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Size of the rare-earth ions: a key factor in phase tuning and morphology control of binary and ternary rare-earth fluoride materials†

In an ionic liquid assisted solvothermal synthesis developed by us for the synthesis of rare-earth (RE) fluorides, it is possible to control the product formation by the choice of the rare earth ion. For rare-earth cations with smaller ionic radii (below 1.075 Å), cubic NaREF₄ with a spherical morphology is obtained, whilst for rare-earth cations with radii between 1.08 Å and 1.13 Å, the formation of hexagonal NaREF₄ with a nanorod-like morphology is observed. For rare earth ions with a larger radius than that of La³⁺ (1.216 Å), instead of ternary fluorides, binary fluorides REF₃ in the trigonal modification is obtained. The growth mechanism behind this morphology change is explained from atomistic origin using electron microscope studies. The lattice strain changes with the rare-earth fluoride phase. For cubic NaREF₄ a tensile strain is observed, whilst for the hexagonal and trigonal binary fluoride a compressive strain is observed. The optical properties of the obtained materials promises use for various optoelectronic applications.

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Introduction

Over the last decade rare-earth (RE) ion doped materials have drawn tremendous attention in the field of photonic and biophotonic applications. 1-10 With respect to their narrow emission bands, large Stokes shifts, weak autofluorescence and decay times typically in the range of milliseconds, RE-doped materials in certain respects are superior to organic dyes and semiconductors.11,12 A judicious choice of the host and dopant ion/ ion pairs is a precondition to obtain the desired optical materials.1 The main criteria in the choice of a suitable host matrix are a low phonon energy, a high refractive index and easy incorporation of dopants together with chemical and thermal stability.13 Based on these criteria, binary and ternary fluorides appear to be effective host materials for RE³⁺ ion doping. ¹⁴⁻²² For rare-earth trifluorides, REF3, two polymorphs are known. Their relative stability depends on the rare earth ionic radius and temperature. For the lighter REF₃ (RE = La-Nd) with larger ionic radius the trigonal tysonite type of structure (LaF₃, $P\bar{3}c1$) is

the stable form, whilst for the heavier REF3 with smaller RE3+ (Dy-Lu, Y, YF₃ type) and the orthorhombic β -YF₃ (Pnma) is the preferred structure. REF₃ with RE³⁺ ions that have a size in between are dimorphic with the orthorhombic form being the room temperature stable phase and the tysonite structure being the high temperature phase. 23-25 Similarly, sodium ternary rare earth fluorides are known to form two polymorphs: a cubic phase and a hexagonal phase.26 Whilst classical high temperature synthesis methods typically lead to the thermodynamically stable form, on the nanoscale by a judicious choice of the reaction conditions, it is possible to selectively obtain the less stable form. For nanomaterials, it is possible to tune the crystal phase by variation of the reaction conditions, such as temperature, pH of the reaction, Ln3+/F- ratio, addition of a surfactant or capping agent, calcination temperature etc. 27-30 Recently a strategy for phase tuning that relies on ion doping has been developed for NaREF4 NCs (nanocrystals).31-33 For optical materials, either the trigonal REF3 form or, even more, the hexagonal modification of NaREF4 is the researcher's choice due to the higher luminescence efficiency. A wide number of NaREF₄ materials doped with various RE³⁺ ions as up and down converting materials have been reported.34-42 By tuning the crystal phase of the host materials, the luminescence dynamics of the RE ion can be tuned. 14,18,26,29 Similarly morphology is an important parameter to tune the luminescence dynamics as illustrated by Mai et al. which had prepared sodium europium fluoride nanospheres, nanopolyhedra and nanorods and compared their fluorescence emission spectra on the basis of different morphologies.37 Similar morphology dependent

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RE3+
NaLnF4

I.132,4

I.132,4

I.132,4

I.132,4

I.1062A

Er3+

Cubic

Scheme 1 Tuning of crystal phase and nature of product (binary/ternary) depending on the size of the RE³⁺ ions for RE-doped NaREF₄.

studies have been reported for other rare-earth doped materials.^{43–46}

In this article we report the phase and morphology selective synthesis of rare earth doped binary and ternary rare-earth fluorides employing an ionic liquid (IL) assisted solvothermal method. 1-ethyl-3-methyl imidazolium bromide (C2mimBr) is used as the structure controlling agent. The RE³⁺/F⁻ ratio is taken as 1:8 in all the cases. Depending on the ionic radius of the chosen host RE3+ ion, a different NaREF4 polymorph for a given RE or even only formation of the binary REF3 is observed. For example, in the ionic radius range of 1.075 Å to 1.042 Å, cubic NaREF₄ particles with sphere-like morphology are obtained (Scheme 1).47 However, for RE"3+ ions with larger ionic radius in the range of 1.13 to 1.095 Å (for Sm³⁺ to Tb³⁺), hexagonal sodium ternary fluoride with nanorod-like morphology is obtained. A drastic change occurs when a RE"3+ ion with quite a large radius is chosen like La3+ (1.216 A). Instead of a ternary fluoride, the binary fluoride with hexagon like morphology is obtained (Scheme 1). Normally phase transition and morphology control of nanomaterials need at least one of the following changes: heating at high temperature, change of the reaction parameters like reactant ratio, pH etc. and other external influences. 48-52 This type of tuning of the crystal phase, lattice strain, morphology and most importantly nature of the product depending on the ionic radius or size of the RE³⁺ ion is novel and not reported earlier to the best of our knowledge.

2. Experimental

2.1 Synthesis

2.1.1 1-Ethyl-3 methyl imidazolium bromide [C₂mim]Br. Modifying a literature procedure, 58 ml of ethyl bromide (0.79 mol, Sigma Aldrich 98%) and 48 ml of *N*-methyl imidazole (0.607 mol, Sigma Aldrich 99%) were heated under reflux under inert gas (Ar) atmosphere at 40 °C for 3 hours in a round bottom flask (250 ml).⁴⁹ After cooling to room temperature, ethyl acetate was added and the product crushed out of the solution. After

filtration, the crude product was washed with ethyl acetate and dried under vacuum at $25~^{\circ}$ C for 10 hours to give a white solid.

2.1.2 NaYF₄:Eu nanoparticles. In a typical synthesis, 10 ml aqueous solutions of 0.265 M NaCl (J. T. Baker) and (0.265 M) $Y(NO_3)_3 \cdot 6H_2O$ (Alfa Aesar) and the required amount of an aqueous solution of Eu(NO₃)₃·6H₂O (99.9%, Alfa Aesar) to give NaYF₄: 1 mol% Eu were added to 60 ml ethanol containing a 20 ml aqueous 5 wt% [C₂mim]Br solution. To the well-stirred solution, the appropriate amount of an aqueous NH₄F (Sigma Aldrich) solution was added producing a Y³⁺/F⁻ ratio of 1 : 8. The mixture was poured into a TeflonTM lined autoclave (Parr Instruments, Moline, Illinois, USA) and subsequently heated at 200 °C for 4 hours. The obtained nanocrystals was washed several times with ethanol, methanol and finally acetone and dried in an oven at 80 °C.

2.1.3 Other rare earth fluoride nanoparticles. Keeping the procedure similar, different RE" precursors were used, which led to the formation of different phase ternary and binary fluorides. La(NO₃)₃·6H₂O; Ce(NO₃)₃·6H₂O; Nd(NO₃)₃·5H₂O; Sm(NO₃)₃·5H₂O; Gd(NO₃)₃·6H₂O; Tb(NO₃)₃·5H₂O; Dy(NO₃)₃·5H₂O and Yb(NO₃)₃·5H₂O are purchased from the Alfa Aesar in a purity of 99.9%. Details on the synthesis condition can be found in Table 1.

2.2 Characterization

PXRD (powder X-ray diffraction) measurements were carried out on a Huber G70 diffractometer (Rimsting, Germany) using Mo-Kα radiation ($\lambda = 0.07107$ nm). The crystallite size was calculated using the Scherer equation $D = K\lambda/\beta \cos \theta$, where K = 0.9, D represents crystallite size (Å), λ is the wavelength of the Mo-K α radiation, and β is the corrected half-width of the diffraction peak. TEM (transmission electron microscopy; a FEI Tecnai STWIN-T30 using 300 kV electron beam source) was used to investigate the shape, size and lattice structure of the nanocrystals dispersed on a carbon coated copper grid from acetone solution. Morphological characterization was also carried out by SEM (scanning electron microscopy) using a NOVA NANO SEM-450, FEI. Excitation, emission spectra and decay time of all samples were recorded on a Fluorolog 3 (HORIBA JOBIN YVON, Germany) luminescence spectrometer equipped with steady and pulsed Xe lamps for sample excitation and a photomultiplier for signal detection.

3. Result and discussion

3.1 Structural characterization by powder X-ray diffraction and phase evolution

Solvothermal conversion of sodium chloride and an appropriate amount of yttrium and europium nitrate in ethanol/water solution in the presence of [C₂mim]Br yielded nanosized NaYF₄:Eu³⁺ (P1). For a reaction time of 4 hours at 200 °C, only cubic NaYF₄:Eu³⁺ (P1) is obtained (Fig. 1a). A detailed study of the effect of the IL, reaction temperature and reactant ratio (Ln³⁺/F⁻) has already been reported by us earlier. Now this synthesis protocol is explored for other elements of the rare earth series. It turned out that, depending on the rare earth ion

Table 1 Summary of the reaction conditions maintained to prepare different binary and ternary fluorides

Name of the sample	Crystal phase	Crystallite size (nm)	Reaction time & temperature	Concentration of NaCl solution/M	Concentration of the rare-earth nitrate solution/M	Concentration of NH ₄ F solution/M
NaYF ₄ :Eu ³⁺ (P1)	Cubic	17.32	4 h & 200 °C	0.265	0.265	2.126
LaF ₃ :Dy ³⁺ (P2)	Trigonal	10.30	4 h & 200 °C	0.21	0.21	1.680
CeF ₃ (P3)	Trigonal	11.8	4 h & 200 °C	0.21	0.21	1.680
CeF ₃ :Tb ³⁺ (P4)	Trigonal	12.03	4 h & 200 °C	0.208	0.209	1.671
NdF ₃ (P5)	Trigonal	14.8	4 h & 200 °C	0.205	0.205	1.644
NaSmF ₄ (P6)	Hexagonal	10.41	4 h & 200 °C	0.200	0.200	1.644
NaGdF ₄ :Eu ³⁺ (P7)	Hexagonal	9.13	4 h & 200 °C	0.196	0.196	1.568
NaTbF ₄ :Ce ³⁺ (P8)	Hexagonal	7.59	4 h & 200 °C	0.193	0.193	1.55
NaDyF ₄ (P9)	Hexagonal (major) + cubic (minor)	25.8	4 h & 200 °C	0.188	0.190	1.526
NaErF ₄ :Yb ³⁺ (P10)	Cubic	16.56	4 h & 200 °C	0.187	0.187	1.502
NaYbF ₄ :Er ³⁺ (P11)	Cubic	9.5	4 h & 200 $^{\circ}\mathrm{C}$	0.184	0.184	1.47

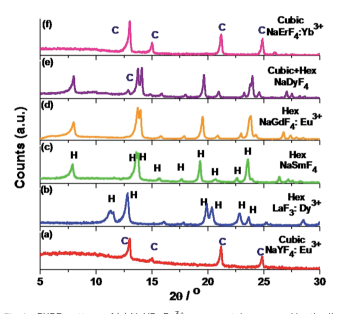


Fig. 1 PXRD pattern of (a) NaYF $_4$:Eu $^{3+}$ nanocrystals prepared by the IL assisted solvothermal synthesis at 200 °C with 4 hours reaction time; (b–e) and (f) PXRD patterns of trigonal LaF $_3$:Dy $^{3+}$, hexagonal NaSmF $_4$, NaGdF $_4$:Eu $^{3+}$, Hex (major) + cubic (minor) NaDyF $_4$ and pure cubic NaErF $_4$:Yb $^{3+}$ prepared with different dopant ions (La $^{3+}$, Sm $^{3+}$, Gd $^{3+}$, Dy $^{3+}$ and Er $^{3+}$) under otherwise similar conditions. In all cases, Ln $^{3+}$ /F $^{-}$ ratio is taken as 1 : 8.

size, different products, either a ternary NaREF₄ polymorph or binary REF₃ in the tysonite type of structure are obtained (Fig. 1 and Scheme 1). In the case of Y³⁺, with an ionic radius of 1.075 Å (using the Shannon radius⁴⁷ for a 9-fold co-ordination) cubic NaYF₄ (P1) (Fig. 1) nanocrystals are obtained. For RE³⁺ ions with an ionic radius in the range of 1.13 to 1.095 Å (for Sm³⁺ to Tb³⁺), hexagonal sodium ternary fluoride (NaLnF₄: Ln = Sm³⁺, Gd³⁺, Tb³⁺ (P6–P8) in Table 1) is obtained (see Fig. 1, S1† and Table 1).

Interestingly, for Dy³⁺ where the ionic radius is in between that of Tb³⁺ and Er³⁺, the cubic modification starts to appear with the majority phase being hexagonal (P9) (as shown in Fig. 1). When the ionic radius is lesser due to lanthanide

contraction as in case of Er³⁺ (1.062 Å) and Yb³⁺ (1.042 Å) only the cubic polymorph (NaErF4 and NaYbF4, P10 and P11) is obtained (Fig. 1, S1[†] and Table 1). At the start of the lanthanide series (for example, for La^{3+} with an ionic radius of 1.216 Å), the formation of the binary rare earth fluoride (P2) instead of the ternary fluoride is observed and this trend is maintained until Nd³⁺ (ionic radius 1.163 Å). It can be anticipated that for smaller size, as for Y³⁺ and for Er³⁺, Yb³⁺, the reaction rate is quite faster which leads to the formation of the kinetic product, the cubic phase. When the size of the RE3+ ion is larger, as for La3+, the reaction rate is slower causing the formation of the binary fluoride instead of the ternary fluoride. For RE³⁺ ions within the range of 1.132 to 1.093 Å, reaction rate is optimum for the generation of thermodynamically stable hexagonal ternary fluoride. This kind of phase tuning depending on the size of the RE³⁺ ion is novel and not previously reported to the best of our knowledge.

3.2 Lattice strain

Generally, the broadening of the diffraction peaks in a PXRD pattern depends upon strain and crystallite size. It is previously noticed that change in the interatomic distance induces the perturbation in the lattice parameter of crystalline nanomaterials leading to stress generation at nanoscale particles. And it is well known that higher the size-volume ratio more will be the strain at the nanoscale level.⁵³ Herein, the lattice strain in the as-prepared nanoparticles can be attributed to the change in ionic radius of the RE³⁺ of the host materials. Lattice strain is changing with changing the ionic radius of RE³⁺ ion in lanthanide series. It means, lattice parameter of host lattice is perturbing in period. It is possible to calculate the lattice strain according to the function derived by Williamson and Hall:⁵¹

$$\beta \cos \theta / \lambda = 1/D + \eta \sin \theta / \lambda \tag{1}$$

where β is the full width at half-maximum (fwhm), θ is the diffraction angle, λ is the X-ray wavelength, D is the effective crystallite size, and η is the effective strain. When plotting $\beta \cos \theta/\lambda$ against $\sin \theta/\lambda$, the strain is the slope and the

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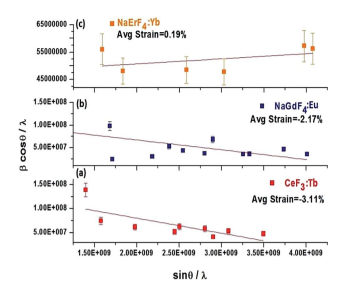


Fig. 2 Plot of β cos θ/λ against sin θ/λ for CeF₃:Tb³⁺ (a) NaGdF₄:Eu³⁺ (b) and NaErF₄:Yb³⁺ (c) prepared solvothermally using [C₂mim]Br at 200 °C.

crystallite size (D) can be derived from the intercept (Fig. 2 and Table 2).52 The lattice strain clearly depends on the size of the rare earth precursor ion. For example, for the binary fluorides where the ionic radius of the precursor RE ion is in the range of 1.216 Å to 1.163 Å (La³⁺ to Nd³⁺), compressive strain is obtained (Table 2). The lattice strain for CeF₃:Tb³⁺ (P4) is compressive (-3.11%) and the crystallite size obtained from the intercept with the x-axis is 17.31 nm which matches with the size obtained from Scherrer equation (Fig. 2a). Likewise, for the RE³⁺ ions with medium ionic radius (1.132 Å to 1.083 Å for Sm³⁺ to Dy³⁺ ions) where the crystal phase is again hexagonal, the obtained lattice strain is compressive (Table 2). For example for $NaGdF_4$ (P7), a compressive strain of -2.17% is obtained and crystallite size is \sim 21 nm (Fig. 2b). However when the size of the rare earth ion is less, for example, for Y3+ and for Er3+ and Yb3+ due to lanthanide contraction, the cubic phase is observed and a drastic change is noticed in lattice strain i.e., tensile strain is obtained. Fig. 3c shows the plot for NaErF₄:Yb³⁺ (P10) showing a tensile strain of +0.19%. Earlier we have reported that cubic sodium yttrium fluoride shows tensile strain however its hexagonal analogue shows compressive strain. Compressive strain is typically observed for smaller crystallite sizes.⁵⁰

3.3 Structural characterizations by scanning electron microscope

To investigate the morphology of the nanoparticles, SEM images are analyzed in detail. Fig. 3a shows the SEM images of CeF3:Tb3+ doped nanocrystals showing nanoparticles with hexagon-like disc shape, whilst for LaF₃:Dy³⁺ (P2) shows cube like morphology (Fig. S2a† inset). A clear change in morphology is observed for NaGdF₄:Eu³⁺ (P7) doped nanocrystals (Fig. 3b). Though the majority of the nanocrystals look spherical, some bunches of cylindrical shape nanostructures are noticed (Fig. 3b). A similar observation is made for NaDyF₄ (P9) nanocrystals (Fig. 3c). PXRD analyses show that for both NaGdF4:-Eu³⁺ and NaDyF₄ the hexagonal modification is the majority phase and the cubic the minority. For NaYF4 nanocrystals doped with Eu³⁺ ions, spherical nanoparticles are obtained (Fig. 3d). Though the crystal phase is same as that of NaYF4 and ionic radius of Eu³⁺ is comparable with Y³⁺, NaErF₄:Yb³⁺ (P10) shows almost monodisperse, oval shaped particles with a higher crystallinity and higher particle size (16.56 nm) which nicely matches with the crystallite size obtained from the Scherrer equation (Fig. 3e). Interestingly, some cubes can be noticed among the oval shaped nanoparticles (Fig. 3e). A similar observation can be made for the NaYbF4:Er3+ (P11) nanoparticles (Fig. 3f) though particle size is little bigger.

3.4 Structural characterizations by transmission electron microscope

To understand the morphology evolution from its atomistic origin, TEM images are analyzed in detail. 10 representative examples, LaF₃:Dy³⁺ (P2); CeF₃:Tb³⁺ (P4); NdF₃ (P5); NaSmF₄ (P6); NaGdF₄:Eu³⁺ (P7); NaTbF₄:Ce³⁺ (P8); NaDyF₄ (P9); NaYF₄:Eu³⁺ (P1); NaErF₄:Yb³⁺ (P10) and NaYbF₄:Er³⁺ (P11) are taken from the three classes of fluoride materials made based on the ionic radius/size of the RE³⁺ ion.

From Fig. 4a and b, it is noticed that hexagonal benzene like particles were found which have narrow range of aspect ratio

Table 2 Lattice strain of binary and ternary rare-earth fluoride nanomaterials prepared by a solvothermal method using [C2mim]Br IL at 200 °C

S. no.	Sample	Ionic radius (Å)	Crystal phase	Average strain ($\pm 5.0\%$)	Crystallite size (nm) ^a	Lattice strain
1.	LaF ₃ :Dy ³⁺ (P2)	1.216	Trigonal	-8.1	$11.83(\pm0.2)$	Compressive
2.	$CeF_3:Tb^{3+}(P4)$	1.196	Trigonal	-3.11	$17.31(\pm 0.2)$	Compressive
3.	NdF ₃ (P5)	1.163	Trigonal	-2.01	$12.3(\pm 0.3)$	Compressive
4.	NaSmF ₄ (P6)	1.132	Hexagonal	-1.245	$19.68(\pm0.2)$	Compressive
5.	NaGdF ₄ :Eu ³⁺ (P7)	1.107	Hexagonal	-2.17	$20.99(\pm0.2)$	Compressive
6.	NaTbF ₄ (P8)	1.095	Hexagonal	-1.87	$23.09(\pm0.2)$	Compressive
7.	NaDyF ₄ (P9) (hexagonal)	1.083	Hexagonal	-0.73	$21.8(\pm 0.2)$	Compressive
8.	NaYF ₄ :Eu ³⁺ (P1)	1.075	Cubic	+0.89	$21.2(\pm 0.2)$	Tensile
9.	NaErF ₄ :Yb ³⁺ (P10)	1.062	Cubic	+0.19	$15.99(\pm0.3)$	Tensile
10.	NaYbF ₄ :Er ³⁺ (P11)	1.042	Cubic	+0.7	$22.5(\pm0.2)$	Tensile

^a Estimated deviation.

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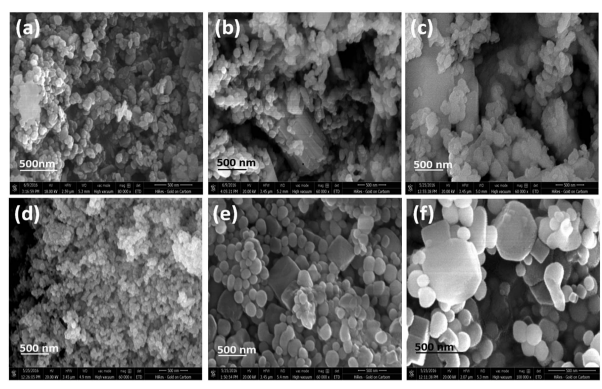


Fig. 3 Scanning electron microscope (SEM) images of (a) CeF₃:Tb³⁺ (b) NaGdF₄:Eu³⁺ (c) NaDyF₄ (d) NaYF₄:Eu³⁺ (e) NaErF₄:Yb³⁺ and (f) NaYbF₄:Er³⁺ nanoparticles prepared solvothermally in the presence of [C₂mim]Br at 200 °C.

(ca. 1.15 to 2.4). Most of the particles are isotropic (length of diagonal is same) in size. Small size particles are usually spherical whereas well matured particles have attended the highly regular hexagonal like shape. From HRTEM images, it can be seen that the growth of these particles are predominantly occurring along the (111) plane (as shown in Fig. 5a and b). And the spacing between the planes is 0.325 nm for the CeF₃. So it can be inferred that the [C₂mim]⁺ ion can serve as a capping agent based on the strong interaction with the (111) facets which favour the directional growth of the hexagonal phase. As the most exposed facets of the CeF₃:Tb³⁺ nanorods are (111) planes, the energy difference between them is minimal, facilitating the oriented attachment via twin boundaries which is noticed in the encircled area of Fig. 5b.54,55

The shape of the particles was also confirmed through the SAED images of LaF₃ and CeF₃, where single crystalline hexagonal diffraction patterns were observed (Fig. S5a and b†). Bright diffracting planes are belonging to (111), (112), (113), (300), (311) and (111), (112), (113), (222) planes of the LaF₃ and CeF₃ respectively. Thus, it is evidently confirmed that (111) plane is the dominating plane for both LaF3 and CeF3 nanoparticles (shown in Fig. S5a and b†). Furthermore, a' b' and b' c' planes are equally spaced (3.16 nm⁻¹) and edge length of the drawn benzene like shape is 5.51 nm⁻¹ in CeF₃:Tb³⁺ single crystal (see Fig. S5b†). However, in the case of NdF₃ significant changes in the shape were noticed (Fig. 4c). Here mixed (hexagonal, spherical and few tubular) morphology is found with wide range of particle size. Aspect ratio (1.020-8.55) of these particles is abruptly increased than the LaF3 and CeF3 nanoparticles.

Analysis reveals that regularity of hexagonal structure is gradually decreased from La to Nd. This is attributed to the contraction of crystal lattice volume of binary rare earth fluorides with decreasing the ionic radius of RE3+ ions. Consequently, anisotropic growth of the nanoparticle would be occurred in distorted hexagonal/trigonal unit cell. Therefore regularity of hexagonal disc is gradually being transformed to distorted hexagonal form.

However, in ternary hexagonal NaREF4 (Sm to Tb), shape of the particles is drastically changed into irregular, branchedtubular shape with higher aspect ratio (ca. 1.14-9.8) that is much higher than the previous binary fluorides i.e. NdF₃ (as shown in Fig. 4c and d). As can be seen from the TEM images, dispersibility of these particles is gradually decreasing from NaSmF₄ to NaTbF₄:Ce³⁺. In the case of NaSmF₄ obtained nanoparticles are highly dispersed whereas in the case of NaTbF₄ particles are agglomerated. Here, (101) plane is highly exposed plane with spacing of 0.298 nm for the hexagonal NaGdF₄:Eu³⁺ nanoparticles (see Fig. 5c). From low magnification TEM images, it can be prominently seen that the however, in HRTEM image of the NaDyF₄, both (111) [cubic] and (101) [hexagonal] planes are appearing (Fig. 5d). The calculated spacing (0.315 nm) between the lattice fringes is belonging to the (111) plane of the cubic NaDyF₄ nanoparticles. SAED image of the same sample indicates that it is polycrystalline in nature and diffracting planes correspond to the (112), (300) and (311) planes (see in Fig. S5c and d†). In present cases, distribution of sodium and RE³⁺ is completely different than that of binary rare earth fluorides. Hexagonal unit cell of ternary rare-earth

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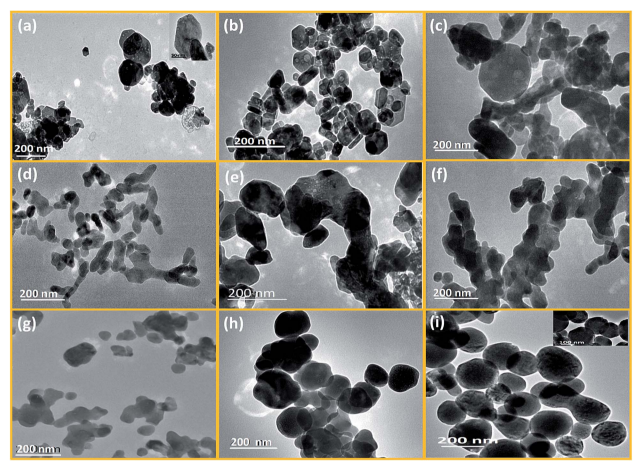


Fig. 4 Low magnification TEM images of $LaF_3:Dy^{3+}$, $CeF_3:Tb^{3+}$, NdF_3 (a-c) $NaSmF_4$, $NaGdF_4:Eu^{3+}$, $NaTbF_4:Ce^{3+}$ (d-f) and $NaYF_4:Eu^{3+}$, $NaF_4:Eu^{3+}$, $NaF_4:Eu^{3$ NaErF₄:Yb³⁺, NaYbF₄:Er³⁺ (g-i) nanocrystals prepared solvothermally at 200 °C.

fluorides are contracting gradually due to contraction of interatomic bond length and causes anisotropic growth leading to the formation of tubular structure. Furthermore, on moving from Sm to Tb, dispersibility of tubular nanoparticles is also decreasing (as shown in Fig. 4d-f), which is attributed to the increased surface energy. In period, charge density (e/r) gradually increases with decreasing the ionic radius of RE³⁺. Consequently, surface energy increases and agglomeration of particles is taken place. On the other hand, mixed phase is obtained in NaDyF₄. Since cubic phase is appearing along with hexagonal phase so irregular spherical as well as tubular both types morphology are obtained (Fig. S3†). In other words, it can be said that Dy3+ is the conduit ion between hexagonal and cubic phase of ternary rare earth fluorides.

In the case of small size RE³⁺ ions (1.075–1.042 Å), nicely dispersed, relatively more spherical-shaped nanoparticles are obtained (Fig. 4g-i). This group of RE³⁺ ions forms cubic ternary rare-earth fluoride nanoparticles under the similar experimental conditions. Significant changes are noticed in the shape of the cubic phase NaYF4 to NaYbF4 nanoparticles. Aspect ratio of these particles is gradually decreasing to the unity and the shape is getting increasingly spherical. It was further confirmed from the HRTEM images, that in NaErF₄:Yb³⁺ major planes are (111) and (200), while (111) is the only dominating plane in case

of NaYbF₄:Er³⁺ (see Fig. 5e and f). Besides this, dispersity of the nanoparticles is also increasing along with shape from NaYF₄ to NaYbF₄. From the diffraction patterns, it is evident that asprepared nanoparticles are polycrystalline and planes can be assigned to (222), (422), (400) and (311), (222), (331) for NaErF₄:Yb³⁺ and NaYbF₄:Er³⁺ nanoparticles respectively (Fig. S5e and f†).

In cubic phase all lattice parameters (a = b = c) of the unit cell are same. So the growth of the as-prepared nanoparticles would be isotropic. Promptness of spherical shape is dependent on the isotropic nature of cubic phase. For example, relatively large ion is forming irregular spherical nanoparticles (for example NaYF₄) while more spherical nanoparticles are found in NaErF4:Yb and NaYbF4:Er. And the average size of nanoparticles are ca. 55.3 nm, 104.95 nm and 151.63 nm for NaYF4:Eu, NaErF4:Yb and NaYbF4:Er respectively. From Fig. S4,† in histogram it can be nicely seen that size distribution of the particles are very much comparable to the calculated average size of the nanoparticles. Moreover, narrow range of size distribution is observed for the NaYF4:Eu nanoparticles whereas wide range of size distribution is occurred in case of NaYbF₄:Er (Fig. S4†). Thus from TEM images it can be said that isotropic nature of cubic phase is increasing with decreasing the ionic radius of RE³⁺ ions in period. Thus from all observations it Paper

Fig. 5 HRTEM images of LaF₃:Dy³⁺, CeF₃:Tb³⁺ (a and b) NaGdF₄:Eu³⁺, NaDyF₄ (c and d) and NaErF₄:Yb³⁺, NaYbF₄:Er³⁺ (e and f) nanocrystals prepared solvothermally at 200 °C.

can be inferred that RE³⁺ ions in period not only governs the phase and type of products but also controlled morphology of the as-obtained products under similar reaction conditions (Fig. 6).

3.5 Optical characterizations

Fig. 7 and S7† depicts the photoluminescence (PL) spectra of eight representative compounds, cubic NaYF₄:Eu³⁺ (P1),

hexagonal NaGdF₄:Eu³⁺ (P7), hexagonal CeF₃ (P3), CeF₃:Tb³⁺ (P4), LaF₃:Dy³⁺ (P2), NaSmF₄ (P6), NaTbF₄:Ce (P8) and NaDyF₄ (P9) respectively. Fig. 7a shows the excitation spectra of 1 mol% Eu³⁺ doped NaGdF₄ nanocrystals measured at room temperature, monitoring the $^5D_0-^7F_2$ electric dipole transition of the Eu³⁺ ion at 615 nm. Narrow transition lines of Gd³⁺ at 272 nm ($^8S_{7/2}-^6I_J$) and 310 nm ($^8S_{7/2}-^6P_J$) along the characteristic ones of Eu³⁺ especially at 393 nm appear. Fig. 7b shows the PL spectrum of hexagonal NaGdF₄:Eu³⁺ nanocrystals obtained upon 393 nm

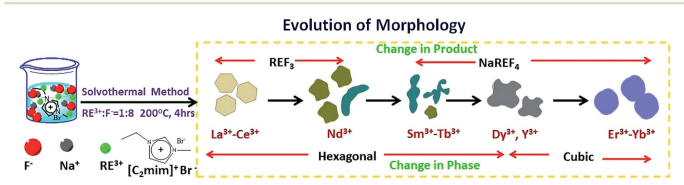


Fig. 6 Schematic presentation of trend of morphology evolution of binary/ternary rare-earth fluorides along the period of lanthanide series.

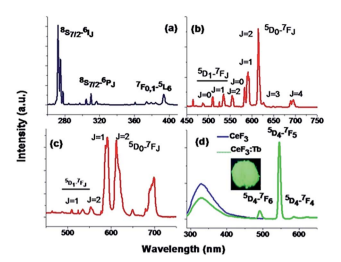


Fig. 7 (a) Excitation spectrum of NaGdF₄:Eu³⁺ nanocrystals ($\lambda_{em} = 615$ nm); (b) emission spectra of NaGdF₄:Eu³⁺; and (c) NaYF₄:Eu³⁺ nanocrystals measured upon excitation at 393 nm (d) CeF₃; as well as CeF₃:Tb³⁺ nanocrystals measured upon excitation at 270 nm.

excitation. It shows the electric dipole transition arises due to ⁵D₀-⁷F₂ at 615 nm which is stronger than the magnetic dipole transitions arise at 591 nm. The asymmetry parameter (Ω_2) using Judd-Ofelt theory was calculated for better understanding [for details see ESI \dagger]. 56,57 Calculated Ω_2 value for the above mentioned sample is $11.6 \times 10^{-20} \text{ cm}^2$. When Eu³⁺ ions doped in cubic NaYF₄ under excitation at 393 nm, the magnetic dipole transition shows stronger emission than the electric dipole transition. This indicates that Eu³⁺ ion is staying in more symmetric environment compared to the hexagonal NaGdF4:-Eu³⁺ sample. This is further confirmed by the lower asymmetry parameter Ω_2 value of 3.75 \times 10⁻²⁰ cm² for cubic Eu³⁺ ion doped NaYF4 nanoparticles. Thus the Eu3+ ion can be used as a spectroscopic probe to monitor phase transition. Another important point which is observed is the presence of the transitions from higher energy levels like ⁵D₁, ⁵D₂, ⁵D₃ etc. in both the cubic NaYF₄ and hexagonal NaGdF₄ doped with Eu³⁺. This hints that these materials can be useful for quantum cutting down conversion purposes which has potential use in energy saving lighting devoid of mercury. 14,50 Fig. S7a† shows the emission spectrum of LaF₃:Dy³⁺ (1%) under 348 nm excitation. The emission peaks appear at 475 nm and 571 nm and are assigned to ${}^4F_{9/2}$ – ${}^6H_{15/2}$ and ${}^4F_{9/2}$ – ${}^6H_{13/2}$ transitions respectively. When NaDyF4 nanoparticles are excited under 348 nm excitation, emission peak at 469 nm appears which is due to ${}^4F_{9/}$ $_2$ - 6 H $_{15/2}$ transition (Fig. S7d†). In the case of NaSmF $_4$, very weak emission at 600 nm is found on exciting the nanoparticles under 365 nm. This transition corresponds to ${}^4G_{5/2}$ of NaSmF₄ (as shown in Fig. S7b†).

Fig. 7d shows the emission spectra of CeF_3 and CeF_3 : Tb^{3+} nanocrystals dispersed in dimethyl sulphoxide (DMSO) solution and excited into the 4f–5d absorption band of Ce^{3+} . A broad emission band ranging from 300 nm to 400 nm with a maximum of 330 nm is observed for CeF_3 . Parity allowed transition of the lowest component of the 2D state to the spinorbit components of the ground state $^2F_{5/2,7/2}$ of Ce^{3+} ion is the

cause of the origin of this particular emission band. A strong excitation peak near 270 nm monitored with the emission wavelength of 330 nm corresponds to the transitions from the ground state ${}^2F_{5/2}$ of Ce^{3+} to the different components of the excited Ce³⁺ 5d states (Fig. S6†).⁵⁸ During the excitation of CeF₃:Tb³⁺ sample (Fig. 7d) a significant decrease in PL intensities of Ce³⁺ ion occurs and the appearance of new peaks at 490, 545 and 586 happens. Appearance of these new peaks is due to the transitions of ${}^5D_4 - {}^7F_{6,5,4}$ for Tb^{3+} ion. Here, Ce^{3+} ions are excited first and then energy transfer took place from Ce3+ to Tb³⁺ ions which non-radiatively decay to the ⁵D₄ excited level of Tb³⁺ ions. Finally radiative decay happens from this level to the various underlying levels of ${}^{5}F_{I=0-6}$ of Tb³⁺ ions. Incidentally the energy levels of Tb3+ ions are suitable for an energy transfer taking place from the Ce³⁺ ion upon excitation with an UV source. When CeF₃:Tb³⁺ powder samples are irradiated with UV light, an intense green colour emission which is due to the magnetic dipole transition (${}^{5}D_{4} - {}^{7}F_{5}$) with $\Delta J = \pm 1$ of at 545 nm which can be observed with the bare eye (inset of Fig. 7d). The energy transfer efficiency (η_{ET}) from donor (Ce^{3+}) to acceptor (Tb³⁺) is calculated using the equation $\eta_{\rm ET} = 1 - I_{\rm d}/I_{\rm d0}$, where $I_{\rm d0}$ and I_d are the luminescence emission intensities of donor in the absence and presence of acceptor (Tb3+) respectively. 40.80% energy transfer efficiency ($\eta_{\rm ET}$) is obtained in DMSO solution. However similar kind of energy transfer from Ce³⁺ to Tb³⁺ occurs in case of NaTbF4:Ce3+ (P8) nanocrystal and presented in Fig. S7c.†

4. Conclusions

In summary, we have succeeded in preparing phase and morphology controlled RE3+ doped binary (LnF3) and ternary rare-earth fluoride (NaREF4) using an IL-based solvothermal method. Crystal phase, lattice strain, morphology and most importantly nature of fluorides (binary/ternary) are tuned by judiciously selecting the size of the RE³⁺ ion, keeping the other reaction parameters same. To the best of our knowledge, the phase and morphology tuning on the basis of ionic radius/size of the chosen RE3+ ion are not reported yet. When size of the RE3+ ions are small like for Y3+ ions or for Er3+ and Yb3+, the reaction rate is quite faster and kinetically controlled cubic polymorph of NaREF₄ (RE = Y^{3+} , Er³⁺ and Yb³⁺) with spherical morphology is obtained. A drastic change in morphology as well as crystal phase is noticed for NaREF₄ with RE = Sm³⁺, Eu³⁺, Tb³⁺, Gd³⁺. Here the sizes of the RE³⁺ ions are in intermediate size and the formation of the thermodynamically stable hexagonal phase occurs. When the same reaction technique is attempted for La³⁺, Ce³⁺ and Nd³⁺, instead of the ternary fluoride, hexagon-shaped particles of the binary fluoride are obtained. This is due to the quite larger size of the RE³⁺ ion which plays a pivotal role in kinetics. HRTEM analysis shows that the [C₂mim]⁺ cation attaches to the (111) plane and favor the directional growth of the hexagonal phase LnF₃. Analysis reveals that lattice strain can be crystal phase dependent. Both the trigonal binary and hexagonal ternary fluoride shows compressive strain whereas cubic ternary fluoride shows tensile strain. Strong emission peaks, higher energy transition of Eu³⁺

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ion for NaYF₄:Eu³⁺ and NaGdF₄:Eu³⁺ ion and efficient energy transfer from Ce³⁺ to Tb³⁺ shows promises in different optoelectronic applications of prepared lanthanide doped materials. Last but not the least, this methodology based on the ionic radius or size of the ions can be applied for any other series of fluorides in the periodic table.

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Notes and references

- 1 F. Wang and X. Liu, Chem. Soc. Rev., 2009, 38, 976.
- 2 J.-C. Bünzli, Chem. Rev., 2010, 110, 2729.
- 3 C. Bouzigues, T. Gacoin and A. Alexandrou, ACS Nano, 2011, 11, 8488.
- 4 G. Chen, T. Y. Ohulchanskyy, S. Liu, W.-C. Law, F. Wu, M. Swihart, H. Agren and P. N. Prasad, ACS Nano, 2012, 6,
- 5 D. A. Chengelis, A. M. Yingling, P. D. Badger, C. M. Shade and S. Petoud, J. Am. Chem. Soc., 2005, 127, 16752.
- 6 E. C. Ximendes, W. Q. Santos, U. Rocha, U. K. Kagola, F. S-Rodríguez, N. Fernández, A. da S. G-Neto, D. Bravo, A. M. Domingo, B. del Rosal, C. D. S. Brites, L. D. Carlos, D. Jaque and C. Jacinto, Nano Lett., 2016, 16, 1695.
- 7 J. M. Meruga, A. Baride, W. Cross, J. J. Kellar and P. Stanley May, J. Mater. Chem. C, 2014, 2, 2221.
- 8 M. Pedroni, F. Piccinelli, T. Passuello, M. Giarola, G. Mariotto, S. Polizzi, M. Bettinelli and A. Speghini, Nanoscale, 2011, 3, 1456.
- 9 Z. Zhang, X. Ma, Z. Geng, K. Wang and Z. Wang, RSC Adv., 2015, 5, 33999.
- 10 S. Kasturi, V. Sivakumar and D. Y. Jeon, Luminescence, 2016, 31, 1138.
- 11 N. Mauser, D. Piatkowski, T. Mancabelli, M. Nyk, S. Mackowski and A. Hartschuh, ACS Nano, 2015, 9, 3617.
- 12 C. Wang, L. Cheng, Y. Liu, X. Wang, X. Ma, Z. Deng, Y. Li and Z. Liu, Adv. Funct. Mater., 2013, 23, 3077.
- 13 K. Krämer, D. Biner, G. Frei, H. U. Güdel, M. P. Hehlen and S. R. Lüthi, Chem. Mater., 2004, 16, 1244.
- 14 P. Ghosh, S. Tang and A.-V. Mudring, J. Mater. Chem., 2011, 21,8640.
- 15 P. Rahman and M. Green, Nanoscale, 2009, 1, 214.
- 16 S. Zeng, G. Ren, C. Xu and Q. Yang, CrystEngComm, 2011, 13,
- 17 D. Yang, Y. Dai, P. Ma, X. Kang, M. Shang, Z. Cheng, C. Li and J. Lin, J. Mater. Chem., 2012, 22, 20618.

- 18 K. A. Abel, J.-C. Boyer and F. C. J. M. van Veggel, J. Am. Chem. Soc., 2009, 131, 14644.
- 19 H. Wang and T. Nann, ACS Nano, 2009, 3, 3804.
- 20 J.-C. Boyer, F. Vetrone, L. A. Cuccia and J. A. Capobianco, J. Am. Chem. Soc., 2006, 128, 7444.
- 21 F. Wang, Y. Han, C. S. Lim, Y. Lu, J. Wang, J. Xu, H. Chen, C. Zhang, M. Hong and X. Liu, Nature, 2010, 463, 1061.
- 22 R. K. Sharma, A.-V. Mudring and P. Ghosh, J. Lumin., 2017, 189, 44.
- 23 X. Sun, Y.-W. Zhang, Y.-P. Du, Z.-G. Yan, R. Si, L.-P. You and C.-H. Yan, Chem.-Eur. J., 2007, 13, 2320.
- 24 S. N. Achary and A. K. Tyagi, Mater. Res. Bull., 2001, 36, 1441.
- 25 I. M. Ranieri, S. L. Baldochi and D. Klimm, J. Solid State Chem., 2008, 181, 1070.
- 26 P. Ptacek, H. Schäfer, K. Kömpe and M. Haase, Adv. Funct. Mater., 2007, 17, 3843.
- 27 D. Chen, P. Huang, Y. Yu, F. Huang, A. Yang and Y. Wang, Chem. Commun., 2011, 47, 5801.
- 28 Q. Zhang and Y. Bing, Chem. Commun., 2011, 47, 5867.
- 29 P. Ghosh and A. Patra, J. Phys. Chem. C, 2008, 112, 19823.
- 30 X. Zhai, S. Liu, Y. Zhang, G. Qin and W. Qin, J. Mater. Chem. C, 2014, 2, 2037.
- 31 A. Aebischer, M. Hostettler, J. Hauser, K. Krämer, T. Weber, H. U. Güdel and H.-B. Bürgi, Angew. Chem., Int. Ed., 2006, 45, 2802.
- 32 M. Wang, Q.-L. Huang, H.-X. Zhong, X.-T. Chen, Z.-L. Xue and X.-Z. You, Cryst. Growth Des., 2007, 7, 2106.
- 33 G. Xiang, J. Zhang, Z. Hao, X. Zhang, Y. Luo, S. Lü and H. Zhao, CrystEngComm, 2014, 16, 2499.
- 34 M. Pang, X. Zhai, J. Feng, S. Song, R. Deng, Z. Wang, S. Yao, X. Ge and H. Zhang, Dalton Trans., 2014, 43, 10202.
- 35 W. Zhang, F. Ding and Y. S. Chou, Adv. Mater., 2012, 24, OP236.
- 36 N. Niu, F. He, S. Huang, S. Gai, X. Zhang and P. Yang, RSC Adv., 2012, 2, 10337.
- 37 H.-X. Mai, Y.-W. Zhang, R. Si, Z.-G. Yan, L.-D. Sun, L.-P. You and C.-H. Yan, J. Am. Chem. Soc., 2006, 128, 6426.
- 38 Y. Liu, D. Tu, H. Zhu, R. Li, W. Luo and X. Chen, Adv. Mater., 2010, 22, 3266.
- 39 L. Wang, R. Yan, Z. Huo, L. Wang, J. Zeng, J. Bao, X. Wang, Q. Peng and Y. Li, Angew. Chem., Int. Ed., 2005, 44, 6054.
- 40 X. Chen, D. Peng, Q. Jua and F. Wang, Chem. Soc. Rev., 2015, 44, 1318.
- 41 G. Chen, H. Ågren, T. Y. Ohulchanskyy and P. N. Prasad, Chem. Soc. Rev., 2015, 44, 1680.
- 42 G.-S. Yi and G.-M. Chow, Chem. Mater., 2007, 19, 341.
- 43 A. Grzechnik, P. Bouvier, W. A. Crichton, L. Farina and J. Köhler, Solid State Sci., 2002, 4, 895.
- 44 D. Yang, Y. Dai, P. Ma, X. Kang, M. Shang, Z. Cheng, C. Li and J. Lin, J. Mater. Chem., 2012, 22, 20618.
- 45 Y. Zhang, X. Li, Z. Hou and J. Lin, Nanoscale, 2014, 6, 6763.
- 46 Y. Zhang, X. Li, X. Kang, Z. Hou and J. Lin, Phys. Chem. Chem. Phys., 2014, 16, 10779.
- 47 R. D. Shannon, Acta Crystallogr., Sect. A: Cryst. Phys., Diffr., Theor. Gen. Crystallogr., 1976, 32, 751.

- 48 R. K. Sharma, Y. N. Chouryal, S. Chaudhari, J. Saravanakumar, S. R. Dey and P. Ghosh, *ACS Appl. Mater. Interfaces*, 2017, **9**, 11651.
- 49 E. R. Parnham, A. M. Z. Slawin and R. E. Morris, *J. Solid State Chem.*, 2007, **180**, 49.
- 50 P. Ghosh and A.-V. Mudring, Nanoscale, 2016, 8, 8160.
- 51 G. K. Williamson and W. H. Hall, Acta Metall., 1953, 1, 22.
- 52 P. Ghosh and A. Patra, J. Phys. Chem. C, 2007, 111, 7004.
- 53 A. M. Smith, A. M. Mohs and S. Nie, *Nat. Nanotechnol.*, 2009, 4, 56–63.
- 54 R. L. Penn and J. F. Banfield, Science, 1998, 281, 969.
- 55 R. L. Penn and J. F. Banfield, Am. Mineral., 1999, 84, 871.
- 56 B. R. Judd, Phys. Rev., 1962, 127, 750.
- 57 G. S. Ofelt, J. Chem. Phys., 1962, 37, 511.
- 58 P. Ghosh, A. Kar and A. Patra, Nanoscale, 2010, 2, 1196.