Method for the Detailed Characterization of Cosputtered Inorganic Luminescent Material Libraries

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Supporting Information

ABSTRACT: Understanding the behavior of combinatorially developed luminescent materials requires detailed characterization methods that have been lacking thus far. We developed a device for directly surveying the luminescent properties of thin-film libraries created through combinatorial gradient sputter deposition. Step-scan recorded excitation-, emission- and luminescence decay spectra of a thin-film library were resolved and combined with EDX measurements on the same film, relating composition to luminescent properties. This technique was applied to a single-substrate gradient thin-film library of NaBr$_{0.75}$I$_{0.25}$ to NaI$_{0.06}$Br$_{0.94}$ doped with 6.5% to 16.5% Eu$^{2+}$. This gradient film closely followed Vegard’s law, with emission fluence shifting from 428 to 439 nm. In comparison, pure NaBr:Eu$^{2+}$ showed emission at 428 nm and NaI:Eu$^{2+}$ at 441 nm. Luminescence decay measurements demonstrated a great degree of concentration quenching in the gradient film. From these measurements we could conclude that an optimized phosphor would most efficiently luminesce when close to NaI:Eu$^{2+}$. This gradient film confirmed that the method presented in this work allows to both study and optimize luminescent behavior in a broad range of host- and dopant systems.

KEYWORDS: inorganic phosphors, combinatorial screening, sputter deposition, luminescence

INTRODUCTION

Combinatorial synthesis finds broad application in many fields, but in the case of phosphor synthesis, only a limited number of reports exist. This lack of past works can be explained by that combinatorial methods, when applied to conventional searches like the search for white light LEDs, usually converge to known phosphors. In addition, the analysis methods used for combinatorially synthesized libraries of luminescent materials are often limited to (color-filtered) charge-coupled device (CCD) photography of a library under UV illumination, substrate-scanning with color filters to isolate emission peaks, or cathode luminescence with no information on low energy excitations. These coarse characterizations still have to be followed by powder synthesis to fully analyze the luminescent properties of a material of interest.

In this Research Article, we present a novel method that enables the detailed characterization of a luminescent composition library, based on combinatorial gradient deposition of thin-films using DC/RF magnetron sputtering. This gradient is realized by taking advantage of the low mobility of sputtered adatoms when depositing without substrate heating. The presented method is capable of resolving position-dependent structural and compositional information, as well as photoluminescent excitation, emission, and decay spectra. This combination allows us to link the position-dependent structural and compositional information to the recorded luminescence properties, thereby retrieving direct composition-property relations.

Determination of the structure and composition is done with step-scan X-ray diffraction (XRD) for crystallinity and phase determination. Energy-dispersive X-ray spectroscopy (EDX) is used to resolve the local composition of the gradient thin-film. Photoluminescence measurements are done using a combination of laser excitation and an xy-stepping stage, with luminescence captured by either a CCD spectrometer, or a photomultiplier tube (PMT) coupled to a digitizer.

In binary halides, substitution of anions can lead to a fluent shifting of emission wavelength between the two mixed compounds, in line with Vegard’s law. For instance, KCl$_{1-x}$Br$_x$ crystals doped with Eu$^{2+}$ have shown this fluent shifting of emission. Therefore, to illustrate the capabilities of the method, we have chosen a continuously varying mixed-phase library of NaBr:Eu$^{2+}$ and NaI:Eu$^{2+}$ as an example. While the individual phosphors have previously been researched, the behavior of the mixed-phase phosphor has not yet been studied. The end points of this library are therefore well-known, while the mixed-phase in between provides new data on the behavior of mixed binary halides. The combination of luminescent, compositional and structural information is employed to see if these two phosphors will also display behavior following Vegard’s law. Furthermore, the possibility of optimizing luminescent behavior in terms of quantum efficiency is explored with this library.

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RESULTS AND DISCUSSION

Library Creation and Method for Characterization.

To provide an illustration of the combinatorial synthesis and characterization route, NaBr, NaI and Eu were magnetron cosputtered at a 90° angle from each other on a 5 × 5 cm² UV fused silica substrate. Figure 1a and 1b shows the layout of the sputtering chamber prior to deposition and during deposition, respectively. UV fused silica was chosen since it does not show luminescence, contrary to other glass types.15 As with Tm,16 Eu sputtered together with binary alkali halides readily forms its divalent state, without the need for any additional oxidizing gases or heat treatments. The library was therefore deposited at room temperature using an inert Ar atmosphere, without further treatment. In addition, being SiO₂ of high purity, it will be stable up to 1500 °C. In the glovebox, the sample (Figure 1c) was transferred to a N₂-filled, airtight sample holder with a UV fused silica or polyimide foil window for optical or XRD measurements, respectively. Figures 1d to 1f show the elemental concentrations based on EDX measurements at 40 locations and an interpolation between these points.

Direct EDX measurements of the Eu concentration carry a high uncertainty. Contrary to Eu, the ratio of Na to Br and I can be repeatedly measured with good certainty across the film. The composition of the film should be charge-neutral. The Eu concentration can therefore be derived from the measured ratio of Na to Br and I by setting the ratio of Eu²⁺ and Na⁺ to Br⁻ and I⁻ equal to unity.

The interpolation is obtained by fitting with the surface-source evaporation equation. The surface-source evaporation equation for a single source is based on the mass deposited on the substrate per unit area dMᵢ/dAₜ, given by

\[
dM_i(x, y) \frac{dA_t}{dA_s} = M_i (n + 1) \cos^n \phi(x, y) \cos \theta \frac{2\pi r^2(x, y)}{S}
\]

Here, \(M_j\) is the total mass exiting the sputtering source, \(n\) determines the sharpness of the elongated sphere of material being sputtered from the source, \(\phi\) is the angle between the source normal and a line extending from the center of the source to a position on the substrate, \(\theta\) is the angle between the source normal and the substrate normal, and \(r\) is the distance between the source and any position on the substrate’s surface. These parameters are illustrated in Figure 1b. Fitting the concentration \(c_i\) of a material \(j \in i\) when using \(N\) target materials \(i\) can be accomplished with

\[
c_i(x, y) = \frac{\sum dM_j(x, y)}{\sum dA_j(x, y)}
\]

where the fitting parameters are the respective \(n\) and \(M_{ij}\) of the sputtering sources.

The resultant \(c_j\) for \(j = \text{I, Br, and Eu}\) are shown in Figure 1d–1f, respectively.

The sputtering yield of NaI is larger than that of NaBr. The iodide concentration therefore follows the distribution yielded by the gun with NaI, which leads to the semicircles seen in Figure 1d. Figure 1g brings all EDX results together and shows that this single substrate comprises a variation of 6.5% to 16.5% Eu, and a variation of NaBr₀.₇₃I₀.₂₇ to NaBr₀.₀₉I₀.₉₁. Across the substrate, this means an average slope of 0.05%/mm from NaBr to NaI and an average slope in Eu-doping percentage of 0.15%/mm (all concentrations mentioned are in atomic percent).

A custom computer-controlled dual xy-translation stage that is synchronized with an optical parametric oscillator (OPO) laser and a data-acquisition system, referred to as the XY-scanner, was developed for mapping the luminescent properties across the thin-film surface. A schematic of the entire characterization system is shown in Figure 2. Light ranging from 193 to 2600 nm with 0.3 nm line width generated by a 100 Hz (3–5 ns fwhm pulses) OPO laser is guided toward the stage to serve as an excitation source. The laser light is passed through a diaphragm to compensate for fluctuations in the position of the laser spot on the thin-film when stepping across wavelengths. A UV fused silica plate placed at a 45° angle reflects 4% of this incident light toward a powermeter that is used to correct for wavelength- and time-dependent laser power fluctuations. Following excitation, luminescence passes through a long-pass filter to remove reflected laser light and is...
focused into an optical fiber that guides the light to a detector. This detector can either be a CCD spectrometer for emission- and excitation measurements, or a monochromator coupled to a PMT (Mono+PMT) connected to a digitizer for decay measurements. Excitation- and emission measurements produced with the XY-scanner show excellent agreement with what is measured with conventional methods, as shown in Figure S1. 

**Luminescent Alkali-Halide Library.** Figure 3a and 3b show the emission (4f65d[t2g] → 4f7) of the NaI−NaBr:Eu2+ thin-film library under laser excitation at 350 nm. Scanning over the film with the laser yields emission spectra per position on the film. This data can be summarized in an RGB color plot, shown in the inset of Figure 3a, which closely resembles the photograph shown in Figure 1c. This color-coded image is obtained by converting the measured emission spectra to CIE-coordinates and translating those to RGB values.

Figure 3a shows that the emissions have a fluent transition from 428 nm at the NaBr-rich side, to 438 nm at the NaI-rich side of the film. A closer look at the emission reveals that it is actually comprised of two bands and a broad (fwhm > 0.8 eV) sideband of low intensity, possibly comprised of multiple smaller bands. In the further discussion, this broad (>0.8 eV) low energy emission is not shown due to its low intensity, and since it is most likely comprised of many higher-order defects. Figure 3b illustrates this presence of multiple bands. We can see a second band at higher wavelengths developing when moving toward increased Eu concentration, while the other band stays centered at the same position. This qualitative analysis can be made more quantitative by deconvoluting the emission into two separate Gaussians. The central wavelengths of these Gaussians are displayed in Figure 3c and 3d, with relative areas in Figure 3e and 3f and bandwidths in Figure 3g and 3h. The emission of the narrow (0.16−0.18 eV) and intense band at 428−438 nm (Figure 3c, 3e, and 3g) closely resembles the semicircles caused by the higher sputtering yield of NaI over NaBr, as seen in Figure 1d. The broader (0.34−0.38 eV) emission’s central wavelength (Figure 3d, 3f, and 3h) follows the increasing Eu concentration, moving toward lower energy and increasing in prominence with higher Eu concentration.

The luminescence quantum yield ηQY is an important measure for the performance of a phosphor, as it indicates the presence of luminescence quenching, for instance due to an overly high dopant concentration. The observed intensity Figures 3e and 3f is not a direct measure of ηQY, because the intensity also depends on the absorption, which in turn depends on the thickness of the film. A measure for ηQY independent of the luminescence intensity is the decay time.

Figure 4a shows individual luminescent decay spectra across the substrate. These spectra were measured with a slit width such that all light from 415 to 445 nm is collected. The exact locations of these measurements are displayed in Figure 4b. Because of the nonexponential nature of the measured decays, a mean decay time

\[ \tau_{\text{mean}} = \frac{\int_0^\infty tI(t) \, dt}{\int_0^\infty I(t) \, dt} \quad (3) \]
is determined. Here, \( I(t) \) is the emission intensity at time \( t \) after a laser pulse. \( \tau_{\text{mean}} \) makes comparison between decay spectra possible, even if the exact behavior of the decay is unknown. Figure 4b shows a mapping of \( \tau_{\text{mean}} \).

The quantum yield of each composition can be estimated using a fitting of the tail of the decay spectrum. Fitting a single exponential to the tail of the decay spectra yields the decay time for an isolated ion \( \tau_{\text{tail}} \). Alkali-halides with a low europium doping are known to have near unity quantum yield at room temperature. It is therefore possible to relate the mean decay time to an estimated quantum yield as

\[
\eta_{\text{QY}} = \frac{\tau_{\text{mean}}}{\tau_{\text{tail}}}
\]

This estimated quantum yield is shown in Figure 4c. Comparison between Figures 4b and 4c shows that the mean decay time does not directly correlate with quantum efficiency. The determined values should therefore not be compared per position, but per composition, as will be discussed in the following.

Using the XY-scanning technique, important luminescence properties of the film at room temperature are determined, while EDX measurements provide the composition of the same film. Since these measurements have the position of the film in common, these measurements can be combined to directly relate the deconvoluted emission, decay times and estimated quantum yields to the composition. These composition-property relationships, which leave out irrelevant position information, are shown in Figure 5.

Figure 5a shows that the high-energy band shifts in concert with the Br to I ratio, from 428 nm at NaBr_{0.73}I_{0.27} to 438 nm at NaBr_{0.5}I_{0.5}, but shifted slightly closer to NaI, at NaBr_{0.43}I_{0.57}. The emission of the substitutional defect in pure NaBr:Eu^{2+} has a bandwidth of 0.13 eV, where in NaI:Eu^{2+} it has a bandwidth of 0.16 eV. This difference leads to the maximal bandwidth in Figure 5b not being displayed at NaBr_{2.5}I_{0.5}, but shifted slightly closer to NaI, at NaBr_{2.0}I_{0.5}.

The emission band at lower energy (Figure 5c) might be attributed to a more complex defect. Previous research attributed a defect emitting at 453 nm in NaBr:Eu^{2+} to a EuBr_{2}-type precipitate, and emission at 461 nm in NaI:Eu^{2+} to a EuI_{2}-type precipitate. The emission shown in Figure 5c, supported by the larger bandwidth shown in Figure 5d, might therefore also be attributed to a EuBr_{2}- or EuI_{2}-type precipitate, or a mixture thereof. The band’s emission center shifts to lower energies with increasing Eu concentration. This redshifting might be a consequence of reabsorption of higher energy emission, which becomes more profound with more absorbing centers present.

Figure 5e shows that for the substitutional defect a decrease in Eu concentration leads to an increase in decay time. Because of the strong correlation with Eu concentration, we can...
conclude that a high degree of concentration quenching is present in our sample. For comparison, the highest $\tau_{\text{mean}}$ of 0.89 $\mu$s is observed for NaBr$_{0.73}$I$_{0.27}$Eu$_{0.09}$, where pure NaBr:Eu$^{2+}$ with a low Eu-doping would show $\tau_{\text{rel}} = 1.05 \pm 0.1 \mu$s and NaI:Eu$^{2+}$ would show $\tau_{\text{rel}} = 1.05 \pm 0.03 \mu$s, as can be found in the Supporting Information. Lowering the concentration of Eu would increase the decay time, and thus the luminescence quantum yield. This relation is confirmed by Figure 5f. In Figure 5f we can also see that the estimated quantum yield for compounds closer to NaI:Eu$^{2+}$ rises more quickly with Eu$^{2+}$ concentration than for those closer to NaBr:Eu$^{2+}$. We can therefore expect that NaI:Eu$^{2+}$-type compounds will yield a brighter phosphor, when fully optimized for photoluminescent yield.

From Figure 6, we see that the fluent shifting behavior is not limited to the emission bands, but also occurs in the excitation bands ($4f^7 [^{8}S_{7/2}] \rightarrow 4f^{6}5d^1$). The crystal field splitting $e_{\text{cb}}$ decreases when moving from NaBr to NaI. This decrease is a result of the larger distance between Eu$^{2+}$ and the surrounding halogen ions due to the larger ionic radius of I ions compared to Br ions. However, the emission for NaI:Eu$^{2+}$ is still at lower energy than for NaBr:Eu$^{2+}$. Hence, we can conclude that the shifting of the emission band from the substitutional defect is mainly dominated by a drop in the centroid energy, because the d-orbitals of the Eu$^{2+}$ ion can expand further when the d-orbitals of the Eu$^{2+}$ ion can expand further when.

As final confirmation of the existence of a well-mixed phase, XRD measurements show that the deposited film exhibited concentration quenching but that, when optimized for photoluminescent yield, compounds close to NaI:Eu$^{2+}$ will yield a brighter luminescent material than compounds close to NaBr:Eu$^{2+}$.

In conclusion, the XY-scanning method presented in this work makes it possible to systematically analyze a broad range of host compositions and dopant concentrations to study the behavior of luminescent materials, using only a single deposition. The results from the thin-film phosphors are highly comparable to those observed with traditional powder-based techniques. Hence, the method may provide a toolbox...
capable of accelerating the discovery and optimization of phosphors for a wide range of applications such as lighting, display and scintillation, but also for solar-conversion, afterglow, and many other types of photoactivated materials.

Experimental Procedures

The NaBr–NaI:Eu2+ gradient thin-film was fabricated in an AJA Orion 5 magnetron sputtering system with a base pressure of $1 \times 10^{-9}$ bar. A 5 cm diameter Eu metal target (99.99%, Demaco) and 5 cm diameter NaBr and NaI pressed-powder targets (both 99.99%, Alfa Aesar, pressed at 36 MPa for 5 min) were simultaneously sputtered at a 90° angle from each other at room temperature in a 4 $\times$ 10$^{-2}$ mbar Ar atmosphere (6N purity, 32 sccm flow rate) onto a 5 $\times$ 5 cm$^2$ UV fused silica substrate (PGO). Prior to deposition, the substrate was cleaned for 15 min in an ultrasonic bath with a solution of soapy water, after which it was rinsed with acetone, ethanol and DI-water, and left to dry in an oven at 200 °C for more than 1 h. The substrate and substrate carrier were loaded into the sputtering chamber while both were >100°C. DC sputter powers were fixed at 25 (0.04 nm/s), 31 (0.08 nm/s), and 10 W ($<1 \times 10^{-3}$ nm/s) respectively. The Eu target’s chimney was masked using a stainless steel mask, with a pattern of concentric holes 1.2 mm in diameter, blocking 99% of the surface of the chimney (shown in Figure 1a). This finally led to a film with an estimated thickness of 0.86 μm in the center, 0.9 μm at the side of the film closest to NaI and 0.8 μm at the side of the film closest to NaBr. Deposition happened at room temperature, without any further heat treatment. Using the same process, two reference films with only (1) NaBr:Eu2+ or (2) NaI:Eu2+ were fabricated: (1) sputter settings fixed at 40 W for NaBr (RF, 0.08 nm/s), 10 W for Eu (DC, $<1 \times 10^{-3}$ nm/s) and (2) NaI at 35 W (RF, 0.13 nm/s) and Eu at 10 W (DC, $<1 \times 10^{-3}$ nm/s). All films were sputtered for 2 h. Sputtering rate determination was carried out using a quartz crystal microbalance.

SEM/EDX analysis was carried out using a JEOL JSM-IT100 operated at 20 kV, with probing current at 70% for 2 min per measurement, leading to a total duration of 80 min. Elemental compositions were quantified at 3000X magnification, corresponding to an area of 31 $\times$ 23 μm$^2$. Low vacuum mode (35 Pa pressure) was used to facilitate quantitative elemental analysis without a conductive coating.

The sample stage of the XY-scanner was made out of two stacked Thorlabs DDSM100 linear translation stages. An Ekspla NT230 OPO Laser was used as excitation source. Two polarizers and a 500 μm diaphragm were placed in the laser beam to reduce intensity of the laser light and to avoid fluctuations of the beam position. The intensity was reduced to prevent saturation of the luminescent centers in the sample. Four % of the incident laser light was split off the main beam by a UV fused silica plate. This split-off light was directed toward a Thorlabs S120VC sensor connected to a Thorlabs PM100D calibrated powermeter for online correction of the laser power. The main laser beam was focused onto the film using a f = 100.3 mm Thorlabs LA4380 lens. Emission passed through a 355 nm long-pass filter (Semrock, BLP01–355R-25), followed by a f = 100.3 mm Thorlabs LA4380 lens. This lens focused the emission into an Ocean Optics QE600–2-SR-BX fiber. Emission was integrated during 100 ms and averaged 5 times using an Ocean Optics QE65000 CCD Spectrometer (100 μm slit, 300 grooves/mm grating, leading to an uncertainty in emission wavelength of ±1.69 nm) and corrected for the quantum efficiency and nonlinearity of the detector. Prior to any emission measurement, the background was measured using identical settings and subtracted from the final measurement, leading to a measurement duration of 1 s. 32 × 32 ($\Delta x = 1.56$ mm, $\Delta y = 1.56$ mm) emission measurements were conducted across the entire substrate, leading to a total measurement duration of 17 min.

For decay studies, the emission passed through an Acton Spect-Pro2300 monochromator coupled to a Hamamatsu R7600U-03 PMT, with the PMT linked to a CAEN DT5730 Digitizer. The digitizer was constantly recording, but started storing information upon reception of a trigger signal from the laser. When a trigger was received by the digitizer, the output of the PMT was stored ranging from ~0.6 μs before the start of a laser pulse and to ~54.6 μs after a laser pulse, with a resolution of Δt = 2 ns. Decays were recorded using 1000 laser pulses, resulting in an error for the calculated mean decay time below 1% (typically below ±7 ns). With the 100 Hz OPO laser, this yields 10 s per measurement. Here, measurements were conducted across the substrate with an 8 × 8 grid, yielding a total duration of 11 min. All devices pertaining to the XY-scanner were controlled using in-house developed Python software.

XRD measurements were performed using a PANalytical X’Pert Pro MPD diffractometer with a Cu Kα anode ($\lambda = 0.1540598$ nm) operating at 45 kV and 40 mA in a Bragg–Breitano geometry measured from $2\theta = 20^\circ$ to $70^\circ$, $\Delta 2\theta = 0.0083731^\circ$ in 1 h. The area illuminated by the X-ray beam was around 1 $\times$ 5 mm$^2$ in size. The film was placed in a hygroscopic sample holder. The measurement locations on the film were set using a manually operated homemade xy-micromanipulator stage.

Associated Content

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscombsci.8b00068.

Comparative excitation and emission measurements of a common Y$_2$Al$_5$O$_{12}$:0.01 Ce$^{3+}$ LED phosphor, photographs (UV/visible), position-dependent decay spectra, and estimated quantum yields of the pure NaI:Eu$^{2+}$ and NaBr:Eu$^{2+}$ films (PDF)

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E.M. and E.K. conceived and designed the experiments, E.M. and E.K. cowrote the manuscript.

Notes

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