Efficient injection of spin-polarized excitons and optical spin orientation of a single Mn$^{2+}$ ion in a CdSe/ZnSe quantum dot

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Circularly polarized optical excitation is used to demonstrate the efficient injection of spin-polarized excitons to individual self-assembled CdSe quantum dots in ZnSe barrier. The exciton spin-transfer is studied by means of polarization-resolved single dot spectroscopy performed in magnetic field applied in Faraday configuration. Detailed analysis of the neutral exciton photoluminescence spectra reveals the presence of exciton spin relaxation during its lifetime in a quantum dot. This process is seen for both nonmagnetic dots and those containing single Mn$^{2+}$ ions. Taking this into account we determine the spin-polarization degree of excitons injected to a dot under circularly polarized below-the-barrier optical excitation at 488 nm. It is found to be close to 40% in the entire range of the applied magnetic field. Exploiting the established spin-conserving excitation channel we demonstrate the optical spin orientation of a single Mn$^{2+}$ ion embedded in a CdSe/ZnSe quantum dot.

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I. INTRODUCTION

Studies of single objects in a solid have recently attracted a lot of research attention. This led to coining the term solotronics – optoelectronics based on solitary dopants\textsuperscript{1–4}. In order to achieve the control over a single dopant, in particular over its spin, different techniques were employed, depending on the studied system. For example, a single phosphorus center on Si surface could be controlled electrically, allowing for realization of a single atom transistor\textsuperscript{5}. In particular over its spin, different techniques were employed, depending on the studied system. For example, a single phosphorus center on Si surface could be controlled electrically, allowing for realization of a single atom transistor\textsuperscript{5}. In particular over its spin, different techniques were employed, depending on the studied system. For example, a single phosphorus center on Si surface could be controlled electrically, allowing for realization of a single atom transistor\textsuperscript{5}. Control of a single N-V center in diamond\textsuperscript{6,7} can be obtained using a combination of internal optical and microwave transitions, which together enabled development of a room-temperature qubit\textsuperscript{8,9} and can be used for nanosensing of local magnetic fields\textsuperscript{10}. Individual lanthanide ions such as Ce, Er or Pr can be addressed using intra-shell 4f, or inter-shell 4f-5d optical transitions\textsuperscript{11–14}.

Transition metal ions with partially filled d-shell embedded in semiconductor offer another attractive possibility of optical manipulation mediated by spin-polarized carriers, which interact with the magnetic ion due to the s-p-d exchange interaction\textsuperscript{15}. Such interaction leads to particularly interesting effects when individual magnetic ion is embedded in a quantum dot (QD)\textsuperscript{16}. In such case, the excitonic states are split due to the interaction with individual magnetic ion, allowing unambiguous readout of the magnetic ion spin state from polarization and energy of photon emitted from such system. So far four systems of QDs with single magnetic dopants have been reported: individual manganese ion in a CdTe\textsuperscript{17}, InAs\textsuperscript{18}, and CdSe QD\textsuperscript{19,20}, and individual cobalt ion in a CdTe QD\textsuperscript{4}. The optical manipulation of the ion spin has been demonstrated only in the case of CdTe and InAs QDs containing manganese ions. It has been achieved either by strictly-resonant excitation of exchange-split excitonic states\textsuperscript{19,20} or through injection of spin-polarized excitons to Mn-doped QD\textsuperscript{21,22}. Such injection typically requires efficient spin-conserving excitation channels. They have been found for CdTe/ZnTe QDs with the use of resonant excitation of either an excited excitonic state in Mn-doped QD\textsuperscript{23–25}, or a ground state of neutral exciton in an adjacent, spontaneously coupled nonmagnetic dot\textsuperscript{26–28}. However, the possibility of such resonant excitation has not been demonstrated for a CdSe/ZnSe QD containing a single manganese ion – a new promising solotronic system with very long Mn$^{2+}$ spin relaxation time\textsuperscript{4}. Nevertheless, self-organized CdSe/ZnSe QDs offer a possibility to employ non-resonant, but still spin-conserving optical excitation channels, as it was shown for QD ensembles\textsuperscript{29–31}.

In this work we show that spin-polarized excitons can be efficiently injected to individual self-assembled CdSe/ZnSe quantum dots by below-the-barrier optical excitation at 488 nm. Utilizing this technique we demonstrate the optical orientation of a single Mn$^{2+}$ ion spin in a CdSe/ZnSe QD. The paper is organized as follows. First, we present the sample growth procedure and describe the experimental setup in section II. Section III contains a discussion of a photoluminescence spectrum of an individual CdSe/ZnSe QD. Next, we demonstrate the efficient optically-induced exciton spin transfer for CdSe/ZnSe QDs, both nonmagnetic (Sec. IV) and containing single Mn$^{2+}$ ions (Sec. V). Finally, the optical orientation of Mn$^{2+}$ ion spin in a CdSe/ZnSe QD is presented and discussed in Sec. VI.

II. SAMPLE AND EXPERIMENTAL SETUP

The sample studied in this work contains a single layer of self-assembled CdSe/ZnSe QDs doped with manganese ions. It is grown using molecular beam epitaxy (MBE) on GaAs (100) substrate. The growth procedure is particularly simple. First, we grow 1 µm ZnSe buffer layer, then 2 monolayers of (Cd,Mn)Se which are immediately transformed into QDs, and finally 100 nm of ZnSe cap.
of 0-10 T. The QDs are excited quasi-resonantly by a 
ducting magnet producing magnetic field in the range 
K. The cryostat is equipped with a split-coil supercon-

cilar, sharp QD emission lines are well resolved in a broad 
energy range micro-photoluminescence spectrum shown 
in Fig. 1(b). In majority of the previ-

ous studies on selenide dots, an observation of isolated 
QD PL typically required additional sample processing, 
such as fabrication of masks or mesas\textsuperscript{30–33}, which might 
afford the optical properties of selected QD (e.g., the 
polarization of emitted light). In contrast, our unstructured 
sample exhibits significantly lower QD density (which is 
apparent from the relatively low number of emission lines 
resent in wide PL spectrum in Fig. 1(b)). In particular, 
it enables us to easily distinguish well-resolved emission 
lines of individual QDs in a long-wavelength tail of the 
PL band.

III. PL SPECTRUM OF A SINGLE CdSe/ZnSe QD

An example PL spectrum of an individual, nonmagnetic 
CdSe/ZnSe QD is shown in Fig. 2(a). It consists of 
three sharp emission lines, which are identified as origin-
ating from recombination of neutral exciton (X), nega-
tively charged exciton (X$^-$) and biexciton (2X). The 
identifcation is based on the in-plane optical anisotropy 
and relative emission energies. The sequence and relative 
energies of observed excitonic transitions are found to 
be similar to ones previously reported for self-assembled 
CdSe/ZnSe QDs\textsuperscript{27,29}. On this basis the highest and low-
est energy emission lines are first tentatively identified as related to the recombination of neutral exciton and biexciton, respectively. This attribution is further con-
formed by a characteristic fine structure of both invoked 
transitions, which is a result of the in-plane optical anisotropy 
of a QD confining potential (typical for self-organized 
dots\textsuperscript{27,29}). More specifically, the anisotropic part of 
exchange interaction between an electron and a heavy 
hole splits two bright states of neutral exciton by $\delta_1$ en-
ergy leading to the presence of two orthogonally linearly 
polarized X emission lines. Since X is a final state of a 
spin-singlet biexciton recombination, the 2X optical tran-
sitions also form a linearly polarized doublet split by the 
same energy $\delta_1$ as X emission lines. These predictions are 
reproduced by the experimental data in Fig. 2(b), which 
shows PL spectra of the analyzed QD measured for dif-
derent orientations of detected linear polarization. Due to 
relatively small $\delta_1$, the QD PL spectrum does not exhibit any noticeable fine 
structure (Fig. 2(b)). This is typical for recombination of
a trion\textsuperscript{29,32,35}, where both the initial state (with two majority carriers forming a closed shell) and the final state (consisting of one residual carrier) are strictly degenerate. Basing on this observation as well as on the comparison of relative energies of QD emission lines with the results of previous studies\textsuperscript{24,25}, we attribute the middle-energy line to a singly charged exciton. The identification of its negative sign (as marked in Fig. 2(a)) is first tentatively based on a typical tendency for selenide materials to be n-type. The sign attribution is unequivocally confirmed by an observation of negative optical polarization transfer of X\textsuperscript{−} transition under quasi-resonant excitation, which is discussed in section IV.B of this paper.

**IV. SPIN-TRANSFER FOR A CdSe/ZnSe QD**

In this section we focus on the ability to control over the spin of excitons injected to a single CdSe/ZnSe QD. In particular, we demonstrate that optical excitation of a CdSe QD with circularly polarized light of photon energy appropriately close to the QD emission energy provides a significant spin-polarization of excitons captured by the dot. In order to investigate the efficiency of spin-transfer we performed polarization-resolved measurements of QD PL under below-the-barrier CW excitation at 476.5 nm or 488 nm in external magnetic field (Faraday configuration). We studied several, randomly selected dots. Here we present data obtained for the representative dot discussed in the previous section.

**A. Neutral exciton**

We first analyze the PL of the neutral exciton. Figures 3(a)–3(i) show the X PL spectra at B = 0, 4 T and 8 T measured in two circular polarizations of detection under circularly or linearly polarized excitation at 488 nm (≈ 2.55 eV). In the absence of the magnetic field no circular polarization of X PL is observed, independently of the polarization of excitation (Figs. 3(a)–3(c)).
This is related to the fine-structure splitting of the neutral exciton state. Even if under circular polarization of excitation a spin-polarized exciton is injected to the QD, its initial state is a coherent superposition of two \( \delta_1 \)-split linearly polarized X eigenstates in the QD. Therefore, the exciton is undergoing oscillations between its eigenstates. The period of these oscillations \( 2\pi \hbar /\delta_1 \approx 40 \text{ ps} \) is several times shorter than the neutral exciton lifetime, which is of the order of a few hundreds of picoseconds in the case of CdSe/ZnSe QDs.\(^{18,31}\) As a consequence, the time-integrated X PL is almost completely unpolarized, as observed in the experiment.

The situation is qualitatively different after application of the magnetic field along the QD growth axis. The field introduces a Zeeman splitting between two spin-polarized states of the neutral exciton. At the sufficiently large field this splitting of \( g\mu_B B \) (where \( g \) is the excitonic g-factor) dominates over the X fine-structure splitting \( \delta_1 \) and exciton eigenstates in the QD become nearly pure spin states. Experimentally, it manifest itself in the presence of almost fully circularly polarized X emission lines. In the case of the studied QD, it is visible even in the field of a few Tesla (Fig. 3(j)) due to relatively small \( \delta_1 \) splitting. Thus, the application of such field enables us to determine the spin-polarization degree of excitons injected to the QD from the X PL spectra. Such spectra measured at \( B = 4 \text{ T} \) are shown in Figs. 3(d)-3(f). The X PL intensities obtained for the same circular polarization of excitation and detection (co-polarized) are clearly larger than those measured in different circular polarizations (cross-polarized). This observation unequivocally confirms the presence of an efficient optically-induced exciton spin transfer, both for \( \sigma^+ \) and \( \sigma^- \) polarized excitation at 488 nm.

Under linearly polarized excitation at \( B = 4 \text{ T} \) we also observe a small, but not negligible degree of X PL circular polarization (Fig. 3(f)). Independently of the orientation of linear polarization of excitation, the lower-energy X emission line (\( \sigma^- \)-polarized) has slightly larger intensity compared to the higher-energy one (\( \sigma^+ \)-polarized). This effect is more pronounced at higher magnetic field of 8 T (Fig. 3(i)). In order to explain its origin, we first notice that excitation with linear polarization is unlikely to induce any spin-polarization of excitons injected to the QD (unimportance of, in principle possible, linear-to-circular polarization conversion\(^{24,26,31}\) will be argued in the latter part of this section). Thus, we relate the observed effect to an exciton spin-flip process most probably assisted by an acoustic phonon. As the phonon emission is strongly favored over the phonon absorption at low temperatures, the spin-flip causes exciton relaxation towards the lower-energy state during X lifetime in the QD. As a consequence, \( \sigma^- \)-polarized emission line of the neutral exciton has enhanced intensity even under excitation providing no spin-polarization of optically-created excitons. Since the spin-flip rate exhibits a strong dependence on the energy of emitted phonon, the efficiency of X relaxation increases with an increasing Zeeman splitting. It underlies the mentioned above increase of the intensity ratio of lower- and higher-energy X emission lines at higher magnetic field under linearly polarized excitation. The influence of exciton spin-flip is also visible in the X PL spectra measured at \( B = 8 \text{ T} \) with circular polarization of excitation (Figs. 3(g), 3(h)). In this case, the intensities of X emission lines are governed by an interplay of the optically-induced exciton spin-transfer and phonon-assisted X spin relaxation. In particular, the excitation with \( \sigma^- \) (\( \sigma^+ \)) polarization results in an increased (decreased) intensity ratio of \( \sigma^- \) and \( \sigma^+ \) polarized X emission lines compared to the ratio obtained under linearly polarized excitation.

In order to perform a quantitative analysis of the exciton spin-transfer efficiency at different values of the magnetic field, we introduce a simple rate-equation model. It includes the probabilities of occupation of three states: the empty QD (\( p_0 \)) and the neutral exciton in lower- or higher-energy state (\( p_- \) or \( p_+ \), respectively). We do not take into account the biexciton state, as the excitation power used in our experiment is sufficiently low to assure that 2X emission intensity is not exceeding a few percent of the X intensity. In general, we expect that the injection of excitons to the dot is a complex process which may involve capture of both single carriers and electron-hole pairs.\(^{24,26,31}\) Since the detailed analysis of this process remains beyond the scope of this paper, we describe it by introducing two effective rates \( \alpha_+ \) and \( \alpha_- \), which account for injection of an exciton to the higher- and lower-energy state, respectively. Moreover, we take into account the exciton radiative lifetime \( \tau_X \), which is assumed to be the same for both considered X states and independent of the magnetic field. Finally, we include the spin-flip process turning a higher-energy exciton into a lower-energy one by introducing its characteristic time \( \tau_J \), which depends on the energy splitting of both invoked states. The scheme of the states and transitions is shown in Fig. 3(a). Within our model, the steady state is given by the following linear equations

\[
\frac{dp_+}{dt} = -\frac{p_+}{\tau_X} - \frac{p_+}{\tau_f} + \alpha_+ p_0 = 0, \tag{1}
\]

\[
\frac{dp_-}{dt} = -\frac{p_-}{\tau_X} + \frac{p_+}{\tau_f} + \alpha_- p_0 = 0, \tag{2}
\]

\[
\frac{dp_0}{dt} = \frac{p_0}{\tau_X} + \frac{p_+}{\tau_f} - (\alpha_+ + \alpha_-) p_0 = 0. \tag{3}
\]

As \( p_- \) and \( p_+ \) probabilities are proportional to the intensities \( I_- \) and \( I_+ \) of lower- and higher-energy X emission lines, their ratio \( I_- / I_+ \) can be easily determined from the above set of equations and reads

\[
\frac{I_-}{I_+} = \left(1 + \frac{\alpha_-}{\alpha_+}\right) \left(1 + \frac{\tau_X}{\tau_f}\right) - 1. \tag{4}
\]

In the case of linearly polarized excitation, the probabilities \( \alpha_+ \) and \( \alpha_- \) of an exciton injection to both considered X states in the QD are equal. Thus the ratio of X intensities under such an excitation is given by
in a QD predicted for a single phonon-mediated X spin relaxation.

power of X energy splitting, as previously theoretically

eigenstates in the QD. Using Eq. (4) it can be expressed

excitation. A quantitative measure of this efficiency is

dependence on the magnetic field we can extract the ex-
sis of the obtained results reveals the power dependence

of X spin-flip time at higher fields. Detailed analy-

curves given by

of neutral exciton states in the dot. In particular, the

transitions considered in the rate-equation model.

FIG. 4: (Color online) (a) A schematic illustration of QD

depends on the magnetic field. In accordance to the above considerations we ob-

the measured

energy splitting of X states. (c), (d) Degrees of spin-transfer

at 488 nm, 476.5 nm with

counting for power dependence of the X spin-flip rate on the

energy splitting. (b) Ratio of lower- and higher-energy X emission line inten-

talities of charged trion. In particular, we focus on the mea-

smeasurements carried out at magnetic field of 4 T (Faraday

power dependence

In such a scenario, the intensities of X emission lines un-

related excitonic fine structure, which prevents precise
determination of the spin-transfer efficiency at lower

fields. Interestingly, the reduction of excitation wave-

length from 488 nm to 476.5 nm (≈ 2.61 eV) results in

an almost complete loss of the exciton spin-transfer in the

entire range of the magnetic field (Fig. 4(d)).

It is noteworthy that our findings do not indicate the

presence of any effective linear-to-circular polarization

conversion, which was observed in the previous studies

on an ensemble of CdSe/ZnSe QD26. Such a conversion

would lead to a non-zero spin-polarization degree of exci-
tons injected to a QD under linearly polarized excitation.

In such a scenario, the intensities of X emission lines un-

der this excitation would not be solely related to the X

spin-flip process. As a consequence, its rate would be im-

possibly determined, which would finally result in a dif-

ference between $P_+ \neq P_-$ degrees obtained for both cir-
cular polarizations of excitation from Eq. (6). However,

almost equal values of both degrees determined experi-

mentally reveals unimportance of the linear-to-circular

polarization conversion in case of the studied QDs.

B. Trion and biexciton

In order to provide a comprehensive description of po-

larization properties of CdSe/ZnSe QD PL under optical

excitation at 488 nm, we briefly analyze the spectra of

two remaining excitonic complexes: biexciton and nega-

tively charged trion. In particular, we focus on the mea-

surements carried out at magnetic field of 4 T (Faraday

Taking into account the determined X spin-flip rate
dependence on the magnetic field we can extract the ex-

citon spin-transfer efficiency under circularly polarized

excitation. A quantitative measure of this efficiency is

degree to which the excitons are injected to one of X

eigenstates in the QD. Using Eq. (4) it can be expressed

as

$$P_\pm = \frac{\alpha_\pm - \alpha_+}{\alpha_+ + \alpha_-} = \pm \left( \frac{1 + \frac{x}{\gamma}}{1 + \frac{x}{\gamma}} - 1 \right),$$

where $P_+$ ($P_-$) corresponds to the excitation with $\sigma^+$ ($\sigma^-$) circular polarization. It should be noted that at

sufficiently large magnetic field (i.e., when nearly pure

spin states are eigenstates of X) the above-introduced $P_\pm$

accounts directly for spin-polarization degree of excitons

injected to the dot. Moreover, for such fields $P_\pm$ would

be equal to the degree of X PL circular polarization in the case of an absence of X spin-flip. On the contrary,

at lower magnetic fields ($B \lesssim 2$ T for the studied QD)

the interpretation of $P_\pm$ is not as straightforward since

X eigenstates are partially linearly polarized.

The evolution of the typical measured $P_\pm$ degree in

the magnetic field is shown in Fig. 4(c). The exciton

spin-transfer efficiency is found to be similar for both

circular polarizations of excitation. At $B > 3$ T we ob-

serve high spin-polarization degree of injected excitons,

which is almost constant and exceeds 40%. This result

unequivocally confirms the pronounced conservation of

exciton spin-polarization under excitation of the studied

CdSe/ZnSe QD at 488 nm. Similar effect is observed

for various quantum dots independently of the excitonic

emission energy. On the other hand, at $B < 3$ T the $P_\pm$

degree drops due to raising influence of the anisotropy-

related excitonic fine structure, which prevents precise
determination of the spin-transfer efficiency at lower

fields. Interestingly, the reduction of excitation wave-

length from 488 nm to 476.5 nm ($\approx 2.61$ eV) results in

an almost complete loss of the exciton spin-transfer in the

entire range of the magnetic field (Fig. 4(d)).

I- / I+ = 1 + 2\tau_X / \tau_f$, allowing a direct determination of

relative neutral exciton spin-flip rate. Fig. 4(b) presents

the measured $I_- / I_+$ ratio as a function of the magnetic

field. In accordance to the above considerations we ob-

serve a clear, superlinear increase of the relative intensity of

lower-energy X emission line resulting from the short-
eening of X spin-flip time at higher fields. Detailed analy-
sis of the obtained results reveals the power dependence

of the spin-flip rate $1 / \tau_f$ on the energy splitting $\Delta E(B)$
of neutral exciton states in the dot. In particular, the

curve given by $I_- / I_+ = 1 + a \Delta E^2$ perfectly reproduces

the experimental data from Fig. 4(b) for $\gamma = 2.95 \pm 0.05$.

This demonstrates that within our experimental preci-

sion, the X spin-flip rate $1 / \tau_f$ is proportional to the third

power of X energy splitting, as previously theoretically

predicted for a single phonon-mediated X spin relaxation

in a QD.39
configuration). The corresponding 2X and X− PL spectra obtained with polarization-resolution under circularly or linearly polarized excitation are shown in Fig. 5.

In case of the biexciton (Figs. 5a−5c), the PL intensities are not affected by switching the polarization of excitation. It is related to the fact that 2X consists of two electron-hole pairs forming closed shells. As such, it cannot carry any spin information and recombines to one out of two Zeeman-split neutral exciton states with the same probabilities. As a result, we observe equal intensities of both 2X emission lines independently of the polarization of excitation.

The physical picture in case of the trion is more complex (Figs. 5d−5f). Under linearly polarized excitation providing no spin-transfer of injected excitons, the lower-energy emission line is slightly more intense (Fig. 5d). Similarly to the case of the neutral exciton, this effect is interpreted as originating from spin relaxation taking place during the trion lifetime in the QD. However, such relaxation has qualitatively different character compared to the neutral exciton spin-flip process analyzed in the previous section. In particular, it is directly related to a spin-flip of a single hole between its Zeeman-split states, as the remaining two electrons forming negatively charged trion are accommodated in a closed shell and cannot participate in the spin relaxation. The presence of the hole spin-flip affects also X− PL spectra measured under circularly polarized excitation increasing the intensity of the lower-energy line (Figs. 5e,5f). Despite this effect, in such a case a pronounced optically-induced spin-transfer is observed. But, in contrast to the neutral exciton, the spin transfer has negative sign since the cross-polarized X− emission is more intense compared to the co-polarized emission. The apparent negative optical orientation for X− was previously observed in different QD systems, both III-V and II-VI. Following Ref. 40, we interpret it as arising due to possible spin mismatch between an electron residing in the dot and the one forming the spin-polarized injected exciton. If both electrons have the same spin, the Pauli blockade prohibits complete energy relaxation of the injected exciton. In such a case, spin-conserving relaxation processes stop at the level of hot trion with one electron in the ground state and the other occupying the first excited state. The hot trion state is typically characterized by relatively strong anisotropic part of the electron-hole exchange interaction, which accompanied by the Fröhlich interaction between electron pair and LO phonon leads to simultaneous spin-flip of the electron and the hole (flip-flop). As a result, the spin of injected exciton is effectively reversed, which finally underlies the negative optical spin-transfer determined experimentally for X−. On the other hand, no such effect has been observed for positively charged trion due to much lower efficiency of the electron-hole flip-flop process in this case. Thus, our findings confirm the negative charge state of the only trion which emission line is visible in the PL spectra of the studied QDs.

V. SPIN-TRANSFER FOR MN-DOPED CDSE/ZNSE QD

The efficient injection of spin-polarized excitons is also demonstrated for CdSe/ZnSe QDs containing single Mn2+ ions. The typical PL spectrum of an example Mn-doped dot is shown in Fig. 6a. Similarly to the case of a nonmagnetic QD (see Fig. 2a), we observe emission lines originating from the recombination of X, X−, and 2X. However, each of them exhibits multiplet splitting related to the presence of s, p-d exchange interaction between confined carriers and Mn2+ ion (total spin $S = 5/2$). Due to heavy hole character of the hole ground state in a QD, the hole-Mn exchange is effectively Ising-like. As a result, the neutral exciton emission line is split into six components as observed in Fig. 6a. Each of them is twofold degenerated and related to a particular spin projection of the X-Mn system on the QD growth axis $z$ (which is the quantization axis of the hole). More specifically, the consecutive energy levels of the neutral exciton with spin projection $J_z = \pm 1$ (coupled to $\sigma^\pm$ polarized light) correspond to subsequent Mn2+ spin projections $S_z$ equal to $\pm 5/2, \mp 1/2, \mp 3/2$, and $\mp 5/2$ (as schematically depicted in Fig. 6b). The same underlying exchange splitting of the neutral exciton affects also the biexciton emission spectrum, since this excitonic complex is a spin-singlet and does not interact with the ion. As a consequence, the 2X state is degenerate (Fig. 6b) and its emission line is split only due to the final state of the recombination process.
exchange interaction with the Mn fine-structure on the neutral exciton states, split by the magnetic field. The states are displayed as a function of their energy and Mn$^{2+}$ ion spin projection on the QD growth axis. Color of the neutral exciton levels denotes the spin projection on the growth axis, as well as the circular polarization of corresponding X optical transitions (red $- \sigma^+$, blue $- \sigma^-$). For clarity, only the transitions related to 3/2 spin projection of the Mn$^{2+}$ ion are shown.

(i.e., the neutral exciton state). Finally, the biexciton PL exhibits similar sixfold splitting as the neutral exciton, which is clearly visible in Fig. 6(a).

The efficiency of exciton spin-transfer under excitation at 488 nm of the Mn-doped QD is determined in a QD under below-the-barrier excitation. The states are displayed as a function of their energy and Mn$^{2+}$ ion spin projection on the QD growth axis. Color of the neutral exciton levels denotes the spin projection on the growth axis, as well as the circular polarization of corresponding X optical transitions (red $- \sigma^+$, blue $- \sigma^-$). For clarity, only the transitions related to 3/2 spin projection of the Mn$^{2+}$ ion are shown.

FIG. 6: (Color online) (a) A PL spectrum of a CdSe/ZnSe QD containing a single Mn$^{2+}$ ion. (b) Simplified diagram of neutral exciton and biexciton energy levels in Mn-doped QD at zero magnetic field. The states are displayed as a function of their energy and Mn$^{2+}$ ion spin projection on the QD growth axis. Color of the neutral exciton levels denotes the spin projection on the growth axis, as well as the circular polarization of corresponding X optical transitions (red $- \sigma^+$, blue $- \sigma^-$). For clarity, only the transitions related to 3/2 spin projection of the Mn$^{2+}$ ion are shown.

The average value of X PL circular polarization degree under circularly polarized excitation is found to be approximately field-independent and equal to about 40% for all studied Mn-doped QDs. However, this value is not necessarily identical to exciton spin-transfer efficiency. More specifically, the quantitative determination of this efficiency requires taking into account possible spin relaxation of X-Mn system occurring during the X lifetime in a QD. Such relaxation may concern both the exciton spin-flip between two states corresponding to the same projection of the ion spin as well as the processes involving simultaneous spin-flip of the X and the Mn$^{2+}$ ion. Their presence is revealed by zero-field X PL spectra.
measured under linearly polarized excitation (Fig. 7(c)). In such a case, no spin polarization of the Mn$^{2+}$ ion is expected, which should manifest itself in equal intensities of six emission lines corresponding to different projections of the ion spin. On the contrary, the intensity of the low-energy emission line is significantly larger compared to the higher-energy lines, independently of the circular polarization of detection. Since the low-energy line corresponds to $±5/2$ ion spin projection in $σ^+$ polarization of detection, this effect cannot be attributed to a steady state polarization of the ion spin. On the other hand, it may originate from X spin-flip process causing the exciton relaxation from $|J_z = ±1, S_z = ±5/2⟩$ state towards $|J_z = ±1, S_z = ±5/2⟩$. As this process does not affect the ion spin, its underlying mechanism is identical as the one entailing exciton spin-flip in a nonmagnetic QD between Zeeman-split states. The latter process is found to be quite efficient for a large energy splitting of the invoked states. Bearing in mind that energy difference between $|J_z = ±1, S_z = ±5/2⟩$ and $|J_z = ±1, S_z = ±5/2⟩$ states corresponds to energy splitting of two extreme X PL lines (equal to about 1.2 meV in the case of the studied QD), the proposed spin-relaxation can play a significant role in Mn-doped QD. In particular, it increases the intensity of the low-energy X emission line. On the other hand, its presence would also lead to a decrease of the high-energy line intensity. However, the experimental results (Fig. 7(c)) do not reflect this prediction. Thus, we expect that other X-Mn spin relaxation processes have to be important. They might be related to a mutual flip of exciton and ion spins or a spin-flip of a single carrier forming the neutral exciton resulting in a creation of a dark exciton. The precise identification and detailed description of these relaxation processes remain beyond of the scope of this paper and require further studies.

The application of the magnetic field modifies the energies of the X-Mn states (presented in Fig. 3(b)) introducing both exciton and ion Zeeman splittings. As a consequence, the symmetry between energy levels corresponding to $J_z = ±1$ excitons is no longer present and X-Mn spin relaxation entails a difference of X PL intensities measured in two circular polarizations of detection under linearly polarized excitation (Figs. 7(f), 7(i)). As the relative energy of $J_z = ±1$ excitons is decreased, the spin-flip processes increase the intensity of the corresponding $σ^−$ polarized X PL. It is clearly visible in Fig. 7(j) presenting the ratio of integrated X PL intensities detected in $σ^−$ and $σ^+$ polarizations under linearly polarized excitation. The determined ratio is however lower compared to the nonmagnetic QD for the same magnetic field (see Fig. 4(b)). The possible explanation of this effect is related to a similar strength of the X-Mn exchange interaction and the excitonic Zeeman effect at high magnetic field. In particular, even at $B = 10$ T the Zeeman effect is too weak to invert the zero-field order of the exchange-split excitonic states corresponding to $±5/2$ ion spin projection. As a result, some of $|J_z = ±1, S_z⟩$ states have lower energy than $|J_z = +1, S_z⟩$ even at high field. It finally leads to a suppression of the overall spin relaxation between $J_z = ±1$ excitons when compared to a nonmagnetic QD. Nevertheless, the influence of the spin-flip is also visible in the X PL spectra measured under circularly polarized excitation. In such case, the spin relaxation results in an increase (decrease) of the circular polarization degree of integrated X PL under excitation with $σ^−$ ($σ^+$) polarization at high magnetic field (Fig. 7(k)). Therefore, the obtained degrees do not exactly correspond to the spin-polarization degrees of excitons injected to the dot. In principle, the latter might be extracted from a detailed analysis of X PL spectra measured under different polarizations of excitation. However, a complicated spin relaxation of the X-Mn system prevents us to perform such an analysis in a similar way as it is carried out in the case of nonmagnetic QD (described in section IV). We only note that the average degree of circular polarization of X PL under $σ^−$ and $σ^+$ spin-polarized excitation of about 40% defines the lower limit of the exciton spin-transfer efficiency in the entire range of the applied magnetic field.

VI. OPTICAL SPIN ORIENTATION OF MN$^{2+}$ ION IN A CdSe/ZnSe QD

Exploiting the ability to inject the spin-polarized excitons to Mn-doped CdSe/ZnSe QD we study their influence on the Mn$^{2+}$ ion spin state. Such an influence so far has been demonstrated for CdTe/ZnTe QDs, in which the single Mn$^{2+}$ ion spin could be oriented towards $±5/2$ state via its exchange interaction with $J_z = ±1$ excitons. In order to examine the presence of this phenomenon in the case of CdSe QDs, we first need to establish a method enabling unambiguous optical readout of the ion spin state. Following the previous studies of CdTe QDs one may think about determining the ion spin basing on the intensities of six emission lines of the neutral exciton detected in a given circular polarization. However, their values provide only information about the probabilities of finding the X-Mn system in a particular state at the moment of exciton recombination. If no spin relaxation is occurring during the X lifetime in a QD, those probabilities are indeed equivalent to the probabilities for the Mn$^{2+}$ ion to have one of its six possible spin projections in the absence of the exciton. On the contrary, the efficient spin-flip processes of X-Mn system present in the studied CdSe QDs make this approach no longer valid. For this reason we monitor the ion spin basing on the intensities of the biexciton emission lines. Since the 2X is a spin-singlet, it is decoupled from the ion and acts as an almost ideal probe of its spin state. Such an influence to a configuration mixing might be neglected here. As a consequence, the intensities of the 2X emission lines reflect the probabilities of finding the Mn$^{2+}$ ion in a spin state defined by the line energy and polarization of detection.
hand, excitation with circularly polarized light results in a direct fingerprint of depolarized ion spin. On the other hand, for sufficiently low $2X$ formation probability the higher-energy lines are almost equal (Fig. 8(a)), which is the case of linearly polarized excitation, the intensities of all $2X$ emission lines are almost equal. This observation unequivocally confirms the presence of optically-induced Mn$^{2+}$ ion spin polarization in a CdSe/ZnSe QD. Since $\sigma^-$ polarized $2X$ emission lines of subsequent energies correspond to Mn$^{2+}$ spin projections ranging from $5/2$ to $-5/2$, our findings indicate that injection of $J_z = \pm 1$ excitons orients the ion spin towards $\pm 5/2$ state, similarly to the case of Mn-doped CdTe QDs\cite{Za97, Za98}. The Mn$^{2+}$ spin can be also independently probed in $\sigma^+$ polarization of detection (Figs. 8(d)-8(f)). In such case, the $2X$ emission energies correspond to subsequent ion spin projections in a reversed order, namely ranging from $-5/2$ to $5/2$. As the ion spin state depends only on the spin of injected excitons, we observe opposite change of $2X$ emission lines intensities under the same circular polarization of excitation compared to the previously used $\sigma^-$ polarization of detection, which is clearly seen in Figs. 8(c)-8(f).

In order to perform a detailed analysis of the optical spin orientation efficiency, we introduce its quantitative measure defined as a difference between mean values of the ion spin projection obtained under $\sigma^-$ and $\sigma^+$ polarized excitation. The Mn$^{2+}$ mean spin ($S_z$) is directly determined from a biexciton PL spectrum detected in a circular polarization as a weighted average of the intensities of consecutive emission lines. The dependences of the above-introduced efficiency on the magnetic field (applied in Faraday configuration), and excitation power at $B = 4$ T are presented in Figs. 8(g) and 8(h), respectively. Since the mean spin of the Mn$^{2+}$ ion can be independently obtained basing on the spectra detected in two circular polarizations, each of the Figs. 8(g), 8(h) contains two sets of points corresponding to $\sigma^+$ and $\sigma^-$ polarized detections. Their mutual agreement confirms the self-consistency of our experimental results.

Excitation power dependence of the ion spin orientation efficiency exhibits a pronounced decrease at high-power regime (Fig. 8(h)). Similarly to the case of CdTe QDs\cite{Za97, Za98}, it is related to the increased probability of the biexciton formation. More specifically, the $2X$ recombination leaves a randomly polarized neutral exciton in the dot. As a consequence, more frequent creation of the biexciton lowers the efficiency of the optically-induced exciton spin-transfer, which finally leads to a smaller efficiency of the ion spin orientation. On the other hand, for sufficiently low $2X$ formation probability assured by the excitation power corresponding to $< 5\%$ of the saturation power, the spin orientation efficiency is almost power-independent. However, under these conditions it clearly increases with the magnetic field (Fig. 8(g)). In the low-field regime such effect can be attributed to a suppression of the mixing between Mn$^{2+}$ spin eigenstates introduced by a hyperfine interaction with nuclear spin. This mixing was previously shown to significantly increase the ion spin relaxation dynamics in the absence of magnetic field\cite{Za97, Za98, Za99}, which finally de-

Having established the procedure of Mn$^{2+}$ spin read-out in a CdSe QD, we verify the possibility of the ion spin manipulation through injection of spin-polarized excitons to the dot. To do so, we measure the biexciton PL spectra under circularly or linearly polarized excitation in a fixed circular polarization of detection. This ensures that each $2X$ emission line corresponds to a specified spin projection of the Mn$^{2+}$ ion. The zero-field spectra detected in $\sigma^-$ polarization are shown in Figs. 8(a)\cite{Za97, Za98} and 8(c). In the case of linearly polarized excitation, the intensities of all $2X$ emission lines are almost equal (Fig. 8(a)), which is a direct fingerprint of depolarized ion spin. On the other hand, excitation with circularly polarized light results in a small, but negligible non-uniform distribution of the intensities between six lines. In particular, under $\sigma^-$ polarized excitation the lower-energy lines are more intense, while the higher-energy lines less (Fig. 8(b)). Switching the polarization of excitation to $\sigma^+$ leads to opposite effect, i.e., stronger higher-energy lines and weaker lower-energy lines (Fig. 8(c)). This observation unequivocally confirms the presence of optically-induced Mn$^{2+}$ ion spin polarization in a CdSe/ZnSe QD. Since $\sigma^-$ polarized $2X$ emission lines of subsequent energies correspond to Mn$^{2+}$ spin projections ranging from $5/2$ to $-5/2$, our findings indicate that injection of $J_z = \pm 1$ excitons orients the ion spin towards $\pm 5/2$ state, similarly to the case of Mn-doped CdTe QDs\cite{Za97, Za98}. The Mn$^{2+}$ spin can be also independently probed in $\sigma^+$ polarization of detection (Figs. 8(d)-8(f)). In such case, the $2X$ emission energies correspond to subsequent ion spin projections in a reversed order, namely ranging from $-5/2$ to $5/2$. As the ion spin state depends only on the spin of injected excitons, we observe opposite change of $2X$ emission lines intensities under the same circular polarization of excitation compared to the previously used $\sigma^-$ polarization of detection, which is clearly seen in Figs. 8(c)-8(f).

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Excitation power dependence of the ion spin orientation efficiency exhibits a pronounced decrease at high-power regime (Fig. 8(h)). Similarly to the case of CdTe QDs\cite{Za97, Za98}, it is related to the increased probability of the biexciton formation. More specifically, the $2X$ recombination leaves a randomly polarized neutral exciton in the dot. As a consequence, more frequent creation of the biexciton in the QD lowers the efficiency of the optically-induced exciton spin-transfer, which finally leads to a smaller efficiency of the ion spin orientation. On the other hand, for sufficiently low $2X$ formation probability assured by the excitation power corresponding to $< 5\%$ of the saturation power, the spin orientation efficiency is almost power-independent. However, under these conditions it clearly increases with the magnetic field (Fig. 8(g)). In the low-field regime such effect can be attributed to a suppression of the mixing between Mn$^{2+}$ spin eigenstates introduced by a hyperfine interaction with nuclear spin. This mixing was previously shown to significantly increase the ion spin relaxation dynamics in the absence of magnetic field\cite{Za97, Za98, Za99}, which finally de-

![FIG. 8: (Color online) (a)-(f) The zero-field PL spectra of biexciton confined in Mn-doped QD. The spectra were excited at 488 nm with indicated circular and linear polarizations and detected in $\sigma^-$ (blue, (a)-(c)) and $\sigma^+$ (red, (d)-(f)) polarizations. The ion spin projections corresponding to the emission lines of extreme energies are indicated. Solid lines representing the direction of change of $2X$ intensities are drawn to guide the eye. (g),(h) The efficiency of optically-induced Mn$^{2+}$ spin orientation measured as a function of: magnetic field (g) and excitation power at $B = 4$ T (h). Circles (squares) represent the efficiencies determined in $\sigma^-$ ($\sigma^+$) polarization of detection. The excitation power is expressed in $P_0$ units, where $P_0$ is a power used to obtain the data presented in (g). The saturation power of this QD corresponds to 30$P_0$.](image)
creases the zero-field efficiency of optical pumping of the ion spin. However, the hyperfine coupling is relatively small\textsuperscript{53} and even a field of the order of few tenths of Tesla is sufficient to almost completely purify the ion spin eigenstates. As a result, it cannot be the underlying reason of the pronounced enhancement of spin orientation efficiency at $B > 1$ T. Taking into account almost constant exciton spin-transfer efficiency in the entire range of the applied field, we interpret this enhancement as reflecting the field-dependence of the intrinsic mechanism leading to the ion spin orientation. Surprisingly, no theoretical model providing such mechanism has been proposed so far, despite the intuitive character of the optical orientation process. In particular, only the optical pumping of the ion spin under strictly resonant excitation\textsuperscript{19,20} was quantitatively understood\textsuperscript{48,54}. The deeper understanding of the ion spin orientation mediated by the injection of spin-polarized excitons would presumably also allow to explain somehow low efficiency of this process in the case of the studied CdSe/ZnSe QDs. In particular, the obtained efficiency at $B = 1$ T is few times lower compared to CdTe QDs\textsuperscript{52}, for even higher efficiency of the exciton spin-transfer in the present case. This observation together with indication of several differences between the two types of dots, including much more efficient exciton spin relaxation for CdSe QDs, may in turn shed some light on the mechanism of the ion spin orientation and help in its further investigation.

VII. CONCLUSION

We have presented a simple all-optical method enabling effective injection of spin-polarized excitons to self-assembled CdSe/ZnSe QDs. Our studies concerned both nonmagnetic dots and those containing single Mn\textsuperscript{2+} ions. Experimentally, the pronounced exciton spin-transfer was observed under below-the-barrier excitation at 488 nm with circularly polarized light. The spin-transfer efficiency was studied in polarization-resolved measurements of a single dot PL performed in external magnetic field applied along the sample growth axis. Experimental results indicated importance of phonon-mediated neutral exciton spin relaxation occurring during the exciton lifetime in a QD. In case of nonmagnetic dots, such relaxation was simply related to X spin-flip between its Zeeman-split states leading to an increased intensity of X emission corresponding to a lower-energy state. On the other hand, the spin relaxation for Mn-doped QDs was exhibiting more complicated character. In particular, our findings revealed that it is not entirely related to X spin-flip occurring without the change of the ion spin. However, basing on the time-integrated spectroscopy we were not able to provide its detailed description.

The straightforward character of X spin relaxation in the case of nonmagnetic QD enabled us to analyze its X PL spectra obtained under circularly and linearly polarized excitation within a frame of a simple rate-equation model. On this basis we established a power-dependence of the exciton spin-flip rate on X energy splitting. Moreover, we determined the spin-polarization degree of excitons injected to QD under $\sigma^+$ and $\sigma^-$ polarized excitation to be similar in both cases and close to 40% at magnetic field of 3-10 T. At lower field, the mixing of exciton spin states imposed by the fine structure prevented exact determination of optically-induced spin-transfer efficiency for a nonmagnetic dot. On the other hand, it was possible for Mn-doped one since in this case the fine structure is dominated by the exciton-ion exchange interaction. Consequently, we found the spin-transfer efficiency under excitation at 488 nm to be almost field-independent and equal to about 40% in the entire range of the applied field 0-10 T. We stress out that similarly large efficiency was obtained for several, randomly selected dots exhibiting different energies of excitonic emission. This finding indicates the robustness of the presented spin-conserving excitation channel in case of the studied CdSe/ZnSe QDs.

Importantly, we observed also a negative spin-transfer for the only emission line in a CdSe/ZnSe QD PL spectrum originating from a trion recombination. Following the previous studies of different QD systems, it enabled us to unequivocally identify the negative charge state of this trion.

Finally, taking advantage of the ability to inject the spin-polarized excitons to Mn-doped CdSe/ZnSe QD we demonstrated the possibility of optical orientation of Mn\textsuperscript{2+} spin. The ion spin readout was based on the intensities of six biexciton emission lines measured in a circular polarization of detection. Our findings revealed that under $\sigma^\pm$ polarized excitation the Mn\textsuperscript{2+} is oriented towards $\pm 5/2$ state. The efficiency of the optical orientation was found to increase with the magnetic field. Simultaneously, it was a few times lower compared to the previously studied Mn-doped CdTe QDs. The origin of both effects is not entirely clear, since the intrinsic mechanism underlying the ion spin orientation remains not fully understood. Nevertheless, the possibility of the optical control over the Mn\textsuperscript{2+} spin embedded in a CdSe QD evidenced in our work may be further exploited to determine the ion spin relaxation dynamics in a low-field regime or even in the absence of magnetic field.

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