Doped Semiconductor-Nanocrystal Emitters with Optimal Photoluminescence Decay Dynamics in Microsecond to Millisecond Range: Synthesis and Applications

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ABSTRACT: Transition metal doped semiconductor nanocrystals (d-dots) possess fundamentally different emission properties upon photo- or electro-excitation, which render them as unique emitters for special applications. However, in comparison with intrinsic semiconductor nanocrystals, the potential of d-dots has been barely realized, because many of their unique emission properties mostly rely on precise control of their photoluminescence (PL) decay dynamics. Results in this work revealed that it would be possible to obtain bright d-dots with nearly single-exponential PL decay dynamics. By tuning the number of Mn²⁺ ions per dot from ~500 to 20 in Mn²⁺ doped ZnSe nanocrystals (Mn:ZnSe d-dots), the single-exponential PL decay lifetime was continuously tuned from ~50 to 1000 µs. A synthetic scheme was further developed for uniform and epitaxial growth of thick ZnS shell, ~7 monolayers. The resulting Mn:ZnSe/ZnS core/shell d-dots were found to be essential for necessary environmental durability of the PL properties, both steady-state and transient ones, for the d-dot emitters. These characteristics combined with high PL quantum yields (70 ± 5%) enabled greatly simplified schemes for various applications of PL lifetime multiplexing using Mn:ZnSe/ZnS core/shell d-dots.

Colloidal semiconductor nanocrystals have been pursued as unique emitting materials in various applications because of their broad yet intensive absorption, high emitting brightness, and size-dependent emission colors. Emission from semiconductor nanocrystals can be classified into two main classes, namely, exciton emission and dopant emission. Exciton is an electron–hole pair generated by photo- or electro-excitation and bonded together through Columba interaction, which delocalizes in the entire nanocrystal in the quantum confinement size regime.1 Recombination of exciton in a semiconductor nanocrystal releases a photon with its energy near the absorption band edge, which is thus also known as either band-edge emission or intrinsic emission. Dopant emission is usually associated with the d-shell orbitals of intentionally doped ions in the lattice of a host semiconductor nanocrystal (d-dot).2-4 Such as the 3d orbitals of Mn²⁺ ions in Mn²⁺ doped ZnSe nanocrystals (Mn:ZnSe d-dots). Different emission mechanisms render these two classes of emitters with complementary luminescence properties. Unlike significant overlapping between emission and absorption in exciton emitters, dopant emission of a d-dot is always noticeably lower in energy than the absorption edge of the host nanocrystal.4,5 Such a large Stokes shift prevents energy transfer and reabsorption for applications using high concentrations of nanocrystal emitters. Different from delocalization of exciton, dopant emission is associated with one atom, implying its low sensitivity to challenging environments.3,6 Opposite to optically allowed exciton emission, dopant emission usually involves optically forbidden transitions of d orbitals of transition metal.7,8 As a result, the typical photoluminescence (PL) decay lifetime for exciton emission is on the order of several tens of nanoseconds,3,6 which is beneficial for applications requiring “nanophosphores”. Conversely, the PL decay lifetime of the dopant emission is in the range between microseconds and milliseconds,7,10-13 which fits those applications requiring “nanophosphors”. Finally, transition metal dopants are often magnetically active, which offers unique magneto-optical properties to d-dots.11-15

The potential of exciton emitters—mostly based on CdSe and their core/shell nanocrystals—have been developed to a level ready for certain real-life applications. Though research of d-dots remains in academic exploration at present, their unique emission properties have been actively studied for a variety of applications. D-dots with pure dopant emission were achieved and reabsorption for applications using high concentrations of

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with decent quantum yield about ten years ago, and color-tunable emission of d-dots was reported by several research groups as non-Cd alternative emitters of CdSe nanocrystals. Taking the advantages of long PL decay lifetime of dopant emission at low energy, various groups reported nanothermometers based on temperature-dependent dual emission of d-dots. The long PL lifetime of d-dot emitters also enabled time-resolved fluorometry to remove biological autoluminescence that possesses a typical PL decay lifetime in a few nanoseconds.

If one could obtain a series of d-dots with single-exponential PL decay dynamics in their typical lifetime window, in principle, it should be possible to realize PL lifetime multiplexing. By labeling each target with an emitter with fixed and single-exponential PL lifetime, PL lifetime multiplexing could potentially read out multiple targets by single excitation. PL lifetime multiplexing possesses numerous advantages over PL color multiplexing and offers a new dimension—time—for PL multiplexing. While time-resolved fluorometry and time-gated imaging only require the labels emitting in a significantly longer time window than the background luminescence, PL lifetime multiplexing is substantially more stringent in terms of PL decay dynamics of the emitters. This is so because PL lifetime multiplexing requires full deconvolution of all emissive components in the system, each of which should possess single-exponential PL decay dynamics. Exciton emission of intrinsic semiconductor nanocrystals was demonstrated as a possible component for PL color multiplexing ~15 years ago, but their nanosecond and environmentally sensitive PL lifetime would prevent them from being ideal for PL lifetime multiplexing. Conversely, the microsecond lifetime of d-dots would potentially overcome these problems.

However, to realize their unique applications, such as PL lifetime multiplexing and magneto-optical applications, there are several challenges facing d-dot emitters. For the most developed Mn:ZnSe d-dots, though several groups reported variation of PL lifetime of the dopant emission—the spin forbidden 4T1 to 6T1 transition within the d-orbitals of a Mn2+ ion, the tunable window of the lifetime was insufficient and the PL decay dynamics was usually multi-exponential. The PL decay lifetime of the 4T1 to 4T1 transitions of Mn2+ ions in bulk InP—VI semiconductor crystals has been known to depend on the dopant concentration, which was considered to be a result of Mn—Mn coupling within the host. Recently, Gamelin’s group found that it was possible to obtain single-exponential PL decay (up to ~2 orders of magnitude of intensity) of the Mn2+ dopant emission with very low Mn2+ concentration (~0.1%, about 1 Mn2+ ion per dot) though an additional fast component became evident upon increase of Mn concentration in the host ZnSe nanocrystals. The authors believed that such multiexponential PL decay dynamics at increased Mn concentration was due to inhomogeneous spatial distribution of Mn2+ ions in each ZnSe nanocrystal. Similarly, with a low Mn2+ concentration (~6 Mn2+ ions per dot), Cao’s group obtained Mn:CsS/ZnS d-dots with single-exponential PL decay. The single-exponential PL decay dynamics should be associated with monodisperse chemical environment of the Mn2+ ions within a sample, including Mn—Mn distance, chemical coordination to Mn2+ ions, surface exposure of Mn2+ ions, uniform concentration among all d-dots in a sample, etc. Although thermal stability of Mn:ZnSe d-dots was well documented, their chemical and photochemical stability was questionable. Similar to the intrinsic semiconductor nano-

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out that, unlike working in the strong quantum confinement regime of the host semiconductor, the sizes of the d-dots were in weak quantum confinement regime of ZnSe. Thus, the influence of dopant concentrations on absorption spectra should be small. PL spectra of these d-dots showed pure dopant emission at ~583 nm, which was confirmed as the $^{4}T_{1} \rightarrow ^{6}A_{1}$ transition of the Mn$^{2+}$ ions in ZnSe nanocrystals. As long as Mn$^{2+}$ ions per dot was higher than ~20 in average, no sign of exciton emission from undoped ZnSe nanocrystals was observed in this new synthetic scheme. The number of Mn$^{2+}$ ions per nanocrystal determined by EDX and TEM in Figure 1, i.e., ~30 to 450 Mn$^{2+}$ ions per nanocrystal of the Mn:ZnSe d-dots, corresponded ~2.5% to 36% of the cations in each initial Mn:ZnSe d-dot (~5 nm in size) for the five representative samples. Both the number of Mn$^{2+}$ ions per nanocrystal and the percentage of cations were quite high in comparison to what has been reported in the literature.

Different from absorption and steady-state PL spectra, the PL decay dynamics of five representative samples were different from each other (Figure 1b). The PL decay curves of all representative Mn:ZnSe d-dots were nearly single-exponential within 3–4 orders of magnitude of intensity. Moreover, by varying the Mn$^{2+}$ ions per nanocrystal from about 500 to 20, the single-exponential PL decay lifetime could be varied from ~50 to 1000 μs (Figure 1c). Reproducibility of the Mn$^{2+}$ ions concentration dependence of the single-exponential PL decay lifetime was rather good (Figure 1c).

The highly tunable and nearly single-exponential PL decay dynamics of the dopant emission in Figure 1 implies several structure features, if we assumed the PL decay dynamics depending on the spin coupling of Mn–Mn ions in a nanocrystal. First, the Mn–Mn distance in d-dots decreased as the number of Mn$^{2+}$ ions per nanocrystal increased. Second, the Mn$^{2+}$ ions in each nanocrystal should be close to homogeneous distribution and should not form MnSe clusters within the d-dots. To further confirm the second point, control experiments revealed that, if MnSe cluster(s) existed within a d-dot, a very short component of PL decay lifetime coupled with a complicated PL decay would be the case (Figure S3). Third, the chemical environments of Mn$^{2+}$ ions in all Mn:ZnSe d-dots in a given sample should be very similar to each other, and there should be no surface exposure of the Mn$^{2+}$ ions. Detailed structure characterization is ongoing to confirm these conclusions.

Mn:ZnSe d-dots were found to be photochemically not sufficiently stable (Figure 2a). Importantly, along with decrease of PL quantum yield shown in Figure 2a (inset), continuous illumination also caused appearance of a noticeable short-lifetime component in the PL decay dynamics (Figure 2a).
Presumably, the new short-lifetime component should be associated with the inefficient surface states created by photobleaching. A similar phenomenon was also reported in the literature, that surface states could influence the PL decay dynamics of the dopant emission of Mn:ZnSe d-dots.37

Epitaxial growth of relatively thick ZnS shell solved the photochemical stability problem. Figure 2a (inset) shows that, under the same conditions, the resulting Mn:ZnSe/ZnS core/shell d-dots were practically resistant to photochemical bleaching. Upon growth of the ZnS shell, the absorption spectrum revealed an apparent increase in the ZnS optical window without noticeable red shift of the absorption edge (Figure 2b). TEM measurements (Figure 2c) revealed that epitaxial growth of the ZnS shell increased the size of the nanocrystal significantly—from 6.5 to 11 nm—without worsening the size distribution. High resolution TEM images (Figure S4) revealed that the resulting core/shell d-dots were single crystalline in nature. X-ray powder diffraction measurements (Figure 2d) indicated that growth of the ZnS shell did not cause a new phase. Instead, the entire pattern shifted from that for ZnSe nanocrystals in standard zinc-blende structure to a pattern similar to that for pure ZnS nanocrystals with the same crystal structure. Given the large volume fraction of ZnS, this shift—corresponding to a lattice compression for ZnSe host—was found to be reasonable. Compression of ZnSe host lattice was consistent with the significant red shift of the dopant PL in Figure 2b.38 PL quantum yield of the Mn:ZnSe/ZnS core/shell d-dots was reproducibly ~70 ± 5% for all dopant concentrations (Figure S5), which was similar to that of the plain core Mn:ZnSe d-dots.

Figure 2e shows that, upon growth of the ZnS shell, PL decay dynamics remained nearly single-exponential with high-count measurements (3–4 orders of magnitude of intensity). The PL decay lifetime decreased from 137 to 114 μs for the specific sample in Figure 2e. For Mn:ZnSe d-dots with different amounts of Mn²⁺ ions, their PL decay dynamics changed in a similar manner after epitaxial growth of the ZnS shell. This was found to be consistent with the compression of ZnSe lattice upon ZnS growth (Figure 2d), which shortened the Mn–Mn distance in a ZnSe host nanocrystal.38 According to the literature,7,39 the PL decay lifetime of Mn²⁺ ions in ZnSe lattice should be about 1 order of magnitude longer than that in ZnSe lattice. Furthermore, literature results revealed that the PL decay lifetime of Mn²⁺ ions in the alloy of ZnSe and ZnS increased steadily as the increase of ZnS content.39 We thus considered that the systematic decrease of the single-exponential lifetime upon ZnS shell growth (see Figure 2e as an example) strongly supported no significant diffusion of Mn²⁺ ions into the ZnS shell and no noticeable alloy formation during the shell growth.

It should be pointed out that epitaxial growth of ZnS shell onto ZnSe (or Mn:ZnSe) nanocrystals with needed thickness for protection of ZnSe host nanocrystals is not well developed, especially for large size core nanocrystals used in this work. Insufficient growth of the shell, formation of pure ZnS nanocrystals, significant blue shift of the absorption spectrum due to alloy, and/or shape irregularity of the resulting core/shell nanocrystals were commonly encountered.28,40–42

Synthesis of the Mn:ZnSe/ZnS core/shell doped nanocrystals (Mn:ZnSe/ZnS d-dots) was realized through a modified SILAR (successive ion layer adsorption reaction) epitaxial scheme using elemental S and Zn stearate as the precursors.43 We varied fatty acids with different chain length and chain structure, and no obvious improvement was observed (data not shown). It was found to be critical to add S into the Mn:ZnSe d-dots solution as the initial step. Furthermore, the S to Zn molar ratio added in consecutive steps in one growth cycle must be S rich, usually with the S/Zn ratio being >1.2. If either the initial step was with the Zn precursor or the S to Zn ratio in one growth cycle was close to 1, appearance of pure ZnS nanocrystals and insufficient shell growth would be inevitable (Figure S6). Presumably, excess S on the surface of the nanocrystals would promote the surface passivation from relatively strong carboxylate ligands to the weak fatty amine ones. The fast ligand dynamics of fatty amines44 would subsequently enhance probability of surface epitaxy of ZnS.

The technical strategies described in the above paragraph would make both epitaxial growth and self-nucleation be extremely fast. This would accelerate formation of pure ZnS nanocrystals as well. In addition, when epitaxy was too fast, size distribution of the resulting core/shell nanocrystals would become broadened upon shell growth (Figure S6). To address these problems, a significant amount of free fatty acid (either 0.5–1 mL of oleic acid or an equivalent amount of stearic acid in 6 mL of reaction mixture) was added. Free fatty acid could reduce the reactivity of metal carboxylate to suppress the extremely fast growth and dissolve newly formed tiny ZnS nanocrystals.

With control of all these key parameters, the resulting Mn:ZnSe/ZnS d-dots retained the size and shape distribution of the Mn:ZnSe d-dots with a desirable thickness of the ZnS shell, i.e., ~7 monolayers (Figure 2c).

Photochemical stability of the d-dots in Figure 2a is one obvious requirement for applications of unique emission properties of d-dots. Epitaxial growth of a relatively thick ZnS shell, i.e., ~7 monolayers, was found to be sufficient to boost the stability of the d-dots against other types of challenging environments. Stability of Mn:ZnSe/ZnS d-dots was studied by comparing with that of CdSe/CdS core/shell nanocrystals with 5 monolayers of CdS shell. This specific reference emitted at a similar PL peak position with PL quantum yield of its exciton emission greater than 90% (see details in Figure S7). The PL decay dynamics of intrinsic emission of the CdSe/CdS core/shell nanocrystals was single-exponential.

One targeted application field of d-dots would be PL lifetime multiplexing for biomedical systems. The homogeneity and stability of the PL decay dynamics of the emitters in aqueous solution with different pH are basic requirements for such potential applications, considering the large variation of pH values in biological systems (e.g., lysosome ~5, cytosol 7.0–7.4, human blood 6.8–7.5, etc.).45 Mn:ZnSe/ZnS d-dots were converted to be soluble in aqueous solution by exchanging the original ligands with mercaptopropanoic acid. After the ligand exchange, the PL quantum yield, peak position, and PL decay dynamics all remained unaltered for the d-dots. Even after dispersion of the resulting water-soluble d-dots in buffer solutions with various pH values, their PL properties were identical to those in the organic solvents (Figure 3a). In comparison, ligand exchange of the CdSe/CdS core/shell nanocrystals and dispersion in the buffer solutions with various pH values changed their emission properties noticeably, which was found to be consistent with the literature.46

Both steady-state PL and transient PL of the dopant emission of Mn:ZnSe/ZnS d-dots in solid thin films—necessary for LEDs,47 solid-sate lighting,48 luminescent solar concentrators,49,50 document security,51,52 and information storage53—
were studied. The PL spectrum of Mn:ZnSe/ZnS d-dots in thin solid film was practically identical to that in the original solution (Figure 3c (inset)), and their PL decay dynamics did not show any wavelength dependence in the solid film (Figure 3c). Conversely, for the reference CdSe/CdS core/shell nanocrystals, significant red shift of the PL spectra was evidenced (Figure 3d, inset), and their PL decay dynamics differed from one wavelength to another in the solid film. Moreover, compared to that in solution, apparent sink (rise) features could be observed for the PL decay curves of the CdSe/CdS core/shell nanocrystals in the solid film (Figure 3d). These changes of the PL properties for intrinsic semiconductor nanocrystals should be mainly caused by Förster resonance energy transfer, which is efficient for the excitation emission of undoped nanocrystals at high concentrations and/or in the solid state.54 In comparison, for d-dots, Förster resonance energy transfer is eliminated by the large Stokes shift (see Figure 1a).

High excitation power results in high emission intensity, which would be beneficial for certain applications. PL lifetime multiplexing is in this category, because PL lifetime multiplexing relies on transient emission, instead of steady-state emission used in PL color multiplexing. The PL decay dynamics and lifetime of the undoped host semiconductor nanocrystals—ZnSe/ZnS core/shell nanocrystals—were confirmed to be closely related to excitation power (Figure S8), which is consistent with the appearance of bi- and multie exciton emission under high excitation power reported in the literature.55 In comparison, under the same excitation conditions, the PL decay dynamics of the Mn:ZnSe/ZnS d-dots apparently did not change by increasing the excitation power up to tens of photons absorbed per pulse for each dot (Figure S8). Moreover, local heating is often a side effect of high power excitation. Results revealed that the PL decay dynamics of the dopant emission in d-dots was found to be much more stable than that of the exciton emission of the undoped semiconductor nanocrystals against local heating (Figure S9). High thermal stability was observed previously in solution and was attributed to the inner atomic emission of the dopant PL.56

With a microsecond excitation source, Figure 4a records a three-dimensional plot of the transient PL of a mixture of a common protein (bull serum albumin, BSA) and two types of d-dots. Colors from red to blue represent the logarithmic PL intensity from high to low. (b) PL decay curve at 570 nm with fitting functions. (c) PL spectra of the solution and three reconstructed components. (d) In vitro image (false color) of a mixture of three groups of cells labeled with d-dots with three different PL lifetimes.
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exponential. In the literature, “perfect” single-exponential requires the goodness-of-fit ($\chi^2$) to be 1.0–1.3.58 Within 3–4 orders of magnitude of intensity (usually 5000 counts), the goodness-of-fit ($\chi^2$) for representative samples in Figure 1 was reproducibly in the range between 1.2 and 2.5. Experimental results (see below) revealed that the extremely large time dimension of the tunable PL lifetime (~50–900 μs) made the degree of single-exponential PL decay for all samples be sufficient for PL lifetime multiplexing.

By fixing the emission wavelength in Figure 4a, one could obtain a regular PL decay curve of the mixture at the given emission wavelength. Figure 4b is such an example, which was extracted from Figure 4a by fixing the emission wavelength at 575 nm. Assuming single-exponential PL decay for all three emitters, eq 1 can be applied to fit the results at a given wavelength.

$$I_I = I_{I_1} e^{-t/\tau_1} + I_{I_2} e^{-t/\tau_2} + I_{I_3} e^{-t/\tau_3}$$

(1)

The terms on the right side depends on the number of emissive components in the system. For the system in Figure 4a, there were three components. Here $t$ is the decay time and $I_{I_1}$, $I_{I_2}$, $I_{I_3}$, $I_{I_1}$, $I_{I_2}$, and $I_{I_3}$ are respectively the PL intensity of total, Component 1, Component 2, and Component 3 at the given wavelength ($\lambda$). The single-exponential PL decay lifetime values of three components are $\tau_1$, $\tau_2$, and $\tau_3$, respectively. For $\lambda$ equal to 575 nm, three lifetime components obtained from the fitting were in excellent accordance with the lifetimes of bull serum albumin (~10 μs, limited by the time resolution of the optical system) and two d-dots (Figure 4b). Applying eq 1 to each wavelength, one would obtain three sets of PL intensity and wavelength, which would result in three separate spectra for three components. Figures 4c and S10 show that the entire emission spectrum reconstructed for each component matched the spectrum of the corresponding pure substance nicely, which quantitatively confirmed suitability of the d-dots emitters for PL lifetime multiplexing.

In imaging applications of PL lifetime multiplexing, PL decay dynamics needs to be deconvoluted in real space, instead of in wavelength dimension discussed above. Three types of Mn:ZnSe/ZnS d-dots with their nearly single-exponential decay lifetime being 220, 550, and 840 μs were in excellent accordance with the lifetimes of bull serum albumin (~10 μs, limited by the time resolution of the optical system) and two d-dots (Figure 4b). Applying eq 1 to each wavelength, one would obtain three sets of PL intensity and wavelength, which would result in three separate spectra for three components. Figures 4c and S10 show that the entire emission spectrum reconstructed for each component matched the spectrum of the corresponding pure substance nicely, which quantitatively confirmed suitability of the d-dots emitters for PL lifetime multiplexing.

In imaging applications of PL lifetime multiplexing, PL decay dynamics needs to be deconvoluted in real space, instead of in wavelength dimension discussed above. Three types of Mn:ZnSe/ZnS d-dots with their nearly single-exponential decay lifetime being 220, 550, and 840 μs were separately loaded into three batches of RAW264.7 mouse macrophage cells. After subsequent mixing of three batches of cells, a series of digital fluorescence pictures were taken under a microscope coupled with a pulsed laser. The PL signals of the pictures were deconvoluted using eq 2.

$$I_{I(x,y)} = I_{I_1(x,y)} e^{-t/\tau_1} + I_{I_2(x,y)} e^{-t/\tau_2} + I_{I_3(x,y)} e^{-t/\tau_3}$$

(2)

The difference between eqs 1 and 2 is the terms for PL intensity. Here, $I_{I(x,y)}$, $I_{I_1(x,y)}$, $I_{I_2(x,y)}$, $I_{I_3(x,y)}$, and $I_{I_3(x,y)}$ are respectively the PL intensity of total, Component 1, Component 2, and Component 3 at a given pixel in the pictures with its spatial position as $(x, y)$. Figure 4d shows the false color image of three batches of cells with good spatial resolution.

Results further verified that the representative d-dots were sufficiently single-exponential to resolve imaging information with spatial overlapping. Figure 5a is a steady-state fluorescence image of an oval overlapped with two letters “M” and “n”, in which each of three objects was printed with one specific type of Mn:ZnSe/ZnS d-dots with nearly single-exponential decay dynamics. Upon exciting the same pattern with a pulsed laser, a series of digital pictures were taken with 67 μs intervals (see three sample pictures in Figure 4b). To obtain the encoded information within one exposure, the average number of photons absorbed per dot per pulse was about 14, which took advantage of the power insensitivity of PL decay dynamics of the Mn:ZnSe/ZnS d-dots (Figure S8). Using eq 2, one could retract the original information as shown in Figure 5c in false colors.

The submillisecond lifetime, nearly single-exponential PL decay dynamics, broad and intense absorption, and high PL quantum yield enabled greatly simplified instrumentation for PL lifetime multiplexing (Figure 5d). It can be as simple as a pulsed light source with ~10 μs pulse width, a commercial high-speed camera with ~100 μs frame rate, and some lens. It is worth mentioning that the exciton emission of intrinsic semiconductor nanocrystals was applied in conjugation with organic dyes in PL lifetime multiplexing.59 However, because of their sensitivity to the environment (Figure 3) and nanosecond lifetime, it would be difficult for intrinsic semiconductor nanocrystals to improve the traditional schemes of PL lifetime multiplexing significantly. For the same reasons, intrinsic semiconductor nanocrystals would not be compatible with the scheme shown in Figure 5d.

PL color multiplexing and PL lifetime multiplexing imply powerful and unique solutions for data storage,53 document security,51,52 and biomedical detection/imaging.60,61 Working in time dimension, PL lifetime multiplexing renders rich coding channels with low sensitivity to excitation and emission intensity.22 However, the nanosecond PL lifetime of common emitters—usually organic dyes—made PL lifetime multiplexing tightly bound with time-correlated single photon counting (TCSPC), whose point-by-point scanning feature dictates the lifetime imaging to be time-consuming and instrumentation intensive. Though time-gated and wide-field imaging techniques shorten detection time, they require highly stable excitation and sophisticated detection schemes, given the short lifetime of common emitters. Up-conversion nanoparticles62 offer a significantly broad lifetime window, i.e., in the microsecond range, but their low absorption coefficient and low PL quantum yield could be challenging for applications with simple scheme shown in Figure 5d.
The magnetic properties at excited states could be different from those at the ground state, at least for the Mn—Mn dimers.\textsuperscript{13,65} Interestingly, it was reported that the PL decay dynamics of d-dots seemed to be closely related to the magnetic properties at excited states of d-dots.\textsuperscript{13} Single-exponential PL decay dynamics in a large time window imply nearly monodisperse magnetic coupling and magnetoluminescence in a given sample described in this work. These monodisperse properties should depend on the Mn\textsuperscript{2+} ion concentration of a given sample. Magnetic properties and magnetoluminescence of these unique emitters are under study for exploring applications in quantum computer and many other spin-based information technologies.\textsuperscript{65}

In conclusion, the dopant emission of Mn:ZnSe/ZnS d-dots could be synthesized not only with high PL quantum yield and desirable excitation properties but also with nearly single-exponential decay dynamics. Furthermore, their single-exponential PL decay lifetime was reproducibly controlled yet tunable in a large time window, approximately between 50 and 900 μs. By epitaxial growth of thick ZnS shell, their PL brightness and decay dynamics were both robust against challenging environments. In comparison to exciton emission of intrinsic semiconductor nanocrystals, these PL properties were found to be unique and would enable interesting applications, such as various schemes of PL lifetime multiplexing, which would be difficult for intrinsic semiconductor nanocrystals. Demonstration of these unique emitters implies that d-dots can be complementary to intrinsic semiconductor nanocrystals to potentially cover a large range of emissive materials in advanced technologies.

\section*{ASSOCIATED CONTENT}

\subsection*{Supporting Information}

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Experimental details and figures (PDF)

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