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Hybrid coordination-network-engineering for bridging cascaded channels to activate long persistent phosphorescence in the second biological window

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We present a novel “Top-down” strategy to design the long phosphorescent phosphors in the second biological transparency window via energy transfer. Inherence in this approach to material design involves an ingenious engineering for hybridizing the coordination networks of hosts, tailoring the topochemical configuration of dopants, and bridging a cascaded tunnel for transferring the persistent energy from traps, to sensitizers and then to acceptors. Another significance of this endeavour is to highlight a rational scheme for functionally important hosts and dopants, Cr/Nd co-doped $Zn_{1-x}Ca_xGa_2O_4$ solid solutions. Such solid-solution is employed as an optimized host to take advantage of its characteristic trap site level to establish an electron reservoir and network parameters for the precipitation of activators Nd^{3+} and Cr^{3+} . The results reveal that the strategy employed here has the great potential, as well as opens new opportunities for future new-wavelength, NIR phosphorescent phosphors fabrication with many potential multifunctional bio-imaging applications.

There is an increasing interest in the use of long persistent phosphorescence in the biologically transparent window to drive the photonic bioprobe for tracing the cancer cells¹. Long phosphorescent phosphors (LPPs) can help avoiding the challenging requirement of high-intensity illumination during the signal collection, which often leads to decreased signal-to-noise ratio and photon-induced deterioration of analytes². This emerging research trend, which incorporates various fields of materials science, biology, chemistry, engineering, physics and pharmaceuticals, follows two main directions: operation waveband and persistent duration, with many relevant crossing points in between^{3,4}. As we know, there are two biologically transparent windows: first one at 650–950 nm and second one at 1000–1350 nm⁵; Near-infrared (NIR) light in the first transparency window can penetrate biological tissues such as skin and blood more efficiently than visible light⁶, yet the second region has even lower absorption and scattering therefore offers more efficient tissue penetration⁷. However, the main researches about the operational waveband of NIR LPPs mainly focus on the short wavelength region, i.e. first NIR window.

In addition to altering the emission center and tailoring the crystal field surrounding the activator, another useful strategy to extend the operational waveband, is to transfer the persistent energy of sensitizers to acceptors⁸. In fact, although the afterglow properties are predominantly controlled by the active traps, more subtle effects, such as topochemical coordination-configuration of dopant ions, can also have a profound role to the spectroscopic features of LPPs, which has long been recognized as a significant issue lying at the heart of doping chemistry and photoluminescent theory⁹. Considering the advanced engineering of cascaded tunnel of energy transfer

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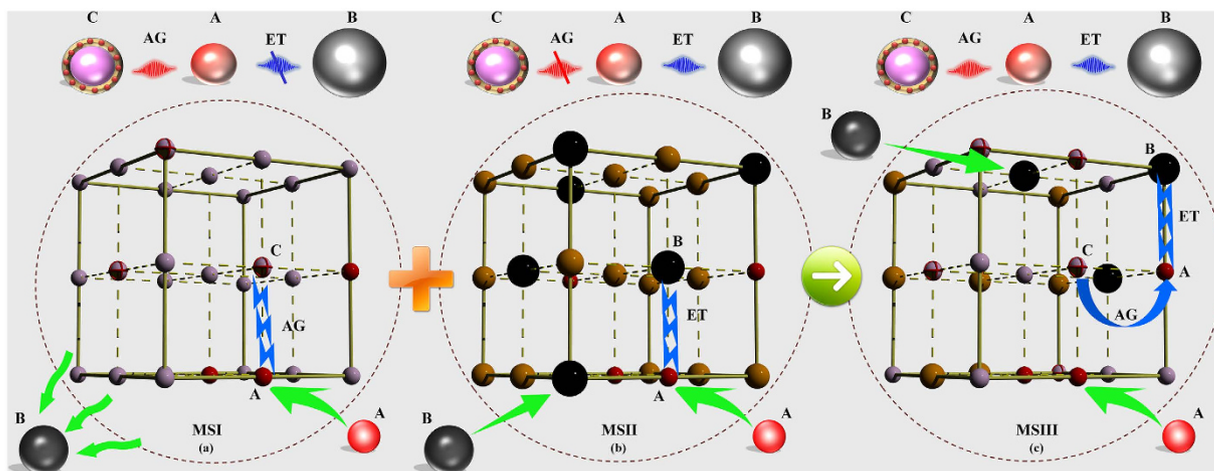


Figure 1. Schematic illustration showing the influence of ions doping-pattern and network-structural motifs on energy transfer process between traps and dopants. (a,b) represent the typical trapping and de-trapping process (AG) (traps (C) \rightarrow activator (A)), as well as energy transfer process (ET) (activator (A) \rightarrow activator (B)) in different material system (MSI and MSII), respectively. Hybrid Materials (MSIII) involves a solid-solution to offer the suitable coordination geometry for activators (A) and (B), and realize the cascaded energy transfer (traps (C) \rightarrow activator (A) \rightarrow activator (B)).

(traps \rightarrow activator(A) \rightarrow activator(B)) and going into the details of it, one has at one's disposal several decades worth of well-established principles in the coincident matching of macroscopical and microscopic features in spectroscopy, coordination chemistry and network connectivity relating to activators and hosts^{10,11}. Traditionally, materials scientists view such network-engineering design accessed via active impurities with a practical eye intent on describing integral architectures in terms of ion types, valency and radius, local coordination geometries, as well as their concomitant implications for electronegativity and chemical bonding¹². However, due to the complex attribute of topological network, there are still remaining grand challenges: to gain better modulation for the local coordination configuration of dopants, to understand the principle linking the indispensable transfer channel of independent individual, and to realize true predictability to the arrangements of traps and dopants (sensitizers, activators, or co-dopants) in coordinated network.

In this work we present a new “Top-down” approach to design and synthesize the long phosphorescent phosphors in the second biological transparent window. The material design approach employed here involves an ingenious engineering for hybridizing the coordination networks of hosts, tailoring the topochemical configuration of dopants, and bridging cascaded channels for transferring the persistent energy from traps, to sensitizers, to acceptors. We present a closed energy transfer channel from Cr^{3+} to Nd^{3+} in ZnGa_2O_4 phosphor and invalid electronic reservoir in CaGa_2O_4 phosphor, respectively. Persistent energy-transfer could occur in $\text{Zn}_{1-x}\text{Ca}_x\text{Ga}_2\text{O}_4$ solid-solution because two dopants were successfully locked in a cage via the efficient crystal packing at an appropriate distance, in addition to the preservation of native electron traps. The hybrid network topologies and structural motifs, thus far will be outlined with particular emphasis on how specific route of energy transfer can be prepared via premeditatedly designing a material system. Such design strategy will notably open a vista of potential avenues for the design of new optical functional materials for the future.

Results and Discussion

Our strategy was inspired by the fundamental spectroscopic theory of energy transfer and local intercalation reaction in inorganic polycrystals (Fig. 1)¹³. In our view, a typical long phosphorescent phosphor (MSI, in Fig. 1a) features a prominent electron reservoir (C, in Fig. 1) with the distinct ability of storing and releasing the captured electrons, as well as a notable photon-emitter (A, in Fig. 1) with higher quantum efficiency under the condition of accurately matching lattice-coordination network and atomic radius¹⁴. A pre-established electronic transfer channel (AG, traps (C) \rightarrow activator (A), in Fig. 1) ensures the long persistent phosphorescence. However, the topological network does not provide an opportunity for another activator (B, in Fig. 1) to embed itself into the suitable lattice site. Such structural constraint thus, closes the possible channel of energy transfer (ET, in Fig. 1) between (A) and (B), leading to the luminescent and phosphorescent quenching. Fortunately, the existing chemical and spectroscopic knowledge offer a far-sighted technique to select another material system (MSII, in Fig. 1b), which allows a synchronous precipitation of activator (A) and (B), as well as engineers a theoretically existent energy-transfer channel (activator (A) \rightarrow activator (B)). But to our surprise, this scheme misses the necessary electron reservoir so as to completely decrease the probability of electrons trapping-de-trapping (Fig. 1b).

The use of solid-solution complexes to engineer predictable, multi-dimensional infinite networks has received ever-increasing attention in the area of chemistry and materials science¹⁵. Solid-solutions have already proven their superiorities in the areas of optical, optoelectronic, electrical and magnetic properties than the single component¹⁶. Pan *et al.* broke new ground in the field by using zinc gallogermanates solid-solution as the system, thereby achieving a super-long NIR afterglow emission time of 360 h³. Kobayashi *et al.* also demonstrated

a state-of-the-art Li_xFePO_4 solid-solution technology, opening the door for lithium ion batteries to take their place in large-scale applications¹⁷. In addition, a series of solid-solution, such as $\text{AgGa}_{1-x}\text{Al}_x\text{O}_2$, $\text{Zn}_{1-x}\text{Cu}_x\text{S}$, $(\text{SrTiO}_3)_{1-x}(\text{LaTiO}_2\text{N})_x$, also have been developed and used as the advanced photocatalysts to enhance the photocatalytic activity of a given semiconductor photocatalyst¹⁸. Therefore, solid-solution highlights hybrid coordination network of host, and is expected to open up a possibility in the visualization of the structural and functional binding process of traps and all activators into an independent system¹⁹. By rationally deploying an indirect intercalation complex comprised by polyhedron ligands of materials (MSI) and (MSII), hybrid coordination-network of novel solid-solution (MSIII, in Fig. 1c) is engineered to steady the activators (A) and (B), modulate the topochemical configuration of activators, and realize the cascaded energy transfers, traps(C) \rightarrow activator(A) \rightarrow activator(B) (Fig. 1c). Such novel structural motif is anticipated to adopt a disturbance to native unit cell and bridging a predictable periodic coordination network.

To validate research idea, a typical NIR long phosphorescent phosphor ZnGa_2O_4 :Cr was pursued as preferential material system, which has been proven capable of supporting high defect densities, thought to be associated primarily with Zn vacancies (V_{Zn}) and O vacancies (V_{O}), as well as some antisite deficiencies (Zn_{Ga})²⁰. Making use of its defect capacity, ZnGa_2O_4 :Cr has been demonstrated as a NIR photo-emitter with surprisingly long persistent phosphorescence in first NIR window (Supplementary Fig. S1). Here, we target the operating waveband in the second NIR window by transferring the persistent energy of Cr^{3+} to Nd^{3+} in Cr/Nd-codoped ZnGa_2O_4 LPPs. Nd^{3+} ion is chosen as the emission center in order to take advantage of the appropriate energy level characteristic, i.e. NIR-absorption (680, 750 and 800 nm) and NIR emission (1064 nm)²¹. The various sharp transitions of Nd^{3+} [$^4\text{I}_{9/2} \rightarrow ^4\text{F}_{5/2}$], [$^4\text{I}_{9/2} \rightarrow ^4\text{F}_{7/2}$], [$^4\text{I}_{9/2} \rightarrow ^4\text{F}_{9/2}$], just overlap the electron transition from metastable state ($^4\text{T}_2$) to ground state ($^4\text{A}_2$) of Cr^{3+} , allowing the potential energy transfer from Cr to Nd²². However, no any NIR phosphorescence in the second NIR window can be observed in Cr/Nd-codoped ZnGa_2O_4 LPPs (Fig. 2a). In fact, the desired phosphorescence is still absent in Nd^{3+} singly doped ZnGa_2O_4 phosphor after ceasing the excitation (Supplementary Fig. S2). It is notable that the diffuse reflection spectrum consists of the characteristic transition bands centered at 530, 588, 688, 748 and 808 nm, respectively, corresponding to Nd^{3+} f-f transition, [$^4\text{I}_{9/2} \rightarrow ^2\text{K}_{13/2} + ^4\text{G}_{7/2} + ^4\text{G}_{9/2}$], [$^4\text{I}_{9/2} \rightarrow ^2\text{G}_{7/2} + ^2\text{G}_{5/2}$], [$^4\text{I}_{9/2} \rightarrow ^4\text{F}_{9/2}$], [$^4\text{I}_{9/2} \rightarrow ^4\text{F}_{7/2}$], [$^4\text{I}_{9/2} \rightarrow ^4\text{F}_{5/2}$] in Nd^{3+} doped ZnGa_2O_4 phosphor (Supplementary Fig. S3)²¹; yet under the excitation at 748 nm, emission peak at 1064 nm, attributed to Nd^{3+} [$^4\text{F}_{3/2} \rightarrow ^4\text{I}_{11/2}$] transition is not identifiable (Supplementary Fig. S4). This attractive optical quenching-phenomenon of luminescence and phosphorescence may be not concerned with the trap distribution, but the microcosmic network architecture.

In ZnGa_2O_4 , a majority of [Ga^{VI}] cations occupy octahedral sites, whereas all of the [Zn^{IV}] cations occupy tetrahedral sites²³. As a preliminary conjecture, Cr^{3+} has proven its strong ability to substitute for Ga^{3+} in distorted octahedral coordination, whereas Nd^{3+} cannot be effectively introduced into this specific network configuration (inset of Fig. 2a). In order to identify this possibility of ion doping, we focus on the intricate topochemical coordination geometry of Cr and Nd ions in zinc gallate spinel. The elucidation is performed in detail by a combination of XRD data and ⁷¹Ga solid state nuclear magnetic resonance (NMR) studies. XRD patterns of ZnGa_2O_4 :xCr (x = 0.5%, 5%, 10% and 20%) and ZnGa_2O_4 :xNd (x = 0.5%, 5%, 10% and 20%) phosphors were measured and shown in Fig. 2b, Supplementary Fig. S5, S6. The peaks in XRD patterns of all Cr-doped samples are well indexed to pure ZnGa_2O_4 spinel structure (JCPDS 86-0848). In stark contrast, the higher doping content (up to 5%) of Nd ion gives rise to an impure phase, NdGa_2O_7 (JCPDS, 70-3810) in Nd-doped samples. Another interesting phenomenon, i.e. XRD dominated peak (PI in Fig. 2b) shifting towards to higher 2 θ value with the increment of Cr content, reveals a small linear variation in ZnGa_2O_4 unit cell lattice parameter with Cr^{3+} substitution, whereas no any shift of same peak is observed in Nd-doped samples, further ensuring the distinct phase splitting (Fig. 2c). In addition, a decline of the peak intensity in Fig. 2c also is present. Nevertheless, the causes of this decline may be different and rooted from either the substitution or the phase splitting.

NMR allows the observation of specific quantum mechanical and magnetic properties of atomic nucleus, as well as provides the detailed information about the structure, dynamics, reaction state, and chemical environment of molecules²⁴. Many scientific techniques exploit NMR phenomena to cover the interplay between the ligands and geometric centers, as well as study the topological network motif in crystals, microcrystalline powders, or anisotropic solutions, etc²⁵. ⁷¹Ga solid-state NMR is famous for the permission of quantitative analyses to different Ga^{3+} central coordination state in inorganic solids²⁶. Figure 2d,e shows the systematical physical investigations of Ga coordination geometry in Cr and Nd singly doped ZnGa_2O_4 , respectively. For the undoped ZnGa_2O_4 samples, ⁷¹Ga NMR spectra exhibit two well-resolved resonances. The relative higher intensive signal at about 31 ppm is characteristic of sixfold coordinated Ga atoms, and the other weaker one ~at 170 ppm corresponds to Ga atoms in the tetrahedral sites of the spinel structure²⁶. It is necessary to mention that with increasing Cr content (from 0.5% to 10%), ⁷¹Ga NMR spectra present a significant broadening of spectral lines (Fig. 2d). In prominent contrast, scarcely any distinct influence on NMR spectral lines can be found by varying the Nd doping content in solid NMR spectra of ZnGa_2O_4 :xNd (x = 0.5% and 10%) phosphors (Fig. 2e). The clear separation of NMR chemical shift at ~31 ppm between the two samples implies the precipitation of Cr^{3+} into the octahedral lattice site and the excludability of local configuration to Nd^{3+} ions. The NMR results are in accordance with XRD data, offering a powerful structural evidence to explain the interesting phenomena of phase splitting and luminescence quenching.

Actually, rare-earth elements generally form complexes which have high coordination numbers (CNs) and weak metal-ligand bonds, because of their large ionic radii and relatively low oxidation states²⁷. Typically transition-metal and main-group elements have coordination numbers 2–6, while rare-earth metals have CNs > 6²⁸. The resulting coordination polyhedra include trigonal prisms (CN = 6) or its variation by stepwise capping of the prism face up to CN = 9, in addition to square antiprisms (CN = 8); Coordination number 3 is realized only under extreme conditions²⁸. Therefore, to supply an ideal dwelling for Nd^{3+} , a suitable material system should be proposed. Alkaline-earth metals have large ionic radii and various coordination-numbers 3–8

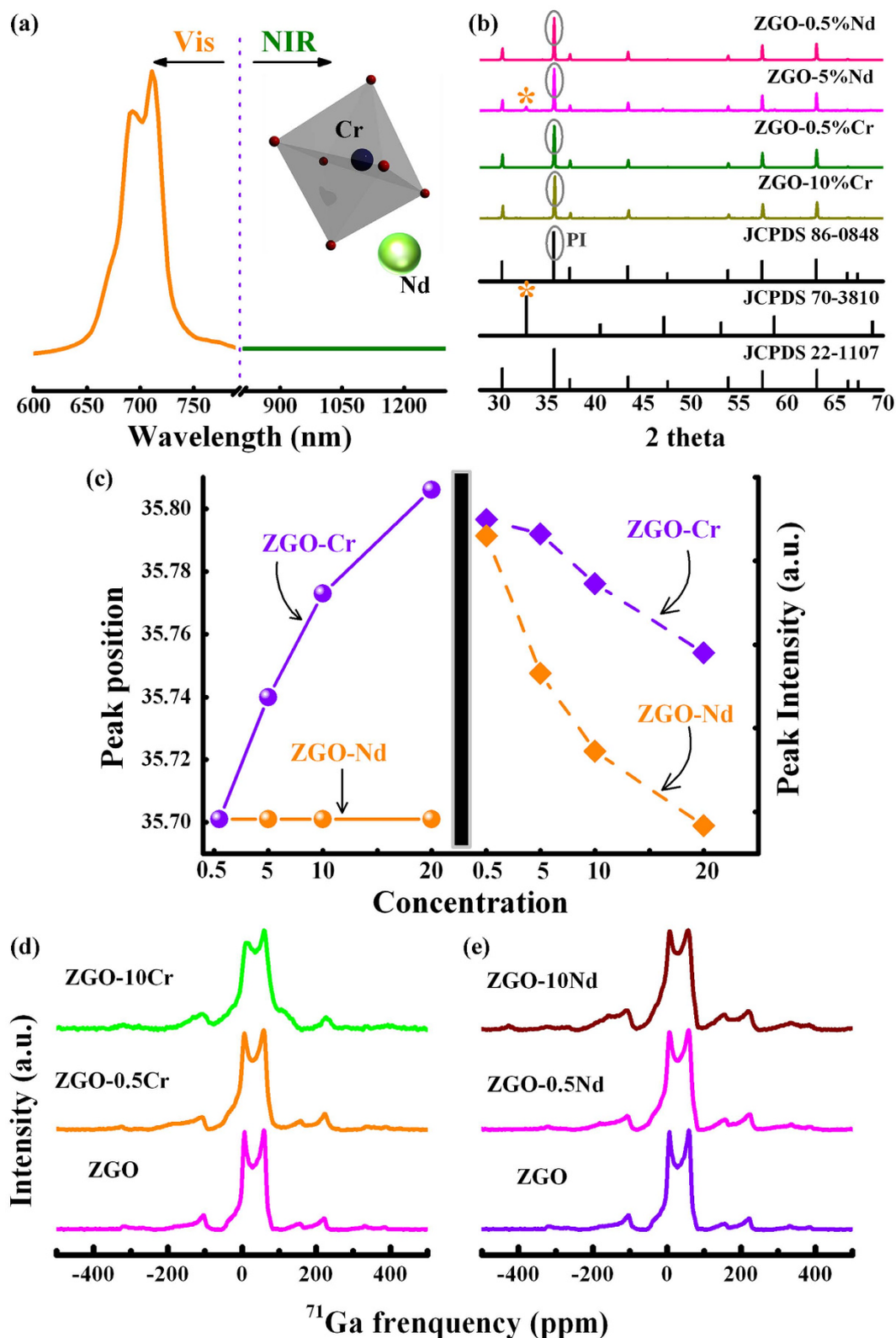


Figure 2. (a) Vis-NIR long persistent phosphorescence spectrum of ZnGa₂O₄: Cr/Nd phosphor. The inset shows a conjectural doping pattern of Cr and Nd. (b) XRD patterns for ZnGa₂O₄: xCr (x = 0.5% and 10%) and ZnGa₂O₄: xNd (x = 0.5% and 5%) phosphors. (c) Dependence of XRD peak ([PI] labeled in Fig. 2b) position and intensity as a function of Cr and Nd concentration in ZnGa₂O₄ phosphor. (d–e) ⁷¹Ga solid state NMR spectra of ZnGa₂O₄: xCr (x = 0.5% and 10%) and ZnGa₂O₄: xNd (x = 0.5% and 10%) samples. All spectra were recorded at a magnetic field of 11.7 T with a sample spinning frequency of 25 kHz.

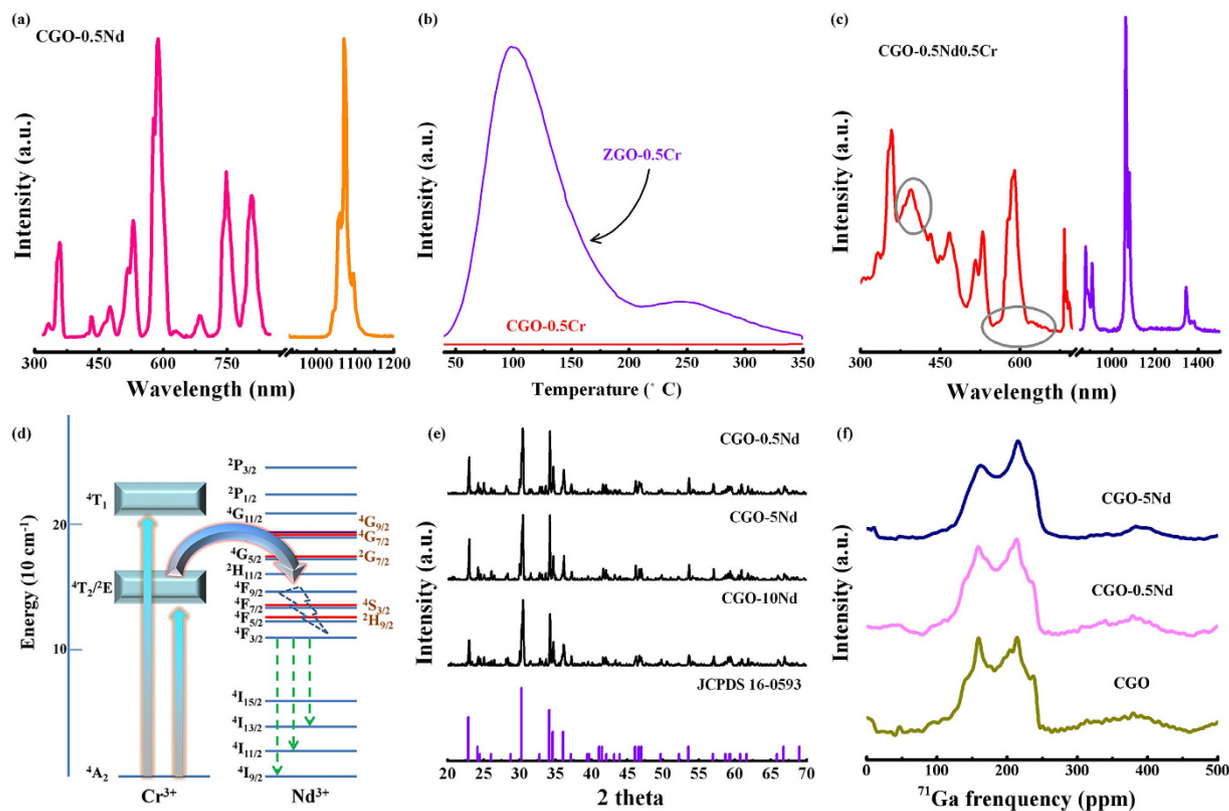


Figure 3. (a) Static photoluminescence spectrum under excitation at 750 nm and the corresponding photoluminescence excitation spectrum monitored at 1064 nm of CGO-0.5Nd phosphor; (b) Thermoluminescence curves of 0.5%Cr-doped ZnGa₂O₄ and CaGa₂O₄ phosphors measured 30 s after irradiation ceased; (c) Photoluminescence spectrum under excitation at 410 nm and photoluminescence excitation spectrum monitored at 1064 nm of CGO-0.5Cr0.5Nd phosphor; (d) Schematic illustration showing the energy-level diagram of Cr³⁺ and Nd³⁺ in CaGa₂O₄ phosphors; (e) XRD patterns for CaGa₂O₄: xNd (x = 0.5%, 5% and 10%) phosphors; (f) ⁷¹Ga NMR spectra of CaGa₂O₄: xNd (x = 0, 0.5% and 5%) samples.

in different hosts, which ensure the selection of alkaline-earth gallates²⁹. CaGa₂O₄ has a similar spinel crystal structure with ZnGa₂O₄. In CaGa₂O₄, [Ca^{VI}] cations occupy octahedral sites²⁹. This configuration thus features a path of easy doping ion precipitation into the octahedral [Ca^{VI}] under the condition of matching geometrical lattice and atomic radius, which occurs with rare earth ion, Nd.

As expected, Fig. 3a exhibits the characteristic transitions of Nd³⁺ in Nd singly doped CaGa₂O₄ phosphor. However, the idealistic and aspirational long persistent phosphorescence is still absence in Cr singly, Nd singly and Cr/Nd doped CaGa₂O₄ phosphors, respectively (Supplementary Fig. S7). A possible cause of this problem is due to the lack of effective traps (Fig. 3b). In sharp contrast to Cr/Nd codoping ZnGa₂O₄, photoluminescence excitation (PLE) spectrum monitored at 1064 nm of Cr/Nd codoping CaGa₂O₄ sample consists of two specific excitation bands centered at ~410 and ~620 nm, in addition to Nd³⁺ characteristic f-f transitions (Fig. 3c), indicating an energy transfer from Cr³⁺ to Nd³⁺. Obviously, the strong one is attributed to the Cr³⁺ [⁴A₂ → ⁴T₁], while the weak one corresponds to Cr³⁺ [⁴A₂ → ⁴T₂]³⁰. Further verification of energy transfer between Cr³⁺ and Nd³⁺ is supplied by emission spectrum and decay curve monitored at 1064 nm under the excitation wavelength at 410 nm (Fig. 3c and Supplementary Fig. S8). A possible channel of energy transfer from Cr³⁺ to Nd³⁺ is Cr³⁺ [⁴T₂ → ⁴A₂]: Nd³⁺ [⁴I_{9/2} → ⁴F_{5/2}], [⁴I_{9/2} → ⁴F_{7/2}], or [⁴I_{9/2} → ⁴F_{9/2}], depending on the overlap between Cr³⁺ emission band and Nd³⁺ absorption band (Fig. 3d)³¹. As discussed above, due to the similar atomic radius and geometric configurations, Nd ions can easily precipitate on Ca lattice site in CaGa₂O₄, enabling the distinct photoluminescence (PL). To probe the lattice configuration and substitution progress in CaGa₂O₄, we performed XRD and solid state NMR experiments. X-ray diffraction pattern first confirms the crystallization of Nd-doped calcium gallate (Fig. 3e). In contrast to Nd-doped ZnGa₂O₄, all Nd-doped CaGa₂O₄ samples can be indexed as standard phase CaGa₂O₄ (JCPDS 16-0593). There is no any apparent observation of phase splitting from XRD data, even under a higher doping content of Nd³⁺, firmly supporting the rational inclusion of Nd³⁺ into an inert matrix, CaGa₂O₄. This result is also supported by ⁷¹Ga solid state NMR spectra. In contrast to ZnGa₂O₄ host, the undoped CaGa₂O₄ sample has a dominant chemical shift at 170 ppm (Fig. 3f). With increasing dopants content, CaGa₂O₄: Nd also has the same effect of NMR resonances' line broadening and the linear increase of NMR resonances integrated intensity, strongly suggesting the successful substitution in substantial amounts of Nd into Ca lattice site.

Seemingly, as the individual backbone, MgGa₂O₄ (Zn and Ca) polymorph is chosen as the prototypical coordination network for its respective ability to engineer the functionally independent tunnel, traps(C) →

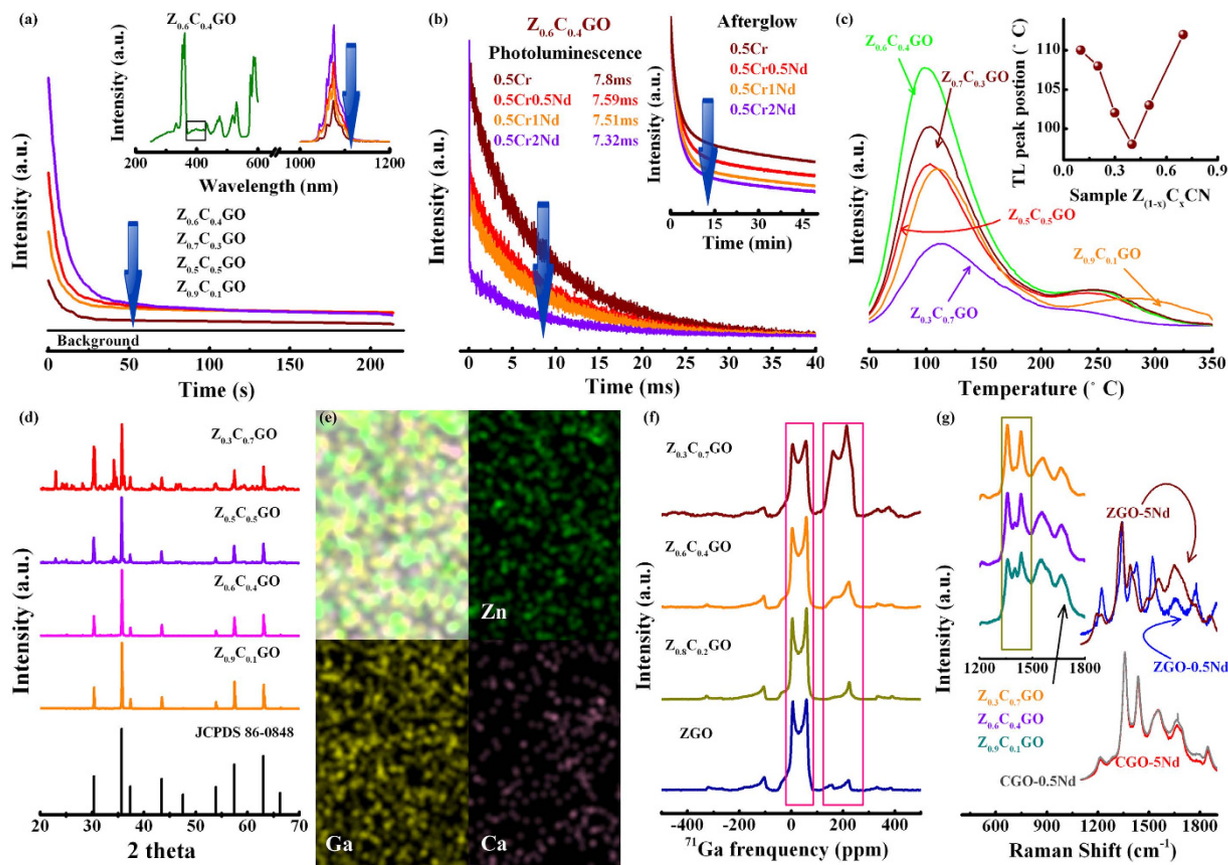


Figure 4. (a) persistence time monitored at 1064 nm as a function of Ca concentration ($x = 0.1, 0.3, 0.4$ and 0.5). The inset shows the long persistent phosphorescence spectra of $Z_{1-x}C_xGO$ ($x = 0.1, 0.3, 0.4$ and 0.5) samples and photoluminescence excitation spectrum monitored at 1064 nm of sample $Z_{0.6}C_{0.4}GO$. (b) Normalized photoluminescent and phosphorescent decay curves of $Z_{0.6}C_{0.4}GO: 0.5\%Cr/xNd$ ($x = 0, 0.5\%, 1\%$ and 2%) samples. The monitored transition is $Cr^{3+} [^4T_2 \rightarrow ^4A_2]$. (c) Thermoluminescence curves of $Z_{1-x}C_xGO$ ($x = 0.1, 0.3, 0.4, 0.5$ and 0.7) phosphors measured 30 s after irradiation ceased. The inset shows the dependence of TL peak position as a function of Ca concentration. (d) XRD patterns for $Z_{1-x}C_xGO$ ($x = 0.1, 0.4, 0.5$ and 0.7) phosphors. (e) EDX mapping of sample $Z_{0.6}C_{0.4}GO$. (f) ^{71}Ga NMR spectra of $Z_{1-x}C_xGO$ ($x = 0, 0.2, 0.4$ and 0.7) phosphors. (g) Normalized Raman spectra of ZGO-0.5Nd, ZGO-5Nd, CGO-0.5Nd, CGO-5Nd and $Z_{1-x}C_xGO$ ($x = 0.1, 0.4$ and 0.7) phosphors.

activator(A), or activator(A) \rightarrow activator(B), used to transfer the required energy. The only regret is the fundamentally missing connection of traps(C) \rightarrow activator(A) \rightarrow activator(B) in a separate material system. To address this issue, we anticipate a novel solid-solution $Zn_{1-x}Ca_xGa_2O_4$ to bridge a new channel for transferring the persistent energy from traps to desired ions, based on the cautious consideration for crystal structure, ion valency and chemical bond relating to hosts and dopants. The desired NIR phosphorescence at 1064 nm is finally present in the afterglow spectra of $Zn_{1-x}Ca_xGa_2O_4$ ($x = 0.1, 0.3, 0.4$ and 0.5) solid-solution (Fig. 4a). Significantly, we also observe a strong dependence (i.e. rising first followed by a decline) of phosphorescent peak intensity and decay dynamics on Ca concentration in Fig. 4a. We attribute this special spectral change of Nd^{3+} to the successful persistent energy transfer from Cr^{3+} to Nd^{3+} , which is supported by the meticulous spectral studies of Nd^{3+} in an optimal $Zn_{0.6}Ca_{0.4}Ga_2O_4: 0.5Cr/0.5Nd$ solid-solution: PLE band at 410 nm should be assigned to Cr^{3+} transition [$^4A_2 - ^4T_2$], while a distinct NIR PL peak at 1064 nm is observed under the excitation at 410 and 600 nm (Fig. 4a and Supplementary Fig. S9). The additional support for the formation of an unrestricted energy tunnel, traps $\rightarrow Cr^{3+} \rightarrow Nd^{3+}$, is the analysis of kinetic processes in $Z_{0.6}C_{0.4}GO: 0.5\%Cr/xNd$ ($x = 0, 0.5\%, 1\%$ and 2%) samples (Fig. 4b). PL decay dynamics study of Cr^{3+} shows a notable shortening in decay lifetime from 7.8 ($Z_{0.6}C_{0.4}GO-0.5Cr$), to 7.59 ($Z_{0.6}C_{0.4}GO-0.5Cr0.5Nd$), to 7.32 ms ($Z_{0.6}C_{0.4}GO-0.5Cr2Nd$), giving clear evidence of successfully simultaneous precipitation of two activators into the corresponding lattice along with the effective energy transfer from Cr^{3+} to Nd^{3+} .

It should be noted that, to the best of our knowledge, this type of NIR long-persistence phosphorescence has not been previously reported to occur in hybrid coordination networks by engineering cascaded energy transfer channels. Such substantial progress is strongly influenced by two key attributes; one is trap distribution and another is network architecture. Apparently, the variation of trap distribution may be not a crucial factor in exploring the nature of transfer channel, because the indispensable electron reservoir is still steadily embedded in all the $Zn_{1-x}Ca_xGa_2O_4$ solid-solutions (Fig. 4c). To probe the evolution of topological network-dependent

topochemical coordination, the systematic characterization, such as, XRD, solid NMR, EDX mapping and Raman spectra should be conducted³². XRD peaks in $Z_{1-x}C_xGO$ ($x = 0.1, 0.4, 0.5$ and 0.7) samples indicate their $ZnGa_2O_4$ spinel solid-solution nature, while the superimposed peaks in samples $Z_{0.5}C_{0.5}GO$ and $Z_{0.3}C_{0.7}GO$ can be well indexed by the diffraction peaks of $ZnGa_2O_4$ and $CaGa_2O_4$ (Fig. 4d, and Supplementary Fig. S10). EDX mapping analysis reveals the solid-solutions have uniform distribution of Ca elements in all of the spinel solid-solutions (Supplementary Fig. S11, S12 and Fig. 4e). EDX experimental composition approximating the theoretical value supports the successful inclusion of Ca elements in spinel crystals (Supplementary Table S2).

⁷¹Ga NMR spectra have provided some insights into the coordination variation of Ga center in $ZnGa_2O_4$ and $CaGa_2O_4$ phosphors, due to the incorporation of Cr and Nd, it is also expected to manifest its ability in resolving the question of topochemical configuration's evolution process, as the addition of Ca element. As shown in Fig. 4f, with increasing Ca content (0, 0.2, 0.4 and 0.7), two resonances at 170 and 31 ppm in ⁷¹Ga NMR spectra increasingly present the linear broadening. In these solid solutions, Zn-O and Ga-O tetrahedron could suppress the intrusion of Ca element due to the mismatch of coordination configuration. In fact, to steady Ca ion, parts of Ga-O octahedron must reorient to form the new polyhedron network Ca-O octahedron along with the transformation from Ga-O octahedron to Ga-O tetrahedron due to the decrease of Zn-O tetrahedron. In detail, for the samples $Z_{1-x}C_xGO$ ($x = 0, 0.2, 0.4$), the motion of local hybrid coordination-networks evolution include: (1) the precipitation of Ca on the lattice site of octahedron Ga, giving rise to the broadening of NMR resonance at 164 ppm; (2) the conversion from Ga-O octahedron to Ga-O tetrahedron, resulting in the enhancement of NMR resonance at 65 ppm. This interesting redeployment of network configuration thus permits the modification of topochemical state of dopants, as well as opens the possibility of bridging cascaded channels to transfer the persistent energy. To further validate the research idea aiming at the network configuration, Raman spectra of the fabricated samples also can be selected as the pertinent tool to further analyze the evolution of network architecture (Fig. 4g and Supplementary Fig. S13)³³. In stark contrast to samples ZGO-0.5Nd and ZGO-5Nd, normalized Raman spectra of samples CGO-0.5Nd and CGO-5Nd do not exhibit the notable Raman peak shift and variation of Raman peak intensity, indicating a strong constraint of topological network to the migration of Nd ions in $CaGa_2O_4$. In fact, only two distinct Raman bands at ~ 1358 and 1434 cm^{-1} are present in the Raman spectrum of CGO-0.5Nd, while the Raman spectrum of ZGO-0.5Nd includes three identifiable Raman peaks at ~ 1341 , 1389 and 1425 cm^{-1} . Thus, Raman spectra of $Z_{1-x}C_xGO$ ($x = 0.1, 0.4$ and 0.7) solid-solutions consequentially show a unit number decrease of Raman peaks with the increment of Ca content (inset of Fig. 4g). The variation of middle peak at 1401 cm^{-1} as a function of Ca doping content ensures the strong signature of the hybrid network structure, which is in accordance with the XRD and solid-state NMR data.

In summary, we report a principle of bridging cascaded energy transfer channels to activate long persistent phosphorescence in the second biological window and fabrication of novel near-infrared phosphorescent phosphor Cr/Nd codoped $Zn_{1-x}Ca_xGa_2O_4$ solid-solutions. Structural studies offer the powerfully fundamental evidences to explain the closed energy transfer channel from Cr^{3+} to Nd^{3+} in $ZnGa_2O_4$ phosphor and invalidation of electronic reservoir in $CaGa_2O_4$ phosphor. We believe that the ingenious solid-solution technology featuring the superiority of engineering a hybrid coordination-network opens new paths for advanced dynamic management of activation energy and gives the inspiration to design future new-wavelength, NIR phosphorescent phosphors by energy transfer.

Methods

Materials. 4N pure $CaCO_3$, Ga_2O_3 , ZnO , Nd_2O_3 and Cr_2O_3 were selected as the raw materials.

Preparation of $ZnGa_2O_4$: xCr/yNd. Phosphors with molar compositions of $ZnGa_2O_4$: xCr/yNd ($x = 0, 0.5\%, 5\%, 10\%, 20\%$; $y = 0, 0.5\%, 5\%, 10\%, 20\%$), (Supplementary Table S1) were prepared by the solid state reaction method. The reaction included a two-step thermal treatment (i.e., initial calcination at 800°C for 5 h, secondary calcination at 1350°C for 3 h).

Preparation of $CaGa_2O_4$: xCr/yNd. Phosphors with molar compositions of $CaGa_2O_4$: xCr/yNd ($x = 0, 0.5\%$; $y = 0, 0.5\%, 5\%, 10\%$), (Supplementary Table S1) were prepared by the solid state reaction method. The reaction included a two-step thermal treatment (i.e., initial calcination at 800°C for 5 h, secondary calcination at 1200°C for 3 h).

Preparation of $Zn_{1-x}Ca_xGa_2O_4$: 0.5Cr/yNd. Phosphors with molar compositions of $Zn_{1-x}Ca_xGa_2O_4$: 0.5Cr/yNd ($y = 0, 0.5\%, 1\%, 2\%$; $x = 0.1, 0.2, 0.3, 0.4, 0.5, 0.7$), (Supplementary Table S1) were prepared by the solid state reaction method. The reaction included a two-step thermal treatment (i.e., initial calcination at 800°C for 5 h, secondary calcination at 1350, 1350, 1300, 1300, 1270, 1250°C for 3 h as a function of x, respectively).

Characterization. The prepared materials were analyzed by X-ray diffraction ($Cu/K\alpha$) to confirm the sole crystalline phase. Room-temperature photoluminescence (PL), photoluminescence excitation (PLE) spectra, afterglow spectra and decay curves were measured with a high-resolution spectrofluorometer (UK, Edinburgh Instruments, FLS920) equipped with a 500 W Xenon lamp as an excitation source, with a Hamamatsu R928P visible photomultiplier (PMT) (250–850 nm) and a liquid nitrogen-cooled Hamamatsu R5509-72 NIR PMT as the detectors. TL glow curves and TL excitation (TLE) spectra were measured with a FJ-427A TL meter (China, Beijing) to characterize defect properties. Unless otherwise mentioned, the samples were pre-annealed at 600 K before testing, and some measurements were taken after pre-irradiating the samples for 10 min by using a xenon lamp. EDX images are characterized by a field emission scanning electron microscopy (FE-SEM), Nova NanoSEM 430. ⁷¹Ga Hahn echo NMR experiments were performed on Bruker Avance III spectrometers operating at magnetic fields of 111.4 T corresponding to ⁷¹Ga Larmor frequencies of 152.54 MHz) using Bruker 2.5 mm triple and double resonance probe heads. The 90° degree pulse length is 1.25 μs with a recycle delay of 8s. ⁷¹Ga chemical

shifts were referenced relative to a 1.0 M aqueous solution of Ga(NO₃)₃. All ⁷¹Ga spectra were fitted using the Dmfit software. Raman spectra were collected with a Renishaw inVia Raman microscope irradiated by a visible laser at 532 nm.

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Author Contributions

J.R.Q. conceived and designed the experiments, and was responsible for the project planning. X.X.Q. prepared the samples. X.X.Q., Y.L. and Y.L.W. investigated the spectroscopic properties. J.J.R. and R.L.Z. carried out the NMR study. Y.L. wrote the manuscript. J.R.Q., M.G., S.F.Z., Z.J.M., K.F. and G.P.D. checked the manuscript. All authors were involved in the discussion of the experimental results.

Additional Information

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