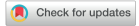
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quasi-equiaxed REPO<sub>4</sub> nanocrystals (RE = La-Dy)

Breaking the strong 1D growth habit to yield

via solvothermal reaction and investigation of

Quasi-equiaxed REPO<sub>4</sub> (RE = La-Dy) nanocrystals of either monoclinic (m-) or hexagonal (h-) structure, depending on the type of RE<sup>3+</sup>, have been successfully synthesized through solvothermal reaction in the presence of ethylene glycol (EG). Detailed characterization of the products was achieved by the combined XRD, FE-SEM, TEM, FTIR, TG/DTA, PLE/PL, and fluorescence decay techniques. The intrinsically strong 1D growth of the phosphates was gradually suppressed through increasing the solution pH and EG dosage, and the formation mechanism was elucidated with h-GdPO<sub>4</sub> as a representative. The differences in the crystal structure, IR response and crystal size of the as-synthesized REPO<sub>4</sub> crystals were interpreted in detail. It was shown that the phase purity and crystal morphology of h-GdPO<sub>4</sub> can be well retained up to 700 °C, followed by a complete transition from the hexagonal to monoclinic phase at 900 °C. The photoluminescence properties of Dy3+-doped h-GdPO<sub>4</sub> nanocrystals, including excitation, emission and fluorescence decay, were thoroughly investigated, and almost pure white-light luminescence was attained with similarly strong yellow (573 nm,  ${}^4F_{9/2} \rightarrow {}^6H_{13/2}$ ) and blue (478 nm,  ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$ ) emissions of Dy<sup>3+</sup> under excitation of the host Gd3+ ions at ~274 nm.

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### 1. Introduction

Rare-earth (RE) orthophosphates (REPO<sub>4</sub>) are attracting wide attention due to their superior advantages such as high thermal stability (melting point ~2300 °C), extremely low water solubility (solubility product  $\sim 10^{-25}$  to  $10^{-27}$ ), and high concentration of doped lasing ions ( $\sim 1.8 \times 10^{21}$  ions per cm<sup>3</sup>).<sup>3</sup> In addition, the unique electronic, optical, magnetic, and catalytic properties of RE elements endow REPO<sub>4</sub> with multiple functionalities. 4,5 As inorganic nanocrystals frequently exhibit many size- and morphology-associated novel properties,6 morphology controllable synthesis of REPO4 has attracted extensive interest, and it embodied great significance in many domains such as lighting, biolabeling, phototherapy, and optical imaging.<sup>7</sup> Till now, REPO<sub>4</sub> nanostructures with a variety of configurations have been successfully fabricated by a number of techniques, including sol-gel,8 the sacrificial template method,9 microemulsion synthesis,10 microwave-assisted crystallization, 11 and so on. Among the developed synthetic strategies, hydro-/solvothermal reaction showed outstanding capacity since it not only directly produces well-crystallized products within a short period of reaction but also allows versatile interior reaction conditions. 12,13 Fang et al., for example, hydrothermally synthesized REPO<sub>4</sub> nanowires (RE = La-Dy) and ill-shaped nanoparticles (RE = Ho-Lu and Y) without the use of any additive.14 Through applying oleic acid (OA) as a surfactant, Huo et al. solvothermally synthesized REPO<sub>4</sub> crystals with multiple morphologies of nanowires, nanorods, and nanoparticles with cuboid and hexagonal shapes in an ethanol/water mixture. 15

It is well documented that REPO<sub>4</sub> may adopt three types of crystal structures depending on the effective ionic radius of RE. Generally, larger La-Tb cations prefer to coordinate with 9 oxygen ions (CN = 9; CN: coordination number) to

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crystallize in the stable monoclinic phase (m-REPO<sub>4</sub>) while smaller Dy-Lu and Y cations tend to crystallize in the stable tetragonal phase (t-REPO<sub>4</sub>, CN = 8). 16,17 Besides, the metastable hexagonal phase (h-REPO<sub>4</sub>, CN = 8) may be adopted by the whole lanthanide spectrum and Y under appropriate conditions. 14,15 Owing to the relatively lower activation energy of growth along the [001] direction (c-axis), m-/h-REPO<sub>4</sub> crystals always present strong 1D growth preference under wet chemical synthesis conditions. 15,18,19 Though a lower solution pH value sometimes proved to be beneficial to 1D growth, 20-22 a systematic account of the role of pH in the phase/shape evolution of REPO<sub>4</sub> can hardly be found.

The intrinsic thermodynamic growth pattern of a crystal is not always likely to be the crucial factor in determining the shape evolution. A variety of strategies, such as manipulation of monomer concentration, 23,24 the use of appropriate capping agents, 25 and control over the anion-exchange rate, 7 have been extensively examined and found to be useful in regulating the growth habit of crystals under hydrothermal conditions. It is worth mentioning that the solubility of reaction products and the reactivity and diffusion behaviors of reactants significantly vary with the type of reaction medium (solvent).26 Accordingly, it is possible to exert precise control over the nucleation and growth patterns of crystals for various materials through solvothermal reaction in an appropriate solvent. You et al., for example, synthesized YBO3 and YVO<sub>4</sub> microstructures via an EG-assisted hydrothermal route, and the effects of the reaction time, pH value, and amount of ethylene glycol on the properties of the products were studied. 27,28 As for REPO<sub>4</sub>, Becerro et al. synthesized intersecting lance-shaped m-GdPO4 nanoparticles with an ethylene glycol/ water mixture as the solvent.29 Despite these successes, breaking the stubborn 1D growth habit of m-/h-REPO4 to produce equiaxed nanocrystals via solvothermal reaction is still hard to reach.

Aside from Eu<sup>3+</sup>, Dy<sup>3+</sup> is a well-known activator frequently used to probe the residential crystal environment of activator because of its hypersensitive yellow emission ( ${}^{4}F_{9/2} \rightarrow {}^{6}H_{13/2}$ transition, ~575 nm).<sup>30</sup> Furthermore, suitably adjusting the intensity ratio of its yellow to blue ( ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$ ,  $\sim$ 480 nm) emissions may realize white luminescence of a phosphor solely doped with Dy3+. By applying solvothermal reaction in an EG/H<sub>2</sub>O mixture and controlling precisely the pH value, the significant 1D growth of REPO<sub>4</sub> (RE = La-Dy) was blocked and quasi-equiaxed nanocrystals of m-/h-REPO<sub>4</sub> (RE = La-Dy) have been successfully synthesized in this work. Almost pure-white emission was also realized through Dy<sup>3+</sup> doping of h-GdPO<sub>4</sub>. In the present study, the morphology evolution of the nanocrystals was thoroughly elaborated by varying the pH value and EG content with h-GdPO4 as a representative, and the effects of lanthanide contraction and REPO<sub>4</sub> solubility on the crystal structure and morphology of REPO<sub>4</sub> were elucidated. Furthermore, the thermal behaviors of the as-synthesized h-GdPO<sub>4</sub> nanocrystals and the phase/ morphology evolution upon calcination were also investigated in detail.

# 2. Experimental

### 2.1. Solvothermal synthesis of REPO<sub>4</sub> crystals

The starting materials (Kanto Chemical Co., Inc., Tokyo, Japan) are  $RE(NO_3)_3.6H_2O$  (RE = La-Dy, >99.99% pure), HNO<sub>3</sub> (ultrahigh purity), sodium hydroxide (NaOH, >99% pure), and ethylene glycol (EG, >99.5% pure). Milli-Q filtered water (resistivity  $\sim$ 18.2 M $\Omega$  cm) was used throughout the experiment. In a typical synthesis, certain amounts of EG and H<sub>2</sub>O were mixed together to make a 60 mL transparent solvent, to which 1 mmol of RE3+ was then added. After vigorous stirring for 15 min, 1 mmol NaH2PO4 was added and the pH was adjusted to a certain value with NaOH/HNO3. The resultant mixture was constantly stirred for 30 min before being transferred to a Teflon-lined stainless steel autoclave of 100 mL capacity. The autoclave was tightly sealed and was put into an electric oven preheated to 200 °C. After 24 h of reaction, the autoclave was left to cool naturally to room temperature and the hydrothermal product was collected via centrifugation, followed by washing with water three times and ethanol once and air drying at 70 °C for 24 h. Calcination of the hydrothermal product was carried out in stagnant air, using a heating rate of 10 °C min<sup>-1</sup> and a duration time of 2 h. After annealing, the sample was cooled to 300 °C at 10 °C min<sup>-1</sup> and then naturally cooled to room temperature.

### 2.2. Characterization

Phase identification was performed via X-ray diffractometry (XRD; Model RINT2200, Rigaku, Tokyo, Japan) at 40 kV/40 mA, using nickel-filtered Cu-Kα radiation (0.15406 nm) and a scanning speed of  $4^{\circ}/2\theta$  per minute. The product morphology was analyzed by field emission scanning electron microscopy (FE-SEM, Model S-4800, Hitachi, Tokyo, Japan) operating at 10 kV and transmission electron microscopy (TEM, JEM-2100F, JEOL, Tokyo) at 200 kV. Fourier transform infrared spectroscopy (FT-IR; Model 4200, JASCO, Tokyo) was performed by the standard KBr pellet method. Thermogravimetry/ differential thermal analysis (TG/DTA; Model STA449F3, Jupiter, NETZSCH, Germany) of the as-synthesized h-GdPO4 was conducted under flowing simulated air (50 mL min<sup>-1</sup>) up to 1000 °C at a heating rate of 10 °C min<sup>-1</sup>. Photoluminescence properties were analyzed at room temperature with an FP-8500 fluorospectrophotometer (JASCO), which has a spectral resolution of ±1.5 nm. Fluorescence decay at room temperature was measured by a time-correlated single photon counting system (TCSPC), using a 370 nm nanoLED as the excitation source (1.2 ns pulse duration, Horiba Jobin Yvon, FluoroCube, Tokyo).

## Results and discussion

### 3.1. The effects of pH on the structure and morphology of GdPO<sub>4</sub>

The phase/morphology evolution of GdPO4 at 200 °C of hydrothermal reaction was firstly investigated against solution pH (Fig. 1). It is seen that hexagonal structured GdPO<sub>4</sub>

100 101 pH=1 pH=3 (a.u.) pH= ntensity pH=10 JCPDS:39-0232 Hexagonal GdPO<sub>4</sub>·nH<sub>2</sub>O JCPDS:38-1042 Hexagonal Gd(OH) 40 50 2Theta (degree)

Fig. 1 XRD patterns of the samples synthesized with different pH values indicated in the figure. The blue triangles and red dots in the pattern of the pH = 13 sample denote monoclinic GdPO<sub>4</sub> and hexagonal Gd(OH)3, respectively.

(h-GdPO<sub>4</sub>; JCPDS no. 39-0232, space group P3<sub>1</sub>21) was obtained at the lowest pH of 1,31 whose phase purity well retained up to pH = 12 though the crystallinity was gradually degraded with increasing pH. Reaction at a higher pH value of 13 led to co-crystallization of small amounts of monoclinic GdPO<sub>4</sub> (m-GdPO<sub>4</sub>; JCPDS no. 32-0386) and hexagonal Gd(OH)<sub>3</sub> (JCPDS no. 38-1042) impurities, as denoted by the blue triangles and red dots, respectively. The morphology variation with pH is displayed in Fig. 2, where it is seen that the h-GdPO<sub>4</sub> crystals all present 1D morphologies of varying aspect ratios. Fig. 2a and b show that the pH = 1 sample consists of microwires with lengths of ~2-6 µm and diameters of ~40-400 nm. The SAED analysis (the inset in Fig. 2b) of a single wire yields a set of well-defined diffraction spots that correspond to the (001), (100), and (101) planes of h-GdPO<sub>4</sub>, indicating that each of the microwires is single crystalline and well crystallized. The HR-TEM analysis of a single wire (Fig. 2c) reveals an interplanar distance of 0.60  $\pm$  0.01 nm that corresponds to the (100) plane of h-GdPO<sub>4</sub> ( $d_{100} = 0.598$ nm), which suggests the preferred growth of the wire along the [001] direction (arrowed). In addition, the (101) and (001) fringes can also be observed with spacings of 0.43  $\pm$  0.01 and  $0.63 \pm 0.01$  nm, respectively ( $d_{101} = 0.435$  nm,  $d_{001} = 0.633$ ). Although the pH = 3 sample is a mixture of micro- and nanowires (Fig. 2d), the pH = 5 product exclusively contains nanowires with lengths of  $\sim$ 90-600 nm and diameters of  $\sim$ 15-20 nm (Fig. 2e). Further increasing the pH value to 10 or 12 did not bring about any obvious change to the wire diameter but significantly shortened the wire length to ~70-200 nm (Fig. 2f-h). The SAED rings (the inset in Fig. 2h) taken from the nanowires can be assigned to the {100}, {101}, and {200} planes of h-GdPO<sub>4</sub>, and the HR-TEM lattice image (Fig. 2i) clearly reveals that the nanowires also grew up along the [001] direction though they are much thinner and shorter than the microwires.

It is known that the liberation of PO<sub>4</sub><sup>3-</sup> from NaH<sub>2</sub>PO<sub>4</sub> is highly dependent on solution pH. At the very low pH value of 1, the PO<sub>4</sub><sup>3-</sup> anions would be almost totally captured by H<sup>+</sup> to form H<sub>3</sub>PO<sub>4</sub>, which prohibited GdPO<sub>4</sub> nucleation, and thus, a clear solution of the reactants was observed before the hydrothermal reaction. As the temperature increases, a small amount of PO<sub>4</sub><sup>3-</sup> would be liberated from H<sub>3</sub>PO<sub>4</sub> to induce GdPO4 nucleation. Under this circumstance, the low nucleation density and hence the high concentrations of remaining solutes facilitated the full growth of GdPO4 crystals, resulting in the microwires with large lengths and diameters (Fig. 2a and b). Increasing the pH value to 3 largely promoted the liberation of PO<sub>4</sub><sup>3-</sup> from the hydrogen phosphate species (H<sub>2</sub>PO<sub>4</sub><sup>-</sup> and HPO<sub>4</sub><sup>2-</sup>), and thus, the precipitation of h-GdPO4 took place prior to hydrothermal reaction. Due to the low activation energy at room temperature, the resultant h-GdPO4 was poorly crystallized and appeared as severely aggregated spiculate particles, as revealed by the broadened XRD peaks and SEM observation (Fig. S1 and its inset†). The room-temperature precipitation would lower the solute concentration, which limited the crystallite growth during the hydrothermal reaction, and thus, averagely thinner microwires were yielded (Fig. 2d). The solutes tend to completely precipitate from the solution generating poorly crystallized h-GdPO<sub>4</sub> at pH ≥5, for which h-GdPO<sub>4</sub> crystals with smaller diameters and lengths were produced by hydrothermal reaction (Fig. 2e-g). It is noteworthy that, although H<sub>3</sub>PO<sub>4</sub> undergoes incomplete dissociation under acidic conditions according to  $H_3PO_4 \leftrightarrow H^+ + H_2PO_4^- \leftrightarrow 2H^+ + HPO_4^{2-} \leftrightarrow 3H^+$ +  $PO_4^{3-}$ , the stronger ability of  $PO_4^{3-}$  than  $H_2PO_4^{-}$  and  $HPO_4^{2-}$ to coordinate with RE3+ leads to the preferential formation of REPO<sub>4</sub> nuclei.<sup>32</sup> This would in turn right-shift the above equilibrium and promote the dissociation of H<sub>3</sub>PO<sub>4</sub>/H<sub>2</sub>PO<sub>4</sub><sup>-</sup>/ HPO<sub>4</sub><sup>2-</sup> species until Gd<sup>3+</sup> is completely consumed. When the pH value reached 13, the large OH concentration led to partial crystallization of hexagonal Gd(OH)3 with h-GdPO4. Owing to the stronger ability of PO<sub>4</sub><sup>3-</sup> than OH<sup>-</sup> to coordinate with RE3+, however, some Gd(OH)3 would be transformed into m-GdPO<sub>4</sub>, rather than h-GdPO<sub>4</sub>, upon hydrothermal reaction, since in the two phases the Gd3+ ions are similarly 9-fold coordinated. 33,34 This accounts for the observed phase mixture of h-Gd(OH)3, m-GdPO4, and h-GdPO4.

The intensity ratio R of (200) to (102) diffraction peaks as a function of solution pH is depicted in Fig. S2.† Combined with the results of SEM analysis, it can be recognized that the R value decreased from  $\sim 3.51$  at pH = 1 to  $\sim 2.43$  at pH = 3 along with the decrease in the overall diameter of the microwires (no obvious change in wire length). The R value then rapidly decreased to  $\sim$ 1.17 at pH = 5 with the decrease in the amount of the microwires and the significant appearance of thinner and shorter nanowires (Fig. 2e), followed by a slow decrease to  $\sim 0.73$  at pH = 12 along with shortening of the nanowires. It can therefore be deduced that the shrinking

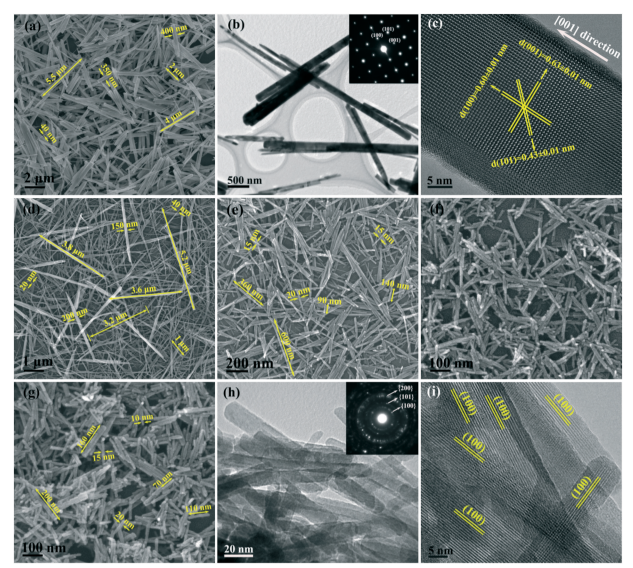


Fig. 2 FE-SEM (a and d-g) and TEM (b and h) micrographs showing the morphologies of the h-GdPO<sub>4</sub> products synthesized with different pH values of 1 (a and b), 3 (d), 5 (e), 10 (f), and 12 (g and h). The insets in (b) and (h) are the selected area electron diffraction (SAED) patterns. Parts (c) and (i) are the HR-TEM lattice images of the pH = 1 and 12 samples, respectively.

diameter, instead of length, of the wires contributed more to the decreasing R value. This is understandable in view that decreasing the wire diameter directly reduces the stacking thickness of {200} facets (Fig. 2c) and lowers the diffraction intensity of the facets.

### 3.2. The effects of EG on the phase and morphology of GdPO<sub>4</sub>

Since the 1D growth of h-GdPO<sub>4</sub> crystals was significantly restricted under alkaline hydrothermal conditions (Fig. 2), the pH value of 12 was chosen to investigate the effects of EG on the phase composition and morphology of GdPO<sub>4</sub> crystals. It is seen from Fig. 3 that the addition of EG did not change the phase purity of h-GdPO<sub>4</sub> but the intensity ratio (R) of the (200) to (102) diffraction peaks gradually increased from  $\sim$ 0.71  $\pm$  0.02 for EG = 0 and 10 mL to  $\sim$ 1.05 for EG = 50 mL ( $\sim$ 70 vol% of the total solvent), as depicted in Fig. S3.† The SEM images of the EG-assisted products are displayed in Fig. 4a-d, where it is seen that increasing EG addition gradually shortened the length of the nanowires and finally led to the crystallization of quasi-equiaxed h-GdPO<sub>4</sub> nanocrystals at EG = 30 and 50 mL (Fig. 4c and d). The TEM observation (Fig. 4e and the inset) found that the nanocrystals possess hexagonal top/bottom surfaces, corresponding well with the hexagonal crystal structure. HR-TEM analysis (Fig. 4f) of a single nanoparticle yields a set of well-resolved lattice fringes that are attributable to the (101) plane of h-GdPO<sub>4</sub>. The lattice fringes spanned the whole particle, indicating that each individual nanocrystal is single crystalline. The "diameter" (shorter side) of the products were measured and labeled in Fig. 4a-d, and the aspect ratio of the quasi-equiaxed h-GdPO<sub>4</sub> nanocrystals was assayed to be below 2. It was found that EG addition slightly yet steadily increased the crystallite diameter from  $\sim$ 10-20 nm (Fig. 2g, EG = 0 mL, R = 0.73) to 30-37 nm

EG-0 ml EG-10 ml ntensity (a.u.) EG-20 m EG-30 ml ≚ 101 JCPDS:39-0232 Hexagonal GdPO, nH, O 40 50 2Theta (degree)

Fig. 3 XRD patterns of the h-GdPO<sub>4</sub> samples synthesized with different EG contents.

(Fig. 4d, EG = 50 mL, R = 1.05), which may account for the increased R value and the gradually sharpened (200) diffraction (Fig. 3) since the (200) diffraction is more sensitive to the particle diameter as inferred from Fig. 2c. It is also interesting to notice that the quasi-equiaxed h-GdPO4 nanocrystals grew from  $\sim$ 25-35 to  $\sim$ 30-37 nm with increasing EG content from 30 to 50 mL (Fig. 4c and d).

The observed shape evolution of h-GdPO<sub>4</sub> crystals can be explained by the selective absorption of EG on specific crystal surfaces and the nucleation kinetics governed by EG. Since the 1D growth was gradually inhibited, it can be deduced that the EG molecules selectively cap the {001} planes of h-GdPO<sub>4</sub> to hinder the crystal growth along the [001] direction. 27,28,35 The intrinsically preferred [001] growth of h-GdPO<sub>4</sub> was almost completely suppressed when the EG volume was 30 mL, and thus, the quasi-equiaxed nanocrystals were obtained. Since the concentrations of Gd and phosphate sources were constant in this work, the significantly prohibited 1D growth by EG capping was expected to yield crystals of substantially larger diameters, but only a slight diameter increase was observed. This could be due to the increased nucleation density as discussed below. For chemical precipitation, the final particle size is known to be closely related to the equilibrium concentration of solutes in saturated solution (the solubility of precipitate), which can be estimated from the following equation proposed by Chen et al.:36

$$C_{\rm s} \approx \exp\{-(z^+z^-e^2)/[4\pi\varepsilon_0\varepsilon_{\rm r}kT(r^{z^+}+r^{z^-})]\}$$
 (1)

where  $r^{z^+}$  and  $r^{z^-}$  are the radii of ions charged by  $z^+$  and  $z^-$ , respectively;  $\varepsilon_0$  is the vacuum permittivity and  $\varepsilon_r$  is the relative dielectric constant of the solvent; k is the Boltzmann's constant, and T is the temperature in Kelvin. Owing to the smaller dielectric constant of EG (37.7) than water (78.5),<sup>37</sup> the solubility  $(C_s)$  of GdPO<sub>4</sub> would successively decrease with

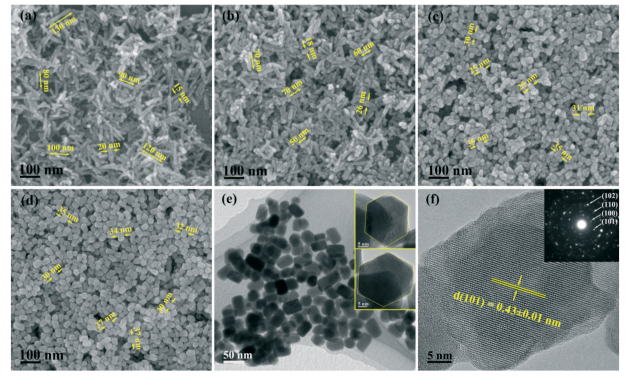


Fig. 4 FE-SEM (a-d) and TEM (e) images showing the morphologies of the h-GdPO<sub>4</sub> samples synthesized with different EG contents of 10 (a), 20 (b), 30 (c), and 50 mL (d and e). The insets in (e) show the hexagonal top/bottom surface of a nanocrystal. Part (f) and its inset are the HR-TEM lattice image and SAED pattern of the EG = 50 mL sample, respectively.

increasing EG addition, thus leading to gradually higher supersaturation for nucleation. On the other hand, the free energy of critical nucleus formation,  $\Delta G_{\text{crit}}$ , can be given by:<sup>38</sup>

$$\Delta G_{\rm crit} = 4\pi \gamma r_{\rm c}^2 / 3 \tag{2}$$

where  $\gamma$  is the liquid/solid interfacial tension and  $r_{\rm c}$  is the critical nucleus radius. EG has a lower surface energy (48.4 N m<sup>-1</sup>) than water (72.8 N m<sup>-1</sup>),<sup>37</sup> and hence, its presence would reduce the energy barrier for stable nuclei formation through lowering the  $\gamma$  term in eqn (2). The above two factors may therefore raise the nucleation density of GdPO<sub>4</sub>, and as a result, only slightly larger crystal diameter was observed despite the significantly prohibited 1D crystal growth. It should also be noted that a high enough EG content may hinder PO<sub>4</sub><sup>3-</sup> dissociation from NaH<sub>2</sub>PO<sub>4</sub> due to the poor accessibility of water molecules, which would in turn lower the nucleation density of GdPO<sub>4</sub>. This may account for the size increment of the quasi-equiaxed nanocrystals upon raising the EG content from 30 to 50 mL.

### 3.3. Extension to the synthesis of other REPO<sub>4</sub> nanocrystals

The EG-assisted approach developed for h-GdPO4 nanocrystals was implemented in the synthesis of other REPO<sub>4</sub> (RE = La-Eu, Tb, and Dy) crystals that similarly present intrinsic 1D growth preference. XRD analysis of the products (Fig. 5) shows that the phosphates of larger La<sup>3+</sup>-Sm<sup>3+</sup> cations crystallized in the monoclinic system while those of smaller Tb3+ and Dy3+, similar to Gd3+, crystallized as hexagonal phases. The Eu product is a mixture of the above two phases, as denoted by the blue and red triangles in Fig. 5, which may

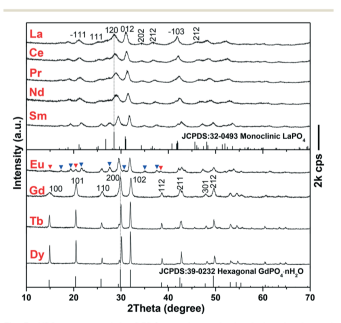


Fig. 5 XRD patterns of the REPO<sub>4</sub> products obtained *via* solvothermal reaction at 200 °C, pH = 12, and EG = 50 mL for 24 h. The blue and red triangles in the XRD pattern of the Eu product denote the monoclinic and hexagonal phases, respectively.

suggest that Eu is the boundary element for phase selection. The above results can be attributed to lanthanide contraction, that is, larger RE cations show stronger ability to coordinate with more oxygen ions and thus favor m-REPO4 crystallization (CN = 9 for m-REPO<sub>4</sub> and CN = 8 for h- and t-REPO<sub>4</sub>). 14-18 The diffraction peaks tend to shift to the highangle region from m-LaPO4 to m-SmPO4 and from h-GdPO4 to h-DyPO<sub>4</sub>, and the cell parameters of both the m- and h-REPO<sub>4</sub> crystals (derived with MDI Jade 5.0 software) linearly decrease with decreasing RE3+ size (Fig. S4†), conforming well to lanthanide contraction. Analysis of the diffraction peak broadening with the Scherrer equation yielded average crystallite sizes of  $\sim 9 \pm 3$  nm for m-REPO<sub>4</sub> (RE = La-Nd),  $\sim 15$ nm for m-SmPO<sub>4</sub>, ~36 nm for h-GdPO<sub>4</sub>, and ~60 nm for h-TbPO<sub>4</sub>.

The morphologies of m-REPO<sub>4</sub> (RE = La-Nd) were analyzed via SEM and TEM with m-LaPO4 as an example (Fig. 6a and b). It was found that m-LaPO<sub>4</sub> consists of nanoparticles with extremely tiny sizes of ~10 nm, which is in agreement with the crystallite size assayed via XRD. The SAED analysis (the inset in Fig. 6b) clearly reveals the diffraction rings assignable to the {200} and {012} planes of m-LaPO<sub>4</sub>. The HR-TEM lattice image (Fig. 6c) exhibits that the nanoparticles are structurally uniform, with the interplanar distance of  $0.32 \pm 0.01$  nm corresponding well to the (120) plane ( $d_{120} = 0.313$  nm). The m-SmPO<sub>4</sub> sample also consists of quasi-equiaxed particles/crystallites, similar to m-LaPO<sub>4</sub>, but the particle size increased to ~15-20 nm (Fig. 6d and its inset). The h-TbPO4 and h-DyPO4 products (Fig. 6e and f) consist of elongated particles with a diameter/length of ~40/ 80 and 60/200 nm, respectively. Firsching et al. reported that the solubility of REPO4 slowly increases from La to Eu and then rapidly rises to the maximum values of Tb and Dy,<sup>2</sup> which well explains the steadily increasing crystal size from La-Dy since higher solubility would lower the nucleation density and thus lead to a larger final crystal size. The elongated phosphate crystals observed in Tb and Dy, particularly Dy, may suggest that the 1D growth preference becomes stronger with decreasing RE3+size. In spite of this, the h-TbPO4 and h-DyPO<sub>4</sub> particles have small aspect ratios of only  $\sim$ 2-4, which implies that the strong 1D growth tendency has indeed been effectively inhibited by the synthetic approach employed in this work.

The FTIR analysis of the as-synthesized m-/h-REPO<sub>4</sub> products is displayed in Fig. 7. For m-REPO<sub>4</sub> (La-Sm), the PO<sub>4</sub><sup>3-</sup> ions present a broad vibration band ranging from ~500 to 1200 cm<sup>-1</sup>, with a sharp peak at ~960 cm<sup>-1</sup> attributable to the P-O symmetric stretching vibration  $(v_1)$ , the broad absorption at ~980-1200 cm<sup>-1</sup> to the P-O antisymmetric stretching  $(v_3)$ , and the splits in the ~500-700 cm<sup>-1</sup> region to the P-O bending mode  $(v_4)$ .<sup>39</sup> The absorptions at ~3438 and 1633 cm<sup>-1</sup> can be assigned to the O-H stretching vibrations  $(v_1 \text{ and } v_3)$  and H-O-H bending mode  $(v_2)$  of water, <sup>40</sup> respectively. Compared with m-REPO<sub>4</sub>, h-REPO<sub>4</sub> (Gd-Dy) exhibited lower resolutions of  $v_1$  and  $v_4$  PO<sub>4</sub>  $^{3-}$  vibrations owing to the higher symmetry of PO<sub>4</sub><sup>3-</sup> in h-REPO<sub>4</sub> (C<sub>2</sub>) than that in

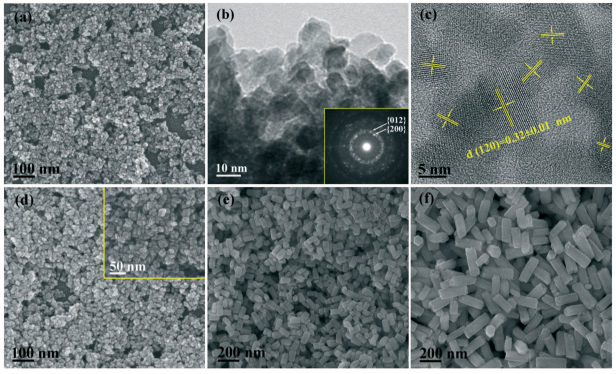


Fig. 6 FE-SEM (a and d-f) and TEM (b) images showing the morphologies of the m-/h-REPO₄ products, where RE = La (a and b), Sm (d), Tb (e) and Dy (f). Part (c) is the HR-TEM lattice image of several m-LaPO<sub>4</sub> nanocrystals. The insets in (b) and (d) are the corresponding SAED pattern and high magnification SEM micrograph, respectively.

m-REPO<sub>4</sub>  $(C_1)$ . The  $v_1$  vibration peak of PO<sub>4</sub>  $^{3-}$  was found to shift toward the higher energy (from ~950 to 975 cm<sup>-1</sup>) with decreasing RE3+ size, owing to the stronger polarizing power of smaller RE3+.42 The vibration frequency increases towards smaller RE<sup>3+</sup> for the  $v_4$  component located at ~620 cm<sup>-1</sup>, as

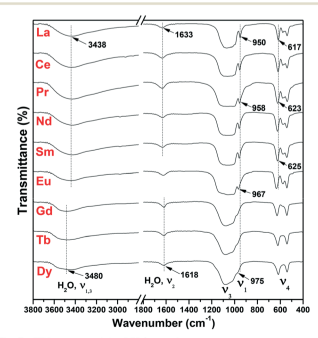


Fig. 7 FTIR spectra of the REPO<sub>4</sub> products

also observed by Ross et al., though the exact reason is yet unknown.<sup>39</sup> It is interestingly noticed that the O-H stretching vibration of H2O has larger frequency in h-REPO4 than in m-REPO<sub>4</sub> while an opposite trend is seen for the H-O-H bending mode. There are two reasons for such a phenomenon: (1) the hydration water in h-REPO<sub>4</sub> has stronger bonding force with the crystal framework than the absorbed water in m-REPO<sub>4</sub>, and thus the stretching vibration needs higher energy, and (2) the hydration effect may enable more lonepair electrons from the 3a<sub>1</sub> orbital of water to take part in chemical bonding in h-REPO<sub>4</sub>, which enlarges the H-O-H bond angle and thus lowers the frequency of bending vibration.43

### 3.4. Thermal behavior of the as-synthesized h-GdPO<sub>4</sub> nanocrystals

The TG/DTA analysis of the h-GdPO<sub>4</sub> nanocrystals (EG = 50 mL) is depicted in Fig. 8A, where it is seen that the sample loses weight via three separate steps upon heating. The first weight loss of ~2.1% up to ~180 °C, accompanied by an endotherm centered at ~63 °C, is largely due to the elimination of adsorbed water. The rapid loss of ~2.4% in the narrow temperature range of  $\sim$ 180-240 °C is ascribed to the removal of water bound to PO<sub>4</sub><sup>3-</sup> through hydrogen bonding, which produces a sharp endotherm at ~222 °C. The subsequent weight loss of ~1.1%, taking place in the wide temperature range of  $\sim$ 240–770 °C, can be attributed to the removal

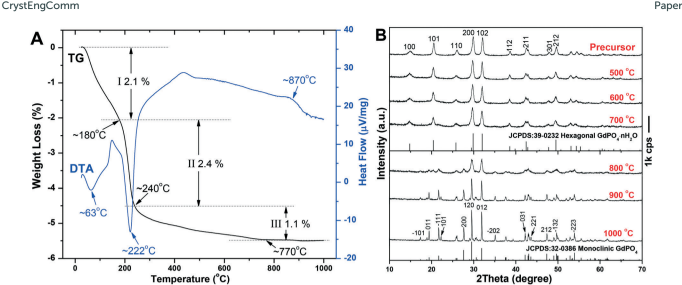


Fig. 8 TG-DTA profiles (A) and the temperature dependent phase evolution (B) of h-GdPO<sub>4</sub>.

of hydration water from the crystal bulk. The number of hydration (n) in each h-GdPO<sub>4</sub>·nH<sub>2</sub>O formula can thus be determined to be  $\sim 0.16$ . The exotherm located at  $\sim 870$  °C, without the occurrence of any weight loss, is associated with the h- to m-GdPO4 phase transition as evidenced by the XRD analysis results (Fig. 8B). It is clear that the hexagonal structure can be well retained up to 700 °C, followed by a complete transition to m-GdPO4 at 900 °C (Fig. 8B). Rietveld refinement of the XRD patterns of the 700, 800, and 900 °C products yielded well acceptable weighted profile R-factors  $(R_{\rm wp})$  in each case (Fig. S5†). It is also clear that the 700 °C (Fig. S5a†) and 900 °C (Fig. S5c†) samples are of pure h- and m-GdPO<sub>4</sub> phase, respectively, while the 800 °C one (Fig. S5b†) is a mixture of the two phases (m-GdPO<sub>4</sub> content:  $\sim$ 93.65 wt%). The results thus further confirmed the phase transition observed in Fig. 8. Calcination at a higher temperature of 1000 °C did not alter the monoclinic crystal structure but significantly enhanced the crystallinity of the product. Analysis of the XRD peak broadening with the Jade 5.0 software yielded crystallite sizes of ~26 and 79 nm for the samples calcined at 700 and 900 °C, respectively. FE-SEM observation (Fig. 9) found slight aggregation and deformation for the particles calcined at 700 °C (Fig. 9c), since the mass transfer via diffusion would bind the neighboring particles together and deform them. Calcination at 900 °C led to severe aggregation and deformation of the particles (Fig. 9d), because of the significant inter-particle sintering, crystallite growth, and re-constructive type hexagonal to monoclinic phase transition.

### 3.5. Luminescence of Dy3+ in h-GdPO4 nanocrystals

Fig. 10A shows the photoluminescence excitation (PLE) and photoluminescence (PL) spectra of h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub> (synthesized with EG = 50 mL) before and after 700 °C calcination, where it is seen that calcination raised both the excitation and emission intensities by a factor of ~750%. Such drastic improvements are largely due to the almost complete

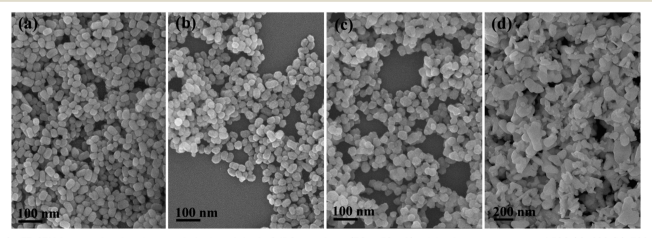


Fig. 9 FE-SEM images showing the morphologies of the h-GdPO<sub>4</sub> products (EG = 50 mL) calcined at temperatures of 500 (a), 600 (b), 700 (c), and 900 °C (d).

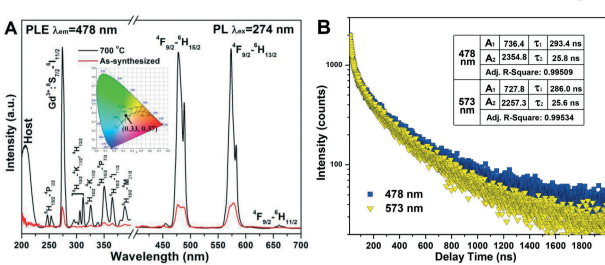


Fig. 10 Photoluminescence excitation (PLE) and emission (PL) spectra of the as-synthesized (EG = 50 mL) and 700 °C calcined h-(Gd $_{0.95}$ Dy $_{0.05}$ )PO $_4$  (A) and fluorescence decay kinetics for the 478 and 573 nm emissions of the 700 °C calcined h-(Gd $_{0.95}$ Dy $_{0.05}$ )PO $_4$  (B). The inset in (A) is the CIE chromaticity diagram of the 700 °C calcined h-(Gd $_{0.95}$ Dy $_{0.05}$ )PO $_4$  phosphor.

elimination of water molecules, which are well known to cause serious luminescence quenching, 9,44 from the surface and crystal interior of h-GdPO<sub>4</sub>, as demonstrated by the TG result (Fig. 8A). The PLE spectra ( $\lambda_{em}$  = 478 nm) exhibit several sharp peaks in the  $\sim$ 230-400 nm region which include the  $^8S_{7/2} \rightarrow ^6I_{11/2}$  transition of  $Gd^{3+}$  at  $\sim$ 274 nm and the weak f-f transitions of Dy3+ within the 4f9 configuration,45 as assigned in the figure. The peak at ~210 nm is due to host absorption. The  ${}^8S_{7/2} \rightarrow {}^6I_{11/2}$  Gd<sup>3+</sup> transition appears to be significantly stronger than any other excitation band, implying an efficient Gd<sup>3+</sup> → Dy<sup>3+</sup> energy transfer. Under UV excitation at 274 nm, two predominant emission peaks were observed to arise from the  ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$  and  ${}^4F_{9/2} \rightarrow {}^6H_{13/2}$ transitions of Dy3+ at 478 nm (blue) and 573 nm (yellow), respectively. Hardly perceptible is the following line at 660 nm associated with the  ${}^4F_{9/2} \rightarrow {}^6H_{11/2}$  Dy $^{3+}$  transition. $^{46}$  The energy transfer from Gd3+ to Dy3+ is schematically displayed in Fig. S6.† Exciting the h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub> phosphor at 274 nm would raise Gd3+ electrons from the 8S7/2 ground state to the <sup>6</sup>I<sub>11/2</sub> energy level, followed by relaxation to the <sup>6</sup>P<sub>7/2</sub> state. Resonant energy transfer from Gd3+ to Dy3+ may then take place upon back jumping of the <sup>6</sup>P<sub>7/2</sub> Gd<sup>3+</sup> electrons, through which the lowest-lying <sup>6</sup>H<sub>15/2</sub> ground-state electrons of the Dy<sup>3+</sup> activator can be excited to the  ${}^4H_{13/2}/{}^4K_{13/2}$  energy level. The <sup>4</sup>K<sub>13/2</sub>/<sup>4</sup>H<sub>13/2</sub> Dy<sup>3+</sup> electrons would relax to the <sup>4</sup>F<sub>9/2</sub> level in a non-radiative way, followed by radiative relaxation to the  $^{6}H_{15/2}$ , <sup>6</sup>H<sub>13/2</sub>, and <sup>6</sup>H<sub>11/2</sub> ground states to produce the observed blue (strong), yellow (strong), and red (very weak) emissions.

The  ${}^4F_{9/2} \rightarrow {}^6H_{13/2}$  emission is a forced electric dipole transition, whose intensity is hyper-sensitive to the crystal field around Dy<sup>3+</sup>, while the  ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$  magnetic dipole transition is not. According to the Judd–Ofelt parity law,  ${}^{47,48}$  the electric dipole transition is allowed only under the condition that the Dy<sup>3+</sup> ions occupy a low symmetry site without an inversion center.  ${}^{49}$  In h-GdPO<sub>4</sub>, Dy<sup>3+</sup> is expected to replace Gd<sup>3+</sup>

to inherit the centrosymmetric  $D_2$  point symmetry. The observed  ${}^4F_{9/2} \rightarrow {}^6H_{13/2}$  transition is, however, as strong as the  ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$  transition. This can be attributed to the high surface area of the nanocrystals arising from their very limited crystallite size ( $\sim$ 40 nm), which allows some of the Dy<sup>3+</sup> ions to be located at or near crystallite surfaces. The loss of point symmetry of these surface region Dy<sup>3+</sup> enhances  ${}^4F_{9/2} \rightarrow {}^6H_{13/2}$  while lowers the  ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$  emission at the same time. The h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub> phosphor calcined at 700  ${}^{\circ}$ C was calculated to obtain the Commission international de l'éclairage (CIE) chromaticity coordinate of (0.33, 0.37), which resides in the white region of the CIE chromaticity diagram (the inset in Fig. 10A).

Fluorescence decay analysis (Fig. 10B) of the 700  $^{\circ}$ C calcined h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub> found that both the 478 and 573 emissions can be fitted with the following second-order exponential equation:<sup>50</sup>

$$I(t) = A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2)$$
(3)

where I(t) is the luminescence intensity at time t,  $A_1$  and  $A_2$  are pre-exponential constants, and  $\tau_1$  and  $\tau_2$  stand for the decay time for the exponential components. The fitting results are summarized in the inset of Fig. 10B together with the adjusted R-squared values of fitting. The goodness of fit can be recognized from the  $R^2$  values of  $\sim 1$  in each case. The results thus indicate that the fluorescence decay can be considered as a superposition of two main components in each case, with the faster one most likely arising from the surface related Dy<sup>3+</sup> and the slower one being related to the innerpart Dy<sup>3+</sup>. The average fluorescence lifetime  $(\tau^*)$  was calculated from the following equation to be  $\sim 234.7$  and 229.4 ns for the 478 and 573 nm emissions, respectively.

$$\tau^* = (A_1 \tau_1^2 + A_2 \tau_2^2) / (A_1 \tau_1 + A_2 \tau_2) \tag{4}$$

The slightly shorter lifetime of the 573 nm emission is due to the involvement of surface-region Dy<sup>3+</sup> ions.

# 4. Conclusion

Quasi-equiaxed nanocrystals of monoclinic (m-) and hexagonal (h-) structured REPO<sub>4</sub> (RE = La-Dy) have been successsynthesized via ethylene glycol (EG)-assisted solvothermal reaction. It was concluded that EG effectively inhibits the 1D growth of REPO<sub>4</sub> by selectively capping the {001} facets and meanwhile affects the nucleation density, leading to substantial size/morphology variation of the resultant nanocrystals. Owing to lanthanide contraction, the orthophosphates of La-Sm and Gd-Dy crystallize in the monoclinic and hexagonal systems, respectively. The solubility of REPO<sub>4</sub> gradually increases from La-Dy, and as a result successively larger phosphate crystals were obtained. Calcination at 700 °C did not affect the phase purity and crystal morphology of h-GdPO4 but greatly enhanced the excitation and emission intensities of photoluminescence. The  ${}^4F_{9/2} \rightarrow {}^6H_{13/2}$ (573 nm) and  ${}^4F_{9/2} \rightarrow {}^6H_{15/2}$  (478 nm) emissions are similarly strong for h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub>, owing to the nanosize effect, though the Dy3+ activators were assumed to have centrosymmetric D<sub>2</sub> point symmetry. The h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub> phosphor has a CIE coordinate of (0.33, 0.37) under 274 nm excitation of the host Gd<sup>3+</sup> ions, which falls in the white light region. The 478 and 573 nm fluorescence of h-(Gd<sub>0.95</sub>Dy<sub>0.05</sub>)PO<sub>4</sub> both decay in a bi-exponential manner and have average lifetime values of ~234.7 and 229.4 ns, respectively.

# Conflicts of interest

There are no conflicts of interest to declare.

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