Resonant optical control of the spin of a single Cr atom in a quantum dot

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A Cr atom in a semiconductor host carries a localized spin with an intrinsic large spin to strain coupling, which is particularly promising for the development of hybrid spin-mechanical systems and coherent mechanical spin driving. We demonstrate here that the spin of an individual Cr atom inserted in a semiconductor quantum dot can be controlled optically. We first show that a Cr spin can be prepared by resonant optical pumping. Monitoring the time dependence of the intensity of the resonant fluorescence of the quantum dot during this process permits us to probe the dynamics of the optical initialization of the Cr spin. Using this initialization and readout technique we measured a Cr spin relaxation time at $T = 5 \text{ K}$ in the microsecond range. We finally demonstrate that, under a resonant single-mode laser field, the energy of any spin state of an individual Cr atom can be independently tuned by using the optical Stark effect.

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Individual spins in semiconductor nanostructures are promising for the development of quantum information technologies [1–3]. Spins trapped in optically active quantum dots (QDs) have attracted strong interest since their coupling to light enables fast spin control, and optical coherent control has been demonstrated for electron [4–6] and hole [7,8] spins confined in QDs. Thanks to their expected longer coherence time, localized spins on individual dopants in semiconductors are also promising media for storing quantum information. Optically active QDs containing individual or pairs of magnetic dopants have been realized both in II–VI [9–13] and III–V [14,15] semiconductors. In these systems, since the confined carriers and magnetic atom spins become strongly mixed, an optical excitation of the QD can affect the spin state of the atom, offering possibilities for a control of the localized spin [16,17]. The variety of magnetic transition elements that can be incorporated in semiconductors gives a large choice of localized electronic spins, nuclear spins, and orbital momentums with optical addressability [9,18–20]. This approach opens a diversity of applications in quantum information and quantum sensing.

Among these magnetic atoms, chromium (Cr) is of particular interest [20]. It incorporates in II–VI semiconductors as Cr$^{3+}$, carrying an electronic spin $S = 2$ and an orbital momentum $L = 2$. Moreover, most of Cr isotopes have no nuclear spin. This simplifies the spin level structure and its coherent dynamics [21]. With biaxial strain, the ground state of the Cr is an orbital singlet with spin degeneracy of 5. The orbital momentum of the Cr connects its spin to the local strain through the modification of the crystal field and the spin-orbit coupling. This spin to strain coupling is more than two orders of magnitude larger than for elements without orbital momentum (Nitrogen Vacancy centers in diamond [22], Mn atoms in II–VI semiconductors [23]). Cr is therefore a promising qubit for hybrid spin-mechanical systems in which the motion of a mechanical oscillator would be coherently coupled to the spin state of a single atom [22,24,25]. The development of these hybrid systems will require efficient control of the Cr spin.

In this article, we first show that the spin of a Cr atom inserted in a CdTe/ZnTe QD can be prepared by resonant optical pumping. The resonant photoluminescence (PL) of the QD is used for the readout of the Cr spin. This spin initialization and readout technique is used to probe the Cr spin relaxation, which remains in the $\mu$s range at low temperature. We finally demonstrate that under a strong resonant laser field, the energy of any spin state of a Cr atom can be tuned by using the optical Stark effect. This ensemble of experiments sets the basis required for a full optical control of this single-spin system and opens the path toward the development of coherent spectroscopy techniques to probe the strained induced coherent dynamics of the magnetic atom.

To optically access an individual magnetic atom, Cr are randomly introduced in CdTe/ZnTe self-assembled QDs during their growth [20,26]. The amount of Cr is adjusted to allow the detection of QDs containing 1 or a few Cr atoms. The PL of individual QDs is studied by optical microspectroscopy at low temperature ($T = 5 \text{ K}$). The PL is excited with a continuous wave (cw) dye laser tuned to an excited state of the QD, dispersed and filtered by a 1-m double spectrometer before being detected by a Si-CCD camera or a fast Si avalanche photodiode. A single-mode dye ring laser can be tuned on resonance with the exciton. In time-resolved pumping experiments, trains of resonant light with variable durations and wavelengths are generated from the cw lasers using acousto-optical modulators with a switching time of about 10 ns.

The circularly polarized PL spectra of an exciton in a Cr-doped QD (X-Cr) is reported in Fig. 1(a). Because of a large magnetic anisotropy of the Cr spin induced by the biaxial strain in the QDs, $D_{0z}S_z^2$, the Cr spin thermalizes to the states $S_z = 0$ and $S_z = \pm 1$, where $z$ is the QD growth axis [Fig. 1(c)]. The exchange interaction with the spin of the bright exciton $|\pm 1\rangle$ further splits the Cr states $S_z = \pm 1$ (lines 1 and 4) [20]. Most of the dots also present a low symmetry and the central line
associated with $S_z = 0$ is split by the electron-hole exchange interaction (lines 2 and 3) and linearly polarized along two orthogonal directions [Fig. 1(b)]. An additional line (labeled line 5) appears on the low-energy side of the PL spectra. It arises from a dark exciton interacting with the same Cr spin state [20]. This mixing is induced by the electron-hole exchange interaction in a confining potential of symmetry lower than $C_{2v}$ [27]. Details about the fine structure of the exciton in a Cr-doped QD can be found in Ref. [20].

To initialize and read out the Cr spin, we developed a two-wavelength pump-probe experiment. A circularly polarized single-mode laser (resonant pump) tuned on a X-Cr level is used to pump the Cr spin (i.e., empty the spin state under resonant excitation). Then, a second laser, tuned on an excited state of the X-Cr (under resonant excitation). Then, a second laser, tuned on a X-Cr level is used to pump the Cr spin (i.e., empty the spin state during resonant excitation by the pump pulse). Under resonant excitation on the high-energy X-Cr line, an exciton spin flip with conservation of the Cr spin can produce a PL on the low-energy line [11]. In this process, the exciton flips its spin.
by emitting (or absorbing) an acoustic phonon. Such spin flip is enhanced by the large acoustic phonon density of states at the energy of the interlevel splitting induced by the exchange interaction with the Cr spin, which act as an effective magnetic field [28,29]. The resulting weak resonant PL signal depends on the occupation of the Cr state $S_z = +1$ and is used to monitor the temperature dependence of the spin selective absorption of the QD.

The time evolution of the PL of the low-energy line of X-Cr under an excitation on the high energy line is presented in Fig. 2(a) for two different pump-probe sequences: probe on and probe off. When the probe laser is on, a large effective Cr spin temperature is established before each pumping pulse. The amplitude of the resonant PL is maximum at the beginning of the pump pulse ($I_1$) and progressively decreases. A decrease of about 80% is observed with a characteristic decay time in the tens of nanoseconds range. As expected for a spin optical pumping process, the characteristic time of the PL transient decreases with the increase of the pump laser intensity [Fig. 2(b)] [11]. When the probe laser is off, the initial amplitude of the PL transient during the pump pulse is significantly weaker. This decrease is a consequence of the conservation of the out of equilibrium Cr spin distribution during the dark time between two consecutive pumping pulses.

The steady-state resonant PL intensity reached at the end of the pump pulse ($I_2$) depends on the optical pumping efficiency, which is controlled by the ratio of the spin-flip rate for the Cr spin in the exchange field of the exciton and the relaxation of the Cr spin in the empty dot. However, even with cross-circularly polarized excitation/detection, this steady-state PL can also contain a weak contribution from an absorption in the acoustic phonon sideband of the low-energy line [30]. Figure 2(c) presents the amplitude of the resonant PL detected on the low-energy line for a detuning of the pump around the high-energy line. A resonance is observed in the initial amplitude $I_1$ of the PL. It reflects the energy and excitation power dependence of the absorption of the QD. A small decrease of the steady-state PL $I_2$ is also observed at the resonance. As displayed in Fig. 2(d), the corresponding normalized amplitude of the pumping transient, $(I_1-I_2)/I_2$, presents a clear resonant behavior, demonstrating the excitation energy dependence of the optical pumping process. The width of the resonance ($\sim 100 \, \mu eV$) is the width of the QD’s absorption broadened by the fluctuating environment [31].

With this resonant optical pumping technique to prepare and read out the Cr spin, we performed pump-probe experiments to observe its relaxation time in the absence of carriers (Fig. 3). A nonequilibrium distribution of the Cr spin population is prepared with a circularly polarized resonant pump pulse on the high-energy X-Cr line. The pump laser is then switched off and switched on again after a dark time $\tau_{\text{dark}}$. The amplitude of the pumping transient observed on the resonant PL of the low-energy line depends on the Cr spin relaxation during $\tau_{\text{dark}}$. As presented in Fig. 3(b), the amplitude of the transient is fully restored after a dark time of about $10 \, \mu s$, showing that after this delay the Cr spin is in equilibrium with the lattice temperature ($T = 5 \, K$). Let us note, however, that the initial amplitude of the pumping transient in this case is weaker than the one observed after a nonresonant probe pulse [Fig. 2(a)]. This means that the nonresonant optical excitation drives the Cr spin to an effective temperature much larger than the lattice temperature.

From the time delay dependence of the amplitude of the transient, we deduce a Cr relaxation time $\tau_{\text{Cr}} \approx 1.7 \, \mu s$ at $B = 0 \, T$ and $T = 5 \, K$. For such neutral QD and in the absence of optical injection of carriers, this spin relaxation is likely to be controlled by the spin-lattice interaction. Despite the large spin-phonon coupling expected for this magnetic atom with an orbital momentum and a strain-induced spin splitting in the meV range [20], the Cr spin relaxation time remains in the $\mu s$ range. This spin memory is long enough for a practical use of Cr in an hybrid spin nanomechanical system and could even be improved in different QDs structures with weaker biaxial strain [32], lower magnetic anisotropy splitting, and consequently less coupling with acoustic phonons [33].

These measurements reveal a significantly different Cr spin-flip times under optical excitation (tens of nanosecond range) and in the dark (microseconds range). The fast Cr spin-flip under optical excitation can be due to the interaction with carriers (exchange induced Cr spin flips [33]) but can also be induced by the interaction with nonequilibrium acoustic phonons created during the energy relaxation of the injected carriers. Both mechanisms probably contribute to the Cr spin heating.

The resonant optical excitation on a X-Cr line can also be used to tune the energy of any Cr spin state through the optical Stark effect [12,34,35]. Such energy shift could be exploited to control the coherent dynamics of the magnetic atom [17,36]. This optical control technique is presented in Fig. 4. When a high-intensity single-mode laser is tuned to the high-energy line of X-Cr in $\sigma^+$ polarization (X-Cr state $|S_z = -1, +1\rangle$), a splitting is observed in $\sigma^-$ polarization in the PL of the two low-energy lines produced by a second nonresonant laser.

At high excitation intensity, a strong coupling with the resonant laser field mixes the states with a Cr spin component
A resonant laser permits us to address any spin state of the Cr and selectively shift its energy. For instance, as presented in Fig. 4(d), a $\sigma^+$ excitation on the dark exciton state (5) induces a splitting of the high-energy line 1 in $\sigma$ polarization (state $|S_z = +1, -1\rangle$) without affecting the central line 2. This shows that such resonant excitation can be used to tune the energy of $S_z = +1$ without affecting $S_z = 0$. The energy tuning induced by a coherent optical driving is particularly interesting for the control of the Cr spin states $S_z = \pm 1$. These states could be efficiently mixed by applied weak anisotropic in-plane strain through a fine structure term of the form $E(S_z^2 - S_y^2)$ [20]. A relative shift of the energy of $S_z = +1$ or $S_z = -1$ by a resonant optical excitation would affect their coupling and consequently the Cr spin coherent dynamics.

Future applications of Cr as a new spin qubit in hybrid nanomechanical systems [38,39] could exploit the efficient mixing of the Cr spin states $S_z = +1$ and $S_z = -1$ induced by anisotropic, in-plane strain. For instance, the resulting mixed spin states, together with an exciton, form an optical three-level system. A coherent optical driving of this level structure opens the possibility of using coherent spectroscopy techniques such as coherent population trapping [40] for a sensitive probing of the splitting of the $S_z = +1$ and $S_z = -1$ induced by the local strain at the atom location. The strain field that could be probed with this technique will depend on the coherence time of the Cr $|+1; -1\rangle$ spin qubit. In addition, dynamical anisotropic in-plane strain, provided for instance by propagating surface acoustic waves [41], whose frequency would be tuned on resonance with the energy splitting of $S_z = \pm 1$ under magnetic field could be used for a direct coherent control of the $|+1; -1\rangle$ spin qubit [42]. Such mechanical coherent control could be probed with the resonant optical pumping technique presented in this paper.

To conclude, we demonstrated that a resonant optical excitation can be used to control the spin of a Cr atom inserted in a QD. The Cr spin can be initialized by optical pumping, readout through the QD resonant PL and the energy of any spin state tuned by optical Stark effect. Despite the large spin-phonon coupling expected for Cr, a magnetic atom with orbital momentum, its spin relaxation time remains in the $\mu$s range at low temperature. These experiments mark an important step toward the development of optical coherent spectroscopy techniques for a precision sensing of externally applied strain on a Cr spin inserted in a nanomechanical system or for a mechanical coherent spin driving of an individual magnetic atom using dynamical strain.

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