Demonstration of diamond nuclear spin gyroscope

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We demonstrate the operation of a rotation sensor based on the nitrogen-14 (14N) nuclear spins intrinsic to nitrogen-vacancy (NV) color centers in diamond. The sensor uses optical polarization and readout of the nuclei and a radio-frequency double-quantum pulse protocol that monitors 14N nuclear spin precession. This measurement protocol suppresses the sensitivity to temperature variations in the 14N quadrupole splitting, and it does not require microwave pulses resonant with the NV electron spin transitions. The device was tested on a rotation platform and demonstrated a sensitivity of $4.7°/\sqrt{s}$ (13 mHz/√Hz), with a bias stability of 0.4°/s (1.1 mHz).

INTRODUCTION

Rotation sensors (gyroscopes) are broadly used for navigation, automotive guidance, robotics, platform stabilization, etc. (1). Various rotation sensor technologies can be subdivided into commercial and emerging. Commercial sensors include mechanical gyroscopes, Sagnac effect optical gyroscopes, and microelectromechanical systems (MEMS) gyroscopes. Among emerging technologies are nuclear magnetic resonance (NMR) gyroscopes. These sensors use hyperpolarized noble gas nuclei confined in vapor cells (2–8); they surpass commercial devices within the next decade in terms of accuracy, robustness, and miniaturization (9).

Recently proposed nuclear spin gyroscopes based on nitrogen-vacancy (NV) color centers in diamond (10, 11) are analogs of the vapor-based NMR devices, constituting a scalable and miniaturizable solid-state platform, capable of operation in a broad range of environmental conditions. An attractive feature is that a diamond sensor can be configured as a multisensor, reporting on magnetic field, temperature, and strain while also serving as a frequency reference (12–14). This multisensing capability is important for operation in challenging environments (15).

In this work, we demonstrate a diamond NMR gyroscope using the 14N nuclear spins intrinsic to NV centers. Its operation is enabled by direct optical polarization and readout of the 14N nuclear spins (16) and a radio-frequency (RF) double-quantum (DQ) pulse protocol that monitors 14N nuclear spin precession. This measurement technique is immune to temperature-induced variations in the 14N quadrupole splitting (16, 17). In contrast to recent work (18, 19), our technique does not require microwave pulses resonant with the NV electron spin transitions. The advantage of this technique is that it directly provides information about the nuclear spin states without requiring precise knowledge of the electron spin transition frequencies, which are susceptible to environmental influences (16, 18, 20). The nuclear spin interferometric technique developed in this work may find application in solid-state frequency references (12, 21) and in extending tests of fundamental interactions (2, 22) at micro- and nanoscale (23, 24) to those involving nuclear spins. With further improvements, it may also find use in practical devices such as miniature diamond gyroscopes for navigational applications.

RESULTS

Experimental setup

Figure 1 shows an experimental setup schematic for demonstration of the diamond gyroscope. The diamond sensor, green diode laser, photodetector, and all optical components are mounted on a 35-cm-diameter rotating platform controlled by a commercial rate table system (Ideal Aerosmith, 1291BL). The diamond is a 400-μm-thick single-crystal plate with an NV concentration of ~4 parts per million (ppm). It is placed in an axial magnetic bias field of 482 G aligned along the NV symmetry axis, providing optimal conditions for optical polarization and readout of 14N nuclear spins (16). The bias magnetic field is produced by two temperature-compensated samarium-cobalt (SmCo) ring magnets (<10 ppm/°C). These magnets were designed and arranged in a configuration that minimizes magnetic field gradients across the sensing volume. An aspheric condenser lens is used to illuminate an ~50-μm-diameter spot on the diamond with 80 mW of green (515 or 532 nm) laser light and collect NV fluorescence. The fluorescence is spectrally filtered with band-pass filter (650 to 800 nm) and focused onto one of the channels of a balanced photodetector. A small portion of laser light is picked off from the excitation path and directed to the other photodetector channel for balanced detection. RF pulses for nuclear spin control are delivered using a 160-μm-diameter copper wire placed on the diamond surface next to the optical focus. To reduce the ambient magnetic field noise, we placed a part of the setup including the diamond and magnets inside low-carbon-steel magnetic shields (see section S1).

Rotation detection principle

Rotation was detected by measuring the shift in the precession rate of 14N nuclear spins intrinsic to NV centers in diamond. Figure 2A shows the ground-state energy level diagram of the NV center. The inset shows the 14N nuclear spin states $|m_I\rangle$ within the $|m_S=0\rangle$ electron spin state manifold. The $|m_I=\pm 1\rangle$ states are degenerate at zero magnetic field and separated from $|m_I=0\rangle$ by ~4.94 MHz because of the nuclear quadrupole interaction. In the presence of an applied magnetic field $B$, the $|m_I=\pm 1\rangle$ states experience a Zeeman shift, and the splitting frequency $\Delta f_{Q}$ between $|m_I=+1\rangle$ and $|m_I=-1\rangle$ is described by the following equation (16)

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**Fig. 1. Experimental setup.** Electrical connections for the laser and the photodetector are wired to the rotating platform through slip-ring lines. RF signals are delivered to the platform via a single-channel RF rotary joint. NA, numerical aperture. ND, neutral density.

**Fig. 2. DQ measurement protocol using $^{14}$N nuclear spins in diamond.** (A) Energy-level diagram of the NV center ground state with and without a magnetic field $B$ applied along the NV axis. The inset depicts the $^{14}$N nuclear spin levels, where the splitting between the $|m_i = ±1⟩$ sublevels depends on the applied field and on the rotation of the sample around the NV axis. Rotation sensing is based on the measurement of this interval. (B) DQ nuclear Ramsey pulse sequence. Inset: 4-Ramsey phase cycling measurement scheme. (C) DQ nuclear Ramsey fringes (R1, R2, R3, and R4) obtained by sequentially alternating phases of the last double quantum pulse as depicted in the inset of (B). The frequency-domain spectrum shows the square of the absolute value of the Fourier transform, which reveals DQ signal at $f_{DQ}$ frequency and residual single-quantum (SQ) signals at $f_1$ and $f_2$ frequencies. Bottom left: DQ nuclear 4-Ramsey fringes obtained by combining four DQ sequential Ramsey measurements $R = R_1 − R_2 + R_3 − R_4$ to cancel residual SQ signals. (D) DQ nuclear 4-Ramsey fringes. Symbols represent experimental data, while the solid line is an exponentially decaying sine wave fit. The oscillation frequency of the signal corresponds to the $f_{DQ}$ while an exponential decay time corresponds to the nuclear DQ spin coherence time $T_2^* = 1.95(5)$ ms. Inset: Zoomed plot of DQ 4-Ramsey fringes near the working point; rotation measurements were performed at a fixed free precession time $t = 1.4$ ms by monitoring changes in the fluorescence signal.
where $\gamma_e$ and $\gamma_n$ are the electron and nuclear $^{14}$N gyromagnetic ratios, respectively; $D$ is the NV electron spin zero-field splitting parameter; and $A_z$ is the transverse magnetic hyperfine constant.

The $^{14}$N nuclear spins are prepared in a $|m_I = \pm 1\rangle$ superposition state and precess about their quantization axis with nuclear precession frequency $f_{\text{DQ}}$. If the diamond rotates about this axis with a frequency $\nu$, then the nuclear precession frequency in the diamond reference frame is $f_{\text{DQ}} + 2\nu$ (see Fig. 2A), where the factor of 2 arises from the fact that each state $|m_I = \pm 1\rangle$ is shifted by $\pm\nu$. The rotation detection presented in this work is achieved by sensitively measuring these frequency shifts using a Ramsey interferometry technique.

**Measurement protocol**

Figure 2B shows a DQ nuclear Ramsey pulse sequence. A green laser pulse (300-µs duration) initializes the $^{14}$N nuclear spins into $|m_I = \pm 1\rangle$ (25–28). Subsequently, an RF $\pi$ pulse (frequency $f_1$; 37-µs duration) is applied to transfer the population into $|m_I = 0\rangle$. Next, to induce a DQ coherence between $|m_I = \pm 1\rangle$ and $|m_I = -1\rangle$, the $f_1$ and $f_2$ transitions are simultaneously irradiated with a two-tone $\pi/\sqrt{2}$ resonant RF pulse. Then, after a free precession interval $\tau$, a second two-tone resonant RF pulse is applied to project the phase between the two nuclear spin states $|m_I = \pm 1\rangle$ into a measurable population difference, which is read out optically. The DQ nuclear Ramsey sequence leads to population oscillations as a function of $\tau$ with a frequency $f_{\text{DQ}} + 2\nu$.

RF pulse imperfections, which arise from RF gradients across the sensing volume, lead to the appearance of residual single-quantum (SQ) signals at frequencies $f_1$ and $f_2$. These SQ transitions are sensitive to temperature fluctuations, and their signals limit the performance and robustness of the technique. To remove these SQ signals, we implemented a “4-Ramsey” measurement protocol, which was recently demonstrated for the NV electron spins in (29); in this work, we extend it to nuclear spins.

The DQ nuclear 4-Ramsey protocol consists of four successive DQ Ramsey sequences with the phase of the second DQ pulse alternated (see Fig. 2B, inset). Figure 2C (top left) demonstrates four optically detected $^{14}$N DQ nuclear Ramsey signals (R1 to R4) measured by sequentially alternating the phase of the second DQ pulse at each $\tau$ delay. The Fourier transform of the individual DQ Ramsey signals reveal residual SQ signals at $f_1$ and $f_2$ frequencies along with DQ signal at $f_{\text{DQ}}$ frequency (top right). Residual SQ signals can be eliminated by combining four DQ Ramsey signals in the following way: $R = R_1 - R_2 + R_3 - R_4$ (bottom left). The Fourier transform of the combined DQ 4-Ramsey signal shows no evidence of residual SQ signal (bottom right).

Figure 2D shows the optically detected $^{14}$N DQ nuclear 4-Ramsey fringes. Experimental data (symbols) are fit to an exponentially decaying sinusoidal function $R(\tau) = A e^{-\alpha T_2} \sin(2\pi f_{\text{DQ}} \tau + \phi)$ (solid line). From the fit to the Ramsey data, we infer the oscillation frequency $f_{\text{DQ}} = 293.332(1)$ kHz and nuclear spin coherence time $T_2 = 1.95(5)$ ms.

Rotation measurements were performed by implementing the DQ nuclear 4-Ramsey protocol at fixed delay $\tau_{\text{wp}} = 1.428$ ms (labeled as working point on Fig. 2D, inset) to detect changes in the phase of the oscillation. The value of $\tau_{\text{wp}}$ was selected to yield the greatest sensitivity (see section S2). Changes in the DQ 4-Ramsey signal were recorded over time and converted into frequency shifts using a calibration coefficient. This coefficient is determined by the values of $\tau_{\text{wp}}$ and the amplitude $A$ of the DQ 4-Ramsey fringes in the vicinity of the working point according to the expression $\alpha = 4\pi \cdot \frac{c}{\tau_{\text{wp}}} \cdot A$ and was found to be $\alpha = 6.56(6) \times 10^{-5}\%/({^\circ}/s)$ (see section S3). The dynamic range for the measurement of the rotation rate is discussed in section S4. The bandwidth of the diamond gyroscope is determined by the duration of the DQ 4-Ramsey sequence $T_{\text{cycle}}$, which, under current experimental conditions, is $1/(2T_{\text{cycle}}) \approx 70$ Hz.

**Diamond gyroscope demonstration and noise characterization**

To demonstrate the capabilities of our diamond gyroscope across a range of rotation rates, we performed a series of test experiments on a rate table. First, we performed a calibration of the gyroscope by varying the rotation rate of the platform while measuring the DQ 4-Ramsey fluorescence signal. This was accomplished by repeatedly sweeping the rotation rate between $-180$ and $180^\circ$/s with a linear sweep rate of $1.8^\circ$/s$^2$ and plotting the resulting diamond gyroscope optical output signal against the rate-table rotation rate output, shown in Fig. 3A. The gyroscope output exhibits a linear response over the measured range with a proportionality factor of $\alpha = 6.58(5) \times 10^{-5}\%/({^\circ}/s)$, which agrees with the expected value $\alpha_0 = 6.56(6) \times 10^{-5}\%/({^\circ}/s)$.
obtained from analyzing the Ramsey fringes without physically rotating the table.

Next, we used the measured value of $\alpha$ to convert the diamond gyroscope fluorescence signal into a calibrated rotation signal. Figure 3B shows the rate-table output signal (red) and the diamond gyroscope signal (black), which were recorded simultaneously for various rotation sequences.

Figure 4 shows the Allan deviation of the nonrotating diamond gyroscope noise measurement as a function of averaging time. The rotation angle random walk sensitivity of a diamond gyroscope at current experimental conditions is $4.7^\circ/\sqrt{s}$ (13 mHz/√Hz) and is near the photoelectron shot noise limit, which is calculated to be $3.5^\circ/\sqrt{s}$ (10 mHz/√Hz) (see section S3). The uncertainty improves until 300 s and reaches 0.4°/s (1.1 mHz) in bias stability. The dashed dotted line shows a $1/\sqrt{t}$ dependence consistent with frequency white noise.

DISCUSSION

We have demonstrated a solid-state NMR gyroscope based on $^{14}$N nuclear spins intrinsic to NV centers in diamond. Key features of the demonstrated technique include the optical polarization and readout of the nuclear spins without the use of microwave transitions and implementation of nuclear DQ Ramsey measurement schemes, which use the entire spin-1 manifold and reduce sensitivity to temperature fluctuations. Through the use of temperature-compensated magnets, magnetic shielding, and robust pulse protocols, we reduce the influence of temperature and magnetic field drift, extending the long-term stability of the gyroscope to hundreds of seconds. The current device demonstrates a sensitivity of $4.7^\circ/\sqrt{s}$ (13 mHz/√Hz) and a bias stability of 0.4°/s (1.1 mHz). These results represent an order of magnitude improvement over those previously achieved with diamond (18, 19). While the current performance of diamond gyroscope is inferior to that of vapor-based NMR gyroscopes, there is still a lot of room for improvement.

The sensitivity can be further improved by at least one order of magnitude by increasing the detected fluorescence, which can be achieved by increasing the laser power and by increasing the photon collection efficiency through improved collection optics and index matching (30, 31). The sensitivity can also be improved by extending the $^{14}$N nuclear spin coherence time. The present sensitivity is limited by the $^{14}$N nuclear spin $T_2^*$, which is ~2 ms. However, we have measured the nuclear spin longitudinal relaxation time $T_1$ to be ~80 s in the same sample. It may be possible to increase the coherence time by up to four orders of magnitude (and improve sensitivity up to two orders of magnitude) using decoupling methods such as spin-bath driving (32, 33).

The long-term stability can be improved by reducing ambient and bias magnetic field drifts. Ambient magnetic field drifts may be reduced with better magnetic shielding, while bias magnetic field drifts can be reduced by temperature stabilization of magnets. Any remaining magnetic field drifts can be corrected using NV electron spin magnetometry (18).

In future research, we will use the $^{13}$C nuclei ensemble in diamond as an alternative sensing spin species. At the natural abundance, these nuclei provide a higher spin density than $^{14}$N intrinsic to NV centers by several orders of magnitude. This in combination with their smaller spin-relaxation rates makes $^{13}$C a promising candidate for rotation sensing. However, using the bulk $^{13}$C requires development of efficient methods for bulk polarization and spin readout, which is an active area of current research (34–39).

MATERIALS AND METHODS

The diamond used in our experiments is a $^{12}$C-enriched diamond (99.99% $^{12}$C) from Element Six. It is a (111)-cut diamond plate (2.5 mm by 1.5 mm by 0.4 mm) grown using chemical vapor deposition with an initial nitrogen concentration of ~13 ppm. NV centers were created by irradiating the sample with 4.5 MeV electrons and subsequent annealing in vacuum at 800°C for 1 hour. The estimated NV concentration is ~4 ppm.

A linearly polarized 515-nm green diode laser light (TOPTICA Photonics iBeam smart 515) was used to excite NV centers in rotation experiments. Laser pulses were generated using an external iBeam smart digital modulation option. In experiments that did not require rotation, we used 532-nm laser (Coherent Verdi G5) due to its superior noise characteristics. Laser pulses for 532-nm laser were generated by passing the continuous-wave laser beam through an acousto-optic modulator. A 0.79–numerical aperture aspheric condenser lens (Thorlabs, ACL25416U-A) was used to illuminate a spot on the diamond with ~80 mW of green light and collect fluorescence. Red fluorescence was separated from the excitation light by a dichroic mirror, passed through a 650- to 800-nm band-pass filter, and detected by a balanced photodetector (Thorlabs, PDB210A), producing ~0.1 mA of photocurrent.

RF pulses with carrier frequencies ($f_1 \approx 5.089$ MHz and $f_2 \approx 4.796$ MHz) were generated by two arbitrary waveform generators (Keysight, 33512B). Each two-channel generator was programmed to output two identical pulses on each channel that are phase shifted by 180° with respect to each other. The pulses with the desired phases were subsequently selected using two transistor–transistor logic (TTL)–controlled RF switches (Mini-Circuits, ZASWA-2-50DR), combined with power combiner (Mini-Circuits, ZFSC-2-6-75), and amplified by an RF amplifier (Mini-Circuits, LZY-22+) (see section S5). RF signals were delivered to the platform via a single-channel RF rotary joint. A TTL pulse card (SpinCore, PBESR-PRO-500) was used to generate and synchronize the pulse sequence. A data acquisition card (National Instruments, USB-6361) was used to digitize experimentally measured signals (see section S4).
SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at https://science.sciencemag.org/content/379/6614/eaax9384.full

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