Synthesis of Nitride Zeolites in a Hot Isostatic Press

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Abstract: The recently introduced nitridophosphate synthesis in a hot isostatic press (HIP) enabled simple access to large-scale product quantities starting from exclusively commercially available starting materials. Herein, we show that this method is suitable for the synthesis of highly condensed functional nitridophosphates, as well. Hence, the syntheses of the nitridophosphate zeolites $\text{Ba}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{X} = \text{Cl}, \text{Br}$) are presented as proof of concept for this innovative approach. Furthermore, samples of unprecedented $\text{Sr}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{X} = \text{Cl}, \text{Br}$) were prepared and characterized to demonstrate the advantages of this synthetic approach over commonly used methods. Luminescence investigations on Eu$^{2+}$-doped samples of $\text{AE}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{AE} = \text{Sr, Ba}; \text{X} = \text{Cl, Br}$) were carried out and characteristics of observed emission bands are discussed.

The versatile compound class of nitridophosphates has been researched in detail for more than 30 years. The structural diversity of nitridophosphates can be derived from their close relationship to oxosilicates, as the element combination Si/O is isoelecronic with P/N, and similar with oxosilicates which are built up from SiO$_4$ tetrahedra, the PN$_4$ tetrahedron is the fundamental building unit in nitridophosphates. However, nitridophosphates can even feature edge-sharing tetrahedra due to the higher covalence (and lower polarity) of P–N bonds compared to Si–O in oxosilicates.[1,2] Additionally, the higher valence of nitrogen allows for higher degrees of condensation (i.e. atomic ratio $\kappa$ of tetrahedra centers to tetrahedra vertices).[1,3] But despite numerous investigations, nitridophosphate synthesis has ever been challenging, as the most common starting material phosphorus(V) nitride (P$_4$N$_3$) is prone to thermal decomposition above 850°C.[4] Therefore, high-pressure high-temperature methods (e.g. multianvil technique) have been employed for their synthesis, since elevated nitrogen pressure suppresses the thermally induced elimination of N$_2$ from P$_4$N$_3$.[1] This high-pressure strategy led to a large number of nitridophosphates with various incorporated electropositive elements.[5–9] However, sample quantities have intrinsically been limited by high-pressure techniques. Investigations on the associated optical and physical properties of nitridophosphates, such as ion conductivity or luminescence, have however quickly revealed the potential of this functional materials class.[10–12] In particular, the intriguing luminescence properties of nitridophosphates like $\text{Ba}_3\text{P}_5\text{N}_{10}\text{X}:\text{Eu}^{2+}$ ($\text{X} = \text{Cl, Br}$, I) and $\text{AE}_3\text{P}_5\text{N}_{10}\text{X}:\text{Eu}^{2+}$ ($\text{AE} = \text{Ca, Sr, Ba}$) clearly underline the quest for a synthetic approach that can be transformed to a large batch scale.[13–15] Here, ambient and medium-pressure methods that have been used for the early syntheses of phosphorus nitrides and nitridophosphates come to mind.[16–23] However, only a limited number of such nitride compounds could be synthesized applying these techniques, as gentle conditions or tailored starting materials had to be used. This limitation has changed only recently, as improved high-temperature ammonothermal techniques and the use of red phosphorus as starting material enabled synthesis of diverse nitridophosphates.[24–27] However, ammonothermal syntheses of nitridophosphates may not be performed industrially due to its demanding handling and the stated goal for a large-scale access remained. Recently, we have reported on the successful synthesis of $\text{Ca}_2\text{P}_3\text{N}_6$ in a hot isostatic press (HIP) under nitrogen pressure, which thus appears as a promising innovative approach for nitridophosphate synthesis.[28] HIPs do not only provide large sample quantities, but also shorten reaction times and facilitate crystal growth under comparatively gentle reaction conditions. Moreover, we have demonstrated that red phosphorus can serve as a starting material in HIPs as well, simplifying nitridophosphate syntheses even more.[29]

Within the scope of this work, this approach is further developed to grant access to highly condensed nitridophosphates with a degree of condensation $\kappa = 1/2$. For this purpose, $\text{Ba}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{X} = \text{Cl, Br}$) were chosen as model compounds, because of their above-mentioned luminescence properties.[11,13] Furthermore, hitherto unknown compounds $\text{Sr}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{X} = \text{Cl, Br}$) were synthesized and investigated with regard to luminescence properties.

All alkaline earth nitridophosphate zeolites $\text{AE}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{AE} = \text{Ba, Sr}; \text{X} = \text{Cl, Br}$) were synthesized under nitrogen atmosphere in a HIP applying hot isostatic conditions (150 MPa N$_2$, 1000°C; Figures S1 and S2). In an initial attempt, the title compounds were synthesized from specially prepared P$_4$N$_3$ and the respective alkaline earth azides and halides following Equation (1) (Table S1).

$$15\text{AE(N)}_3 + 3\text{AE}_2\text{X}_2 + 10\text{P}_4\text{N}_3 \rightarrow 6\text{AE}_3\text{P}_5\text{N}_{10}\text{X} + 40\text{N}_2$$

(1)

Subsequently, all variants of $\text{AE}_3\text{P}_5\text{N}_{10}\text{X}$ ($\text{AE} = \text{Ba, Sr}; \text{X} = \text{Cl, Br}$) were prepared using commercial red phosphorus.
Starting material for laboratory synthesis of P₃N₅ and may the structure of Sr₃P₅N₁₀Cl as starting model (Figure S5, Table S7). Both compounds crystallize homeotypically to the 8; Sr₃P₅N₁₀Cl:

Subsequent oxidation into P⁵⁺ is coupled to the disproportionation of azide ions into network-forming nitride anions and elemental nitrogen. Thereby, it is conceivable that activated P₃ is gradually oxidized and present as P⁴⁺ in form of molecular “PN” after initial reaction with N₂. In a second oxidation step, halide-containing intermediates, such as (P⁴⁺NCl₂), might be formed. (PNCl₂)₃ in turn serves as starting material for laboratory synthesis of P₃N₅ and may facilitate the in situ formation of the latter, leading to the desired zeolites AE₃P₅N₁₀X. An experimental evidence for this hypothesis by in situ measurements is still pending.

To investigate luminescence properties of AE₃P₅N₁₀X, Eu²⁺-doped samples were prepared by adding 3 mol % EuCl₃ (with regard to alkaline earth ions) to the mixture of starting materials.

The undoped samples are yielded as colorless cube-like crystals, while sinter cakes of Eu²⁺-doped products exhibit yellow (Ba compounds) to orange body colors (Sr compounds). All products have been washed with de-ionized water and are not sensitive towards air or moisture. SEM imaging of the products reveals the microcrystalline character of Ba compounds (edge length up to ≈3 μm, Figure 1). Sr compounds form slightly larger crystals with edge lengths up to 15–20 μm (Figure 1). Phase purity of Ba₃P₅N₁₀Cl (X = Cl, Br) was confirmed by Rietveld refinements, using literature known structures as starting models.[11,13] Detailed information on the refinements is provided in the Supporting Information (Figures S3 and S4, Table S2).

The structures of Sr₃P₅N₁₀X (X = Cl, Br) were elucidated from a single-crystal XRD measurement of Sr₃P₅N₁₀Cl. Sr₃P₅N₁₀Br was refined using the Rietveld method and using the structure of Sr₃P₅N₁₀Cl as starting model (Figure S5, Table S7). Both compounds crystallize homeotypically to the Ba compounds in the JOZ zeolite structure type (Pnaa; Z = 8; Sr₃P₅N₁₀Cl: a = 12.240(3); b = 12.953(3); c = 13.427(3) Å; Sr₃P₅N₁₀Br: a = 12.297(1); b = 12.990(1);

![Figure 1. SEM images of obtained AE₃P₅N₁₀X samples (AE = Sr, Ba; X = Cl, Br) containing single crystals.](image1)

![Figure 2. Projection of the crystal structure of Sr₃P₅N₁₀Cl along [100]: P₅N₅ tetrahedra blue, Cl atoms pink, Sr₁–Sr₄ atoms gray, split position of Sr5 black. All atoms are displayed with 95% probability.[29](c = 13.458(1) Å)[29–31] The crystal structure of Sr₃P₅N₁₀X is shown exemplarily for X = Cl in Figure 2. The as-refined crystallographic data is summarized in the Supporting Information (Tables S3–S5) and a more detailed description of the crystal structure is provided in literature.[11,13] Phase purity of Sr₃P₅N₁₀Cl has been confirmed by Rietveld refinement (Figure S6, Table S7). In contrast to the Ba compounds, Sr₃P₅N₁₀X (X = Cl, Br) show split positions of the Sr5 site. Although the split position is not present in the Ba compounds the corresponding Ba5 site in Ba₃P₅N₁₀X, however, shows the most elongated ellipsoids and Ba–X distances (Figure S7).[11,13] This observation can be explained by the significantly smaller space filling of Sr²⁺ compared to Ba²⁺ and the associated increasing displacement of ions.[32,33] Moreover, owing to the larger radius of Br⁻, the distance between the split position is reduced in Sr₃P₅N₁₀Br (Figure S8).

These observations may explain the reason for Sr₃P₅N₁₀X (X = Cl, Br) being exclusively accessible at medium-pressure conditions, as the pressure has to be well balanced. Although increased pressure is necessary to prevent thermal decomposition, it must not be chosen too high for Sr₃P₅N₁₀X syntheses to prevent collapsing of the large cages, considering the reduced space filling of the Sr²⁺ ions. In line with this hypothesis, the Ba compounds are rather difficult to access at ambient pressure, as well, and phase pure samples have only been obtained at 1–5 GPa, thus far.[11,13]

The interatomic P–N distances and N–P–N angles of Sr₃P₅N₁₀X (X = Cl, Br) are in very good agreement with values reported for other nitridophosphates (Table S9).[11–13] In line with previous refinements, the observed AE–N/X distances differ significantly depending on the coordination number of the alkaline earth metal ion (Figure S8). A detailed list of the interatomic AE–N/X distances is provided in literature for Ba₃P₅N₁₀X and in the Supporting Information for Sr₃P₅N₁₀X (Table S8).[11,13]

The chemical compositions of the title compounds were confirmed by energy dispersive X-ray spectroscopy (EDX),
The two emission maxima likely correspond to the different Eu\(^{3+}\) coordination spheres that are provided by the host lattice. The sites \(AE1\), \(AE4\), and \(AES\) are coordinated by eight \(N\) and two \(X\) ions (\(CN=10\)) and thus, feature rather elongated \(AE-X\) and \(AE-N\) distances, causing a weak crystal field (Figure S8, Table S8). Therefore, the higher energetic emission bands can be assigned to Eu\(^{3+}\) ions occupying these sites according to the parity-allowed transition \(4f^7 \rightarrow 4f^5 5d^1\). Consequently, the second emission band is assigned to Eu\(^{3+}\) ions located on \(AE\) sites with lower coordination number \((AE2, AE3: CN=8, six \(N\) and two \(X\) ions), which feature shorter \(AE-X\) and \(AE-N\) distances and thus a strong crystal field (Figure S8, Table S8). A detailed illustration of the emission and excitation spectra for each element combination \(AE-X\) is provided in the Supporting Information (Figures S10–S13).

When comparing the emission spectra, two trends are particularly striking. First, emission bands of \(AE\), \(P\), \(N_{10}X\), and \(AE\), \(P\), \(N_9\), \(Br\) are shifted comparing \(AE=\) Ba and \(AE=\) Sr. This effect is attributable to the different \(AE\) radii and the associated \(AE-N/X\) and Eu\(^{3+}-N/X\) distances, leading to an increased crystal field splitting for Eu\(^{3+}\) on the Sr site and a red-shift in emission. The second trend describes the influence of the halide ions on the position of the emission maxima. While the higher energetic bands are shifted red with increasing size of \(X\), lower energetic bands are shifted blue. A detailed discussion of these observations is provided in the Supporting Information.

Furthermore, measurements of the internal quantum efficiency have been carried out for non-optimized powder samples of the title compounds. The IQE of Sr\(_3\)P\(_5\)N\(_{10}\):Eu\(^{3+}\) was determined to 29\% \((X=\text{Cl})\) and 32\% \((X=\text{Br})\). Measurements on Ba\(_3\)P\(_5\)N\(_{10}\):Eu\(^{3+}\) yield quantum efficiency of 12\% each, which shows potential for improvement of the investigated samples, as values of >60\% have been reported for Ba\(_3\)P\(_5\)N\(_{10}\):Eu\(^{3+}\) \((X=\text{Cl}, \text{Br})\), previously\([11,13]\).

The presented results suggest that the pressure range generally applied under HP/HT conditions may exceed the actually required synthesis pressure for nitridophosphates by far. Since the minimum pressure to suppress thermal decomposition is not known, numerous published and some novel nitridophosphates are already accessible under medium-pressure conditions. Consequently, future investigations may focus on synthesis of Ca\(_3\)P\(_5\)N\(_{10}\)X, other P/N based zeolites (e.g., \(NPO\), \(N\)), and even higher condensed nitridophosphates (e.g., \(AE\), \(P\), \(N\)). In particular, the successful activation of red phosphorus as starting material could contribute to a considerable acceleration of these investigations. It should also be examined whether other starting materials, such as nitrides, can be replaced by precursors, like metals or alloys. The fact that \(N_2\) can serve as necessary redox partner has already been shown in the synthesis of Ca\(_3\)P\(_5\)N\(_{10}\)\(_2\),\([28]\). These significant simplifications in synthesis, the wide range of achievable degrees of condensation and the large sample volumes may allow that nitridophosphates could find their way into industrial application as phosphor materials.

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Conflict of interest

The authors declare no conflict of interest.

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[29] Crystal data of Sr$_3$P$_5$N$_{10}$ (orthorhombic, Pnma (no. 62)): $X = \text{Cl}$: $M = 593.26$ g mol$^{-1}$, $a = 12.241(2)$, $b = 12.953(3)$ and $c = 13.427(3)$ Å, $V = 2126.9(8)$ Å$^3$, $Z = 8$, $\rho = 3.702$ g cm$^{-3}$, $\mu = 15.980$ mm$^{-1}$. Mo-K$_{\alpha}$ ($\lambda = 0.71073$ Å, Bruker D8 Venture), $T = 293$ K, 33698 observed reflections, 3522 independent reflections, 116 parameters, $R_{int} = 0.036$, $R_1 = 0.072$, $R_{wp} = 0.069$, $GoF = 1.065$, residual electron density $3.360$, $-1.277$ e Å$^{-3}$; $X = \text{Br}$: $M = 637.70$ g mol$^{-1}$, orthorhombic, Pnma (no. 62), $a = 12.2970(2)$, $b = 12.9896(2)$ and $c = 13.4585(2)$ Å, $V = 2149.76$ ($5$) Å$^3$, $Z = 8$, $\rho = 3.942$ g cm$^{-3}$, $\mu = 30.520$ mm$^{-1}$. Cu-K$_{\alpha 1}$ ($\lambda = 1.5406$ Å, Stoe StadiP), $T = 293$ K, 993 observed reflections, 101 parameters, $R_1 = 0.035$, $R_{wp} = 0.046$, $R_{int} = 0.020$, $R_{ave} = 0.022$, $GoF = 2.257$, Deposition Numbers 2011982 (for Cl) and 2012398 (for Br) contain the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service www.ccdc.cam.ac.uk/structures.
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