Exploration of New Phosphors Using a Mineral-Inspired Approach in Combination with Solution Parallel Synthesis

Masato Kakihana¹, Jihae Kim¹, Tetsufumi Komukai², Hideki Kato¹, Yasushi Sato¹, Makoto Kobayashi¹, Yuji Takatsuka²

¹Institute of Multidisciplinary Research for Advanced Materials, Tohoku University, Sendai, Japan
²Ichikawa Research Laboratories, Sumitomo Metal Mining Co. Ltd., Ichikawa, Japan

Email: kakihana@tagen.tohoku.ac.jp

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ABSTRACT

The concept, as well as the methodology of using a mineral-inspired approach in combination with solution parallel synthesis (SPS) for exploration of new phosphors among Na/Sr(Ba)/Al-silicate and Zr(Ti)-silicate minerals, is reported. By employing the proposed approach, we have discovered new phosphors of NaAlSiO₄:Eu²⁺ and BaZrSi₃O₉:Eu²⁺, that emit green-yellow (553 nm) and blue-green (480 nm) light, respectively, when excited by radiation of 290 to 420 nm.

Keywords: Mineral; Solution Parallel Synthesis; Exploration of New Phosphors; Silicate

1. Introduction

Research aimed at the discovery of new phosphors has been dependent on the knowledge of solid-state physics and crystal chemistry. Generally, the exploration of new phosphors involves the repetition of a sequential process: 1) devising the composition of a chosen candidate substance, 2) its synthesis, 3) characterization of the fluorescence of the synthesized material, 4) modification of the original composition, 5) its re-synthesis, and so on. However, it should be noted that the aforementioned process needs to be repeated several times to develop a new phosphor. Therefore, the process of discovering a new phosphor requires an enormous amount of time and effort. As a consequence, the search for new phosphors was confined to a limited composition range.

One of the effective ways of addressing the challenge is to employ generic algorithm-assisted combinatorial chemistry (GACC), which was applied to the development of heterogeneous catalysts in the recent past [1-3]. GACC has been successfully utilized by Kee-Sun Sohn’s group for the exploration of new phosphors for the use in plasma displays and light-emitting diodes (LEDs) [4-8]. More recently, the same research group employed high-throughput combinatorial chemistry in combination with an advanced computational approach based on particle swarm optimization [9], and executed a so-called particle swarm optimization-assisted combinatorial materials search (PSOCMS) to discover novel phosphors such as Ce₂.6Ca₁.₂Si₁₂O₄₄N₁₆.₆-Eu²⁺ and La₂.₅₄Ca₁.₄₂₅Si₁₂O₄₄N₁₆.₅₄-Eu²⁺ [10]. Considerable attention has been paid to the following in both GACC and PSOCMS: 1) development of computational algorithms to make the high-throughput combinatorial screening efficient and 2) shortening of the time needed for the final identification of promising compositions. On the other hand, lesser attention has been devoted to the method of the material fabrication itself although it is crucial to establish the methodology for the reliable synthesis of phosphors, as this ensures homogeneity of compositions and uniform distribution of activators in given host compounds. In most previously reported studies based on GACC/PSOCMS as well as the synthetic studies on phosphors, researchers have primarily relied on the traditional method based on solid-state reactions. As a result, precise control of the compositions of phosphors and achievement of a homogeneous distribution of small amounts of activators in the host compounds were not possible. Therefore, we propose an alternative methodology involving a mineral-inspired approach in combination with the so-called solution parallel synthesis (SPS) method [11].

In the SPS method, tens of types samples are synthesized at once under the same conditions by employing the “polymerizable complex (PC) method” [12-14] or the “amorphous metal complex (AMC) method” [12-14]. This results in relatively simple and efficient screening of
the phosphor candidate substances, which was difficult through conventional techniques. The PC and AMC methods are best suited for homogeneous synthesis of ceramics with complex compositions. The scheme for the SPS method is summarized in Figure 1. We explored new phosphors containing silicon by focusing on silicon of Clarke number 2 and employing the SPS method. There are more than 18,000 inorganic crystalline compounds containing silicon, a common element [15]. However, since most of them are found in natural minerals, their artificial synthesis is difficult. As a result, their application as functional ceramics, e.g., phosphors, is uncommon.

The obstacle in the artificial synthesis of compounds containing silicon imitating minerals comes from the lack of an appropriate silicon source that is required when an advanced solution technology has to be employed to obtain high-quality Si-containing ceramic materials. In fact, the only silicon sources that can be used in a given solution process are “volatile” tetraalkoxysilanes (TAS). However, the use of TAS is not recommended, since TAS evaporates when subjected to repeated heat treatment, as a result of which the composition changes greatly from the original. For this reason, the discovery of new phosphors from silicon-containing compounds regarded as a cornucopia of functions is challenging. We systematically synthesized silicon-containing phosphors using nonvolatile and nonflammable glycol-modified silane (GMS) [16], which was uniformly dispersed in water.

The accomplishment of the reliable synthesis of silicon-containing phosphors from GMS has heralded the development of new silicon-containing phosphors by the SPS method.

In this paper, we report a process for exploring new phosphors by the SPS method based on a mineral-inspired approach. Specifically, we detail the discovery of new phosphors of silicate minerals, nepheline, i.e., NaAlSiO4:Eu2+ (green-yellow) [21] and benitoite, i.e., BaZrSi3O9:Eu2+ (blue-green) [22], by the SPS method utilizing water-dispersible GMS. We have first described the method for the synthesis of GMS, followed by the importance of the use of GMS in the SPS method.

2. Glycol-Modified Silane (GMS): Water-Dispersible Silicon Compound

Until now, we reported on the synthesis of silicon-containing phosphors [16-22] through the aqueous solution method involving GMS obtained by reacting tetraethoxysilane (TEOS)—a typical silicon alkoxide—with glycol. For example, silane modified with propylene glycol (PG) (propylene-glycol-modified-silane; PGMS) can be synthesized in the presence of hydrochloric acid as a catalyst by reacting TEOS with PG at 80°C for a couple of minutes. GMS has a distinct property of reacting with water to form silicon clusters [23], such as Si(OH)4 and (OH)3-Si-O-Si(OH)3, and of distributing uniformly in water, unlike other conventional Si compounds such as TEOS. While GMS is important in the synthesis of silicon-containing phosphor, it is indispensable in the discovery of new silicon-containing phosphors by the SPS method.


We describe herein the conditions employed for the search of new phosphors by the SPS method. The success or failure of the search for new substances hinges on the effectiveness of the composition library that is established. We chose to use “minerals as models” in our artificial composition library. By employing the SPS method, we recently discovered NaAlSiO4:Eu2+ phosphor, which emits a strong green-yellow light (553 nm) when excited by near ultraviolet radiation of 290 to 420 nm. We first describe the sequence of steps implemented in the discovery of the aforementioned phosphor below and, then, introduce BaZrSi3O9:Eu2+ phosphor discovered by the same method.

Figure 1. Scheme of solution parallel synthesis scheme for exploration of new phosphors in A,B2AlSi2O6:Eu2+ (A = Li, Na, K; B = Sr, Ba) systems. GMS stands for “glycol-modified silane,” which was used as the silicon source. Visual observation of emission light by illumination with near-UV light is also shown as the image.
3.1. NaAlSiO₄:Eu²⁺ Phosphor

Step 1: Establishment of a library of artificial compositions using minerals as models.

First, we extracted alkaline aluminum silicate-based substances of mineral origin from the database. Eleven minerals that were specifically chosen are listed in Table 1. Additionally, two more minerals based on alkaline-earth/alkali metals and aluminum silicate, which include banalsite and stronalsite, are listed in Table 1. Of the artificial compositions in Table 1, the standard composition was created by removing water of crystallization (‘‘•nH₂O’’) from original mineral compositions. We have taken into consideration artificial solid solutions of Na and Sr in sodium-aluminum silicates assuming that a certain portion of sodium in a given sodium-aluminum silicate can be replaced by Sr. For instance, in the case of jadeite NaAlSi₂O₅, (1st column in Table 1), two possible solid solutions have been created artificially, i.e., Na₀.₉₅Sr₀.₀₅Al₁.₀₅Si₁.₉₅O₆ and Na₀.₉Sr₀.₁Al₁.₁Si₁.₉O₆. Owing to the mismatch in the charges between Na⁺ and Sr²⁺, the charge neutrality in the artificial composition was maintained by adjusting the molar ratio of Al to Si. In the case of banalsite, BaNa₂Al₅Si₇O₂₈·₆H₂O, two possible solid solutions, i.e., Ba₀.₆Sr₀.₄Na₂Al₅Si₇O₂₈ and Ba₀.₈Sr₀.₂Na₂Al₅Si₇O₂₈, have been artificially created as shown in the penultimate row in Table 1. Similarly, we have constructed 27 artificial compositions, including the standard compositions. Thus, the list in Table 1 is referred to as a “composition library” inspired by minerals.

Step 2: Preparation of a mixed solution in accordance to the library of compositions.

An aqueous solution of each element that appeared in the library was prepared. The concentration of each metal was analytically determined prior to its use. In addition, an aqueous solution of europium was also prepared. The mixed solution with a composition corresponding to that mentioned in the library was prepared in a test tube, and Eu, the activator, was then added to the test tube so that its amount is 1 mol% for Na/Sr or Ba/Sr (when necessary, charge compensation was achieved by adjusting the ratios of Al and Si). Finally, citric acid was added as a metal stabilizer and glycol was added as a polymerizing agent.

Table 1. Library of artificial compositions inspired by natural minerals of AₓBᵧAlₓSiₘO₁₆ (A = Li, Na, K; B = Sr, Ba) systems.

<table>
<thead>
<tr>
<th>No.</th>
<th>Name of Mineral</th>
<th>Original compositions of minerals</th>
<th>Artificial compositions inspired by natural minerals</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Standard compositions</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5% for Na and 20% for Ba</td>
</tr>
<tr>
<td>1</td>
<td>Jadeite</td>
<td>NaAlSi₂O₅</td>
<td>NaAlSi₂O₅</td>
</tr>
<tr>
<td>2</td>
<td>Albite</td>
<td>NaAlSi₃O₈</td>
<td>NaAlSi₃O₈</td>
</tr>
<tr>
<td>3</td>
<td>Nepheline</td>
<td>NaAlSiO₄</td>
<td>NaAlSiO₄</td>
</tr>
<tr>
<td>4</td>
<td>Natrolite</td>
<td>Na₂[Al₂Si₃O₁₀]·2H₂O</td>
<td>Na₂Al₅Si₇O₁₈</td>
</tr>
<tr>
<td>5</td>
<td>Barrerite</td>
<td>Na₃[Al₂Si₇O₁₄]·6H₂O</td>
<td>Na₃Al₇Si₉O₂₈</td>
</tr>
<tr>
<td>6</td>
<td>Gobbinsite</td>
<td>Na₅[Al₅Si₁₁O₃₂]·12H₂O</td>
<td>Na₅Al₇Si₇O₂₈</td>
</tr>
<tr>
<td>7</td>
<td>Orthoclase</td>
<td>KAlSi₅O₁₆</td>
<td>KAlSi₅O₁₆</td>
</tr>
<tr>
<td>8</td>
<td>Kalsilite</td>
<td>KAlSi₅O₁₆</td>
<td>KAlSi₅O₁₆</td>
</tr>
<tr>
<td>9</td>
<td>Leucite</td>
<td>KAlSi₅O₁₆</td>
<td>KAlSi₅O₁₆</td>
</tr>
<tr>
<td>10</td>
<td>Lithosite</td>
<td>K₅AlSi₅O₁₆·2·H₂O</td>
<td>K₅AlSi₅O₂₅</td>
</tr>
<tr>
<td>11</td>
<td>Petalite</td>
<td>LiAlSi₅O₁₀</td>
<td>LiAlSi₅O₁₀</td>
</tr>
<tr>
<td>12</td>
<td>Banalsite</td>
<td>BaNa₂Al₅Si₇O₂₈·₆H₂O</td>
<td>BaNa₂Al₅Si₇O₁₆</td>
</tr>
<tr>
<td>13</td>
<td>Stronalsite</td>
<td>SrNa₂Al₅Si₇O₂₈</td>
<td>SrNa₂Al₅Si₇O₁₆</td>
</tr>
</tbody>
</table>
**Step 3:** Parallel, simultaneous chemical processes, such as polyesterification (fabrication of polyester resin).

These test tubes were placed in an oven and heated, and the mixture in them was concentrated at 120°C for 12 h to promote polyesterification.

**Step 4:** Heat treatment processes for production of precursors and inducing fluorescence by final heat treatment.

The test tube containing polyester resin was heat-treated in a sand bath at 500°C for 2 h, for degreasing. The degreased substances were transferred to crucibles and fully baked in the electric furnace in the presence of graphite as a reducing agent at 1200°C for 2 h for simultaneous synthesis of many kinds of samples.

**Step 5:** Visual examination of light emission.

The appearance of luminescence of oxides thus obtained, through illumination by ultraviolet light (254 nm) or near-ultraviolet lights (365 nm and 400 nm), is shown in Figure 2. The image clearly indicates that many of the Na(K)/Sr(Ba)/Al/Si-oxide systems are hosts for blue-to-yellow luminescence originating from Eu²⁺.

**Step 6:** Choice of promising phosphors: Re-synthesis based upon rational inspection.

Phase identification by X-ray diffraction measurement and the evaluation of photoluminescence properties by excitation and emission-spectrum measurement were performed to choose promising phosphors. In the specific case of Eu²⁺-doped Na(K)/Sr(Ba)/Al/Si-oxide systems, further reduction of a stable trivalent europium (Eu³⁺) to a less stable divalent europium (Eu²⁺) was found to be necessary. In order to reduce Eu³⁺ to Eu²⁺ completely, additional heat-treatment under a strongly reducing atmosphere was carried out by passing N₂/NH₃ (90/10 mol%) gas at 1200°C for 2 h. As expected, the luminescence characteristics were significantly improved due to the increased amount of available Eu²⁺. Figure 3 shows images depicting luminescence of some representative samples after reduction by N₂/NH₃. The nepheline sample with the nominal composition of NaAlSiO₄:Eu²⁺, the natrolite sample with the nominal composition of Na₂Al₂Si₃O₁₀:Eu²⁺, and their Sr-substituted derivatives emit strong green-to-yellow lights when illumination with UV/near-UV lights between 254 nm and 400 nm. Careful phase identification by X-ray diffraction indicated that while NaAlSiO₄ was found in the nepheline sample, the same was not true for the natrolite sample; instead a mixed phase of NaAlSiO₄ and SiO₂ formed. These observations led us to the conclusion that it is nepheline that emits strong green-to-yellow lights in samples with the nominal compositions of NaAlSiO₄:Eu²⁺ and Na₂Al₂Si₃O₁₀:Eu²⁺.

The excitation and emission spectra of the NaAlSiO₄:Eu²⁺ phosphor discovered in the aforementioned manner are shown in Figure 4. While this phosphor displays weak emission when excited by visible light, it emits strong green-yellowish light with excitation by near ultraviolet light of 320 nm. In addition, the emission spectrum is comprised of one simple broad component with a peak maximum at 553 nm.

Our future goal is to achieve a redshift (long-wavelength shift) in an excitation spectrum. In oxynitride, wherein a portion of oxygen was replaced with nitrogen, a redshift in the excitation spectrum was observed along with a slight increase in intensity at 400 nm [21].

### 3.2. BaZrSi₃O₉:Eu²⁺ Phosphor

Like in the case of NaAlSiO₄, we extracted zirconium (titanium) silicate-based substances of mineral origin from the database. Of the elements present in original mineral compositions, we replaced those that were un-
Figure 3. Appearance of luminescence in some representative samples upon illumination of the samples with ultraviolet light (254 nm) or near-ultraviolet light (365 nm and 400 nm) before and after reduction by N\textsubscript{2}/NH\textsubscript{3}. Figures on each image indicate the wavelength (nm) used for excitation.

Figure 4. Excitation and emission spectra of the nepheline-origin NaAlSiO\textsubscript{4}:Eu\textsuperscript{2+} (1 mol% for Na) discovered by employing the SPS method.

suitable for phosphors. For instance, Fe and Ti were replaced with elements having similar ionic radii such as Mg and Zr, respectively. We replaced the removed artificial composition for water of crystallization (“•nH\textsubscript{2}O”). In addition, we took into consideration the solid solutions comprising alkaline earth metals Ca, Sr and Ba and alkali metals Li, Na, and K. Thus, as depicted in Table 1, 49 artificial compositions were established using each mineral as a model, and a part of the artificial composition library established in this way is shown in Table 2. As in the case of NaAlSiO\textsubscript{4}, we employed Eu\textsuperscript{2+} as an activator for all the 49 artificial compositions (typically 3 mol% Eu\textsuperscript{2+} for Na or Ba).

After careful inspection of the result obtained by the SPS experiment, the BaZrSi\textsubscript{3}O\textsubscript{9}:Eu\textsuperscript{2+} phosphor, emitting blue-green light (480 nm), was found. Its emission intensity was comparable to the maximum intensity of a commercially available BAM (BaMgAl\textsubscript{10}O\textsubscript{17}:Eu\textsuperscript{2+}) phosphor. The light emission when irradiated with 405 nm of near ultraviolet light is shown in Figure 5, and the excitation and emission spectra of the same sample are shown in Figure 6 in comparison with that of BAM. It is worthwhile to note that in contrast to BAM, a flat excitation spectrum was observed for BaZrSi\textsubscript{3}O\textsubscript{9}:Eu\textsuperscript{2+} over the range of 290 nm to 420 nm. The intensity of luminescence due to excitation with 405 nm was more than twice that of BAM. In conclusion, BaZrSi\textsubscript{3}O\textsubscript{9}:Eu\textsuperscript{2+} is considered to be useful as a blue-green phosphor for use in near-ultraviolet LEDs [22,24].

4. Concluding Remarks

We believe that research involving exploration for new phosphors is still in infancy. The key features that are common to the currently known useful phosphors are cations of elements such as silicon, phosphorus, aluminum, gallium, alkaline earth metals, and alkali metals, and anions of elements such as oxygen, nitrogen, sulfur, and halogens. The possible combinations of these elements result in astronomical options. As a result, no one can deny the possibility of the existence of new substances and new phosphors. The principle of synthesis of inorganic phosphors, which are categorized as ceramics, differs fundamentally from organic chemical synthesis on the basis of the reaction between molecules. Further, due to slow movement of cations in the solid lattice, inorganic phosphor synthesis is not simple and generally spans a long duration. Moreover, while in organic synthesis, it is possible to refine the final product by techniques such as chromatography, in phosphor synthesis, it is almost impossible to refine only the target phase separately, if a mixed phase sample is obtained. Unless these fundamental challenges are redressed in phosphor syn-
Table 2. Library of artificial compositions inspired by natural minerals of zirconium (titanium) silicate systems.

<table>
<thead>
<tr>
<th>No.</th>
<th>Name of Mineral</th>
<th>Original compositions of minerals</th>
<th>Artificial compositions inspired by natural minerals</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Standard compositions</td>
</tr>
<tr>
<td>1</td>
<td>Armstrongite</td>
<td>CaZrSi₆O₁₅·2.5H₂O</td>
<td>CaZrSi₆O₁₅</td>
</tr>
<tr>
<td>2</td>
<td>Benitoite</td>
<td>BaZrSi₃O₉</td>
<td>BaZrSi₃O₉</td>
</tr>
<tr>
<td>3</td>
<td>Calciohilarite</td>
<td>CaZrSi₃O₉·H₂O</td>
<td>CaZrSi₃O₉</td>
</tr>
<tr>
<td>4</td>
<td>Cataplelite</td>
<td>Na₂ZrSi₃O₉·2H₂O</td>
<td>Na₂ZrSi₃O₉</td>
</tr>
<tr>
<td>5</td>
<td>Elpidite</td>
<td>Na₂ZrSi₁₅O₉·3H₂O</td>
<td>Na₂ZrSi₁₅O₉</td>
</tr>
<tr>
<td>6</td>
<td>Parakeldyshite</td>
<td>Na₂ZrSi₅O₉</td>
<td>Na₂ZrSi₅O₉</td>
</tr>
<tr>
<td>7</td>
<td>Terskite</td>
<td>Na₂ZrSi₁₅O₉·3H₂O</td>
<td>Na₂ZrSi₁₅O₉</td>
</tr>
<tr>
<td>8</td>
<td>Aenigmatite</td>
<td>Na₂Fe₅TiSi₆O₂₀</td>
<td>Na₂Mg₅ZrSi₆O₂₀</td>
</tr>
<tr>
<td>9</td>
<td>Titanite</td>
<td>CaTiSiO₅</td>
<td>CaZrSiO₅</td>
</tr>
<tr>
<td>10</td>
<td>Natisaite</td>
<td>Na₂TiSiO₅</td>
<td>Na₂ZrSiO₅</td>
</tr>
</tbody>
</table>

Figure 5. Appearance of the powder of BaZrSi₃O₉:Eu²⁺ (a) and its emission by illumination of 400 nm light (b).

In this paper, we described the process parameters involved in the exploration of new phosphors using the SPS method based on a mineral-inspired approach. We would like to reiterate the importance of glycol-modified silane (GMS) in the implementation of the SPS method for preparing new phosphors from silicates. Using this method, we recently found Eu²⁺-activated oxide phosphor that emitted red-orange light (625 nm) when excited with blue light [25]. Moreover, we are currently developing new phosphorus-containing phosphors using new phosphate compounds applicable to the SPS method [26]. It is expected that the SPS method will become popular in the near future and this will inject new vigor into the field of phosphor exploration.

Figure 6. Excitation and emission spectra of the benitoite-origin BaZrSi₃O₉:Eu²⁺ (3 mol% for Ba) discovered by employing the SPS method.

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