $B_{\rm NV}$ enhanced) by local fields. This mode of bit-imaging allows for spatial resolutions $\approx 20 - 30$ nm (Fig. ??c). At the same time, a magnetic image recorded with the technique described in the main text shows no appreciable imaging contrast (Fig. 2.7.2b). Only exceedingly long integration times on the order of hours allowed us to reveal weak magnetic features with dimensions on the order of 20 nm (Fig. 2.7.2d).

The rates of the two effects which lead to a disappearance of ESR contrast, i.e. spin-flip optical transitions and shelving of $m_s = 0$ electronic states into the metastable singlet, scale approximately as $\left(\frac{B_{\perp}}{D_{GS}-D_{ES}}\right)^2$ and $\left(\frac{B_{\perp}}{D_{ES}}\right)^2$, respectively, with $D_{GS(ES)}$ the ground- (excited-) state zero-field spin-splitting of 2.87 GHz and 1.425 GHz [25], respectively. Given that $D_{GS} \approx 2D_{ES}$, the scaling of the two mechanisms with B_{\perp} will be very similar. The characteristic scale of D_{ES} ($D_{GS}/2$) for the disappearance of ESR contrast thus allows us to estimate B_{\perp} close to the sample to be $B_{\perp} \approx D_{ES}/\gamma_{NV} \approx 514$ Gauss. We note however that this simple argument likely gives and over-estimation of B_{\perp} as smaller values can already significantly effect ESR contrast and NV fluorescence intensity due to the complex dynamics of NV spin pumping. Indeed, strong reductions of NV fluorescence rates for B_{\perp} less than 100 G have been observed in the past [23]. Transverse magnetic fields on this order were consistent with the largest *on-axis* magnetic fields observed on our experiments as well as with the calculations of



Figure 2.7.3: Contamination of diamond tips. (a) AFM image of the end of a scanning diamond nanopillar after contamination during scanning. The image was acquired by scanning the diamond nanopillar over a sharp diamond tip as shown in Fig. 2.7.4. (b) AFM Image of the same nanopillar as in (a) after cleaning of the pillar's end-face by repeated "scratching" over the sharp diamond tip.

magnetic field profiles presented in Sect. 2.7.4 (for the parameters used in Fig. 2.7.1, we obtain maximal values of $B_{\perp} \approx 200$ Gauss for an NV-to-sample distance of 20 nm).

2.7.6 Limitations to NV-sample distance

As mentioned in the main text, NV-sample distance is an essential parameter for the performance of our microscope as it determines the overall resolving power with which weak magnetic targets can be imaged. We identified three critical parameters that can affect NV-sample distance:

 Implantation-depth of NV centers in the diamond nanopillars: The depth of the NV centers created using ion implantation can be controlled by the energy of the ions used for NV creation. However, the stopping of ions in matter is a random process [98] and the depth of the created NV centers therefore not perfectly well-defined. This straggle in ion implantation poses an intrinsic uncertainty to the distance between the scanning NV and the end of the diamond nanopillar. For implantation energies of 6 keV (with nominal implantation-depths of 10 nm) as used in this work, NV straggle has recently been shown to be as large as 10 - 20 nm [29, 90]. We note that since straggle in NV implantation is hard to circumvent it is essential for the future to develop techniques to precisely pre-determine the depth of a given sensing NV in a diamond nanopillar. This could be performed using recently developed nanoscale imaging methods for NV centers [29], or by scanning the NV sensor over a well-defined magnetic field source.

• Contamination of scanning diamond nanopillars:

During scanning-operation, the scanning diamond nanopillar can gather contamination from the sample or environment. An example for such a contaminated diamond-tip is shown in the AFM image shown in Fig. 2.7.3a (which was acquired with the scanning protocol employed in Fig. 2.5.1, using the a sharp diamond tip as shown in Fig. 2.7.4). Such contamination can artificially increase the distance of the scanning NV center to the sample by several 10 's of nm (see Fig. 2.7.3a). To undo contamination of the diamond-tip after excessive scanning over dirty samples, we developed a "tip-cleaning technique" that allowed us to revert a contaminated tip to its initial, clean state (as illustrated by the transition from Fig. 2.7.3a to b). Tip cleaning is performed by repeated scanning of the diamond nanopillar over a sharp diamond tip (Fig. 2.7.3a) in the absence of AFM feedback. Such feedback-free scanning can partly remove contamination from the diamond pillar, which after repeated operation leads to a clean device as the one shown in Fig. 2.7.3b. We note that with proper sample-cleaning, control over environmental conditions and occasional "tip-cleaning" runs, adverse effects of tip-contamination can be essentially eliminated. This, together with the excellent photo-stability of NV centers, then allows for long-term operation of the scanning NV sensor.



Figure 2.7.4: Sharp diamond tip for FQI. Image of a sharp diamond tip similar to the one used for the experiments presented in Fig. 2.5.1a of the main text. Typical tip-radii are on the order of 10nm.

• AFM control:

Proper AFM control is necessary to assure close proximity of the NV center to the sample surface. It has been shown in the past that bad mounting or improper AFM feedback control can lead to AFM tip-sample distances in excess of 20 nm [40]. Careful mounting of AFM tips and proper setup and tuning of AFM feedback (here provided by an Attocube ASC500 controller) was therefore essential to observe, for instance, the fluorescence quenching features discussed in Fig. 2.5.1 of the main text.

2.7.7 FABRICATION OF SHARP DIAMOND TIPS

For the experiment presented in Fig. 2.5.1a of the main text, we fabricated sharp diamond tips which were metal coated for in order to localize the NV in the scanning nanopillar through fluorescence quenching. Diamond tip fabrication was based on the nanofabrication techniques [34] that we already employed for the production of the scanning diamond nanopillars presented in Fig. 2.2.1. A type Ib diamond (Element six) was patterned with circular etch-masks (flowable oxide, FOx XR, Dow Corning) of 100 nm diameter. Here, in order to obtain sharp diamond tips instead of cylindrical diamond nanopillars, we modified the RIE etching recipe we had previously used: While we kept the (oxygen) etching

chemistry identical to pillar fabrication, we significantly increased the etching time, such as to completely erode the etch mask on the diamond substrate. As a result, the etched diamond structures acquired the form of sharp tips as shown in the representative SEM image in Fig. 2.7.4. Typical tip-radii were in the range of 10 nm and tip lengths were on the order of 200 nm.

For the experiments described in the main text, we then coated the sharp diamond tips with a thin metallic layer using thermal metal evaporation. To avoid oxidation of the metal, we chose gold as the quenching metal and used a chrome adhesion layer between the gold and the diamond. For the tips employed in this work, we deposited 5 nm of gold and 5 nm of chrome.

2.7.8 Explanation of topographic features in Fig. 2.5.1

The features observed in Fig. 2.5.1a of the main text were governed by direct fluorescence quenching through metallic objects (as highlighted by the red square in the figure) and by a confluence of the distance-dependence of the NV fluorescence with topographic features on the sample (bright, ring-shaped feature in the figure). When approaching the NV to our metallic sample-surface, the total NV fluorescence collected in the far-field through the pillar changed as shown in the measurement in Fig. 2.7.5b. This well-known $\begin{bmatrix} 21 \end{bmatrix}$ variation of NV fluorescence is a result of the variable electromagnetic density of states in the vicinity of metallic or dielectric interfaces, which influences the NV radiative lifetime as well as the total effective laser excitation intensity impinging on the NV center. During our scanning experiments, the topography causes the mean distance between the scanning NV center in the nanopillar and the metallic substrate to vary, which in turn causes variations in the collected NV fluorescence rate. Assuming to first order that the metallic tip does not itself affect NV fluorescence (so long as it is not placed in direct contact to the NV center as in the "red-square region"), one can understand most features observed in Fig. 2.5.1a as a pure effect of topography. Based on this principle, in Fig. 2.7.5 we reconstruct the data in Fig. 2.5.1a from a measurement of sample topography (a)





quenching. (a) AFM topography recorded during the experiment presented in Fig. 2.5.1 (same data as shown in Fig. 2.5.1a). (b) "Approach-curve" of the far-field NV florescence rate as the nanopillar with the NV center was approached to the sample. "z=0" was defined as the point of AFM contact (leftmost data-point) (c) Total fluorescence image reconstructed form the datasets in (a) and (b): Looking up the NV fluorescence intensity in (b) for every tip-sample displacement measured in (a) yields the reconstructed topographic features shown in the panel. We note that here, fluorescence quenching is solely induced by the sample surface, while the tip merely acts as a "spacer" between NV center and sample. Fluorescence-quenching that is directly induced by the tip (additional dark spot in d) is not reproduced here. (d) Original data (same data as Fig. 2.5.1a). The features common to (c) and (d) are attributed to effects of sample topography. The additional, dark feature in the center of (d) (red square in Fig. 2.5.1a) has no correspondence in topography and stems from direct fluorescence quenching of the NV center on the sharp metallic tip.

and an independently acquired fluorescence "approach-curve" (b), characteristic for fluorescence quenching of an emitter approaching a metallic surface [10]. The reconstructed image (Fig. 2.7.5c) was obtained by taking the value of the AFM z-displacement for each point in the scan in Fig. 2.7.5a and looking up the corresponding fluorescence-rate obtained in the approach-curve. The resulting image shows striking similarity with the actually measured data (Fig. 2.7.5e; same data as Fig. 2.5.1a) and supports the validity of our explanation.

3 Nanoscale magnetic imaging of a single electron spin under ambient conditions

3.1 INTRODUCTION

The detection of ensembles of spins under ambient conditions has revolutionized the biological, chemical, and physical sciences through magnetic resonance imaging [54] and nuclear magnetic resonance [7, 77]. Pushing sensing capabilities to the individual-spin level would enable unprecedented applications such as single molecule structural imaging; however, the weak magnetic fields from single spins are undetectable by conventional far-field resonance techniques [28]. In recent years, there has been a considerable effort to develop nanoscale scanning magnetometers [12, 41, 55, 99], which are able to measure fewer spins by bringing the sensor in close proximity to its target. The most

sensitive of these magnetometers generally require low temperatures for operation, but measuring under ambient conditions (standard temperature and pressure) is critical for many imaging applications, particularly in biological systems. Here we demonstrate detection and nanoscale imaging of the magnetic field from a single electron spin under ambient conditions using a scanning nitrogen-vacancy (NV) magnetometer. Real-space, quantitative magnetic-field images are obtained by deterministically scanning our NV magnetometer 50 nanometers above a target electron spin, while measuring the local magnetic field using dynamically decoupled magnetometry protocols. This single-spin detection capability could enable single-spin magnetic resonance imaging of electron spins on the nano- and atomic scales and opens the door for unique applications such as mechanical quantum state transfer.

To date, the magnetic fields from single electron spins have only been imaged under extreme conditions (ultralow temperatures and high vacuum), with data integration times on the order of days [83]. Magnetometers based on negatively charged nitrogen-vacancy (NV) centers in diamond have been proposed as sensors capable of measuring individual spins [4, 18, 58, 88] because they can be initialized and read-out optically [30] and have long coherence times [5], even under ambient conditions. Moreover, since NV centers are atomic in size, they offer significant advantages in magnetic resolution and sensing capabilities if they can be brought in close proximity of targets to be measured. Recent advances in diamond nanofabrication have allowed for the creation of robust scanning probes that host individual NV centers within roughly 25 nm of their tips [52]. Here, we employ such a scanning NV center to image the magnetic dipole field of a single target electron spin.

3.2 Scanning NV magnetometer

As described in Chapter 2, our scanning NV magnetometer (Fig. 3.2.1a) consists of a combined confocal and atomic force microscope (AFM), which hosts a sensing NV center embedded in a diamond nanopillar scanning probe tip [52].

The sensor NV's spin-state is initialized optically and read out through spin-dependent fluorescence, while its position relative to the sample is controlled through atomic-force feedback between the tip and sample. Microwaves (MWs) are used to coherently manipulate the sensor NV spin. Magnetic sensing is achieved by measuring the NV spin's optically detected electron spin resonance (ESR), either by continuously applying near-resonant MW radiation (Fig. 3.2.1b) or through pulsed spin-manipulation schemes [58, 88], (Fig. 3.2.1c), where the sensor NV spin precesses under the influence of its local magnetic field (projected along the NV center's crystallographic orientation). We measure the contribution of the magnetic field from a target electron spin to this precession. The entire system, including both the scanning NV magnetometer and the target sample, operates under ambient conditions.

3.3 ISOLATED NV SPIN AS A TARGET SINGLE ELECTRON SPIN

To verify the single-spin detection and imaging, we choose our target to be the spin associated with an additional negatively charged NV center in a separate diamond crystal (so that the sensor and target NV centers can be scanned relative to one another). The advantage of using an NV target is that both its location and spin state can be independently determined by its optical fluorescence. As discussed below, we can thus compare the target NV's magnetically measured location to its optically measured location and ensure that the magnetic image is from a single targeted spin. Additionally, we can guarantee that the target spin is initialized and properly modulated, as is useful for optimizing AC magnetic sensing.

To isolate single NV targets for imaging, NV centers are created in a shallow (< 25 nm) layer of a bulk diamond through established implantation and annealing techniques (see Supp. 3.7.2). The target diamond surface is structured to create nanoscale mesas, whose diameters ($\sim 200 \text{ nm}$) are chosen to contain,



Figure 3.2.1: Scanning NV magnetometer. (a) Conceptual schematic of the scanning NV magnetometer. The sensor NV is hosted within a scanning diamond nanopillar [52], where its spin is initialized and read-out optically from above (532 nm excitation laser spot shown). Coherent NV spin manipulations are performed via a nearby microwave (MW) coil, in this work operating near resonance with the $|0\rangle$ or $|-1\rangle$ transition, in the presence of a static applied magnetic field (not shown). The sensor NV is scanned over target spins of interest to construct magnetic field images. (b) By continuously applying the excitation and sweeping the MWs across the $|o\rangle$ to $|-1\rangle$ transition, optically detected magnetic resonance provides a measure of the static magnetic field at the NV center, with a DC sensitivity of $\sim 2\mu T/\sqrt{Hz}$. (c) By dynamically decoupling the sensor NV spin from its environment, the sensor's magnetic field sensitivity is dramatically improved for AC magnetic fields. Plotted are a spin-echo (1-pulse) magnetometry sequence with a $40 - \mu$ s total evolution time, and a 512-pulse XY8 [31] magnetometry sequence (see Supp. 3.7.1) with $_{330} - \mu$ s total evolution time, which achieve magnetic field sensitivities of 56 nT/ $\sqrt{\text{Hz}}$ and 18 nT/ $\sqrt{\text{Hz}}$, respectively.

on average, a single NV spin. Mesas with single NV centers (as determined through photon auto-correlation experiments, Supp. 3.7.4) are chosen for our measurements. In order to individually control the target and sensor NV spins, we choose a target NV center with a different crystallographic orientation (which determines the spin quantization axis) from the sensor NV, so that their spin transitions can be spectrally separated in ESR measurements by applying a uniform static magnetic field.

Spatial features in the collected fluorescence from scanning the NV magnetometer over target diamond mesas allow us to independently determine the relative positions of the sensor and target NV spins (Fig. 3.3.1). Firstly, the scanning diamond nanopillar acts as a waveguide [3] which, when centered precisely above the target NV, provides efficient collection of fluorescence from the target NV (in addition to the sensor NV). Also, the sensor NV's fluorescence is efficiently coupled into the target bulk diamond when it is centered on a mesa, due to the diamond's high refractive index. The intersection of these two near-field fluorescence features indicates where the sensor NV spin is closest to the target NV spin. This spatial location is later used to confirm the location of the magnetically imaged target NV spin.

3.4 SINGLE SPIN DETECTION SCHEME

Near the expected location of the target, the local magnetic field is measured with a magnetometry pulse sequence performed on the sensor NV using a combination of dynamic decoupling [16] and double electron-electron resonance [47]. The sensor NV spin is prepared in a superposition of spin states, where it accumulates phase proportional to the local magnetic field, including contributions from the target NV spin. To optimize magnetic field sensitivity, the sensor NV is dynamically decoupled from fluctuating magnetic fields in its environment (Fig. 3.4.1, upper panel) through the repeated application of MW π -pulses. Normally, this pulse sequence would also remove any magnetic signal



Figure 3.3.1: Independent determination of a target spin's location.

The sensor NV's diamond nanopillar is scanned over a target nanostructure ("mesa") containing a single target NV center. The combined NV fluorescence is recorded as a function of position (center panel). The fluorescence has a strong spatial dependence because (i) sensor NV fluorescence can partially couple into the target bulk diamond when the sensor NV is close to the sample surface, and (ii) target NV fluorescence can couple into the nanopillar waveguide when the nanopillar is located above of the target NV. When the nanopillar is located away from the target NV, only fluorescence from the sensor NV is collected, as indicated by ESR measurements showing two spectral peaks corresponding to the sensor NV spin $m_S = o \leftrightarrow \pm 1$ transitions (left panel). For ESR measurements taken with the nanopillar located above the target NV (right panel) there are four observable spectral peaks that correspond to both the sensor and target NV spin transitions (blue and red, respectively), with reduced ESR contrast due to collecting fluorescence from both NV spins. The center of the target-coupling circle (red dashed circle around bright fluorescence spot) indicates the lateral location of the target NV spin relative to the center of the nanopillar. Similarly, the center of the sensorquenching circle (blue dashed circle around dark fluorescence spot) indicates the lateral location of the sensor NV spin. With both NV spins' lateral locations known, the position of sensor-target closest approach can be ascertained (green dot).

from a static target spin, but we also simultaneously invert the target NV spin in phase with the π -pulses applied to the sensor NV spin (Fig. 3.4.1, lower panel) to maintain the sign of phase accumulation by the sensor spin due to the target NV spin. The total acquired phase is converted to a population difference, which is measured via NV spin-dependent fluorescence.

With the sensor NV in close proximity to the target diamond surface, the field sensitivity of the sensor NV is reduced, because sensor NV fluorescence is partially emitted into the target bulk diamond (due to its high refractive index), and the target NV adds background fluorescence to magnetic measurements. Because of these effects, our sensor NV's magnetic field sensitivity at closest approach to the target NV is somewhat reduced to approximately 96 nT/ $\sqrt{\text{Hz}}$ (with a 32- π -pulse XY8 [31] decoupling scheme and a 40- μ s total phase accumulation time; Supp. 3.7.8). Since the target NV is embedded in bulk diamond, the sensor-to-target vertical separation is roughly twice the distance between the sensor NV and the diamond surface. Thus, for our magnetic field imaging of a single target NV spin, we expect a \sim 50-nm sensor-target vertical separation, which results in a magnetic field of about 10 nT at the sensor NV location.

3.5 NANOSCALE IMAGING OF THE MAGNETIC FIELD FROM A SINGLE ELECTRON SPIN

A magnetic field image centered at the expected target spin location is acquired by averaging the sensor's NV fluorescence in multiple scans of the NV magnetometer across a $\sim 200 \times 200$ -nm field-of-view (taken using a lateral drift correction scheme detailed in Supp. 3.7.6). A normalization scheme is applied to the magnetometry, where we alternately initialize the target NV spin in the $|0\rangle$ state and the $|-1\rangle$ state and measure the equal and opposite phase shifts induced during the sensor NV's magnetometry sequence (Supp. 3.7.5). We subtract the measured NV fluorescence rates for these two initial target NV spin polarizations,



Figure 3.4.1: Single spin detection scheme and target spin modulation verification. To detect the magnetic field from the target NV spin, the sensor NV spin (top panel, blue arrow) is placed in a superposition of spin states with a MW $\frac{\pi}{2}$ pulse (around the X axis). It then evolves under the influence of the magnetic field from the target spin (red arrow), accumulating phase (shaded blue region, whose extent is visually exaggerated for visibility). To optimize the sensor spin's magnetic sensitivity, it is dynamically decoupled from its environment (see Supp. 3.7.1) [16] by the repeated application of MW π -pulses using an XY8 sequence [31]. In order to magnetically measure the target NV spin, it is inverted, synchronously with the π -pulses applied to the sensor NV, so that phase shifts induced on the sensor by the target spin constructively accumulate. (The target NV spin is modulated between the $m_S = o$ and $m_S = -1$ states to isolate an effective spin- $\frac{1}{2}$ system from the target NV's spin triplet.) To invert the target NV spin with high fidelity, we employ adiabatic fast passages (see Supp. 3.7.3). Plotted is the measured fluorescence for pulses 1, 2, 127, and 128, indicating that the target NV spin can be modulated many times without substantial polarization decay. The sensor NV's accumulated phase is converted to a population difference using a final $\frac{\pi}{2}$ pulse, whose axis (Y) is chosen to maximize sensitivity to small magnetic fields.

which isolates the magnetic field signal from the target spin (Supp. 3.7.7).

Near the center of the magnetometry scan, we observe a drop in the normalized fluorescence from the magnetometry sequence that is well beyond the uncertainty set by the measurement's noise level and is consistent with the effect of a single target NV spin's magnetic field on the sensor NV. The complete magnetic field image clearly indicates the presence and location of the target NV spin (Fig. 3.5.1a). This single electron spin detection is confirmed by repeating the measurement with a spatial linecut of magnetometry measurements (Fig. 3.5.1b), with a resulting magnetic response that fits well to a vertical separation of 51.1 \pm 2.0 nm between the sensor and target NV centers. (Errors are determined from the χ^2 of the fit as a function of distance, where the sensor-to-target displacement is the only free-parameter and the orientations of both NV spins are independently measured using ESR.) The measured fluorescence difference is converted to a magnetic field at the sensor NV (peak value of 8.6 nT, Fig. 3.5.1b) by using the sensor NV spin's independently calibrated magnetic field response and fluorescence rate. Both scanning magnetometry measurements are in good agreement with simulations of the sensor NV's response to the magnetic field from a single electron spin at a vertical distance of 51 nm (Fig. 3.5.1c). Thus the above measurements are consistent and confirm the detection and nanoscale imaging of the single target spin.

In the demonstrated magnetic field imaging, single-spin measurements with a signal-to-noise ratio (SNR) of one can be acquired in 2.3 minutes. The data for both single-spin measurements presented in Fig. 3.5.1 have been integrated for a total time of 42 minutes per point, yielding an SNR of 4.3. This integration time is consistent with the measured target NV spin magnetic field (8.6 nT) and sensor NV magnetic field sensitivity calculated by assuming the noise is dominated by photon shot noise (96 nT/ \sqrt{Hz}).

Figure 3.5.1: Single-spin magnetic imaging. (a) Magnetic field image of a target NV spin near the surface of a diamond mesa, acquired with the scanning NV magnetometer. While repeatedly running an AC magnetometry pulse sequence (here with a 32-pulse XY8 sequence, with 40 μ s of total evolution time), the sensor NV is laterally scanned over the target, and the fluorescence rates for the target spin starting in the $|o\rangle$ state as well as the $|-1\rangle$ state are independently recorded. Plotted is the difference between these measurements, which depends only on the sensor NV's magnetic interaction with the target spin and not on background fluorescence variations (see Supp. 3.7.7). The pronounced drop in fluorescence near the center of the image indicates a detected single electron spin. (b). An independent magnetometry linecut taken along the green arrow confirms the single spin imaging, which has an intensity and width consistent with the recorded image. The measured fluorescence difference is converted to the measured magnetic field using the sensor NV's calibrated field sensitivity and fluorescent rate (see Supp. 3.7.8). (c) Simulated fluorescence due to a target spin. With only the sensor-target displacement as a free parameter, the spin signal is simulated, which agrees well with both the spin image and the linecut for a vertical distance of 51 nm (the fit in (b) and the image in (c) have the same parameters). (d) If the sensortarget vertical distance can be moderately reduced, the quality of single-spin imaging will be dramatically improved. Plotted are simulated lateral magnetic field contours from a single target electron spin for different sensor-target vertical separations where each contour indicates an increase of signal-to-noise by one for a 100 second integration time. At 50 nm (the current condition), there is only one contour, indicating single-spin imaging with a signal-to-noise of one; however at 10 nm, a signal-to-noise of roughly 100 is possible, such that many contours and the dipole lobes of the target spin are clearly observable.



Figure 3.5.1 (Continued)

3.6 Discussion and outlook

By successfully measuring the magnetic field from a single target NV spin, our spin-sensing protocols have been confirmed, enabling otherwise undetectable "dark" electron spins to be detected with confidence. Note that imaging dark spins rather than an NV target could potentially be performed with higher sensitivity, because optical fluorescence from the sensor NV could be better collected and isolated. In addition, for dark spins on or near a sample surface rather than embedded beneath it, the sensor-to-target separation would be reduced by a factor of about two: thus the SNR for magnetic field imaging would increase by nearly an order of magnitude (Fig. 3.5.1d) because dipolar fields decay as $1/r^3$. Moreover, the required measurement time for a given SNR scales with the sixth power of sensor-to-target separation for a shot-noise-limited measurement: e.g. at a separation of 25 nm, a target surface spin would be detectable in two seconds (with our instrument's demonstrated sensitivity and a SNR of one).

For target spins of interest that cannot be initialized, the variance of the magnetic field at the sensor NV could instead be measured, which is detectable with nearly the same sensitivity as the field itself if an appreciable amount of phase can be acquired [88]. For instance, at a 25 nm sensor-to-target distance, with a phase evolution time of 100 μ s, an uninitialized, driven spin could be detected within two seconds of integration time (SNR of one, for the same sensor NV spin-dependent fluorescence rate and contrast as in the demonstrated spin imaging; Supp. 3.7.9). For the phase-evolution time used in the demonstrated single-spin imaging (40 μ s), the integration time would be 20 s.

If the coherent sensor-target coupling is strong enough for more than 2π of sensor NV phase to be accumulated during magnetic field measurements, then phase-estimation techniques can be employed, thus allowing the measurement noise to decrease linearly in time [67, 92], and potentially offering a great boost in speed to magnetic imaging. Moreover, if a target spin can be initialized and has a coherence time as long as the sensor NV, then the target and sensor spins could

be entangled. Combined with long-lived storage techniques for quantum states [57], the ability to entangle a scanning sensor and target spins could allow for mechanical transfer of quantum information between solid-state spins.

3.7 SUPPLEMENTARY SECTION

3.7.1 Dynamic decoupling of the sensor NV spin

The sensor NV spin coherence time is prolonged by dynamically decoupling it from its noisy environment [8, 16, 17]. This is achieved by the repeated application of microwave (MW) π -pulses, which causes the effects of slowly fluctuating magnetic fields to re-phase and cancel out. To apply a large number of pulses without scrambling the sensor NV spin state, the control pulses are carefully calibrated to within 2% using a boot-strap tomography scheme [?]. For the dynamic decoupling scheme and magnetometry, we employ an XY8 sequence [31], which uses π -pulses around two orthogonal axes on the equator of the sensor NV's Bloch sphere to minimize the accumulation of pulse errors. This sequence ($\pi_x - \pi_y - \pi_x - \pi_y - \pi_y - \pi_x - \pi_y - \pi_x$) is repeated as many times as possible to maximize the sensor NV's magnetometry sensitivity, which – as a function of the number of pulses – is a compromise between the extended NV coherence from the decoupling and the reduced contrast from accumulated pulse errors.

MW fields are supplied from a Rhode and Schwarz SMB100A signal generator. MW phase control is achieved using an IQ mixer (Marki-1545) with pulsed analog inputs on the I and Q ports supplied by an arbitrary waveform generator (Tektronix AWG5000). NV spin Rabi frequencies in this work are 15-20 MHz, with typical π -pulse durations of 30 ns.

3.7.2 NV CENTER TARGETS

NV centers were created through implantation of ¹⁵N ions [76] into ultrapure diamond (Element Six, electronic grade diamond, < 5 ppb nitrogen). The

implantation was done at an energy of 6 keV with a dose of $2.5 \cdot 10^{11}$ cm⁻² to give a nominal nitrogen depth of 10 nm [98]. To form NV centers, the sample was annealed in vacuum at 800 °C, where existing vacancies are mobile and can pair with the implanted nitrogen atoms. The resulting density of NV centers corresponds to one center every ~ 200 nm, forming a layer within ~ 25 nm from the surface. To isolate NV centers, we selectively etched [48] away most of the shallow diamond surface layer, leaving individual NV-containing nanostructures. This was done using electron-beam lithography to define an etch mask from a flowable oxide [34] (Dow Corning, XR-1541). A reactive-ion etch then removed any exposed diamond surfaces, resulting in shallow diamond nanostructures (200 nm across) containing, on average, single NV centers. The spacing between the structures are chosen to be 3 μ m, sufficiently larger than the size of focused laser spot, which ensures optical measurements of only one structure at a time, We then, perform photon autocorrelation measurement (see Supp. 3.7.4) with each fluorescent diamond structure to find the ones with single NV centers.

3.7.3 Adiabatic fast passages for controlling the target NV spin

To control the target NV spin with high fidelity over numerous spin inversions, we employ adiabatic fast passages. The spin-state is prepared optically in the mS = |0> state, and microwaves (MW) with bare Rabi frequency ω_R are applied and detuned by $\delta(t = 0)$ from the target NV transition. The detuning is ramped through zero to $-\delta(t = 0)$ over a pulse time, T_p . At any point in time, the target NV spin in the rotating frame prececess around an effective magnetic field Ω_R , which is the sum of the MW field and the remaining static magnetic field in the rotating frame resulting from the non-zero MW detuning. If the angular velocity, $d\theta/dt$ of Ω_R is slow compared to ω_R , then the NV spin-state is effectively locked to the motion of this effective magnetic field as it moves from $|0\rangle$ to $|-1\rangle$. In general, it is advantageous to sweep the detuning non-linearly in time and spend most of TP when the NV spin is near the equal population state where it is most susceptible to dephasing [27]. To achieve this, we ramp the detuning to keep the rate of change of the spin's angle with respect to the $|0\rangle$ state constant, so that:

$$\delta(t) = \omega_R \tan(\beta(\frac{t}{2T_p} - 1)) \tag{3.1}$$

where β is chosen to achieve the desired sweep range. For the adiabatic fast passages presented in Fig. 3.4.1 of the main text, $T_p = 300$ ns, $\delta(t = 0) = 100$ MHz, and $\omega_R = 17$ MHz.

The detuning ramping is implemented by using an arbitrary waveform generator to output a sinusoid at a frequency of the desired detuning, which is mixed with a continuous-wave MW source (all are the same make and model as the sensor-addressing MW equipment). By setting the phase of this sinusoid to be the integral of the detuning as a function of time, the mixed MW frequency can be continuously varied.



Figure 3.7.1: Photon-autocorrelation measurements for the target NV center.

3.7.4 Photon-autocorrelation measurements for the target NV center

Photon-autocorrelation measurements for the target NV in the absence of the sensor NV (Fig. 3.7.1) give $g^2(\tau = 0) < 0.5$, ensuring that a single target NV spin lies in the diamond mesa on which magnetic field imaging is performed with the scanning NV magnetometer. No background subtraction was performed on the data and normalization was performed based on count-rates on the individual detectors as well as the time-binning in the photon-correlation hardware.

3.7.5 MAGNETOMETRY NORMALIZATION SCHEME

To isolate the sensor NV's spin-state-dependent fluorescence from spatially varying background fluorescence, we employ a normalization scheme that involves alternatively performing two slightly different magnetometry pulse sequences, and then subtracting the measured NV fluorescence rates (Fig. 3.7.2). These two sequences are similar to the double-electron-electron resonance



Figure 3.7.2: Magnetometry normalization scheme.

scheme presented in Fig. 3.7.3, except that one target NV spin-inversion (performed via an adiabatic fast passage) is removed from each sequence: in Sequence 1, the last spin inversion is removed, while in Sequence 2 it is the first spin inversion. By removing spin inversions in this manner, we ensure that the sensor NV (blue arrow) acquires an equal and opposite target-spin-induced phase shift (Φ , shaded blue region) during the two pulse sequences, because the target spin-state is inverted in the two sequences for the majority of the phase evolution time. Crucially, the target NV spin ends in the same state for the two pulse sequences (here, $|-1\rangle$). Thus subtracting the measured NV fluorescence rates for the two pulse sequences removes the contribution of background fluorescence from the NV target, which has a non-trivial spatial dependence (Fig. 3.3.1). Moreover, both pulse sequences have the same number of target spin inversions, which alleviates unavoidable spin-polarization losses associated with flipping the target spin and cross-talk between the applied MW sources. In this scheme, a small amount of integration time is unused for sensor NV phase accumulation (the portions next to the $\frac{\pi}{2}$ pulses at the beginning and end of each sequence cancel, equal to one delay period τ between π pulses). However in the limit of a large number of spin inversions, this loss of integration time is negligible.



Figure 3.7.3: Magnetic field image acquisition protocol.

3.7.6 MAGNETIC FIELD IMAGE ACQUISITION PROTOCOL

Scanning an AFM for long periods of time with nanometer precision can be difficult to achieve under ambient conditions because of thermal-induced drifts. Temperature fluctuations on the order of a fraction of a degree can lead to tens of nanometers of relative motion between the sensor and target NV spins, which would considerably smear out our magnetic field imaging. These drifts generally occur on long time-scales, with a few nanometers of drift every hour. To minimize their effect, we employ an image acquisition protocol that periodically corrects for sensor-to-target drifts, as described in Figure 3.7.3.

The protocol has three major components: (1) Drift-correction using the sample topography to determine the target's location (top panel); (2) Taking a relatively quick magnetic scan over the target spin location (lower right panel); and (3) Checking to make sure that the MW pulsing has not degraded over time, and that the sensor NV's magnetic sensitivity has not been significantly compromised (lower left panel).

Drift correction is performed by scanning over the target-containing diamond

mesa, and the measured topography is the convolution of this mesa (\sim 200 nm in diameter) with the diamond nanopillar scanning tip (\sim 200 nm in diameter). From this topography, and the simultaneously measured fluorescence (as in Fig. 3.3.1), the target NV spin can be located (green dot) and an appropriate scan range can be defined (green square). The topography of successive scans (taken after both magnetometry and diagnostic measurements), can be compared to the first reference scan by cross-correlation, and thus drifts can be corrected between scans. In the single-spin measurements presented in Fig. 3.5.1, we observe a mean variation of 5 nm between successive scans (limited by the pixel size of the reference scan), indicating that magnetic field images can be overlapped with roughly 5-nm precision.

After zooming into the appropriate scan region, where the expected target spin NV lies in the center of the scan range, magnetic field images are acquired while simultaneously alternating between the two magnetic detection pulse sequences (Fig. 3.7.2) and monitoring their fluorescence rates (only one sequence is illustrated). Each scan is integrated for roughly 30 minutes to minimize the drifts between scans.

When a magnetic scan is finished, the sensor NV is placed at the approximate measurement position to measure the optimal sensitivity to the target NV and as a function of time. In general, the sensor NV can slowly drift in and out of the green laser confocal spot, causing variations in the overall detected NV fluorescence. Additionally, the power of the MW source can drift, which can decrease the performance of dynamic decoupling and magnetometry pulse sequences. Magnetic field sensitivity is experimentally determined by running the magnetometry sequence, with the phase of the last $\frac{\pi}{2}$ pulse set at $\pm \frac{\pi}{2}$ (red and brown data points, respectively) to measure the $\langle -\sigma_y \rangle$ and $\langle \sigma_y \rangle$ projections of the sensor NV. The difference between these measurements gives the contrast and counts of the sensor NV's magnetic response, and when combined with the phase accumulation time, determines the magnetic field sensitivity of the sensor NV. To differentiate overall NV fluorescence rate changes from pulsing performance changes, we also measure fluorescence counts for the $|-1\rangle$ state



Figure 3.7.4: Magnetic field image data processing.

(black data points), which should overlap with the $-\sigma_y$ measurement in the case of no pulse errors or dephasing.

After these pulse diagnostics, we zoom out to measure the topography of the sample again, completing a measurement cycle. This procedure is repeated until a desired signal-to-noise in the magnetic field image has been achieved.

3.7.7 MAGNETIC FIELD IMAGE DATA PROCESSING

The multiple images taken during a magnetic field scan for each magnetometry sequence (Fig. 3.7.4, left column) are averaged (without any further spatial correction), to yield the average fluorescence map (Fig. 3.7.4, center left column). In these averaged measurements, there are large variations in fluorescence due to near field coupling into and out of the diamond nanopillar for the target and sensor NV centers, as described in Fig. 3.3.1. These variations are quite large (\sim 150 CPS) compared to the expected effect of a single target NV's magnetic field on the sensor NV signal (fluorescence change \sim 4 CPS under inversion of

the target NV spin). Subtracting the average fluorescence maps of the two magnetometry sequences yields a difference fluorescence signal free of the large background fluorescence (Fig. 3.7.4, center right). In general, the difference in fluorescence between the two magnetometry sequences has a small remaining offset, and so it has a mean of a few counts per second, even in the absence of the target NV. This is likely due to a small amount of cross-talk between the target-addressing MW and the sensor NV, which is slightly different between the two sequences. During our pulsing diagnostics in the measurement acquisition, we measure this fluorescence offset with the sensor NV very far from the sample $(> 1\mu m)$, and we subtract this value from the difference fluorescence, which yields a mean of zero counts per second away from the target NV spin. As long as this remainder fluorescence (4.3 CPS for this image) times the percent fluorescence variations across the scan region (15%) is smaller than the target NV spin signal - as is the case here - then the target NV spin signal will be the largest feature in the difference fluorescence map. To increase the signal-to-noise ratio (SNR) of magnetic field imaging, we average multiple pixels together to coarse-grain the scan (Fig. 3.7.4, right panel), yielding a scan with 64 pixels across a field-of-view of \sim 200 imes 200 nm, with 42 minutes of integration time per pixel providing average SNR of 4.3.

3.7.8 Simulation of single-spin magnetic field imaging

The response of the scanning NV magnetometer to a single electronic spin is simulated by considering an electron spin at the origin of a coordinate system with a quantization axis oriented along the $[x, y, z] = [o, -\sqrt{2}, 1]$ direction (to match the orientation in the sample that is measured through ESR measurements using a three-axis Helmholtz coils). Equi-field contours of this target spin's magnetic field are plotted as a function of three-dimensional space (Fig. 3.7.5, displayed here are 2 nT, 4 nT, and 6 nT). Because the sensor NV is first-order sensitive only to magnetic fields along its quantization axis $([x, y, z] = [o, \sqrt{2}, 1])$, the plotted field contours from the target spin have been



Figure 3.7.5: Simulation of single-spin magnetic field imaging.

projected along the sensor NV quantization axis, which yields the dipole field lobe pattern shown here. Experimental magnetic field scans are taken as plane-cuts of this dipole field pattern above the location of the target spin (at an a-priori unknown distance; plotted is the best fit value of 51 nm). The simulated magnetic field profile is converted into a spatial map of sensor NV fluorescence rate using measured values of the sensor NV fluorescence rate, contrast, and phase evolution time, giving a magnetic field sensitivity conversion factor of -1.8 nT per count per second.

3.7.9 Measuring the variance of a non-initialized spin

If target electron spins cannot be initialized (unlike the target NV spin measured in this work), then the spin's magnetic field will average out to zero over multiple measurements, as at the start of a given measurement ("shot") the target spin has an equal probability of being either up or down. However, if the target's spin's



Figure 3.7.6: Measuring the variance of a non-initialized spin.
longitudinal relaxation time is much longer than the magnetometry phase-evolution time τ , within a single shot, then the target spin maintains its statistical polarization, and a net phase shift will be accumulated by the sensor spin. By choosing the axis of rotation of the final $\frac{\pi}{2}$ pulse to match the axis of the first $\frac{\pi}{2}$ pulse, when the accumulated phase shifts from multiple measurement shots are converted to a net population difference, the effect of the target spin's magnetic field no longer cancels out and can be measured via the sensor NV's spin-dependent fluorescence. This scheme effectively measures the variance of the target spin polarization ($\langle \sigma_z^2 \rangle - \langle \sigma_z \rangle^2$, for the thermal state of a target spin) instead of its mean polarization ($\langle \sigma_z \rangle$)

Plotted in Figure 3.7.6 is the sensor NV's response for $\tau = 100\mu s$, plotted is the sensor's NV response to a (driven) target electron spin with random polarization (either up or down) at the measurement's start. (The sensor NV's fluorescence and spin-dependent contrast used are those demonstrated in spin-imaging; Supplementary Fig. 3.7.5.). The magnetic field profile for this driven target spin is a square wave with amplitude B_{max} , which is synchronized to the sensor NV's decoupling scheme. For a sensor-to-target distance of 25 nm (and the same sensor and target spin-quantization axes used in the present work; Supplementary Fig. 5), $B_{max} = 74$ nT, which gives a signal of 25 CPS with respect to the $|0\rangle$ state. Within two seconds of integration time, this signal divided by the measurement's shot noise gives a signal to noise ratio of one.

4

Quantum control of proximal spins using nanoscale magnetic resonance imaging

4.1 INTRODUCTION

Quantum control of individual spins in condensed matter systems is an emerging field with wide-ranging applications in spintronics [1], quantum computation [65], and sensitive magnetometry [13]. Recent experiments have demonstrated the ability to address and manipulate single electron spins through either optical [37, 95] or electrical techniques [24, 33, 61]. However, it is a challenge to extend individual spin control to nanoscale multi-electron systems, as individual spins are often irresolvable with existing methods. Here we demonstrate that coherent individual spin control can be achieved with few-nm resolution for proximal electron spins by performing single-spin magnetic

resonance imaging (MRI), which is realized via a scanning magnetic field gradient that is both strong enough to achieve nanometric spatial resolution and sufficiently stable for coherent spin manipulations. We apply this scanning field-gradient MRI technique to electronic spins in nitrogen-vacancy (NV) centers in diamond and achieve nanometric resolution in imaging, characterization, and manipulation of individual spins. For NV centers, our results in individual spin control demonstrate an improvement of nearly two orders of magnitude in spatial resolution compared to conventional optical diffraction-limited techniques. This scanning-field-gradient microscope enables a wide range of applications including materials characterization, spin entanglement, and nanoscale magnetometry.

Magnetic field gradients allow for spatially distinguishing spins in ensembles, as fields locally modify the spins' resonance frequencies. Spatially separated spins can therefore be addressed selectively, allowing for magnetic resonance imaging (MRI), which has revolutionized the medical and biological sciences by yielding micron-scale imaging of nuclear spins [49, 54]. Performing MRI on single spins with high spatial resolution is attractive both for determining structure on the molecular scale and for achieving individual spin quantum control in ensemble systems. With conventional MRI techniques, however, it is difficult to improve the spatial resolution to the nanoscale due to insufficient readout-sensitivity and inadequate magnetic field gradients [28]. Recently, magnetic field gradients introduced via scanning probe techniques have enabled single spin detection with few-nm resolution [4, 83]; however, control and characterization of individual spins in nanoscale clusters has not been demonstrated thus far.

Here we perform scanning field-gradient MRI on proximal electron spins in nanoscale ensembles and demonstrate a spatial resolution < 10 nm under ambient conditions. We show that scanning field-gradient microscopy not only allows for imaging but further provides quantum spin control for characterization and manipulation of individual spins on the nanoscale. By pushing the spatial resolution to few-nm length scales, our results illustrate that quantum control of individual spins can be maintained in dense ensembles of

spins (with spins separated by a few tens of nanometers), where the mutual coupling between adjacent spins can exceed their decoherence rates $[64](1/T_2)$. Thus, scanning-field-gradient MRI will help facilitate the creation of entangled spin-states with applications for quantum information processing and sensitive, nanoscale magnetometry. Although our approach is applicable to any spin system where spins can be initialized and read out, we focus here on the electronic spins associated with NV centers in diamond, where spin initialization and readout can be performed optically [30]. In addition, NV spins are attractive for performing quantum information processing [22, 64] and sensitive magnetometry [4, 18, 58, 88]. Individual NV spin control in a nanoscale ensemble is a key advance towards the implementation of these applications.

4.2 MAGNETIC RESONANCE IMAGING OF A SINGLE NV CENTER US-ING THE SCANNING-FIELD-GRADIENT

Our scanning-field-gradient MRI system (Fig. 4.2.1a) is comprised of an atomic force microscope (AFM) with a magnetic tip and integrated into an optical confocal microscope (see Supp. 4.6.2), all operating under ambient conditions. The magnetic tip used in the experiments is a laser-pulled quartz tip coated with a 25 nm magnetic film (see Supp. 4.6.1). Small ensembles of shallowly implanted NV centers, nominally 10 nm below the surface of a diamond sample, are placed in the confocal spot (volume $< 1 \, \mu m^3$) where an excitation laser at 532 nm is used to initialize and read out the NV centers' spin states. Because of the NV centers' spin-dependent fluorescence ($\approx 630 - 800$ nm), optically detected electron spin resonance (ESR) can be observed by sweeping the frequency of a driving radio-frequency (RF) field through the spin resonance and measuring the corresponding variation in fluorescence (Fig. 4.2.1b) [30], in this case on a single NV center. As has been demonstrated by Balasubramanian et al. [4], a magnetic tip in proximity to an NV center shifts the energy of the NV spins, particularly for fields along the NV axis. By selectively detuning the applied RF field and scanning the magnetic tip over a single NV center, a magnetic field map

corresponding to the applied detuning can be acquired (Fig. 4.2.1c). Using the AFM to regulate the magnetic tip's height allows for precise nanoscale control of the tip's location in three dimensions: the height of the tip is maintained via a conventional feedback loop on the force from the sample, while several-nm precision in lateral positioning is achieved by using the sample's topography as a marker.

4.3 Three-dimensional nanoscale MRI of proximal NV spins

We first demonstrate how this technique can be used for imaging proximal NV spins. The magnetic response shown in Figure 4.2.1c provides a direct means for determining the relative location of proximal NV centers. For NVs with an identical orientation, the indistinguishable nature of their spins leads to an identical but spatially shifted magnetic field map for each NV spin. Thus, magnetic resonance images of single-spins (such as in Figure 4.2.1c) can serve as the point spread function for multi-spin imaging. To demonstrate the performance of this technique, we executed scanning-field-gradient imaging on three closely spaced NV centers (Fig. 4.3.1). As the magnetic tip is scanned laterally across the sample, a magnetic field contour is observed for each center. Here, we selected an ensemble of NVs where all three have a common quantization axis orientation, so that every spin responds in the same way to the presence of the magnetic tip. Thus, the relative distances between NV spins can be obtained by quantifying the spatial shift between the observed resonance rings through a deconvolution procedure (see Supp. 4.6.4). The resulting image (Fig. 4.3.1a) indicates the relative positions of the NV centers, which we find are spaced by 50 nm and 75 nm with respect to NV II. A spatial resolution of 9 nm can be extracted from the width of the resonance ring along the vector connecting NV centers II and III; a precision of 0.2 nm can be determined from a two-dimensional Gaussian fit to the deconvolved peak. This estimate of precision only accounts for variance induced by random noise and does not account for



Figure 4.2.1: Two-dimensional imaging of a single NV center using the scanning-field-gradient MRI microscope. (a) Schematic of the experimental approach. A magnetic tip is scanned by an AFM over a diamond sample containing multiple shallow NV centers separated by distances smaller than the optical excitation wavelength. Each NV center experiences a different magnetic field from the tip. Thus, spin transitions for individual NV centers can be selectively driven by tuning the frequency of an externally applied radio-frequency (RF) field. A confocal microscope provides NV spin-state preparation via optical pumping, and detection via spin-state dependent fluorescence. (b) Optical fluorescence measurement of the electron spin resonance (ESR) of a single NV center in the absence of the magnetic tip. The NV-spindependent fluorescence rate leads to a drop in emitted (and detected) photons when the external RF source is swept onto the NV spin resonance (here at 2.47 GHz instead of the zero-field value of 2.87 GHz because of an applied static magnetic field). (c) Two-dimensional magnetic resonance image of a single NV center, created by scanning the magnetic tip across the surface and fixing the RF frequency off resonance from the NV ESR transition (see arrow in (b)). A reduction of fluorescence is observed for positions of the magnetic tip relative to the NV that put the NV spin on resonance with the RF field, creating a dark "resonance ring". The plotted fluorescence magnitude is normalized to the NV fluorescence when no driving RF field is applied near the ESR transition (see Supp. 4.6.3).

systematic deviations caused by effects such as scanner non-linearity or deviations from the peak shape to the Gaussian fit.

Our single-spin MRI technique can be extended to three dimensions by performing magnetic tomography, where the magnetic tip is retracted from the surface in few-nm steps and scanned laterally (Fig. 4.3.1b). The size of the resonance rings evolves quickly as a function of z-distance as the magnetic field from the tip becomes too weak to bring the spin-transitions into magnetic resonance with the detuned RF driving field. The height differences between the NVs can be seen here as NV I vanishes roughly 15 nm before the other two, which indicates that NV I lies roughly 15 nm further below the surface than both NV II and III. A vertical scan across NVs II and II shows the z-resolution of this measurement to be 10 nm, which is extracted from the width of the resonance line in the z-direction. Since the lateral separation of these two NVs is larger than our spatial resolution, we can determine the relative height of the two NVs with high precision, which we find to be 3 nm.

The NV centers in our demonstration experiments were created through implantation of nitrogen ions forming a layer \approx 10 nm below the diamond surface. Modeling of this implantation procedure [98] predicts a spatial variation of \pm 3 nm; whereas we observe a larger variation in the distribution of NV depths (> 10 nm) using our scanning-field-gradient MRI technique. This discrepancy may arise from ion channeling or surface effects. Nanoscale precision in measuring the distance between proximal NV spins also allows determination of the mutual dipole coupling between adjacent spins (see Supp. 4.6.5), a key component for creating entangled spin-states.



Figure 4.3.1: Three-dimensional magnetic resonance imaging of proxi**mal NV spins.** (a) Scanning the magnetic tip over a cluster of three NV centers with the same crystallographic orientation yields multiple dark-resonance rings in the observed fluorescence, one for each NV center (upper image). The relative locations of the NV centers are extracted from the magnetic resonance image through a deconvolution process (see Supp. 4.6.4), yielding adjacent NV-NV distances of 50 nm and 70 nm (bottom, left image, which plots the normalized deconvolved signal). ESR spectral linecuts (top, right) give a spatial resolution of roughly 9 nm with 0.2 nm precision taken from a twodimensional Gaussian fit to the deconvolved peak (bottom, right). (b) The relative depths of the NV centers below the diamond surface are determined by taking magnetic resonance images for different magnetic tip heights above the sample surface. Comparing the evolution of the three NV resonance rings as a function of tip-to-sample distance, we determine that NV I lies roughly 15 nm below NV II and NV III (stack of images on left). A vertical cut further resolves NV II and NV III, showing that they are 3 nm apart in depth (right image).

4.4 INDIVIDUAL AND SELECTIVE QUANTUM CONTROL AND CHARAC-TERIZATION OF PROXIMAL NV SPINS

Scanning-field-gradient MRI provides not only a method for nanoscale mapping of the spatial locations of proximal spins but also allows for individual spin transitions to be resolved in frequency space with high precision. In the absence of magnetic field gradients, identical spins sharing a quantization axis are indistinguishable, making selective control over proximal spins impossible. Performing scanning-field-gradient MRI on proximal spins differentiates their transition frequencies, allowing for coherent manipulation and characterization of individual spins in the ensemble. This can be done simultaneously on several NV spins while preserving independent control of each spin. Such selective control over proximal spins can be maintained as long as the magnetic field gradient separates the transition frequencies of neighboring spins by more than their resonance linewidth, which is the same condition that determines the spatial resolution in MRI. Therefore, spins separated by more than the achieved MRI spatial resolution (\approx 9 nm for the NV spin experiments reported here) can be addressed independently using our technique.

To demonstrate such selective nanoscale characterization and control of proximal spins we examined a pair of proximal NV centers (NV IV and NV V, separated by 135 nm, see Supp. 4.6.6), which share the same NV axis orientation. In the absence of the magnetic tip, we performed a continuous-wave ESR measurement on both NV spins simultaneously (Fig. 4.4.1a, left panel), finding no difference in their spectra, as expected. After tuning the RF frequency to the pair of NV spins, we drove Rabi oscillations and observed the free induction decay of the two NV spins via a Ramsey sequence (Fig 4.4.1a, right panels). The measured Ramsey fringes show a pronounced beating pattern due to the hyperfine structure of the NV spin transitions and are damped due to inhomogeneous dephasing of the two NV centers. However, information

distinguishing the individual coherence-properties of the proximal NV centers cannot be extracted from these measurements. In contrast, when the magnetic tip is brought in close proximity to the NV spins, it splits their resonance frequencies due to the differing magnetic fields applied to each NV, thereby allowing each NV spin to be addressed and characterized individually and coherently (Fig. 4.4.1b) without modifying the spin of the neighboring NV.

Such coherent individual spin control was realized by tuning the RF frequency to the ESR resonance of the target NV center. Pulsing the RF with a variable duration induced coherent Rabi oscillations of either NV IV or NV V, depending on the tuning of the RF frequency. Similarly, characterization of individual spin coherence properties was achieved by measuring the spin's free-induction decay via a Ramsey sequence using the appropriate RF frequency for the target NV spin. We observed that NV IV has a faster free-induction decay rate than NV V, indicating that the two spins have different inhomogeneous dephasing times, T_{2}^{*} . The measured free-induction decay rate for NV V with and without the magnetic tip are comparable, indicating that additional decoherence induced by the tip is small compared to that spin's ambient dephasing $1/T_2^*$. In order to prevent tip-induced spin-decoherence, we employed a three-dimensional spatial feedback scheme, which ensured that variations in the applied tip-field were smaller than the intrinsic NV inhomogeneous dephasing rates (see Supp. 4.6.7). Thus, the applied magnetic field gradient can be used to characterize individual spin coherence properties, which is a direct consequence of achieving single-spin control.

Figure 4.4.1: Individual quantum control and characterization of proximal NV centers. (a) With the magnetic tip pulled far away from the sample, two NV centers (IV and V) separated by 135 nm (see Supp. 4.6.6) and sharing the same spin-quantization axis cannot be resolved by their ESR spectra $(m_S = o \rightarrow m_S = 1$ transition of the NV triplet groundstate, observed by spin-state dependent fluorescence), as they experience the same static magnetic field (left panel). By driving both NVs at once, Rabi oscillations (middle panel) and the NV spins' collective free-induction decay (right panel) can be observed using the indicated Rabi and Ramsey RF pulse sequences. (b) With the magnetic tip in close proximity to the sample, the two NV spin resonances are spectrally distinguishable by their differing Zeeman shifts (left panel; \approx 30 MHz splitting), allowing each NV to be addressed independently, so that Rabi oscillations (middle panel) and Ramsey free-induction decay (right panel) of each NV spin can be individually measured (NV IV in red and NV V in green). For these measurements, the probability of the spin being in the mS=0 state is plotted.





Figure 4.4.2: Selective, independent RF control of proximal NV spins in the presence of the tip magnetic field gradient. (a) Fluorescence measurements were performed for the same two proximal NV centers as in Figure 4.4.1, while undergoing simultaneous, near-resonant driving by fields RF 1 and RF 2; and with the tip magnetic field gradient inducing a large $(\approx$ 30 MHz) Zeeman frequency shift between the ESR frequencies of the two NV spins. The frequency of RF 1 was set on resonance with NV IV and was pulsed with varying duration (τ_{RF_1}) to induce Rabi oscillations. Simultaneously, RF 2 was continuously applied while its frequency (ω_{RF_2}) was swept through the spin resonance of NV V to measure its ESR spectrum. (b) Twodimensional data set sweeping over τ_{RF_1} and ω_{RF_2} for the two NV spins (base of cube). Individual behavior of each NV spin is extracted by subtracting the mean measured fluorescence for each row (i.e., fixed value of ω_{RF_2}) from the data for that row; and similarly for each column (fixed value of τ_{RF_1}). The resulting extracted data sets (shown in the sides of the cube) are found to be independent of row or column number, showing that varying the Rabi pulse duration τ_{RF_1} for NV IV does not influence the results of sweeping ω_{RF_2} through the spin resonance of NV V, and vice versa. Summing up the measurements for all rows and columns yields the resulting averaged Rabi oscillation measurement for NV IV (top of left wall) and ESR spectrum from NV V (top of right wall), respectively.



Figure 4.4.2 (Continued)

Under the influence of the tip's magnetic field gradient, manipulating one NV spin does not perturb the state of a neighboring NV spin. To experimentally verify this selective, independent spin control, we simultaneously drove Rabi oscillations on NV IV while measuring the ESR spectrum of NV V (Fig. 4.4.2a) by applying two separate RF fields (RF 1 and RF 2), each used to drive one NV spin-resonance. To illustrate the independence of the two measurements, we performed a two-dimensional sweep over the RF 1 pulse duration (τ_{RF1}) and the frequency of RF 2 (ω_{RF2}) (Fig. 4.4.2b). We then subtracted out the mean of each row or column to show that the two measurements are independent. Rows or columns were summed to reconstruct the resulting Rabi oscillations of NV IV or the ESR curve of NV V, respectively. The only deviations from simultaneous, independent NV spin measurements are observed when ω_{RF2} is swept close to the resonance of NV IV, yielding a slight damping in the Rabi oscillations. These deviations do not persist through the ESR transition of NV V, which shows that both NVs can be manipulated independently when they are driven on resonance.

4.5 DISCUSSION AND OUTLOOK

There are several promising avenues in applications of field-gradient MRI. First of all, we expect that the demonstrated magnetic field gradient method can be used to improve spatial resolution of the scanning NV magnetometer introduced in Chapter 2 and, 3 [88]. The field-gradient-induced spectral separation of identical spins at different positions provides an orthogonal means to attain spatial resolution not limited by the sensor-to-target separation, and could potentially be pushed to the sub-nm scale with experimentally achieved field-gradients [53]. Realizing such nano- or atomic scale resolution in imaging single paramagnetic spins under ambient conditions would enable diverse applications such as imaging magnetic point defects in solid-state systems [62] and tracking individual spin-labels in biological systems [2].

In addition, scanning-field-gradient MRI of spins yields precise determination of their relative locations with nanometric spatial resolution, which will be crucial for optimizing the functionality of spin-based quantum bit ensembles. Once suitable spin ensembles are identified and spatially mapped, individual spin control both allows for the determination of individual spin properties and when combined with magnetic dipole coupling between adjacent spins provides a method for achieving complete control of the quantum state of spin ensembles. For our demonstrated spatial resolution (9 nm) in resolving nearby spins, coupling rates between proximal spins can be as strong as 36 kHz for optimally oriented spins. Thus, so long as spin coherence times exceed 28 μ s, such spins can be entangled. As NV spin coherence times up to 1.8 ms have been demonstrated [5], our scanning-field-gradient MRI is immediately applicable from sensitive nanoscale magnetometers to scalable quantum information processors [96].

The control and manipulation of individual spins using magnetic field gradients is independent of the method used for spin readout, therefore can be extended to other spin qubit systems. For optically addressable spins, such as NV spins, integrating far-field, sub-diffraction schemes - such as stimulated emission depletion (STED) [81] and reversible saturable optical linear fluorescence (spin-RESOLFT) [56] - with a scanning magnetic field gradient would allow for both robust individual spin control and readout with nanometric resolutions. In addition, selective optical control of such systems is possible via the incorporation of an electric field gradient to the scanning tip [35], which would allow both spin and electronic degrees of freedom to be both addressed individually. Alternatively, using demonstrated single-shot electrical readout of individual spins [6, 61] would allow for MRI to be performed rapidly and efficiently, as acquisition times would not be limited by the readout integration time.



Figure 4.6.1: Scanning electron micrograph of a quartz tip with magnetic coating. The metal coating is overlain in red (false color).

4.6 SUPPLEMENTARY SECTION

4.6.1 MAGNETIC TIPS

Magnetic tips were created by evaporating a magnetic layer onto quartz tips with the aspect ratio of 5 and the diameter of roughly 40 nm, which were fabricated using a commercial laser-pulling system (Sutter Instrument, P-2000). Using a thermal evaporator, a 25 nm layer of cobalt–iron was deposited on the side of the pulled quartz tip. A 5 nm chrome layer was then evaporated, which serves as a capping layer to prevent oxidation of the magnetic material. These tips result in magnetic field gradients of roughly 1 G/nm at distances of roughly 100 nm.

4.6.2 Combined AFM and confocal microscope

Two stacks of piezo coarse positioners and scanners (Attocube ANPxyz101/RES ANSxyz100) are used for positioning a magnetic tip and a bulk diamond sample containing NV centers. NV centers are placed at the focus of a confocal microscope using one set of piezo scanners (XYZ1), while the tip's distance to the sample is regulated via AFM feedback on the other set of stacks (XYZ2). Atomic force detection is done electrically by measuring changes in the resonant frequency of an oscillating tuning-fork operating in shear mode. The tip can be



Figure 4.6.2: Combined AFM and confocal microscope.

scanned using XYZ₂ to produce magnetic resonance images, while NV spins on the sample are read out confocally. A green excitation laser (532 nm wavelength) is focused on the NV centers using a long working distance objective (Mitutoyo, M Plan Apo NIR 100X) and excites NV centers non-resonantly, which then fluoresce at longer wavelengths. An acousto-optical modulator (not illustrated), is used to pulse the laser allowing for measuring the time-dependent fluorescence of NV centers. A dichroic mirror (DM) separates the collection and excitation laser paths to avoid unwanted photon counts from the excitation laser. Emitted light passes through a telescope containing a pinhole (PH), making the optical detection confocal. A longpass filter (LP) is used to further suppress green excitation counts before the emitted light is collected via an avalanche photo diode (APD) (Perkins and Elmer, SPCM-AQRH-13).



Figure 4.6.3: Normalized fluorescence for magnetic imaging. (a) Pulse sequence for magnetic resonance imaging. (b) The frequency of RF 1 and RF2 relative to ESR frequency of an NV center in the absence of the magnetic tip. (c) Normalization of counter 1 to counter 2 for background-free magnetic image.

4.6.3 NORMALIZED FLUORESCENCE FOR MAGNETIC IMAGING

This section describes the procedure to acquire background-free magnetic resonance imaging. Figure 4.6.3a is the pulse sequence used in the imaging. Green light is continuously applied to the NV center. For the first half of the sequence, one radio-frequency (RF 1) field is applied, and the corresponding NV counts are collected in counter 1. For the second half of the sequence, the same is done for RF 2 and counter 2. The length of the total sequence is roughly 50 μ s, much longer than the spin-pumping time and much shorter than the pixel integration time of our MRI scans. The frequency of RF 1 is tuned close to the NV resonance, while RF 2 is detuned by more than 10 linewidths so that the tip never brings the NV spin into resonance with RF 2 (Fig. 4.6.3b). When a tip is scanned over the sample (same data as Figure 4.2.1c), counts are collected in both counter 1 and counter 2. Counter 1 contains a magnetic contour showing the location where the tip brings the NV into resonance with RF 1 (Fig. 4.6.3c, left panel). However, there are also significant non-magnetic features due to artifacts such as tip shadowing and the influence of topographic features (Fig. 4.6.3c, middle panel). These non-magnetic features can be removed by normalizing counter 1 to counter 2; the latter is affected by the same artifacts as the former but contains no magnetic signal. This normalization results in an image that has only magnetic-field induced fluorescence variations (Fig. 4.6.3c, right panel). We note that by modulating between two RF sources, the overall RF power can be kept constant in time, which is crucial for maintaining AFM performance.

4.6.4 Deconvolution to extract spin-spin distances

A deconvolution process is used to extract the relative distance between NVs from the magnetic contours in Figure 4.3.1 of the main text. As the magnetic response of each NV to the tip is identical, but spatially shifted, we note that images of multiple NV spins are simply the magnetic response of one NV convolved with a map of delta functions corresponding to the spins' relative locations. Thus, we can use deconvolution of the observed magnetic image with

the response of one of the NV spins to retrieve relative spin locations. Doing so requires the point spread function of a single NV spin, which can be found by first measuring the response of a single NV spin before moving to ensemble systems. Alternatively, for ensembles where there is not significant overlap between magnetic resonance curves, one can isolate a single ring and use that as the point-spread function for the deconvolution. We use the latter approach here by manually cropping out data not associated with NV II's magnetic resonance (shown in Fig. 4.3.1a in the main text).

Once the point-spread function of a single spin is determined, deconvolution can be readily performed. Because of finite signal to noise ratios in our data, applying a filter during the deconvolution process is necessary to avoid artificial divergences. We accomplish this by using a Wiener filter, provided in the MATLAB Image Processing Toolbox, which minimizes the contribution from noise at frequencies which have low signal-to-noise ratios (Gonzalez, R. C. & Woods, R. E. Digital Image Processing. (Addison-Wesley Publishing Company, Inc., 1992)). Wiener deconvolution has no free parameters as the optimal amount of filtering is dictated by the noise power spectrum of the image. For NV spin states read out optically, the dominant source of noise is photon shot noise, and consequently, the power spectrum is flat. Thus, the only relevant parameter is the signal to noise ratio of the image, which we find to be 7. Using this value yields the deconvolved image in Figure 4.3.1 which results in an image where each NV spin's location corresponds to a single peak. We note that the peak corresponding to NV I is significantly skewed (as compared to NV III) because that NV lies at a different depth (15 nm further into the surface) than NV II, which slightly modifies its response to the magnetic tip. In principle, to deconvolve spins separated in three dimensions it is necessary to perform a deconvolution on the three-dimensional response of the NV spins to the tip. For this work, because spins are separated laterally more than in the z-direction, this is unnecessary for measuring relative spin distances.

Performing deconvolution is more involved if NV spins in an ensemble are not identically oriented. In bulk diamond, there are four possible quantization axes

for NV centers, dictated by the crystallographic vector connecting the nitrogen atom and neighboring carbon vacancy. As spins' resonant frequencies predominantly shift to energies along their quantization axes, up to four different magnetic responses can be observed upon scanning arbitrary NV spin ensembles with a magnetic tip. While simple deconvolution can be used to find the relative distances between spins sharing an axis in the ensemble, finding the relative distances between the four NV types requires additional consideration.

One method for measuring the distance between NV spins of different axes is to magnetize the tip-field to be perpendicular to the diamond surface, which for cylindrical tip geometries will yield a cylindrically symmetric magnetic field. Then for [001] diamond crystals, where all four NV axes have the same projection along the z-direction, the tip induces an identical, but rotated, magnetic response from each NV axis direction. Thus deconvolution can again be performed after rotating the magnetic response by 90, 180, or 270 degrees depending on the NV axis. Alternatively, for tips without cylindrical symmetry, the sample can be rotated in 900 increments to have each NV direction respond identically to the tip. Relative distances between these four measurements can be determined by using AFM topography, which should not change upon rotation of the sample and can thus be used to correlate the magnetic measurements. Additionally, this technical challenge is unique to spins associated with NV centers, where the quantization is defined by a zero-field splitting along a crystallographic vector. For many other systems, the quantization axis for spins can be dictated by an externally applied field, and consequently all spins will respond identically to the local tip field.

4.6.5 Theoretical NV coupling for the measured NV locations

NVs I, II, and III (Fig. 4.3.1), are plotted in real space with the relative distances extracted through our scanning-magnetic field gradient imaging. Arrows indicate the NV orientation determined by measuring the Zeeman splitting along different directions with a three-axis Helmholtz coil (data not shown). All three



Figure 4.6.4: Theoretical NV coupling for the measured NV locations.



Figure 4.6.5: Distance between NV IV and NV V.

NVs have the same NV axis. The dipolar magnetic field strength from NV II along the NV axis is plotted as a function of position. We plot equal field-strength contours that intersect NV I (red) and NV III (blue), yielding their theoretical dipolar coupling strengths of 1.8 nT and 4.0 nT respectively. The shapes of the contours indicate the strong angular dependence of the coupling strength, illustrating the importance of three-dimensional imaging for a priori determination of NV coupling frequencies.

4.6.6 Distance between NV IV and NV V $\,$

Analogous to Figure 4.3.1, magnetic imaging is done on NVs IV and V to determine their relative lateral distances. When the tip is scanned over of the pair of NVs, four contours are seen for the two NVs. There are two contours per NV because the tip field is strong compared to the applied static field, which splits the $m_S = -1$ and $m_S = +1$ in the NV triplet ground state. Unlike the imaging done in Figures 4.2.1 and 4.3.1, where the tip is only strong enough to bring one spin

transition into resonance, now the tip can bring either $m_S = -1$ or $m_S = +1$ into resonance with $m_S = 0$, depending on the sign of the field. The distance between NVs is given by the offset of the magnetic contours for the two NVs, which is the same for both spin transitions and is 135 nm (length of blue arrows). We note that this configuration of field-strength and detuning is advantageous in this situation, since the detection of the two ESR transitions ($m_S = 0 \rightarrow m_S = -1$ and $m_S = 0 \rightarrow m_S = 1$) allows for accurately determining both x and y displacements of the NV centers while maintaining a small scan range.

4.6.7 ACTIVE AFM DRIFT CORRECTION

This section describes the active drift correction scheme used to ensure the tip position with few-nm resolution during repetitive NV-control experiments. Figure 4.6.6a shows AFM topographic image of an NV-containing nanostructure (here a triangle). This triangle contains the two NV centers studied in Figures 4.4.1 and 4.4.2 in the main text. Magnetic resonance imaging is performed on the two NV spins, and their magnetic responses are observed in the normalized fluorescence (plotted is the same region as Fig. 4.6.5). In order to achieve coherent control of each of the two spins, it is necessary to position the tip where it splits the two NV resonances and maintain the tip location with few-nm resolution to overcome NV spin-decoherence induced by tip-drifts. Here, we fix the tip at the green marker, but after performing experiments on each NV spin, the tip may have changed location due to temperature-induced drifts or piezo-electric creep. As an example of drifts, whose magnitude is exaggerated here, we examine the situation where the tip has drifted from the green marker to the blue marker. The real-space location of the drifted tip is unknown. To determine its location efficiently, we take a single AFM line-scan which intersects the edge of the AFM triangular structure (green line in Fig. 4.6.6a). In Figure 4.6.6b, we plot the measured line-scan topography (green line), and cross-correlate it to individual lines of a topographic scan of the triangle edge (blue and grey lines). The AFM line yielding the highest cross-correlation (blue



Figure 4.6.6: Active AFM drift correction. (a) AFM topographic image of an NV-containing nanostructure. (b) Lateral positioning feedback via line-scan correction. (c) Lateral positioning feedback performance.

line), measures the tip's drift in the y-direction (dy). Drifts in the x-direction (dx) can additionally be extracted from the spatial offset between the green and blue panels, which is given by the location of the peak in the performed cross-correlation. Positioning is performed intermittently, and the resulting drifts in the y-direction (dy) are plotted (Fig. 4.6.6c, left panel). Magnetic field-gradients are predominantly directed along the y-direction (measured from the MRI image of the NVs in Fig. 4.6.5), and consequently achieving high-precision in y-positioning is necessary for maintaining the NV spins resonance. Here we show positioning can be achieved with a standard deviation of 2.3 nm, only limited by pixelation of the topographic data. Spatial variations can then be converted to spectral uncertainties using the measured magnetic field gradient of our tip (0.1 G/nm here), indicating that the magnetic resonance can be maintained with 500 kHz precision. This sets a limit for tip-drift induced decoherence of NV spins which is smaller than the inhomogeneous dephasing rate $(1/T_2^*)$ of the NV spins studied here. Integrating the measured drifts between line-scans gives the overall tip motion (Fig. 4.6.6c, right panel), which indicate that the tip location would drift considerably more without the use of line-scan feedback.

4.6.8 RAMSEY FREE-INDUCTION DECAY MEASUREMENTS

The Ramsey free-induction decay measurements in Fig. 4.4.1 are taken with the radiofrequency driving field detuned by 5 MHz from the target NV center's nominal ESR frequency in the presence of the magnetic tip; the observed oscillations are due to beating of this detuning with the ¹⁵N hyperfine splitting (3.1 MHz). The measured Ramsey data are fitted to the sum of two exponentially damped sinusoids, whose phases are fixed by the relative strength between the net detuning (the radiofrequency field detuning plus or minus half the hyperfine splitting) and the Rabi frequency (5.5 MHz), governed by the strength of the applied radiofrequency field.

5 Coherent, mechanical control of a single electronic spin

5.1 INTRODUCTION

The ability to control and manipulate spins via electrical [24, 33, 66], magnetic [37, 74] and optical [75] means has generated numerous applications in metrology [13] and quantum information science [65] in recent years. A promising alternative method for spin manipulation is the use of mechanical motion, where the oscillation of a mechanical resonator can be magnetically coupled to a spin's magnetic dipole, which could enable scalable quantum information architectures [78] and sensitive nanoscale magnetometry [4, 58, 88]. To date, however, only population control of spins has been realized via classical motion of a mechanical resonator [53, 83, 93]. Here, we demonstrate coherent mechanical control of an individual spin under ambient conditions using the driven motion of a mechanical resonator that is magnetically coupled to the electronic spin of a single nitrogen-vacancy (NV) color center in diamond. Coherent control of this hybrid mechanical/spin system is achieved by synchronizing pulsed spin-addressing protocols (involving optical and radiofrequency fields) to the motion of the driven oscillator, which allows coherent mechanical manipulation of both the population and phase of the spin via motion-induced Zeeman shifts of the NV spin's energy, thus demonstrating full control of the spin's quantum state via mechanical means. We demonstrate applications of this coherent mechanical spin-control technique to sensitive nanoscale scanning magnetometry.

5.2 COHERENT CONTROL SCHEME OF A COUPLED MECHANICAL-SPIN SYSTEM

The magnetic coupling between spins and mechanical resonators has been recently investigated for imaging the locations of spins via magnetic resonance force microscopy [53, 83]. as well as for sensing nanomechanical resonator motion [68]. A scrarcly explored resource in such coupled spin-resonator systems is the coherence of a driven resonator's motion, and here, we demonstrate that this motion can be used to fully control the quantum state of an individual electron spin. We employ an electronic spin associated with an NV center in diamond as the target spin due to the efficient optical initialization and readout of the NV spin [30], as well as its long coherence time [5]. A single NV center is prepared near a bulk diamond surface (at a nominal depth of 10 nm, see Supp. 3.7.2), and its spin-state is read out optically through spin-dependent fluorescence $\begin{bmatrix} 30 \end{bmatrix}$. The mechanical resonator is a quartz tuning fork, with a micro-fabricated quartz tip attached at the end of one of its prongs. The tuning fork operates in a transverse oscillation mode with a resonant frequency of 41.53 kHz and oscillation amplitude that can be controllably varied up to 250 nm (see Fig. 5.2.1 and Supp. 5.6.1). Magnetic coupling between the NV spin and the

tuning fork resonator is provided by a 25 nm CoFe magnetic film evaporated onto the quartz tip (see Supp. 4.6.1). Consequently, the transverse mechanical motion of the tip generates an oscillatory magnetic field at the spin's location, which modulates the Zeeman splitting between the NV's energy levels (Fig. 5.2.1b, c). We note that the tip is laterally positioned 500 nm from the NV spin, which keeps the NV outside from the range of the tip's motion. This helps to achieve linear coupling between the motion and the spin (see Supp. 5.6.2). To achieve coherent control of the target NV spin using the resonator's mechanical motion, we synchronize spin manipulation protocols to the driven oscillation of the mechanical resonator (Fig. 5.2.1d). With this synchronization, the relative timing between applied radiofrequency (RF) pulses and the resonator motion is fixed, allowing for the resonator's motion to coherently and deterministically influence the NV spin state.

The pronounced coupling between the resonator's mechanical motion and the NV spin is measured with optically detected electron spin resonance (ESR). The ESR of the target NV center is acquired by sweeping the frequency of an applied RF field through the NV spin's magnetic dipole transition and collecting the resulting NV's spin-state-dependent fluorescence, with lower (higher) fluorescence when RF field is on (off) resonance with the NV spin transition $\begin{bmatrix} 30 \end{bmatrix}$. As the oscillation amplitude of the tuning fork increases, we observe a broadening in the ESR spectrum (Fig. 5.2.2a), resulting from the larger magnetic field modulation associated with the greater range of motion of the resonator's magnetic tip. As the driving amplitude of the resonator is further increased, the ESR lineshape becomes bimodal, which reveals the distribution of dwell-times of the magnetic tip as a function of its position during the oscillatory motion, with the two peaks representing the turning points of the oscillation trajectory (see Supp. 5.6.3). To deconvolve this time-averaged spectral broadening in the coupled dynamics of the NV spin and resonator, we perform stroboscopic ESR measurements synchronized to the resonator's motion (Fig. 5.2.2b), which reveal the magnetic tip's position as a function of time via the



Figure 5.2.1: Coherent dynamics of a mechanical resonator and an electron spin. (a) Schematic of an electron spin of an NV center next to a tuning fork resonator. A magnetic tip provides coupling between the tuning fork resonator and the NV spin. Readout and addressing of the NV spin is provided through a confocal microscope and an RF coil. The details of the experimental setup are described in Grinolds et al. [29]. (b, c) Coupling of the motion of the resonator to the electronic spin of the NV center. The local magnetic field at the position of the NV spin changes as a function of the resonator's motion due to the magnetic field gradient of the tip. This resonator-induced magnetic field oscillation, $\delta B_{\rm res}(t)$, with the oscillation period, $T_{\rm res} = 24.1 \ \mu s$, modulates the spin's dipole transition energy, $\delta \omega(t)$, through a Zeeman shift, $\delta \omega(t) = \gamma \delta B_{res}(t)$, where γ is the gyromagnetic ratio of the spin (Inset). (d) Scheme for coherent coupling of the resonator's motion to the NV spin. The resonator's motion is synchronized to the NV addressing sequences. After this synchronization, the motion of the resonator influences the NV spin coherently with respect to the standard optical/RF pulse sequences which addresses the NV spin.

tip-induced Zeeman shift of the NV's ESR resonance frequency. Figure 5.2.2b shows two example stroboscopic ESR spectra with shifted resonance frequencies consistent with the magnetic tip's position in its oscillatory cycle. Such measurements are acquired with an acquisition time (1 μ s) much shorter than the resonator's oscillation period (24.1 μ s), which enables stroboscopic readout of the NV spin's resonance frequency for a well defined position of the magnetic tip. The observed remaining broadening of the ESR linewidth (full width at half maximum) is within 20% of the value we find for an undriven magnetic tip (6.62 ± 0.49 MHz), which is due to the resonator's remnant motion during the non-zero acquisition time (see Supp. 5.6.3). The near-restoration of the zero-tip-motion ESR linewidth at arbitrary tip position confirms that the resonator motion is coherent with the spin addressing protocols.

5.3 QUANTUM CONTROL OF A SPIN WITH THE MECHANICAL MOTION OF A RESONATOR

This synchronization of the resonator's motion with respect to the external optical/RF control can be used for coherent control of both the population and phase of the target spin states. Population control (Fig. 5.3.1) is achieved via an adiabatic fast passage [86]. The oscillation of the magnetic field induced by the magnetic tip motion modulates the NV ESR frequency. By fixing the applied RF frequency (ω_{RF}) to the center of this frequency modulation range, the target spin's population is adiabatically inverted each time the resonator-induced magnetic field brings the NV ESR frequency onto resonance with ω_{RF} . This spin population inversion occurs twice for each period of the resonator oscillation (here 24.1 μ s), resulting in periodic population inversion (i.e. spin flips). Note that such adiabatic population inversion can have a higher fidelity than conventional RF π -pulses, since it is robust against inhomogeneous variation in ESR frequency. It is especially useful for flipping ensembles of spins, where differing local environments for individual spins can otherwise limit control



Figure 5.2.2: Spin-resonator coupling as observed by electron spin resonance (ESR). (a) Motion-induced broadening of ESR. As the amplitude of the resonator's oscillation is increased, the ESR linewidth broadens beyond its initial value (6.62 ± 0.49 MHz, here power-broadened by the applied RF field). The shape of ESR evolves to a bimodal form, owing to the simple harmonic oscillation of the resonator (see Supp. 5.6.3). (b) Stroboscopic ESR measurement. Measurements are taken synchronously to the motion of the resonator with 1 μ s acquisition time. Having this short acquisition time compared to the oscillation period of the resonator ($24.1 \ \mu$ s) results in stroboscopic ESR snapshots, which capture the applied tip field - and the corresponding shift in ESR - as a function of the resonator's position relative to the spin. The actual data (blue and red) are taken with the timings of $-1.8 \ \mu$ s for the blue and o.8 μ s for the relative to the node of the resonator's oscillation. The linewidth of the blue colored ESR is 7.98 \pm 0.43 MHz, which is nearly recovered to the original linewidth of 6.62 MHz.



Figure 5.3.1: Population control of a spin with the mechanical motion of a resonator. The motion of the resonator periodically modulates the local magnetic field at the position of the target spin, and therefore modulates its spin resonance. Adiabatic spin inversions occur when the resonator sweeps the spin resonance through the frequency of an applied RF field (inset, right). The corresponding range of frequency sweep is $6_{5.1}$ MHz, larger enough than the RF Rabi frequency, 6.2_5 MHz to achieve high fidelity of spin inversion. Plotted is the population of the NV spin in the $m_s = 0$ state ($P(m_s = 0)$) as a function of delay between initialization and readout (τ). The left inset shows a periodic population modulation with multiple cycles and a period of $24.1 \ \mu$ s in accordance with the measured resonator frequency. Fits to the adiabatic passages take into account a finite RF Rabi frequency of 6.25 MHz (see Supp. 5.6.4). The decay in the inversion amplitude is determined by the spin-lifetime, T_1 .

fidelity, provided that the tip-induced modulation of the field is larger than such inhomogeneous broadening.

Furthermore, synchronizing the resonator motion with an appropriate RF pulse sequence allows for high fidelity control of the target NV spin's phase. As a demonstration of such phase control, the NV spin is first prepared in a superposition state, using optical pumping followed by an RF $\pi/2$ -pulse. Next, the time-varying magnetic field induced by the resonator's motion results in a

detuning of the NV ESR frequency from $\omega_{\rm RF}$, causing the spin to acquire a differential phase relative to its precession at ω_{RF} . In order to accumulate the phase with high fidelity, and to minimize the effect of other, incoherent magnetic field fluctuations, a Hahn echo sequence is employed and synchronized to the driven motion of the resonator (Fig. 5.3.2a). By placing the pulse-sequence symmetrically with respect to the node of the resonator oscillation, the acquired phase is maximized for a given tip amplitude. For a fixed duration of the echo sequence (here, $\tau = 12 \ \mu s$), the amount of accumulated phase is proportional to the amplitude of the magnetic field oscillation, which is given by the oscillation amplitude of the magnetic tip displacement multiplied by the tip-induced magnetic field gradient along the direction of the tip motion. To read out the accumulated phase, a final RF pulse (here, $3\pi/2$) is applied to project the accumulated phase of the NV's spin onto a distribution of the state populations, which is then measured via spin-state-dependent fluorescence. Figure 5.3.2b shows an example of such resonator-induced spin phase accumulations, controlled by the amplitude of the resonator's oscillation.

In these demonstrations of coherent, mechanical control of a single NV spin, the rate of spin manipulation is set by the 41.53 kHz resonance frequency of the quartz tuning fork. However, nano-mechanical resonators can have resonance frequencies ranging up to nearly 1 GHz [50, 51], which would allow rapid control of target spins. For example, a resonator with 1 MHz frequency could perform more than 1000 coherent spin manipulations within the demonstrated NV spin coherence time of a few milliseconds [5].

5.4 MOTION-ENABLED AC MAGNETOMETRY

With the demonstrated population and phase control, we have achieved coherent, mechanical control over the state of a single electron spin. This coherent mechanical spin-control technique enables new applications in sensitive, nanoscale metrology. In particular, electronic spins in NV centers have

Figure 5.3.2: Phase control of a spin with the mechanical motion of a resonator and applications to magnetic imaging. (a) Phase accumulation scheme based on a Hahn echo sequence. The evolution time, τ , is fixed to 12 μ s in this experiment. The echo sequence is symmetrically placed at the node of resonator's oscillatory motion, allowing for robust phase accumulation, which is only sensitive to the motion of the resonator, while canceling out slowly varying background magnetic fluctuations. (b) Phase accumulation of a spin as a function of the oscillation amplitude of the resonator. The accumulated phase is proportional to the magnetic field modulation induced by the resonator, which scales linearly with the oscillation amplitude of the resonator. When the phase is converted to the population of the NV spin in the $m_s = o$ state $(P(m_s = o))$ by a RF $3\pi/2$ -pulse, we observe a sinusoidal oscillation of the population. (c) Scanning DC magnetometry with the NV spin. The external RF frequency is set on resonance with ESR in the absence of the magnetic tip. The change in local magnetic field at the position of NV results in ESR frequency shifts, and then changes the spin-state-dependent fluorescence. The larger scan image in the inset reveals a dipole-like pattern of the tip-induced magnetic field. The NV's position is marked as a black dot. The scan is zoomed to a region depicted by the dotted square box. The darker region is the region where the spin is near resonance with the external RF frequency. The field variation across the band is $_{220} \mu T$. (d) Scanning AC magnetometry with the NV spin. The magnetic tip is scanned laterally with 20 nm oscillation amplitude. The modulation of the NV position relative to the magnetic target (here, the magnetic tip) produces a modulation of the magnetic signal from the source, providing a general method for adapting optimally sensitive AC magnetic imaging scheme. In comparison to DC magnetic image, we observe additional features, multiple fringes, which correspond to contours of constant magnetic field-gradient along the direction of the resonator's oscillation.




been recently identified as excellent magnetic field sensors because of their long coherence time and efficient optical readout [4, 58, 88], even under ambient conditions. Previous work [58, 88] showed that optimal magnetic field sensitivity is achieved when the target field is modulated with a period comparable to the NV's spin coherence time ('AC magnetometry'). However, such AC magnetometry cannot be applied a priori to targets with static magnetizations. Our coherent mechanical spin-control scheme provides a solution as the motion of the tip transforms a spatially varying, static magnetic field of a magnetic sample into a time-varying magnetic field at the position of the NV center. We demonstrate such mechanical-resonator-enabled AC magnetometry by performing scanning, nanoscale magnetic field imaging of our magnetized tip (Fig. 5.3.2c,d). At first, we acquire a 'DC magnetic image' [4, 29] by scanning the magnetic tip laterally near a single NV center, where the external RF frequency is fixed on resonance with the NV ESR frequency in the absence of the tip, and the fluorescence change due to Zeeman shifts is monitored (Fig. 5.3.2c). In this mode, the change in signal directly reflects the change in the local magnetic field at the position of NV. From the ESR spectrum (Fig. 5.2.2a, top), the corresponding DC magnetic field sensitivity can be deduced to be 14 μ T/ $\sqrt{\text{Hz}}$ (see Supp. 5.6.5). A resonance band with decreased fluorescence is formed where the tip's magnetic field along the NV axis is within one linewidth of the ESR. The width of this resonance band corresponds to the magnetic field variation along the NV axis of 220 μ T. Next, we perform resonator-motion-enabled AC magnetometry (Fig. 5.3.2d), where the same optical/RF-pulse sequence is used as for the phase-control experiment described above, while the driven motion of the magnetic tip is synchronized to the NV addressing protocols with fixed tip oscillation amplitude of 20 nm. We calculate our NV spin's AC magnetic field sensitivity to be 0.92 μ T/ \sqrt{Hz} (see Supp. 5.6.6), which constitutes a factor of 15 improvement over our DC experiment. In contrast to the 'DC magnetic image' (Fig. 5.3.2c), additional structures in the form of multiple parallel fringes in the resonance region are revealed. In this configuration, the NV center senses magnetic field variations along the direction

of the tip oscillation, and the observed fringes (Fig. 5.3.2d) correspond to contours of constant magnetic field gradients, with neighboring fringes differing by a gradient of 0.29 μ T/nm, or a field variation of 5.8 μ T which is a factor of 40 smaller than that of DC image (see Supp. 5.6.7).

A particularly appealing application of our motion-enabled AC magnetometry could be sensitive nanoscale imaging of static, weak, and spatially varying magnetic features. Our magnetic imaging technique optimizes magnetic field sensitivity through AC Magnetometry and should thereby be capable of detecting the magnetic moment of a single Bohr magneton within few seconds of data acquisition (see Supp. 5.6.8) at a 10 nm distance (which is the nominal depth of the prepared NV centers, see Supp. 3.7.2). It therefore offers the sensitivity required to detect local magnetic features down to a few Bohr magnetons in magnitude, provided that the NV spin can be brought within a few tens of nanometers of the sample. For example, this could be used to study much-debated graphene ferromagnetism [70, 79, 85, 87, 97], where scanning nanoscale AC magnetometry can potentially provide local information of its magnetic origin. While in our demonstration of motion-enabled AC magnetometry, the NV spin-sensor is located in a fixed bulk diamond sample, our scheme can be readily applied to studying arbitrary samples in a scanning geometry. Such a scannable diamond device $\begin{bmatrix} 52 \end{bmatrix}$ has recently been developed, where an NV spin-sensor with 56 nT/\sqrt{Hz} sensitivity can be brought within a few tens of nanometers to the sample, making these applications of motion-locked AC magnetometry readily achievable [52].

An additional application of our coherent mechanical spin-control technique is motion sensing for nanoscale mechanical resonators. Such detection of motion, while routinely performed for microscale mechanical resonators using optical interferometry, remains challenging for nanoscale objects. Our mechanical spin-control scheme employs a single, atomically localized NV spin, thereby allowing nanoscale displacement and motion sensing. In principle, our demonstrated phase control scheme can be used to measure the amplitude and the phase of the motion of a resonator of interest. Using the measured magnetic tip field gradient of 18.4 μ T/nm combined with the NV's AC field sensitivity, the oscillator amplitude sensitivity in our setup is estimated to be 49.8 pm/ $\sqrt{\text{Hz}}$, which is already comparable to the sensitivity (\approx 10 pm/ $\sqrt{\text{Hz}}$) achieved by optical inteferometry of sub-micron sized resonators [43]. Similarly, the same scheme can be applied to measure the phase of the resonator's oscillatory motion, with an estimated sensitivity of 5.1 mrad/ $\sqrt{\text{Hz}}$ at an oscillation amplitude of 10 nm.

5.5 DISCUSSION AND OUTLOOK

The demonstrated sensitivity for metrology applications can be further optimized through a variety of experimentally demonstrated improvements. Such schemes of improvements include extending NV's spin coherence time [4, 16] and enhancing photon collection efficiency [3, 32, 52]. Our displacement sensing scheme can also be improved by engineering higher magnetic field gradients. For example, by using state-of-the-art magnetic tips with a field gradient of 1 mT/nm [53], an NV spin with T_2 of few ms, and enhanced photon collection efficiencies, the displacement amplitude sensitivity of 22.8 fm/ $\sqrt{\text{Hz}}$ could potentially be achieved. Finally, we note that the techniques demonstrated here could be integrated with related methods for scanning-field-gradient spin MRI [29] and super-resolution optical imaging and coherent manipulation of proximal spins [56], with an ultimate goal of combined magnetic field sensitivity and spatial resolution to achieve real-time atomic-scale mapping of individual electron and nuclear spins in physical and biological systems of interest.

5.6 SUPPLEMENTARY SECTION

5.6.1 QUARTZ TUNING FORK

In our experiment, a commercially available quartz tuning fork resonator (DIGI-KEY, X801-ND) is used. Its resonance frequency and quality factor are 41.53 kHz, and 1400, respectively. The excitation of the tuning fork is performed by mechanically dithering the tuning fork using a piezoelectric actuator. Additionally, the response of the tuning fork is read out electrically through piezoelectric detection.

5.6.2 Magnetic field profile and the measurement location of the tip

In our experiments, we carefully positioned the tip near the NV center to achieve pronounced and linear coupling for mechanical spin-control. To find such a position, we acquired the magnetic image [4, 29] of the tip (Fig. 5.6.1a). It is acquired by scanning the tip around the NV center while monitoring the NV's fluorescence. The RF frequency was set on resonance with NV's ESR transition in the absence of the tip. As a result, a resonance band appeared where the tip's magnetic field along the NV axis is within the linewidth of the ESR. Across the band, the on-axis magnetic field, or NV's frequency shift, is monotonic, which assures a quasi-linear magnetic coupling. It is desired to locate the tip on the narrower side of the band to have stronger field gradient. Therefore, we position the tip in the middle of the disappearing end of the resonance band. To qualitatively understand the field gradient, we performed a simulation of the tip's magnetic field (Fig. 5.6.1b-d). We assumed the tip was a single dipole, since the tip is far enough from the NV center. The direction of NV was measured to be (o, $\sqrt{2}$, 1), and the free parameters for the simulation were the dipole strength, direction, and the height from the plane, which were $7 \cdot 10^9$ Bohr magnetons, (0, -0.4, 1), and 600 nm. Indeed, the contours of the NV-axis field gradients along the tip motion (Fig. 5.6.1d) implies the magnetic field is monotonic along the direction of the tip motion. The simulated field gradient at the position of the tip was 15.4 μ T/nm, which is in a good agreement with the measured value of $18.3 \,\mu T/nm.$



Figure 5.6.1: Magnetic image of the tip. (a) Measured image. The dashed line with arrows indicates the tip's oscillation direction (with a drawn exaggerated amplitude for visual clarity). (b) Simulated magnetic field map. (c) Simulated field gradient map (along the tip's direction of motion). (d) Simulated magnetic image. The contours of field gradients along the direction of the motion were presented.

5.6.3 Resonator-induced magnetic field fluctuation

The ESR spectra of NV center can be described as

$$ESR(f) = 1 - c \frac{(\Gamma/2)^2}{(f - f_o)^2 + (\Gamma/2)^2},$$
 (5.1)

where *c* is the normalized fluorescence-reduction in the signal, f_o is the spin resonance frequency in the absence of the mechanical resonator, and Γ is the linewidth (full width at half maximum). The magnetic field variation at the position of the NV induced by the resonator's motion $\delta B_{res}(t)$ can be approximated as,

$$\delta B_{res}(t) = \frac{\partial B}{\partial x_{res}} \cdot A_{res} \cdot \sin(2\pi f_{res}t), \qquad (5.2)$$

where f_{res} , A_{res} , and $\frac{\partial B}{\partial x_{res}}$ are the frequency, the amplitude, and the magnetic field gradient along the direction of the resonator's oscillation, respectively. The corresponding modulation of the spin resonance frequency due to Zeeman effect is,

$$f_{spin}(t) = \gamma \cdot \delta B_{res}, \qquad (5.3)$$

where γ is the gyromagnetic ratio of the spin. Given the time interval (t1, t2), the resulting time-averaged ESR spectrum due to the resonator's motion, $ESR_{res}(f)$, is,

$$ESR_{avg}(f) = f_{res} \cdot \int_{t_1}^{t_2} dt (1 - c \frac{(\Gamma/2)^2}{(f - f_o - \delta f_{spin}(t))^2 + (\Gamma/2)^2})$$
(5.4)

In Figure 5.2.2a in the main text, The two broadened ESR are numerically fitted using the model above with the full time interval $(0, 1/f_{spin})$, with the measured values of c, Γ , f_o , f_{res} , and A_{res} . The resulting gradients, $\frac{\partial B}{\partial x_{res}}$, of the two broadened ESR are 18.3 μ T/nm (Fig. 5.2.2a, middle) and 18.6 μ T/nm (Fig. 5.2.2a, bottom). The small difference in the obtained value can be ascribed to drift in the relative distance between the magnetic tip and the NV, and higher-order terms of the gradients. We also fitted the stroboscopic ESR (Fig. 5.2.2b) with the time interval $(-2.3 \ \mu s, -1.3 \ \mu s)$ and $(0.3 \ \mu s, 1.3 \ \mu s)$ respectively, and reproduced additional broadening of ESR due to the resonator's motion during the acquisition time. For instance, The linewidth of a stroboscopic ESR (Fig. 5.2.2b, blue) was 7.98 MHz, while the simulated ESR linewidth with the model was 7.07 MHz.

5.6.4 Population control of spin via resonator's motion

To calculate the population of the state $|o\rangle$, $P_{|o\rangle}(t)$, we consider the ideal case of adiabatic fast passage. In this case, the spin state is locked to the effective field, $\overrightarrow{B_{eff}} = \frac{\Omega_o}{\gamma} \overrightarrow{x} + \frac{\delta f_{spin}}{\gamma} \overrightarrow{z}$, in the rotating frame, where Ω_o is the bare Rabi frequency (given by an external RF source) and δf_{spin} is the detuning due to the resonator's motion (Fig. 5.6.2). As a result, $P_{|o\rangle}(t)$ is

$$P_{|\mathsf{o}\rangle}(t) = \frac{1}{2}(1 + \cos(\theta)), \tag{5.5}$$

Where θ is the angle between $\overrightarrow{\Omega_{eff}}$ and \overrightarrow{z} (Fig. 5.6.2a), and $cos(\theta)$ is,

$$\cos(\theta) = \frac{\delta f_{spin}}{\sqrt{\delta f_{spin}^2 + \Omega_o^2}}.$$
 (5.6)

The fit in Figure 5.3.1 in the main text is obtained by plugging in $\delta f_{spin} = (32.55 \text{ MHz}) \cdot sin(2\pi f_{res}t)$ (calculated in the previous section, data based on the bottom panel in Fig. 5.2.2a), and $\Omega_o = 6.25 \text{ MHz}$ (measured by a Rabi experiment, data not shown). To better understand the spin-dynamics in adiabatic fast passage, we numerically solved the Schrodinger equation for our experimental situation. Fig. 5.6.2b shows the calculated trajectory of the NV spin for the experimental parameters in the main text. Note that, due to the moderate field gradient (18.6 μ T/nm), there is an uncertainty of 13 % in the final spin state. This is because the maximal detuning induced by the oscillating tip (32.55 MHz) is only moderately large compared to the applied Rabi frequency (6.25 MHz), i.e. spin rotations are induced even at the extrema of the tip motion. This problem



Figure 5.6.2: Simulation of adiabatic fast passage.

can be solved by engineering tips with higher magnetic field gradient. Fig. 5.6.2c shows such a situation, using a gradient of 74.4 μ T/nm which results in an uncertainty of only 1.5 % in the spin-state after adiabatic fast passage. (Note that even for these large gradients, the adiabatic condition is easily fulfilled)

5.6.5 MAGNETIC FIELD SENSITIVITY OF 'DC MAGNETOMETRY' SCHEME WITH A SINGLE NV SPIN

The magnetic field sensitivity of the 'DC magnetometry' scheme can be derived from the electron spin resonance (ESR) experiment given in the main text (Fig. 5.2.2a, top). Provided that the frequency of the external RF source is fixed within the linewidth of ESR, any additional magnetic field results in the shift in



Figure 5.6.3: Electron spin resonance spectrum. (same as Fig. 5.2.2a, top)

the resonance frequency, which then causes change in the NV's fluorescence. The ESR spectrum can be described as

$$ESR(f) = C_{o} - (C_{o} - C_{1}) \cdot \frac{(\Gamma/2)^{2}}{(f - f_{o})^{2} + (\Gamma/2)^{2}},$$
 (5.7)

where C_0 is the fluorescence far from the resonance, C_1 is the fluorescence on resonance, f_0 is the resonance frequency, and Γ is the linewidth (full width at half maximum). When the external RF frequency is set around Γ from the resonance frequency, the fluorescence change, C, as a function of the external magnetic field, B, is

$$C \approx \frac{C_{o} - C_{i}}{\Gamma} \cdot (\gamma B),$$
 (5.8)

where γ is the gyromagnetic ratio of the spin. The corresponding photon-shot-noise of the measured signal per shot is given by, $\delta C \approx \sqrt{(C_{o} + C_{1})/2}$, due to the Poissonian statistics of photon counting. The resulting minimal detectable magnetic field amplitude per shot is $\delta B = \frac{\delta C}{\partial C/\partial B}$. The parameter-values for the experimental data shown in Figure 5.6.3 are $C_{o} = 19068/s$, $C_{1} = 16753$, and $\Gamma = 6.62$ MHz. Using these numbers, we obtained $\delta B \approx 14 \,\mu T/\sqrt{\text{Hz}}$.



Figure 5.6.4: Bloch sphere representation of spin dynamics during AC magnetometry.

5.6.6 MAGNETIC FIELD SENSITIVITY OF 'AC MAGNETOMETRY' SCHEME WITH A SINGLE NV SPIN

The magnetic field sensitivity of the 'AC magnetometry' scheme can be derived from the mechanical spin-phase control experiment given in the main text (Fig. 5.3.2a, b). The total phase accumulated after the measurement sequence is illustrated in Figure 5.6.4. The spin state is first brought into the equator of the Bloch sphere along the -y axis $by\frac{\pi}{2}$ -pulse around the x axis. During the first half period of a Hahn echo sequence, phase is accumulated due to the oscillating magnetic field, $B_{AC}(t)$ (Fig. 5.3.2a). Consequently, the phase is equal to the integral of the detuning, $\gamma B_{AC}(t)$, during that time interval.

$$\varphi_{1} = \int_{-\tau/2}^{\circ} dt 2\pi \gamma B_{AC}(t) = \int_{-\tau/2}^{\circ} dt 2\pi \gamma Bsin(2\pi f_{res}t) = \frac{\gamma}{f_{res}}(cos(\pi f_{res}\tau) - 1)B$$
(5.9)

The above equation describes the proportionality between the phase, φ , and the oscillation amplitude of the field, *B*, which is the variable to be detected. After the

spin is flipped by a RF π -pulse around the x axis, the accumulated phase becomes $(\pi - \varphi_1)$. During the second half interval of the pulse-sequence, the same amount of a phase with the opposite sign as the first interval is accumulated, since the area of integration is the same while the sign of the AC magnetic field is changed with respect to the DC component of the magnetic field, resulting in the total phase of $(\pi - 2\varphi_1)$. Finally, the phase information is converted into the population difference by applying a $\frac{\pi}{2}$ -pulse around the y axis. The resulting population of the state $|o\rangle$ as a function of the oscillation amplitude of the magnetic field, $P_{|o\rangle}(B)$, is

$$P_{|0\rangle}(B) = \frac{1}{2} (1 + \cos(\frac{\pi}{2} + 2\varphi_1)) = \frac{1}{2} (1 - \sin(2\varphi_1))$$
(5.10)

This population is measured through spin-state-dependent fluorescence, and the above expression of population is converted into the fluorescence signal,

$$S(B) = S_{mean} - \frac{1}{2}S_{cont}sin(2\varphi_1), \qquad (5.11)$$

where S_{mean} is the number of photons collected per measurement shot when the spin-state is in the equal superposition of $|0\rangle$ and $|1\rangle$, and S_{cont} is the peak-to-peak amplitude of the signal variation. Ideally, S_{cont} corresponds to the difference between the numbers of photons collected per measurement shot when the spin-state is $|0\rangle$ and $|1\rangle$, respectively. However, dephasing of the spin during the measurement results in a reduction of S_{cont} . When the amplitude of the magnetic field is nearly zero, the photon-shot-noise of the measured signal per shot is given by $\delta S = \sqrt{S(B = 0)}$, due to the Poissonian statistics of photon counting. The resulting minimal detectable magnetic field amplitude per shot is $\delta B = \frac{\delta S}{\partial S/\partial B}$. When averaged for 1 second, the signal-to-noise ratio is improved by $\sqrt{N_{meas}}$, where N_{meas} is the number of measurements in 1 second. In the demonstrated experiments, the duty cycle of the measurement is given by the frequency of the mechanical resonator, f_{res} . Consequently, the magnetic field sensitivity, or the

minimal detectable magnetic field amplitude per 1 second, δB_{sec} , is,

$$\delta B_{sec} = \frac{\delta B}{\sqrt{N_{meas}}} = \sqrt{S_{mean}} \cdot \frac{1}{S_{cont}} \cdot \frac{\sqrt{f_{res}}}{\gamma} \cdot \frac{1}{(1 - \cos(\pi f_{res}\tau))}$$
(5.12)

The parameter-values for the experimental data shown in Figure 5.3.2b are $S_{mean} = 0.0199$, $S_{cont} = 0.0011$, $f_{res} = 41.53$ kHz, and $\tau = 12.16 \ \mu s$. Using these numbers, we obtained $\delta B_{sec} = 0.92 \ \mu T / \sqrt{\text{Hz}}$.

5.6.7 Imaging of the magnetic field from the magnetized tip

The magnetic field oscillation due to the tip motion, $B_{AC}(t)$, is given by the equation (2), and by using equation (9), the resultant fluorescence map, M(x, y), can be expressed as,

$$M(x,y) = S_{mean} - \frac{1}{2} \cdot S_{cont} \cdot sin(2\frac{\gamma}{f_{res}}(cos(\pi f_{res}\tau) - 1) \cdot A_{res} \cdot \frac{\partial B(x,y)}{\partial x_{res}}) \quad (5.13)$$

Note that it is a sinusoidal function of the magnetic field gradient of the tip along the direction of the tip oscillation, $\frac{\partial B(x,y)}{\partial x_{res}}$, where all the other parameters are fixed during the image acquisition. Therefore, the modulation of the signal can be acquired as $\frac{\partial B(x,y)}{\partial x_{res}}$ is varied monotonously. Consequently, fringes in the image appear as $\frac{\partial B(x,y)}{\partial x_{res}}$ changes by a certain step, which is 0.29 μ T/nm in the demonstrated data. To better understand the image Figure 5.3.2c, we performed a numerical simulation of tip-induced, scanning AC magnetometry. The magnetic field from the tip is modeled as a single magnetic dipole with the dipole strength and direction as the fit parameters. The simulation takes account of the tip during RF pulses, and phenomenological dephasing during the phase evolution. The simulated magnetic image (Fig. 5.6.5a) with the appropriate choice of fitting parameters agrees qualitatively with the actual magnetic image (Fig. 5.6.5b), containing multiple of fringes with similar direction and spacings.



Figure 5.6.5: Simulation of magnetic field-gradient image of the magnetized tip.

5.6.8 MAGNETIC IMAGING WITH TIP MOTION, SMALL TARGET SAMPLE (SINGLE ELECTRONIC SPIN) CASE

The resonator-motion-enabled AC magnetometry demonstrated in the main text (Fig. 5.3.2d) could be especially useful in probing weakly magnetized static materials, since our technique combines optimal magnetic field sensitivity with the ability to modulate the target's field, free from the choice of materials. To illustrate this, we conducted following simulations, where a magnetic dipole moment with a single Bohr magneton is the imaging target. All the parameters – resonator's frequency (41.53 kHz), echo period (12 μ s), average photon counts during the measurements per second (822/s), the fluorescence signal contrast (5.34 %) - for AC magnetic field sensitivity calculations were taken from the actual data presented in Figure 5.3.2a. The dipole direction of the target dipole is arbitrarily set as (x, y, z) = (0, 1, 0), where as the NV sensor's direction is chosen to be one of its possible crystallographic direction, (x, y, z) = (0, $\sqrt{2}$, 1). The distance between the NV sensor and the sample surface is assumed to be 10 nm, as the NV used in the experiment is implanted with the nominal depth of 10 nm.

In the first set of simulations (Fig 5.6.6a, b), the target is resonantly driven with an additional RF source while the tip doesn't oscillate, as typical of conventional AC magnetometry proposals [88]. We assumed that, while driving the target, its dipole direction can be fully inverted. The results show that the signal contrast compared to the background is around 23 counts per second. Figure 5.6.6b simulates the effect of photon-shot-noise, and demonstrates that the signal from a single electron spin can be clearly detectable with the integration time of 60 seconds per pixel. To compare the performance of the resonator-motion-enabled AC magnetometry, a second set of simulations is conducted where the tip with NV sensor is oscillating with an amplitude of 10nm. The results show that the signal contrast is around 36 counts per second (Fig 5.6.6c), which is even higher compared to the conventional AC magnetometry configuration. Figure 5.6.6d additionally shows that, in this mode, single electron spin detection can still be possible with the pixel integration time of 60 seconds. We ran multiples of simulation with different choices of NV/dipole orientations and NV-to-sample distances (data not shown), and confirmed that these two different AC magnetometry modes result in similar signal strengths. In conclusion, our resonator-motion-enabled AC magnetometry can be as good as target-modulation AC magnetometry, and small magnetic targets such as single electrons can be detected within a few tens of seconds. Note that our magnetic field sensitivity can be improved further by employing higher quality diamond [5] and introducing more efficient photon collection techniques [3, 32].





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Colophon

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