

Engineering bright sub-10-nm upconverting nanocrystals for single-molecule imaging

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1 Imaging at the single-molecule level reveals heterogeneities
2 that are lost in ensemble imaging experiments, but an
3 ongoing challenge is the development of luminescent probes
4 with the photostability, brightness and continuous emission
5 necessary for single-molecule microscopy^{1–6}. Lanthanide-
6 doped upconverting nanoparticles overcome problems of
7 photostability and continuous emission^{7–12} and their upcon-
8 verted emission can be excited with near-infrared light at
9 powers orders of magnitude lower than those required for con-
10 ventional multiphoton probes^{13,14}. However, the brightness of
11 upconverting nanoparticles has been limited by open questions
12 about energy transfer and relaxation within individual nano-
13 crystals and unavoidable tradeoffs between brightness and
14 size^{15–18}. Here, we develop upconverting nanoparticles under
15 10 nm in diameter that are over an order of magnitude brighter
16 under single-particle imaging conditions than existing compo-
17 sitions, allowing us to visualize single upconverting nanoparti-
18 cles as small ($d = 4.8$ nm) as fluorescent proteins. We use
19 advanced single-particle characterization and theoretical mod-
20 elling to find that surface effects become critical at diameters
21 under 20 nm and that the fluences used in single-molecule
22 imaging change the dominant determinants of nanocrystal
23 brightness. These results demonstrate that factors known to
24 increase brightness in bulk experiments lose importance at
25 higher excitation powers and that, paradoxically, the brightest
26 probes under single-molecule excitation are barely luminescent
27 at the ensemble level.

28 Lanthanide-doped upconverting nanoparticles (UCNPs) absorb
29 multiple photons in the near-infrared (NIR) and emit at higher
30 energies in the NIR or visible (Fig. 1a). They demonstrate significant
31 advantages over other luminescent reporters, and have therefore
32 generated much excitement^{7,8}. These include an absence of on-off
33 blinking, single-molecule multiphoton NIR excitation at powers
34 approaching those used for standard one-photon confocal
35 imaging (Fig. 1b)¹⁹, no overlap with cellular autofluorescence, and
36 no measurable photobleaching under prolonged single-particle
37 excitation^{9–12}. Recent synthetic efforts have established control
38 over UCNP size to produce smaller nanocrystals more compatible
39 with many imaging applications^{15,16}, but this size reduction also sig-
40 nificantly reduces brightness because surface losses increase in
41 importance while the number of sensitizer and emitter ions per parti-
42 cle are reduced as r^3 . UCNPs make use of energy transfer upcon-
43 version (ETU), in which sensitizer ions with relatively large
44 absorption cross-sections sequentially transfer absorbed energy to
45 emitter ions, which luminesce (Fig. 1a, Supplementary Fig. 1).
46 The most common upconverting nanocrystal composition is

β -phase NaYF₄ doped with 20% Yb³⁺ sensitizer and ~2% Er³⁺ 47
emitter; these concentrations, in both bulk materials^{20,21} and nano- 48
crystals^{3,22}, have been suggested to optimize brightness by increasing 49
photon absorption and minimizing luminescence quenching. 50

Recent work on larger nanocrystals ($d \approx 40$ nm) has shown 51
improvements in brightness with higher emitter concentrations at 52
high excitation irradiance¹⁸. The photophysical processes leading to 53
luminescence quenching in larger nanocrystals and in bulk are 54
related primarily to cross-relaxation between dopants and energy 55
migration to defects, but it is less clear how these kinetics are modi- 56
fied as nanocrystal sizes drop to single-digit diameters. To under- 57
stand the efficiency of the ETU process in these UCNPs and the 58

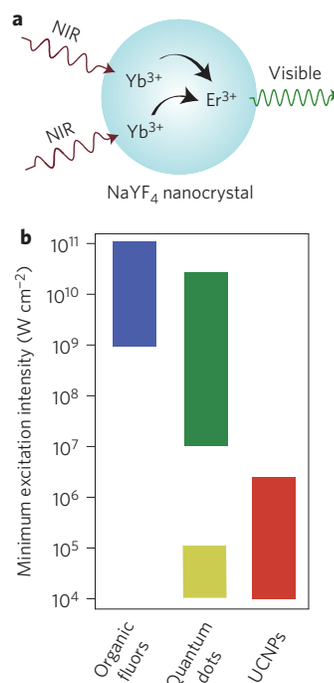


Figure 1 | Luminescence of UCNPs. **a**, Schematic of energy transfer upconversion with Yb³⁺ as sensitizer and Er³⁺ as emitter. **b**, Minimum peak intensities of NIR light needed for multiphoton single-molecule imaging of various classes of luminescent probes. The peak intensity ranges shown are required to detect signals of ~100 c.p.s. for core-shell quantum dots (green, ref. 13), 40 nm colloidal double dot-rods (yellow, ref. 34), organic fluorophores¹⁴ and UCNPs (this study and refs 15, 16).

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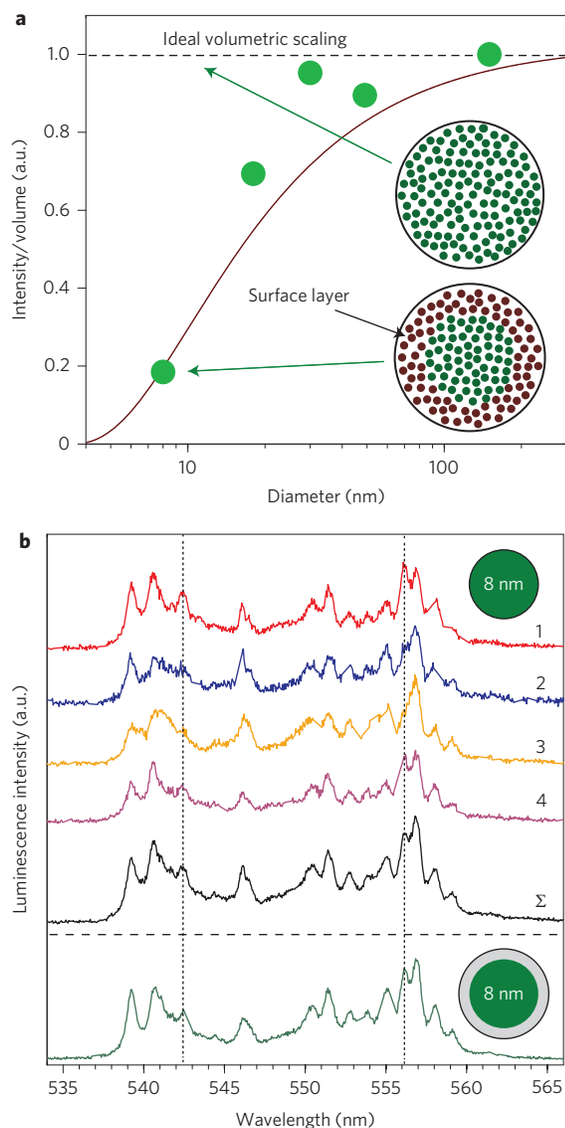


Figure 2 | UCNP size-dependent luminescence intensity and heterogeneity.

a, Deviation of single UCNP luminescence intensity normalized to particle volume ($n = 300$ total) from ideal volumetric scaling. The curve represents calculated intensity normalized to volume for UCNPs with a non-luminescent surface layer of 1.7 nm. Only intensities from single, unaggregated nanocrystals, as determined by Supplementary Fig. 5, are used. **b**, Fine spectra of the green emission bands collected from four single 8 nm UCNPs and their averaged spectra. Vertical dotted lines highlight peaks exhibiting heterogeneity between individual UCNPs. The green emission spectrum of an 8 nm UCNP with epitaxial 1.8 nm undoped shell is shown below the horizontal dashed line.

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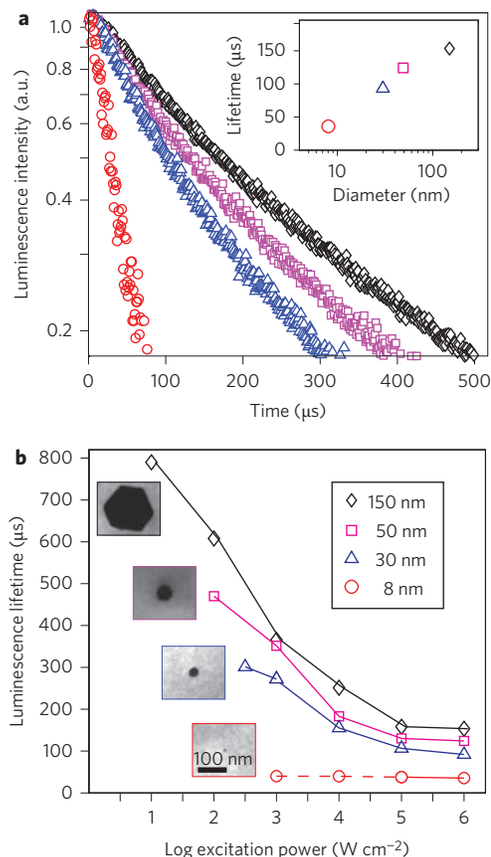


Figure 3 | Single UCNP luminescence lifetime as a function of particle size and excitation power.

a, Luminescence decay (normalized) plotted for various UCNP diameters. See Supplementary Fig. 10 for full lifetime curves. Inset: lifetime values plotted as a function of UCNP diameter. Lifetime values were determined by fitting luminescence decay curves to a double exponential and plotting only the dominant decay value. **b**, Single UCNP lifetime values for various diameters plotted as a function of excitation power. In these plots, emission from all wavelengths between 532 nm and 700 nm was used for the fit because the trends were the same for all emission bands in this range. Separate fits for just green and red emission were also collected and are discussed in Supplementary Figs 7 and 8 and Supplementary Table 1. For simplicity, only dominant lifetime decay values are plotted. Dashed line represents data collected from 8 nm UCNP clusters.

reducing brightness below that predicted for ideal volumetric scaling (Fig. 2a). This trend is consistent with recent ensemble measurements on sub-10-nm UCNPs^{23,24}, and these data can be analysed using a simple calculation in which the UCNP is divided into two regions: a dark surface region and a luminescent core region. In the context of this calculation, our data indicate the dark surface radius is ~ 1.7 nm (ref. 25), because the observed intensity of the 8 nm particles is equivalent to the extrapolated intensity of an ~ 4.6 -nm-diameter particle assuming ideal volumetric scaling.

To understand the origins of these surface-related losses, we collected visible emission spectra from approximately 40 individual 8 nm UCNPs. Unlike homogeneous room-temperature spectra of larger UCNPs³, these high-resolution spectra are heterogeneous, with particle-to-particle variations in peak intensities at 541 and 557 nm (Fig. 2b; compare curves 1 and 4 with curves 2 and 3). One explanation for such heterogeneity would be that emission from these UCNPs is dominated by only a few of the approximately 70 Er^{3+} emitters present in each UCNP, but this is not supported by photon antibunching studies of single UCNPs (Supplementary Fig. 6). The addition of undoped $NaYF_4$ shells to these nanocrystals

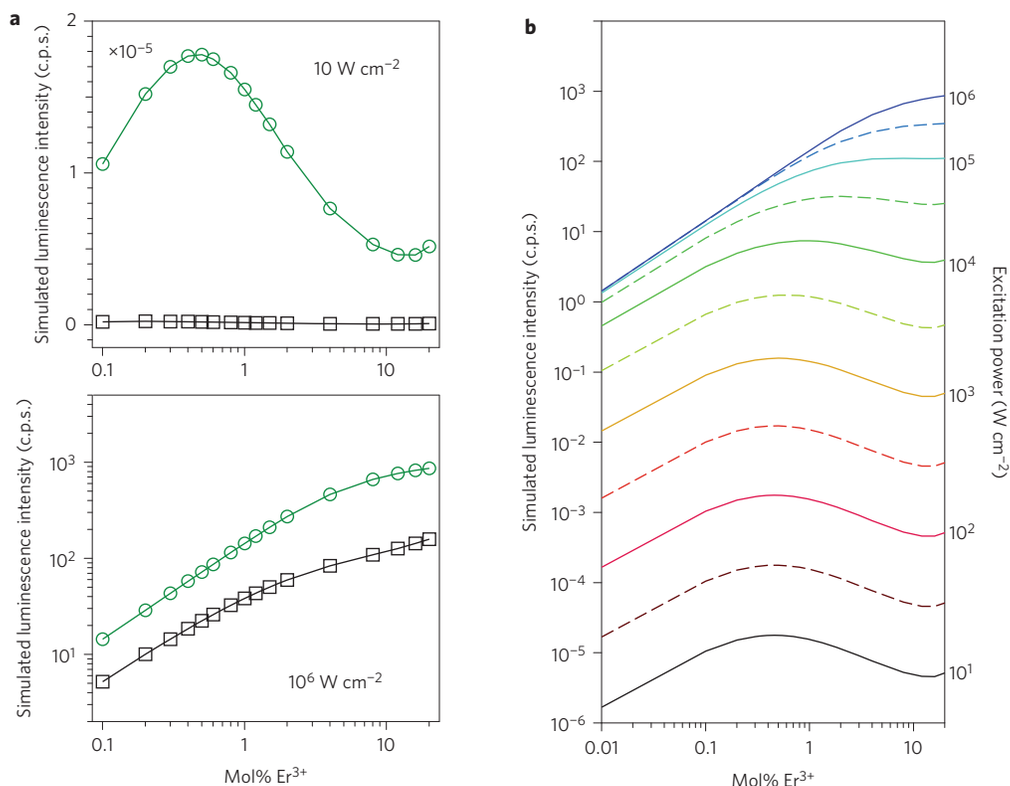


Figure 4 | Simulated UCN emission intensity. **a**, Theoretical models of integrated 8 nm UCN emission (500–700 nm) as a function of Er^{3+} mol% for 20% Yb^{3+} (green circles) and 2% Yb (black squares) for low (10 W cm^{-2} , upper panel) and high (10^6 W cm^{-2} , lower panel) power excitation. **b**, Simulated luminescence intensity of UCNPs with 20% Yb^{3+} as a function of Er^{3+} doping and excitation power. Solid lines are calculated UCN emission following excitation at powers on the right axis. Dashed lines are half logarithmic spacings.

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1 eliminates this heterogeneity (Fig. 2b, bottom trace) suggesting a
 2 region within the nanocrystal in which the lanthanides may be
 3 emissive but are energetically coupled to the surface. The observed
 4 spectral differences may arise from either variations in lanthanide
 5 distributions between nanocrystals, or from subtle variations in
 6 surface defects, surface reconstruction, or faceting. This identifica-
 7 tion of losses from energy transfer to the surface suggests one
 8 means for improving emission from small UCNPs.

9 We next measured the luminescence lifetimes of individual
 10 UCNPs of various sizes to probe the balance between the energy
 11 transfer pathways that lead to radiative and non-radiative relaxation.
 12 As UCNP size decreases, fast and presumably non-radiative recombi-
 13 nation dominates (Fig. 3a). To determine whether surfaces are a
 14 primary source of the non-radiative relaxation, we measured the life-
 15 times of 8 nm UCNP cores with added undoped NaYF_4 shells.
 16 Emission intensities and lifetimes both increase as shell thickness
 17 increases, up to a shell thickness of $\sim 1.8 \text{ nm}$ (Supplementary
 18 Fig. 11), suggesting that the increased luminescence is due to
 19 improved quantum yields for core-shell nanocrystals^{16,26}. This satura-
 20 tion of lifetime and brightness at a shell thickness of $\sim 1.8 \text{ nm}$ is con-
 21 sistent with a model (Fig. 2a) in which the emitters within this surface
 22 radius are quenched by energy transfer to vibrational modes at the
 23 nanocrystal surface or in organic capping ligands. Because dopant
 24 excited states can be coupled to surface vibrations directly or via reso-
 25 nant, donor-to-donor energy migration to directly coupled states,
 26 this $\sim 1.8 \text{ nm}$ distance can be interpreted physically as a convolution
 27 of the critical distance for direct coupling and the average energy
 28 migration length (Supplementary Discussion 1). Critically, for
 29 UCNPs with $d < 8 \text{ nm}$, this dark surface layer occupies a large
 30 majority ($>80\%$) of the total nanocrystal volume.

31 Previous work has shown that UCNP lifetimes are roughly inde-
 32 pendent of excitation power for powers $< 100 \text{ W cm}^{-2}$ (ref. 25), but

the low power densities used in those experiments are not useful for
 imaging small, single UCNPs. At higher single-nanocrystal powers,
 we observe a pronounced lifetime dependence on excitation power
 density for all UCNPs with $d > 30 \text{ nm}$ (Fig. 3b). We considered the
 possibility that these higher powers may generate enough heat to
 significantly affect lifetimes, although we observed no change in life-
 time for any UCNPs as pump pulse widths were increased from
 250 μs to 2.5 ms, indicating that the particles reach a steady-state
 temperature in less than 250 μs . Rather, the lifetime dependence
 on excitation power suggests that the higher fluences increase the
 spatial density of populated Er^{3+} excited states with longer lifetimes
 (such as $^4\text{S}_{3/2}$), increasing rates of ETU and cross-relaxation out of
 these states. This leads to shorter lifetimes for states that emit visible
 photons and increases the population of higher-energy excited states
 (such as $^4\text{G}_{11/2}$)²⁷. However, for the sub-10-nm UCNPs, the lumi-
 nescence lifetime is short and remains constant for all excitation
 powers studied here (Fig. 3b), due to the dominance of surface-
 related non-radiative recombination in UCNPs of this size. This
 suggests that the entire 8 nm particle is energetically coupled to
 the surface—although the nanocrystal radius is larger than the
 dark radius of 1.7 nm—and is consistent with the presence of
 the sub-surface region containing emissive lanthanides that are
 nonetheless influenced by the surface.

Surface-related non-radiative recombination greatly shortens the
 lifetime of excited emitters, which suggests an opportunity in that
 emitter concentrations could be increased substantially beyond 2%
 before self-quenching becomes a major factor. In this case, the
 surface energy losses change the relative balance between energy
 transfer pathways in smaller UCNPs. In addition, the higher flu-
 ences of single-molecule imaging push the nanocrystals into the
 excitation saturation regime and further modify the balance of
 energy transfer between states¹⁸. These findings suggest that the

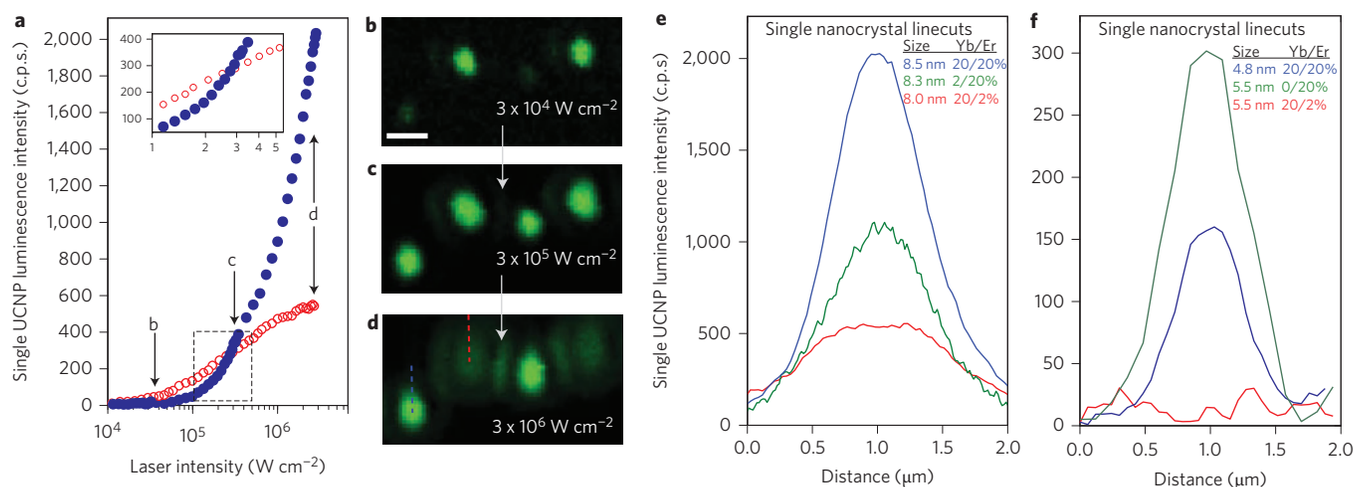


Figure 5 | Luminescence intensity of single UCNPs as a function of Er^{3+} and Yb^{3+} doping. **a**, Luminescence intensity of single UCNPs with 20% (red curve) and 2% (blue curve) Er^{3+} , each with 20% Yb^{3+} , plotted as a function of excitation power. **b–d**, Confocal luminescence images of a region containing single UCNPs in **a**, collected at increasing excitation powers. Dashed lines indicate regions from which luminescence intensity was collected for data in **a**. Scale bar, $1\ \mu\text{m}$. **e,f**, Linecuts of single-particle luminescence scans of $\sim 8\ \text{nm}$ UCNPs and $\sim 5\ \text{nm}$ UCNPs with varying Er^{3+} and Yb^{3+} doping levels. Size is diameter (nm) and Yb and Er levels are in atomic %. Emission from $5.5\ \text{nm}$ UCNPs with 20% Yb^{3+} and 2% Er^{3+} are indistinguishable from our sensitivity limit determined by instrument noise. These data were collected using $3 \times 10^6\ \text{W cm}^{-2}$ excitation power and a $\times 100$, $1.4\ \text{NA}$ oil-immersion lens (see Methods and Supplementary Fig. 2 for details).

1 optimal design has not yet been achieved for sub-10-nm UCNPs
 2 intended for single-molecule applications, where the goal is to maxi-
 3 mize emission over background and noise levels.

4 We used these observations to refine computational models of
 5 UCNP energy transfer to design UCNPs that are brighter under
 6 single-molecule imaging conditions. Emission intensity was calcu-
 7 lated as a function of Er^{3+} and Yb^{3+} dopant concentrations using
 8 a population balance model that has successfully predicted the
 9 steady-state luminescence spectra of various lanthanide-doped
 10 UCNPs²⁸. Based on the single-nanocrystal intensity and lifetime
 11 data, we modified this model to include a third, non-emissive
 12 surface species that can accept energy from excited lanthanide
 13 states (Supplementary Methods 1). At the low excitation powers
 14 typical of ensemble measurements ($10\ \text{W cm}^{-2}$), the simulated
 15 emission intensity is maximized at $\sim 0.5\%$ Er^{3+} (Fig. 4a, top)—
 16 close to the 1–2% Er^{3+} maximum typically observed for Er^{3+} -
 17 containing nanocrystals²⁹. On the other hand, increasing the Yb^{3+}
 18 concentration from 2 to 20% increases the luminescence by two
 19 orders of magnitude at all Er^{3+} concentrations. However, when we
 20 apply the refined model to the higher powers (10^5 – $10^7\ \text{W cm}^{-2}$)
 21 needed to observe single small nanocrystals, we find that the sensitizer
 22 $\text{Yb}^{3+}\ ^2\text{F}_{5/2}$ excited manifold has its population approach 70% of
 23 the overall Yb^{3+} ion concentration, well into the saturation regime
 24 (Supplementary Fig. 14). At these higher fluences, the incremental
 25 Yb^{3+} absorption cross-section decreases significantly¹⁹, thereby
 26 resulting in a reduced dependence of emission on Yb^{3+} sensitizer
 27 concentration (Fig. 4a, bottom). The number of Er^{3+} emitters
 28 becomes the fundamental bottleneck for visible emission (Fig. 4b
 29 and Supplementary Discussion 2) because radiative relaxation
 30 rates for parity-forbidden $4f^*4f$ transitions are significantly slower
 31 than photon absorption rates in this regime. Luminescence
 32 increases linearly as a function of Er^{3+} doping percentage
 33 (Fig. 4b), because the emission intensity is proportional to the
 34 number of Er^{3+} states that can emit visible photons (for example,
 35 $^4\text{S}_{3/2}$, $^4\text{F}_{9/2}$)²⁸, as well as the number of Er^{3+} ions that can absorb
 36 incident photons. This model predicts that a UCNP with
 37 20% Er^{3+} will be three to five times brighter at higher excitation flu-
 38 ences than a conventional 2% Er^{3+} UCNP of the same size and that
 39 Yb^{3+} doping levels are of less importance than at lower fluences.

This points to a radically different design strategy for nanocrystals to be used for ensemble measurements versus those to be used for single-molecule studies: for single-molecule studies, emitter concentrations should be as high as possible without compromising the structure of the nanocrystal, while sensitizer content becomes less significant at higher powers and can potentially be eliminated for single-molecule imaging applications. Based on these calculations, we synthesized a series of 8 nm and 5 nm nanocrystals with higher emitter or lower activator content, and imaged them at single-particle powers (Fig. 5). At lower powers ($\sim 100\ \text{W cm}^{-2}$), these new compositions have vanishingly low quantum yields (Supplementary Table 4), indicating that they would behave poorly in ensemble imaging experiments. However, comparing β - NaYF_4 nanocrystals doped with 20% Yb^{3+} and 20% Er^{3+} (as well as 25% Gd^{3+} added to maintain β -phase morphology at high lanthanide doping levels⁷) to conventional 8 nm UCNPs (β - NaYF_4 with 20% Yb^{3+} and 2% Er^{3+}) dispersed on the same glass substrate, we observe that the conventional UCNPs are visible at lower powers, but the high- Er^{3+} UCNPs are not (Fig. 5a–d). As excitation powers are raised, the conventional UCNPs saturate in brightness while the high- Er^{3+} UCNPs become visible and continue to increase in brightness, surpassing the conventional UCNPs. The excitation intensity at which the 20% Er^{3+} UCNPs become brighter than their 2% Er^{3+} counterparts is $\sim 3 \times 10^5\ \text{W cm}^{-2}$ (Fig. 5a). The diverging brightness trends of these UCNPs, which agree well with simulated data (Supplementary Fig. 14), indicate one advantage of tailoring dopant compositions specifically for higher-flux imaging. This advantage is explicitly illustrated by simultaneously imaging the two classes of UCNPs at increasing excitation intensities and also may be potentially useful in optical encoding applications^{18,30} or in conjunction with surface modifications that shift UCNP absorption or emission³¹.

This strategy for increasing single-nanocrystal brightness suggests that even smaller UCNPs may be viable as single-molecule probes. We tested this idea by synthesizing 5.5-nm-diameter β - NaYF_4 UCNPs with 20% Er^{3+} and no Yb^{3+} sensitizer, as well as 4.8 nm UCNPs with $\sim 20\%$ each of Er^{3+} , Yb^{3+} and 25% Gd^{3+} (Fig. 5e,f and Supplementary Discussion 3). These nanocrystals are significantly smaller than other UCNPs imaged at the single-

particle level^{16,18} and are the approximate size of monomeric genetically encoded fluorescent proteins. We find that each of these compositions is significantly brighter than the canonical β - $\text{NaYF}_4\text{:}20\% \text{Yb}^{3+} 2\% \text{Er}^{3+}$ nanocrystals. We measured signals of ~ 150 c.p.s. for single 4.8 nm UCNPs doped with 20% Yb^{3+} and 20% Er^{3+} and note that these nanocrystals have quantum yields of $< 0.001\%$ at lower excitation powers (Supplementary Table 4). In comparison, we were unable to image single 5.5 nm β - $\text{NaYF}_4\text{:}20\% \text{Yb}^{3+} 2\% \text{Er}^{3+}$ nanocrystals because their signal falls below our sensitivity limit of ~ 25 c.p.s. (Fig. 5f, Supplementary Fig. 12). The 4.8 nm UCNPs with 20% Yb^{3+} and 20% Er^{3+} are over 500-fold smaller in volume than nanocrystals optimized with higher emitter concentrations for single-particle excitation irradiance and imaged as single nanocrystals in suspension¹⁸. These protein-sized nanocrystals have significant advantages over the larger UCNPs previously used in cellular imaging applications, including increased accessibility to small subcellular structures, greater tissue penetration and reduced interference with biomolecule function, trafficking and binding events¹⁶.

These new rules for designing small, bright UCNPs address key obstacles for optimizing nanocrystals as single-molecule probes and suggest a single-molecule probe development strategy involving iterative rounds of kinetic modelling and detailed nanocrystal characterization. We find that factors known to increase brightness at low powers are unimportant at single-molecule powers and that the brightest single-molecule probes may be non-luminescent at the ensemble level. For the most efficient nanocrystals, we find that 5 nm UCNPs are bright enough to be used in single-molecule detection. We anticipate further gains in brightness through iterative rounds of modelling and nanocrystal characterization, as well as surface modifications that alter the balance between various energy transfer pathways^{31,32}. Together, these advances open the door to a range of applications, including cellular and *in vivo* imaging^{12,33}, as well as reporting on local electromagnetic near-field properties of complex nanostructures.

Methods

β - $\text{NaYF}_4\text{:Yb}^{3+}, \text{Er}^{3+}$ nanocrystals were synthesized as reported previously¹⁶ and characterized by analytical transmission electron microscopy (TEM), dynamic light scattering and X-ray diffraction. UCNPs were dispersed in hexane to ~ 0.1 nM before dropcasting onto silicon nitride TEM grids (Ted Pella, #21569-10). Laser scanning confocal imaging was performed under ambient conditions using a 980 nm continuous-wave laser (Thorlabs TCLDM9, 300 mW diode) (see Supplementary Fig. 2 for instrument details). Because the diffraction-limited beam spot was larger than individual nanoparticle size, single particles were confirmed on SiN TEM-grid samples by subsequent SEM imaging (Zeiss Ultra-55, operating in transmission mode).

Single-particle luminescence intensity histograms were compiled from ~ 50 –300 individual particles for each size (Supplementary Fig. 5), and single-particle spectra were obtained using the same confocal imaging system. To obtain lifetime data, a time-correlated single-photon counter (TCSPC, Picoquant) was used to tag the photon arrival times of the collected luminescence with respect to the laser operating in pulsed mode. The resulting time-resolved luminescence plots were fitted to a double exponential, because the non-radiative recombination rates from the surface and core regions of a UCNP are generally different²⁵. For clarity, only the dominant decay value was plotted in Fig. 3 (Supplementary Figs 7 and 8 show more complete information).

In our optical set-up, the excitation laser was pre-focused with a 500 mm lens before entering the back-aperture of either a 0.95 NA, $\times 100$ air objective (used for the data in Figs 2 and 3) or a 1.4 NA, $\times 100$ oil objective (Fig. 5). Emitted light was collected with the same objective and filtered by two 700 nm short-pass filters (Chroma) to remove residual laser light. Emission was then routed either through a spectrometer, or through additional 532 nm long-pass filters, and onto a single photon-counting avalanche photodiode. To collect data from just the green or red spectral band, a 540 ± 20 nm (green) or 650 ± 20 nm (red) bandpass filter was used in place of the 532 nm long-pass filters. A TCSPC (Picoquant) was used for luminescence lifetime measurements.

The single-particle imaging shown in Fig. 5 was performed using a 1.4 NA, $\times 100$ oil-immersion objective. Equal dilutions of 20/2% (Yb/Er) and 25/20/20% (Gd/Yb/Er) UCNPs were dropcast onto a glass coverslip and imaged at various powers. For mixtures of nanocrystals with different dopant compositions, the compositions were identified by comparison with the optical behaviour of each

composition imaged individually (Fig. 5d). Single-particle power-dependent plots were constructed by scanning the laser beam over an isolated particle and dividing the collected luminescence curve by the laser beamspot profile, assuming Gaussian shapes. Consecutive linecut scans at increasing excitation powers were compiled to produce the plots shown in Fig. 5a. The linecuts shown in Fig. 5e,f were collected from single particles at a fixed excitation power of $3 \times 10^6 \text{ W cm}^{-2}$.³⁴

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Author contributions

P.J.S., B.E.C., D.J.M., D.J.G., E.M.C. and A.D.O. developed the idea for the project. A.D.O.,
27 P.J.S. and B.E.C. synthesized the nanocrystals. E.M.C. performed the kinetic modelling
28 and simulations. D.J.G. performed the single-UCNP optical imaging, lifetime and
29 spectroscopy measurements, as well as single-UCNP scanning TEM imaging. S.A. and
30 M.V.P.A. provided TEM and analysis. J.J.U. provided in-depth discussions. E.S.B. and B.S.
31 provided in-depth discussions and performed data analysis. D.J.G., E.M.C., B.E.C. and
32 P.J.S. wrote the paper. All authors reviewed the paper. 33 34

Additional information

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Competing financial interests

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