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Scalable Spin Amplification with a Gain Over a Hundred

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We propose a scalable and practical implementation of spin amplification which does not require individual addressing nor a specially tailored spin network. We have demonstrated a gain of 140 in a solid-state nuclear spin system of which the spin polarization has been increased to 0.12 using dynamic nuclear polarization with photoexcited triplet electron spins. Spin amplification scalable to a higher gain opens the door to the single spin measurement for a readout of quantum computers as well as practical applications of nuclear magnetic resonance spectroscopy to infinitesimal samples which have been concealed by thermal noise.

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A magnetic moment of a single nuclear spin is so small that its induction signal is buried under thermal noise. The minimum detectable number of perfectly polarized nuclear spins is still in the order of 10^6 with a cryogenic inductive detection circuit. If information of a single nuclear spin is quantum-logically copied to a large number of spins, a spin component can be amplified and hence its signal may be detected with sufficient signal-to-noise ratio (SNR). This scheme is called spin amplification, which can be realized by the quantum circuit shown in Fig. 1(a) [1]. Although copy of an arbitrary, unknown state itself is prohibited by the no-cloning theorem [2], the information about whether the state is in $|0\rangle$ or $|1\rangle$ can be copied. Figure 1(a) can be used for a readout of quantum computers.

We propose other potential applications of spin amplification. To experimentally determine an arbitrary, unknown quantum state $|\psi\rangle$ of a spin, quantum state tomography is used, in which the state $|\psi\rangle$ is repeatedly prepared and measured [3]. If the number of repetitions is N and the measurement is limited by the thermal noise of the detection apparatus, then the SNR is increased by a factor of \sqrt{N} . If the state is prepared and accumulated elsewhere to make N identical copies of $|\psi\rangle$, as shown in Fig. 1(b), and measured at once, then the signal becomes N times as large and the SNR is increased by a factor of N, which is nontrivial because it is much more efficient than \sqrt{N} attained by the simple repetition. This scheme can also be regarded as amplification in a broader sense because it improves the SNR against the noise entering afterwards. The spin amplification [Fig. 1(b)] can be applied to quantum metrology which estimates a small unknown parameter ϕ contained in a quantum evolution U from the initial state $|0\rangle$ to the final state $|\psi\rangle$ [4]. This procedure is regarded as spectroscopy as well when the parameter ϕ is given by an internal Hamiltonian of a spin system of interest. The minimum detectable value of ϕ is improved by a factor of N as long as it is limited by the thermal noise of the detection apparatus.

In the *S*- I^m spin system (a rarely existing *S* spin and *m* abundant *I* spins), the absolute maximum gain of spin amplification shown in Figs. 1(a) and 1(b) is limited by *m*. To increase the maximum gain, *m* must be increased. Figures 1(a) and 1(b) are, in principle, scalable, but they hardly scale up to larger *m* in practice because they require individual addressing to all spins or a special spin network with tailored couplings such as a spin chain and a star topology [5–8]. Until now, a gain as low as four has been demonstrated with nuclear spins in molecules in solution [8,9]. Building a spin amplifier with a high gain, say, 100, seems as daunting as building a 100-qubit quantum computer.

In this Letter, we propose a new scalable implementation of spin amplification shown in Fig. 1(c), which utilizes spin diffusion [10] and magnetic-field cycling [11]. The total Hamiltonian of an *S-I^m* nuclear spin system with a dipolar coupling network, naturally existing in a bulk solid, is given by



FIG. 1 (color online). (a) Spin amplification referred to by DiVincenzo [1]. (b) Spin amplification of the *S* spin response signal using selective SWAP gates and (c) that using the hetero-SD process. (d) Scalable version of (a). The *I* spins are initialized to $|0\rangle$ for simplicity, although they do not have to be in practice.

$$\mathcal{H} = \omega_S S_Z + \omega_I \sum_i I_{iZ} + \sum_{i,j} \mathcal{H}_{i,j} + \sum_i \mathcal{H}_{i,S}, \quad (1)$$

where ω_S and ω_I are the resonance frequencies of the rare spin S and the abundant spins I. $\mathcal{H}_{i,i}$ and $\mathcal{H}_{i,S}$ represent the homonuclear and the heteronuclear dipolar interactions, respectively. The dipolar interaction, $\mathcal{H}_{i,j}$, between the *i*th spin I_i and the *j*th spin I_j (or S for the heteronuclear case, $\mathcal{H}_{i,S}$) is given by $\mathcal{H}_{i,j} = \mathcal{H}_{i,j}^{ZZ} + \mathcal{H}_{i,j}^{XY}$, where $\mathcal{H}_{i,j}^{ZZ} = d_{i,j}I_{iZ}I_{jZ}$ (for the heteronuclear case, the dipolar coupling strength $d_{i,j}$ is replaced with $d_{i,S}$) generates mutually controlled rotations about the z axis and $\mathcal{H}_{i,i}^{XY}$ = $-d_{i,j}(I_{iX}I_{jX} + I_{iY}I_{jY})/2$ generates fractional power of iSWAP operations. Both are always on between homonuclear spins regardless of the static magnetic field strength. Between heteronuclear spins, the former is always on and the latter is turned off in high static magnetic fields by the Zeeman energy difference being much larger than the dipolar interaction strength $(|\omega_s - \omega_I| \gg |d_{i,s}|)$. In low fields $(|\omega_S - \omega_I| \leq |d_{i,S}|)$, the latter is turned on. Under free time evolution of the total Hamiltonian in the high field, the z components of only the I spins are shuffled by simultaneous fractional powers of iSWAP operations, and this behavior is known as spin diffusion [10]. In the low field, the S spin is also involved in the spin diffusion among the *I* spins via its coupling to at least one *I* spin: we call it the heteronuclear-spin-involved spin diffusion (hetero-SD) process, where the total z spin component is preserved. The hetero-SD process can be switched on and off by magneticfield cycling between the low and the high fields [11–13]. Therefore the quantum circuit of Fig. 1(c) can be easily realized by the simple repetition of synchronized field cycling and a rf pulse irradiation for the quantum operation U to the S spin alone. The process requires only a reasonably dense network of dipolar-coupled spins without requiring the tailored coupling strength $(d_{i,j})$, and, therefore, is quite general, robust, and practical in real samples which may have some irregularities or randomness.

By exciting the z component of the S spin with a rf pulse as U and subsequently switching on the hetero-SD process for a sufficiently long duration, as shown in Fig. 1(c), the excited z component of the S spin is diffused in the reservoir of the abundant I spins. By repeating the sequence, the excitations are accumulated in the abundant Ispins. The information about whether the S spin is inverted by U is converted into the decrease in the z component of the total magnetization of *m* abundant *I* spins, which is proportional to the I spin polarization. Assuming that the polarizations of the S spin and the I spins are uniformly mixed in the hetero-SD process, the I spin polarization after the *N*th repetition of the U = NOT and the hetero-SD process decreases to $\epsilon_0 [(m-1)/(m+1)]^N$ from the initial polarization ϵ_0 of the S and the I spins. Compared with the difference between the directly observed signal of the S spin of polarization ϵ_0 and that of the inverted S spin, the



FIG. 2 (color online). The top dashed line shows the gain G of the SNR of ideal spin amplification [Figs. 1(a) and 1(b)] with respect to the number of SWAP gates N. The solid lines show the expected gain [Eq. (2)] of the proposed one [Fig. 1(c)] with respect to the number of the hetero-SD process gates N for different system sizes. The bottom dashed line shows the gain of the direct simple repetitive detection with respect to the number of repetitions N.

expected gain of the signal obtained from the polarization difference of m abundant I spins at the Nth step of spin amplification with and without U is given by

$$G = \frac{m\{\epsilon_0 - \epsilon_0[(m-1)/(m+1)]^N\}}{\epsilon_0 - (-\epsilon_0)}$$
$$= \frac{m}{2} \left[1 - \left(\frac{m-1}{m+1}\right)^N \right].$$
(2)

In an *S*-*I*^{*m*} spin system, the gain of the spin amplification using the hetero-SD process increases with *N* but is saturated to m/2 for $N \gg m/2$, as shown in Fig. 2. The proposed implementation using the hetero-SD process gives better SNR improvement with respect to *N* than the simple *N* repetitive detection. When *m* and *N* are increased simultaneously while keeping N = m/2, the amplification gain scales up linearly as $m(1 - e^{-1})/2$ (the dotted line) with respect to $m (\gg 1)$.

The quantum circuit of Fig. 1(d) is a scalable version of Fig. 1(a) and could be used for a readout of quantum computers [14]. While there is a functional difference between Figs. 1(d) and 1(c), the most difficult part, the scalability of implementation to achieve a higher gain, is common in Figs. 1(d) and 1(c). In Fig. 1(d), the information about whether the state of a rare spin S is $|0\rangle$ or $|1\rangle$ is nondestructively copied to a buffer spin I_b by the CNOT gate and accumulated in abundant I spins with hetero-SD processes. The field cycling between a high field ($|\omega_I|$ – $|\omega_{Ib}| \gg d_{I,Ib}$ and $|\omega_S - \omega_{Ib}| \gg d_{S,Ib}$ and a low field $(|\omega_I - \omega_{Ib}| \sim d_{I,Ib} \text{ and } |\omega_S - \omega_{Ib}| \gg d_{S,Ib})$ enables the hetero-SD process in Fig. 1(d). Our implementations, both Figs. 1(c) and 1(d), using the field cycling do not require rf irradiation to the abundant I spins, of which a subtle imperfection causes the large degradation of the spin amplification gain.





FIG. 3 (color online). (a) An experimental procedure of spin amplification using the hetero-SD process [15]. (b) A DNP sequence with integrated solid effect [16]. (c) A schematic diagram of an experimental setup where the sample position at each stage is indicated.

In this work, we have performed experiments to amplify the response of a rare spin with respect to a rf pulse and demonstrated a spectroscopic application in an $S-I^{799}$ spin system in a bulk solid with the quantum circuit shown in Fig. 1(c). We used a sample of a single crystal of naphthalene doubly doped with $\sim 0.005 \text{ mol }\%$ pentacene and $\sim 1 \mod \%$ 2-fluoronaphthalene. If the sample is uniform, it is approximately regarded as the ensemble of $S-I^{799}$ spin systems because a ¹⁹F spin is surrounded by 799¹H spins on average. The experimental procedure is shown in Fig. 3(a) [15]. In the first stage of every experiment, we enhanced the ¹H spin polarization to ~ 0.12 at 233 K by dynamic nuclear polarization (DNP) with photoexcited triplet electron spins of pentacene molecules [16–19]; the sequence of the photoexcitation and the integrated solid effect [16] are shown in Fig. 3(b).

The key stage in this experiment, the hetero-SD process, was realized by field cycling (4000 G \rightarrow 100 G \rightarrow 4000 G) using sample shuttling, as shown in Fig. 3(c). In 100 G, the resonance frequency difference between the ¹H and the ¹⁹F spins is of the same order as the strength of the nearest neighbor ¹H-¹⁹F dipolar coupling. After repeating a rf pulse to ¹⁹F spins as the quantum operation *U* and the sample shuttling *N* times, we observed the ¹H magnetization.

We investigated the behavior of the ¹H spin polarization with respect to the number of steps *N* of spin amplification [Fig. 4(a)]. In the case of U = I with no rf pulse, the ¹H spin polarization decayed with η of ~99.91% per cycle, which gave close agreement with the decay rate calculated from T_1 measured at each sample position. In the case of an on-resonance π pulse (i.e., U = NOT), the result was close to the theoretical decay $\epsilon_0\{(m-1)/(m+1)\}^N \eta^N$ (the dashed line) calculated from the decreased polarization in Eq. (2) and the decay η . The *S* spin response to *U* is converted into the difference between the *I* spin polarization with and without *U*. The polarization difference was amplified 37 times for N = 40 and 136 times for N = 200, compared with that for N = 1. The amplification gain



FIG. 4 (color online). (a) The behaviors of the ¹H spin polarization with respect to the number of steps N of spin amplification without rf pulse (U = I) and with on-resonance π pulses (U = NOT). The dashed line is the theoretical decay. (b) The circles show the spin-amplified frequency response spectra obtained for N = 40 and 200. They were obtained with the rf pulse with a peak amplitude of 45 kHz (the pulse length is ~3 times as long as that of 140 kHz). The squares show the spin-amplified frequency response spectra obtained for N = 200 with the rf pulse with a peak amplitude of 140 kHz. Directly detected ¹⁹F spectrum vertically magnified is also shown for comparison.

anticipated from Eq. (2) is 38 for N = 40 and 157 for N = 200. By taking into account the frequencies for ¹⁹F and ¹H spins, the signal obtained from the polarization difference of the ¹H spins with and without U = NOT was estimated to be ~140 times as large as that of the ¹⁹F spin with the polarization 0.11. It is noteworthy that the 140-time amplified signal is more than 10 times as large as the signal of the perfectly polarized ¹⁹F spins. Furthermore, a factor of 140 spin amplification in conjunction with the modern magnetic resonance force microscopy [20] technology, which is expected to detect 100 nuclear spins in the near future [21], may open the door to the detection of a faint signal from a single nuclear spin.

Amplified responses to the rf pulses with various carrier frequencies for N = 200 give the excitation spectrum of the ¹⁹F spin, as represented by the squares in Fig. 4(b). The spin-amplified response spectra with the weaker rf pulse (the circles) were narrower than that but still broader than the directly observed ¹⁹F NMR spectrum. The pulse intensity should be lowered in order to avoid power broadening. The depth of the spectral response was increased with *N*.

All spin-amplified ¹H NMR signals were reduced by a factor of 100 on purpose to avoid saturation of the detection circuit [15]. With the reduction, the directly observed ¹⁹F NMR signal was completely concealed with the thermal noise. The ¹⁹F signal observed without the reduction is shown in Fig. 4(b). If the sample size is 100 times as small, the ¹⁹F signal cannot be detected even without the reduction. On the other hand, the spin-amplified ¹H signals of the small sample can be detected with the approximately

same SNR as the present result without the reduction. Spin amplification will have a distinct advantage over the simple repetitive detection in a smaller or a more diluted sample. If we use a sample with a lower 2-fluoronaphthalene concentration, we could get a higher gain by increasing N in the same manner. The maximum achievable gain is inversely proportional to the concentration. It has been reported that T_1 of the ¹H spins in naphthalene doped with \sim 0.01 mol % pentacene in \sim 7 G at 77 K is \sim 166 min [18], and the result implies that T_1 of a sample with a lower 2-fluoronaphthalene concentration at a lower temperature is longer. In addition, there is room for speeding up field cycling [11,22], and hence earning the number of steps Nwell before the ¹H polarization vanishes by T_1 relaxation. Scalable spin amplification will enable practical applications of NMR spectroscopy to rarely existing nuclear spins in infinitesimal bulk samples which have been concealed by thermal noise. For higher sensitivity and spectral resolution, the quantum operation U and the NMR detection should be performed in a higher magnetic field, which is realized by the field-cycling system described in [23].

In the proposed implementation using field cycling for switching the hetero-SD process, the nuclear species are modestly restricted to those with the resonant frequency close to that of the abundant ¹H spins. For other nuclear species to be involved in spin diffusion process, the double resonance irradiation satisfying the Hartmann-Hahn condition [24], as in the cross polarization experiment [25], may be used, although it is subject to the rf pulse imperfection and the much faster decay with $T_{1\rho}$. State-of-the-art high fidelity pulse engineering [26,27] may be required to implement high-gain spin amplification.

The demonstrated spin amplification [Fig. 1(c)] can be straightforwardly modified to Fig. 1(d). It can be realized, for example, by doping 2-¹³C 2-fluoronaphthalene, in which the ¹³C, ¹⁹F, and ¹H spins play the roles of the *S*, the I_b , and the *I* spins, respectively. Recently, optically detected magnetic resonance [28,29] has accomplished single-event state readout of a single nuclear spin [30], which is attracting attention in quantum information science. The breakthrough of the gain shown in this Letter is an important step for state readout of a single nuclear spin in bulk solids with more versatile detection such as magnetic resonance force microscopy and the inductive detection.

In this Letter, we have proposed a scalable and practical implementation of spin amplification and successfully demonstrated a gain as large as 140. A faint magnetization of a rare spin has been quantum-logically transferred beyond the molecular boundary to abundant spins in the bulk solid sample through spin diffusion process which does not require individual addressing nor a specially tailored spin network. The amplified signal obtained from the abundant spins is more than 10 times as large as that of the completely polarized rare spins. A spectroscopic application has been demonstrated by 140-time amplified frequency response spectra of the rare spin. It will enable practical applications of NMR spectroscopy to infinitesimal samples which have been concealed by thermal noise. A higher gain is possible in the sample with a lower concentration of rare spins. Spin amplification scalable to a higher gain opens the door to a single spin measurement and a readout of spin-based quantum computers.

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