Crystalline phase analysis and phosphorus availability after thermochemical treatment of sewage sludge ash with sodium and potassium sulfates for fertilizer production

Hannes Herzel1 · Zeynep Aydin1 · Christian Adam1

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Abstract
Phosphorus rich sewage sludge ash is a promising source to produce phosphorus recycling fertilizer. However, the low plant availability of phosphorus in these ashes makes a treatment necessary. A thermochemical treatment (800–1000 °C) with alkali additives transforms poorly plant available phosphorus phases to highly plant available calcium alkali phosphates (Ca,Mg)(Na,K)PO4. In this study, we investigate the use of K2SO4 as additive to produce a phosphorus potassium fertilizer in laboratory-scale experiments (crucible). Pure K2SO4 is not suitable as high reaction temperatures are required due to the high melting point of K2SO4. To overcome this barrier, we carried out series of experiments with mixtures of K2SO4 and Na2SO4 resulting in a lower economically feasible reaction temperature (900–1000 °C). In this way, the produced phosphorus potassium fertilizers (8.4 wt.% K, 7.6 wt.% P) was highly plant available for phosphorus indicated by complete extractable phosphorus in neutral ammonium citrate solution. The added potassium is, in contrast to sodium, preferably incorporated into silicates instead of phosphorus phases. Thus, the highly extractable phase (Ca,Mg)(Na,K)PO4 in the thermochemical products contain less potassium than expected. This preferred incorporation is confirmed by a pilot-scale trial (rotary kiln) and thermodynamic calculation.

Keywords Phosphorus recovery · Recycling fertilizer · Calcium alkali phosphate · Silicate

Introduction
Phosphorus (P) and potassium (K) are essential elements for all life forms, and thus main compounds of fertilizers. The supply of raw materials to produce K bearing fertilizers is classified as uncritical because of large K deposits around the world and K recovery from the ocean [1]. This is different for P. Conventional P fertilizers are based on phosphate rock. These deposits are located in a few countries [2]. Europe has negligible deposits and depends nearly completely on imports [3]. Thus, countries in Europe are searching for other P sources [4]. Sewage sludge ashes (SSA) which are currently mainly landfilled [5, 6] are prominent alternative resources for P fertilizer production. P-rich SSA contains up to 12 wt.% P in Germany [5]. Recovery of P from suitable SSA types that are rich in P and low in pollutants (10,900 t/a P based on Krüger and Adam [5]) could substitute more than 10% of imported conventional fertilizers in Germany (86,100 t/a P based on Industrieverband Agrar [7]). The P recovery from SSA and other waste streams from wastewater treatment will become mandatory in Switzerland [8] and in Germany [9]. A direct application of SSA as fertilizer is avoided because it contains heavy metals which are partly above limit values of fertilizer legislation [10] and the plant availability of P is low [11]. Phosphorus in the SSA is mainly bonded in the crystalline phase whitlockite type (Ca3-x(Mg,Fe)x(PO4)2) [12] a poorly plant available P phase [13]. Thus, P recovery technologies target recycling fertilizers that comply with fertilizer legislation and contain highly plant available P phases.

Thermochemical treatment of SSA is a promising method to produce highly plant available P fertilizers and to reduce the content of toxic heavy metals [4, 10]. This process transforms the poorly plant available P from whitlockite and AlPO4 to highly plant available calcium alkali phosphates due to thermochemical reactions with alkali additives.
such as sodium sulfate (Na₂SO₄) in the temperature range 800–1000 °C [14, 15]. Previous plant growth studies confirmed that this product has plant availability comparable to triple super phosphate [16, 17]. The plant availability of P can be estimated using the chemical extraction in neutral ammonium citrate, calculated as a fraction (P_{NAC,rel}) of the total P-content [11].

The thermochemical treatment was successfully demonstrated on technical scale using Na₂SO₄ as additive during a pilot trial in 2014 [10]. The maximal P_{NAC,rel} value of 80 wt.% was satisfying to use this material for greenhouse pot experiments and a field trial [18].

In general, the formed calcium alkali phosphate is the crystalline phase CaNaPO₄ [10, 14]. Current studies postulate that calcium could be partly substituted by Mg [15, 19–21] and forms a phosphate phase comparable to Ca₁₃Mg₅Na₁₈(PO₄)₁₈ [22].

Compared to sodium (Na) containing additives, the thermochemical treatment with K-containing additives is rarely investigated. The addition of K-containing additives could be used to produce a PK- or NPK-fertilizer. The additional costs for the more expensive K additives might be compensated by the value of K in the fertilizer. Potassium hydroxide (KOH) [23] and potassium carbonate (K₂CO₃) [24] were successfully tested to convert P compounds to highly plant available CaKPO₄. Potassium sulfate (K₂SO₄) could be also a suitable additive because it is a by-product of biofuel production [25] and can be available at a reasonable price. However, first investigations at 1000 °C with K₂SO₄ as an additive showed that the reaction of the phosphate phases with K₂SO₄ is not complete due to its high melting point of 1070 °C [10]. To reduce the melting point of the additive, K₂SO₄ can be mixed with Na₂SO₄. The composition 3 Na₂SO₄*K₂SO₄ has the lowest melting point (823 °C) in this system [26].

The added Na or K-containing additives react with phosphates as well as with silicates. The reaction with silicates is thermodynamically favored and takes place first. Thus, at first alkali aluminosilicates are formed and later calcium alkali phosphates. Depending on silicon (Si) and P contents of SSA, a certain fraction of the alkali additive is consumed for the reaction with silicates and must be taken into account for total additive dosing and thus additive costs for the process [14].

In this study, we focus on the possibilities to partly substitute the alkali additive Na₂SO₄ by K₂SO₄ to increase the content of the valuable nutrient K in the recycling fertilizer. Processing with different ratios of Na₂SO₄/K₂SO₄ was investigated in laboratory-scale experiments (crucible trials) as well as in pilot-scale trials in a rotary kiln. The thermochemical products are evaluated by analyzing extractable P as well as by crystalline phase analysis. The formation of phases is compared with thermodynamic calculations.

**Materials and methods**

**Materials**

Sewage sludge ash for laboratory scale calcination experiments and the demonstration trial was taken from an industrial mono-incineration facility in Germany that was already investigated in a SSA monitoring project [5]. Dried sewage sludge (SS) originating from a large-scale wastewater treatment plant in Germany was pulverized to a particle size < 500 µm for muffle furnace experiments. Elemental compositions of SSA and SS are displayed in Table 1. Na₂SO₄ and K₂SO₄ (technical grade) was acquired from Cordenka GmbH & Co KG Obernburg, Germany and from company Kemira, Finland, respectively.

**Laboratory scale calcination experiments**

The set up for laboratory crucible trials was similar to Herzel et al. [10] and Stemmann et al. [14]. SSA was mixed with dry SS as a reducing agent in a weight ratio 5 to 1. This mixture of SSA and SS was mixed with milled Na₂SO₄ and/or K₂SO₄ (vibrating cup mill made of tungsten carbide, Siebtechnik GmbH, Mülheim/Ruhr, Germany) targeting on a molar ratio (Na + K) / P of 2 mol/mol. This molar ratio was chosen

| Component | SSA | SS | CT-Na-1000 | CT-K25-1000 | CT-K50-1000 | CT-K75-1050 | CT-K100-1150 |
|-----------|-----|----|------------|-------------|-------------|--------------|---------------|
| Al        | 6.7 ± 0.5 | 0.8 ± 0.1 | 6.2 ± 0.1 | 5.9 ± 0.2 | 6.4 ± 0.1 | 5.9 ± 0.2 | 5.7 ± 0.1 |
| Ca        | 11.4 ± 0.8 | 3.8 ± 0.1 | 9.0 ± 0.1 | 9.3 ± 0.4 | 9.6 ± 0.1 | 9.9 ± 0.4 | 8.3 ± 0.1 |
| Fe        | 5.9 ± 0.3 | 7.5 ± 0.2 | 4.8 ± 0.3 | 4.8 ± 0.2 | 4.8 ± 0.1 | 4.7 ± 0.2 | 5.1 ± 0.1 |
| K         | 1.5 ± 0.1 | 0.2 ± 0.1 | 1.1 ± 0.1 | 4.7 ± 0.3 | 8.4 ± 0.2 | 11 ± 0.6 | 13.5 ± 0.2 |
| Mg        | 1.6 ± 0.1 | 0.4 ± 0.1 | 1.2 ± 0.1 | 1.3 ± 0.1 | 1.5 ± 0.1 | 1.3 ± 0.1 | 1.2 ± 0.1 |
| Na        | 0.5 ± 0.1 | 0.2 ± 0.1 | 10.8 ± 0.4 | 10.6 ± 0.3 | 6.5 ± 0.1 | 4.8 ± 0.1 | 0.5 ± 0.3 |
| P         | 9.4 ± 0.3 | 4.2 ± 0.1 | 7.2 ± 0.3 | 7.6 ± 0.2 | 7.6 ± 0.1 | 7.3 ± 0.2 | 7.2 ± 0.1 |
| S         | 0.9 ± 0.1 | 0.9 ± 0.1 | 2.7 ± 0.1 | 3 ± 0.2 | 1.6 ± 0.1 | 1 ± 0.2 | 1.4 ± 0.4 |
| Si        | 11.4 ± 0.8 | 3.1 ± 0.4 | 8.9 ± 0.3 | 9.9 ± 0.1 | 9.8 ± 0.3 | 9.8 ± 0.5 | 9.4 ± 0.4 |
to achieve complete extractability in neutral ammonium citrate (PNAC,rel) as shown in a previous study by Herzel et al. [10]. Experiments with additive mixtures of molar ratios of $K_2SO_4$ and $Na_2SO_4$ were labelled CT-Na (Crucible trial with $Na_2SO_4$, $K_2SO_4$), CT-K25 ($Na_2SO_4$/$K_2SO_4$ = 75/25), CT-K50 ($Na_2SO_4$/$K_2SO_4$ = 50/50), CT-K75 ($Na_2SO_4$/$K_2SO_4$ = 25/75) and CT-K100 ($Na_2SO_4$/$K_2SO_4$ = 0/100). The $Na_2SO_4$/$K_2SO_4$ mixtures were pre-processed and homogenized by heating at 600 °C over night and subsequent milling in a vibrating cup mill before utilization in the thermochemical trial.

10–13 g of the above described mixtures of SSA, SS and alkali additive were calcined in crucibles of 75 mL volume. The material mixtures in crucibles were thermochemically treated in a preheated muffle furnace at different temperatures from 750 °C to 1150 °C for 30 min. After thermochemical treatment, the crucibles were removed from the furnace and were allowed to cool at ambient conditions. During calcination and cooling, the crucibles were covered with a corundum lid to maintain reducing conditions. The samples were each pulverized in a vibrating cup mill made of tungsten carbide to prepare the samples for subsequent analytics.

Pilot-scale trial

The influence of the composition of the alkali additive ($Na_2SO_4$/$K_2SO_4$) on the thermochemical process was investigated with a bench scale rotary kiln (RT1700, Thermal technology GmbH). The kiln consists of a temperature resistant ceramic tube that rotates in an oven heated by electric heating elements. The trial was operated at a co-current nitrogen gas flow of 300 L/h.

The feed material was a mixture of 3.0 kg SSA, 0.75 kg SS and varying amounts and composition of alkali additive ($Na_2SO_4$/$K_2SO_4$). The materials were mixed in an Eirich (R02, Eirich) mixer for 5 min before processing in the rotary kiln. The composition of the alkali additives was (i) 1.39 kg $Na_2SO_4$ (starting material mixture: L-Na), (ii) 0.66 kg $Na_2SO_4$ and 0.77 kg $K_2SO_4$ (starting material mixture: L-K50) and (iii) 0.31 kg $Na_2SO_4$ and 1.21 kg $K_2SO_4$ (starting material mixture: L-K75). The names of the trials with different composition of the alkali additive were chosen according to the molar ratio of K:Na that was used. In opposite to the laboratory-scale experiment, the additive mixtures (K50, K75) were not preheated. The three mixtures were fed after each other into the rotary kiln without any break in the order L-Na, L-K50 and L-K75 (feeding rate 2.6 kg/h). Consequently, the ratio of K:Na was increased along the rotary kiln trial.

The trial was done at a constant kiln set up temperature of 1000 °C. The temperature in the tube was recorded by a manually held thermocouple of 3 m length (Type K, electronic sensor, Heilbronn, Germany). The maximum temperature in the material bed was lower than the set kiln temperature and reached 950–970 °C in the center of the heating zone.

The trial lasted 9 h. The first product could be sampled 45 min after start of feeding material. Subsequently, the product was collected in different charges. A new product sampling charge was started after every approx. 30 min. The twelve product samples was collected after feeding stop and for a time period of 2 h and 10 min. Each sample was milled in a cross-beater mill (SK300, Retsch GmbH, Haan, Germany). The milled samples were divided with a sample splitter to achieve a representative sub-sample of 100 g. This was pulverized in a vibrating cup mill made of tungsten carbide and was used for subsequent analytics.

Analysis

Total digestion

Approximately 0.1 g of SSA or thermochemically treated ash was mixed with 4 mL of concentrated nitric acid ($HNO_3$), 1.5 mL perchloric acid ($HClO_4$), and 0.5 mL hydrofluoric acid (HF) and was digested in a microwave (mikroPrepA, MLS GmbH, Leutkirch, Germany). Excess HF was complexed with 2.5 mL cold saturated boric acid ($HBO_3$). Additional information on the analysis and quality assurance can be found elsewhere [27, 28].

Extraction in neutral ammonium citrate solution

Extraction of P in neutral ammonium citrate solution was determined according to EU [29]. The amount of extractable P is defined as $P_{NAC,rel}$. The ratio of soluble P and total amount of P is defined as $P_{NAC,rel}$. The elements Ca, K, Mg and Na were also measured in the extraction solution (Table 2).

Determination of element content with ICP-OES

The element concentrations of digestions solutions and extract in neutral ammonium citrate were measured by ICP-OES (Thermo iCAP 7400, Dreieich, Germany). The total digestion and extraction solution were diluted by 1:100 and analyzed with a six-point-calibration, including the blank. Reference material CTA-FFA-1 [30] was used to evaluate precision of chemical analysis for Al, Ca, Fe, K, Mg and Na. Reference material U826-1 [31] was used for P and extractable P in citric acid [29]. The recovery was 89% for Al, 82% for Ca, 93% for Fe, 95% for K, 97% for Mg, 100% for Na, 93% for P, 98% for Si and 94% for extractable P in citric acid.
Crystalline phase analysis by X-ray diffraction (XRD)

Powder X-ray diffraction (XRD) analyses were done for fertilizier samples (Table 3) as well as for fertilizer samples after extraction in neutral ammonium citrate to verify reflexes disappearing due to extraction. Analyses were performed in Bragg–Brentano geometry over a 2θ range from 5° to 80°, with a step size of 0.02° (D8 Advance, Bruker AXS, Germany). The diffraction patterns were collected using Cu Kα1/Kα2 (\(\lambda_1 = 1.54056 \, \text{Å} / \lambda_2 = 1.54443 \, \text{Å}\)) radiation. The diffraction patterns were recorded with a Lynxeye detector. Qualitative identification of the crystalline phases was performed using the MATCH! Software (version 3.6) [32] in combination with the PDF2 database [33].

Different thermal analysis (DTA)

Different thermal analysis (DTA) were carried out on a Netzsch STA 449 F3 Jupiter for mixtures of sodium and potassium sulfates (Fig. S2). The samples were heated with a rate of 10 Kelvin per minute from 30 to 1200 °C in an oxidizing atmosphere adjusted with synthetic air (80% nitrogen and 20% oxygen).

Table 2 Extracted element mass fractions after extraction of selected thermochemical products with neutral ammonium citrate solution (NAC) displayed in wt.-% with standard deviation (triplicates) and on the molar ratios normalized to 1 P atom per formula unit. The calculated molar ratios are used for estimating P-phase composition

| Series | Extracted chemical composition of P-phases After extraction in NAC | Classification of crystalline P-phases according to literature | Estimated P-phase composition after extraction in NAC |
|--------|---------------------------------------------------------------|---------------------------------------------------------------|-----------------------------------------------------|
| Ca     | Mg | Na | K | P | CaNaPO₄ [37] | (Ca₀.₇₂Mg₀.₂₈)NaPO₄ [22] | Two phases |
| molar ratio | 0.87 | 0.09 | 1.23 | 0.00 | 1.00 |
| CT-K25 | 1000 | 8.1 ± 0.3 | 0.6 ± 0.01 | 5.6 ± 0.1 | 0.7 ± 0.0 | 7.2 ± 0.1 | CaNaPO₄ [37] | (Ca₀.₇₂Mg₀.₂₈)NaPO₄ [22] | Two phases |
| molar ratio | 0.87 | 0.1 | 1.06 | 0.08 | 1.00 |
| CT-K50 | 1000 | 7.2 ± 0.1 | 1.0 ± 0.01 | 5.1 ± 0.1 | 1.6 ± 0.2 | 7.3 ± 0.1 | CaNaPO₄ [37] | (Ca₀.₇₂Mg₀.₂₈)NaPO₄ [22] | Two phases |
| molar ratio | 0.77 | 0.18 | 0.94 | 0.18 | 1.00 |
| CT-K75 | 1050 | 7.1 ± 0.2 | 0.5 ± 0.04 | 2.9 ± 0.1 | 3.8 ± 0.1 | 6.4 ± 0.1 | Ca(Na₀.₆K₀.₄)NaPO₄ [37] | (Ca₀.₉₀Mg₀.₁₀)(Na₀.₅₇K₀.₄₃)PO₄ |
| molar ratio | 0.88 | 0.62 | 0.47 | 1.00 |
| CT-K100 | 1150 | 7.0 ± 0.1 | 0.7 ± 0.01 | 0.8 ± 0.1 | 6.3 ± 0.2 | 6.3 ± 0.1 | Ca(Na₀.₃₅K₀.₆₅)NaPO₄ [37] | (Ca₀.₉₀Mg₀.₁₄)(Na₀.₁₈K₀.₈₂)PO₄ |
| molar ratio | 0.85 | 0.14 | 0.18 | 0.78 | 1.00 |

Table 3 Semi-quantification of phase composition of one selected thermochemical product of each series at reaction temperature required for high P_{NAC,rel}

| Series | T [°C] | Phosphate phases | Other phases |
|--------|-------|-----------------|--------------|
|        |       | CaNaPO₄ | (Ca,Mg)NaPO₄ | Ca(Na,K)PO₄ | Lazurite* | (Na,K)AISiO₄ | KAlSiO₄*** | KAlSi₂O₆ | Fe₃O₄ | Magnetite |
| CT-Na  | 1000  | ●     | ● | ● | ● | ● | ● | ● | ● |
| CT-K25 | 1000  | ●     | ● | ● | ● | ● | ● | ● | ● |
| CT-K50 | 1000  | ●     | ● | ● | ● | ● | ● | ● | ● |
| CT-K75 | 1050  | ●     | ● | ● | ● | ● | ● | ● | ● |
| CT-K100| 1150  | ●     | ● | ● | ● | ● | ● | ● | ● |

Meaning of symbols: ‘–’ not detected, ○ low ● medium ● high amount. This quantification is a rude semi-quantification with comparison of reflex intensity and chemical composition. Assumed chemical composition for (Ca,Mg)NaPO₄ and Ca(Na,K)PO₄ are listed in Table 2. *Na₃Ca₃Si₃Al₃O₁₂S **Two modifications of KAlSiO₄
Results

Thermochemical treatment in laboratory scale crucible experiments

Characterization of additives

Three mixtures of Na₂SO₄ and K₂SO₄ (not pre-heated) in the molar ratios 75:25 for CT-K25; 50:50 for CT-K50 and 25:75 for CT-K75 were analyzed with DTA. During heating, the characteristic phase transition points of Na₂SO₄ at approx. 253 °C and of K₂SO₄ at 584 °C are visible (Fig. S2). The temperatures of melting points are comparable to the temperatures of desired composition of mixtures of Na₂SO₄ and K₂SO₄ according to Rowe [26] (Fig. S2, Table 4).

These three mixtures were afterwards pre-heated at 600 °C and analyzed by XRD analysis after cooling. They contain different amounts of crystalline phases K₃Na(SO₄)₂, Na₂SO₄, NaKSO₄ and K₂SO₄ related to the Na and K portion in mixtures of Na₂SO₄ and K₂SO₄ (Fig. S3).

Correlation between temperature and extractable phosphorus (PNAC,rel)

Sewage sludge ash was thermochemically treated with Na₂SO₄, K₂SO₄ and their mixtures (series CT-Na to CT-K100) in corundum crucibles at different temperatures. The aim was to identify the required temperature to achieve a high extractable P in neutral ammonium citrate, indicated as PNAC,rel in Fig. 1. The data show a clear correlation of PNAC,rel and treatment temperature for all tested additives and are described with regression curves of a dose–response four-parameter logistic function [34]. The lowest measured PNAC,rel in each series were between 40–50%. The PNAC,rel increased at different temperatures in these experiments. The point of increase of PNAC,rel correlated with melting points in the phase diagram Na₂SO₄-K₂SO₄ [26].

Sewage sludge ash treated with Na₂SO₄ has the strongest extractability increase in the temperature range 800–900 °C around the melting point 894 °C of Na₂SO₄. The maximum level of 90–100% were above 925 °C and is confirmed by experiments by Herzel et al. [10] and Herzel et al. [15].

PNAC,rel for a treatment with K₂SO₄ by Herzel et al. [10] correlates also with current results. In this previous study, approximately half of P was soluble in neutral ammonium citrate for a K/P-molar ratio of 2.0. We observed a PNAC,rel of 55% at 1000 °C. The maximum PNAC,rel was observed at the maximum temperature in the experiment of 1150 °C with approximately 90% and correlates with the melting point of K₂SO₄ of 1070 °C. Higher temperatures were not investigated as SSA started to melt at these temperatures. The melt would react with the corundum crucible changing the chemical system. A thermochemical treatment with K₂SO₄ is not suitable for a technical process due to the high temperature required for the reaction of the phosphate phases in SSA with K₂SO₄.

Nevertheless, a K-containing additive is desirable to increase the K content of the recycling fertilizer. Therefore, reaction temperatures were investigated when K₂SO₄ is

| Series  | K₂SO₄ in additive [mol%] | Treated ash | Measured melting points of additives ¹ | Temperature of melting point ² |
|---------|--------------------------|-------------|----------------------------------------|-----------------------------|
| CT-Na   | 0                        | 925         | 888                                   | 894                        |
| CT- K25 | 25                       | 900         | 838                                   | 823                        |
| CT- K50 | 50                       | 925         | 865                                   | 910                        |
| CT- K75 | 75                       | 1000        | 948                                   | 1000                       |
| CT-K100 | 100                      | 1150        | 1073                                  | 1071                       |

Fig. 1 PNAC,rel of products thermochemically treated in crucibles at different temperatures of five series with different additive mixtures of Na₂SO₄ and K₂SO₄ (number in series name represent molar portion of K₂SO₄ in additive). Total ratio of (Na + K)/P was 2 mol/mol in starting material. The horizontal dashed line represents 90% PNAC,rel. Colored dashed curves are regression curves of a dose–response four-parameter logistic function [34] (color figure online)

Table 4 Sufficient reaction temperature to achieve PNAC,rel ≥ 90% for thermochemical treatment compared with melting points of alkali additives: ¹ measured temperature of melting point of used Na₂SO₄/K₂SO₄ with differential thermal analysis (DTA) (Fig. S2) and ² melting point according to Rowe [26]
partly substituted by Na$_2$SO$_4$ in an additive mixture. K$_2$SO$_4$ was substituted by 25 mol% (CT-K75), 50 mol% (CT-K50) and 75 mol% (CT-K25) Na$_2$SO$_4$.

One quarter substitution results in a significant temperature reduction to achieve high $P_{\text{NAC,rel}}$. It increased from 50 to 90% in the temperature range from 950 °C to 1000 °C and is constant at higher temperatures. In case of substitution of 50 mol% (CT-K50), P was nearly completely extractable in neutral ammonium citrate after treatment at 925 °C. Series CT-K25 (substitution 75 mol%) has the lowest reaction temperature of all five series. More than 90% of P was extractable in neutral ammonium citrate after thermochemical treatment at 900 °C (Fig. 1).

**Phase analysis in laboratory-scale calcination experiments**

The phase analysis of thermochemical products was done with samples in which the highest $P_{\text{NAC,rel}}$ levels were achieved (1000 °C for all samples, except 1050 °C for CT-K75 and 1150 °C for CT-K100). The sample names include the reaction temperature (e.g., CT-Na-1000). The added Na$_2$SO$_4$ and K$_2$SO$_4$ react with phosphate phases as well as with quartz in the SSA. Furthermore, the amount of amorphous phase is decreased after thermochemical treatment. This means more crystalline phases are available in the thermochemical products because additives react with e.g., Al$_2$O$_3$ and SiO$_2$ from the amorphous phase [35] and form crystalline phases.

The different content of Na and K in series CT-Na-1000 to CT-K100-1150 changed the Na and K contents in the silicates formed (Table 3). Na-containing nepheline ((Na,K)AlSiO$_4$) was identified in samples CT-Na-1000 and CT-K25-1000 and was replaced by KAlSiO$_4$-phases in the other samples (CT-K50-1000, CT-K75-1050 and CT-K100-1150). Furthermore, the fraction of leucite (KAlSi$_2$O$_6$) decreases with increasing K content and is replaced by KAlSiO$_4$. Only, CT-Na-1000 contains a sulfur bearing silicate phase which belongs to the sodalite group. The assumed composition of Na$_3$CaAl$_3$Si$_3$O$_{12}$S belongs to the lazurite group but many isostructural types exist. Sulfur containing silicates were previously observed by Stemmann et al. [14] using Na$_2$SO$_4$ as additive.

The P phases whitlockite and AlPO$_4$ in SSA (Fig. S1) are transformed due to thermochemical treatment to calcium alkali phosphates (Ca(Na,K)PO$_4$). Furthermore, Calcium (Ca) could be partly substituted by Magnesium (Mg). Sodium-rich samples (CT-Na-1000, CT-K25-1000 and CT-K50-1000) contain the low temperature modification of CaNaPO$_4$ (ICDD PDF entry 00–029-1193) and (Ca,Mg)NaPO$_4$ (Table 3). The latter correlates to the reflexes of the data base entry for (Ca$_{0.72}$Mg$_{0.28}$)NaPO$_4$ (ICDD PDF entry 01–088-1548). A reflex shift in the diffraction pattern indicated that this phase contained less Mg compared to Ca (Fig. 2). Thus, the composition is most likely (Ca$_{0.8}$Mg$_{0.2}$)NaPO$_4$. CT-Na-1000 and CT-K25-1000 mainly contain CaNaPO$_4$ and less (Ca,Mg)NaPO$_4$. Only (Ca,Mg)NaPO$_4$ was detected in CT-K50-1000 (Fig. 2 and Table 3) and is confirmed by increased content of Mg in solution after extraction in neutral ammonium citrate (Table 2). The chemical composition of P phase in CT-K50-1000 is assumed as (Ca$_{0.81}$Mg$_{0.19}$)Na$_{0.84}$K$_{0.16}$PO$_4$ (Table 2).

The identifications of crystalline P-phases in samples CT-K75-1050 and CT-K100-1150 were difficult because no data base entry for P phases correlates very well. To identify the P-bearing crystalline phases, the ash was analyzed by XRD before and after extraction with neutral ammonium citrate (Fig. 3). The reflexes which are disappeared after extraction are located between reflexes of high-temperature modification of CaNaPO$_4$ and CaKPO$_4$ [36] (Table 3). Additionally, reflex positions were compared with pure Ca(Na,K)PO$_4$-phases with different portion of K and Na (Fig. 3). The reflexes in CT-K75-1050 are comparable to synthesized phase Ca(Na$_{0.4}$K$_{0.6}$)PO$_4$ [37] but shifted to higher angles indicating a lower K content (Fig. 3). This is confirmed by an estimated phase composition of (Ca$_{0.99}$Mg$_{0.10}$)Na$_{0.5}$K$_{0.4}$PO$_4$ (Table 2). The reflexes in CT-K100-1150 are located between samples Ca(Na$_{0.35}$K$_{0.65}$)PO$_4$ and high temperature modification of CaKPO$_4$ [37] and correspond to the estimated phase composition (Ca$_{0.86}$Mg$_{0.14}$)(Na$_{0.18}$K$_{0.82}$)PO$_4$.
Thus, phase modification of CaKPO₄ incorporated Na and has different reflexes (Fig. 3) as previous reported phase modification of CaKPO₄ [24, 37].

### Pilot trials in a rotary kiln

#### Pilot-scale observations

The pilot-scale trial (L) in the rotary kiln was conducted at kiln set up temperature of 1000 °C. The measured temperature in the material bed in the tube was 950–970 °C. Adhesion of the material was observed at the hot walls of the tube. Thus, a material ring continuously formed had to be removed every 20 min. The clogging effect was even more pronounced compared to Herzl et al. [15] because the kiln has a smaller diameter and its corundum tube has a porous ceramic surface. The material adhesions were unchanged over the whole trial including variation of starting materials (L-Na, L-K50, L-K75).

A mass flow at the outlet of about 2.1 kg/h was expected (material feeding rate of 2.6 kg/h, total mass yield of 80% mainly due to combustion of organic compounds in SS). The measured mass flows were approx. 1.5 kg/h. Thus, a certain storage of material in the kiln was observed.

#### Extractable phosphorus and change of alkali content

The ratios of alkali additives Na₂SO₄ and K₂SO₄ were varied to evaluate how the portion of K₂SO₄ in the additive influences \( P_{NAC,rel} \). Laboratory-scale experiments show that the mixture of Na₂SO₄ and K₂SO₄ needs minimal 25 mol% \( \text{Na}_2\text{SO}_4 \) (series CT-K75) to achieve a high \( P_{NAC,rel} \) at 1000 °C (Fig. 1). Based on this data, the same amounts of SSA, SS and alkali additives were used to conduct a rotary kiln trial using the starting materials mixtures (L-K50 and L-K75).

The first material charging was starting materials mixtures with additive \( \text{Na}_2\text{SO}_4 \) (L-Na, in total 5.14 kg). After 45 min, the first material product was collected for half an hour. The treatment increased the \( P_{NAC,rel} \) from 35% in SSA to 73% in the thermochemical product. The content of Na in the product was 11 wt-% in agreement with the adjusted additive dosing (Fig. 4).

The second materials charging (L-K50, in total 5.18 kg) was immediately after feeding with first charging was finished after 1.5 h. Consequently, the K content increased continuously and was constant at 6 wt-% in sample L-5, L-6 and L-7, in parallel the Na content decreased to 7–8 wt-%. \( P_{NAC,rel} \) varied between 65 to 73% for samples L-3 till L-7 (Fig. 4).

Feeding of the third starting material L-K75 (in total 5.27 kg) after 4 h increased the K content up to 10 wt-% and \( P_{NAC,rel} \) decreased at the same time to 50%. The last sample 12 represents the product material after feeding stop resulting in decreasing material bed and is therefore not representative for any defined process condition (Fig. 4).

This rotary kiln trial shows very well how the Na and K contents adjusted by the additive mixtures influence the fraction of P extractable in neutral ammonium citrate and thus its availability to plants. The \( P_{NAC,rel} \) varied between 65

![Fig. 3](https://example.com/image.png)

**Fig. 3** Sections of diffractograms of samples CT-K75-1050 (left) and CT-K100-1150 (right). The dashed diffractograms after NAC-extraction. Arrows mark and dotted vertical lines marked reflexes vanished after NAC-extraction. Diffractogram of synthesized sample Ca(Na₃₅K₀₆₅)PO₄, Ca(Na₀₄K₀₆)PO₄, CaKPO₄ (high temperature modification) [37] are displayed for comparison of reflex positions (color figure online)
and 75% during the L-Na and L-K50 campaigns and continuously decreased during the L-K75 campaign to values below 50% (Fig. 4).

**Phase analysis in pilot trial**

$P_{\text{NAC,rel}}$ in the products of pilot-scale trial (Fig. 4) were significantly lower compared to the products of the corresponding laboratory-scale crucible experiments (Fig. 1). This suggests that the transformation from less soluble whitlockite and AlPO$_4$ in the SSA (Fig. S1) [11, 38] to highly NAC-extractable calcium alkali phosphates was not complete. This is confirmed by X-ray diffraction analysis done with the eleven samples from the pilot trial (sample names L-1 to L-11) (Table S1).

The identified highly NAC-extractable phases are mainly (Ca,Mg)NaPO$_4$ and minor CaNaPO$_4$. In the pilot trial, portions of these highly NAC-extractable phases decreased to the expense of whitlockite (Table S1) resulting in a drop of $P_{\text{NAC,rel}}$ after 4.5 h (L-8) (Fig. 4). There was no detectable K-containing calcium alkali phosphate, while the K content in the products increased (Fig. 4).

Sodium-rich silicates phases changed to K-rich phases due to increased K addition over time. Lazurite [Na$_3$CaAl$_3$Si$_3$O$_{12}$S] and sodium nepheline [NaAlSiO$_4$] is substituted by (Na,K)AlSiO$_4$ and Kalsilite [KAISiO$_4$] and...
for heat capacity up to 950 °C are available. An exception is KAlSiO$_4$ (Kalsilite) as heat capacity data are only available up to 537 °C in HSC Chemistry. Therefore, these data were substituted by similar data for Kalsilite (25–537 °C) and by data for high kaliophillite (537–1527 °C) from Robie and Hemingway [40].

These calculations resulted in six main element species Al$_2$O$_3$*SiO$_2$, CaNaPO$_4$, CaKPO$_4$, CaNaPO$_4$, KAlSiO$_4$ and NaAlSiO$_4$. Further element species were below 0.03 kmol and were thus negligible. Amounts of the same chemical element species for KAlSiO$_4$ (kaliophilite, kalsilite/high kaliophilite) and NaAlSiO$_4$ (kalsilite, nepheline) were summed up in Fig. 5 and S4.

The calculation shows that the formation of the element species does not further change above 500 °C (Fig. 5). The amounts of CaNaPO$_4$ and KAlSiO$_4$ are at the same level of around 0.65 kmol. The amount of CaKPO$_4$ (0.35 kmol) is in the same level as the sum of the silicate species NaAlSiO$_4$, Na$_2$SiO$_3$ and Al$_2$O$_3$*SiO$_2$. Thus, the element species distribution can be divided into two groups. The dominant group contains CaNaPO$_4$+ KAlSiO$_4$ and the subordinated group CaKPO$_4$, NaAlSiO$_4$, Na$_2$SiO$_3$ and Al$_2$O$_3$*SiO$_2$. Thus, the element species distribution can be divided into two groups. The dominant group contains CaNaPO$_4$+ KAlSiO$_4$ and the subordinated group CaKPO$_4$, NaAlSiO$_4$, Na$_2$SiO$_3$ and Al$_2$O$_3$*SiO$_2$ in case Na$_2$O and K$_2$O were added in equal amounts of 0.5 mol (HSC-K50). Further calculations with other Na$_2$O and K$_2$O ratios resulted in the same six main element species (HSC-K25 and HSC-K75). The portion is shifted to higher and lower contents of Na and K correlating to the added amounts of alkali oxides (Fig. S4).

Furthermore, we tested calculations with extra addition of 2 mol SiO$_2$ as well as 2 mol SiO$_2$ and 1 mol Al$_2$O$_3$ (data not shown). Phosphates species do not change and the content was quite similar. KAlSiO$_4$ and KAlSi$_2$O$_8$ were additionally formed by adding 2 mol SiO$_2$. Addition of 2 mol SiO$_2$ and 1 mol Al$_2$O$_3$ increased the content of Al$_2$O$_3$*SiO$_2$.

Finally, we can conclude the thermodynamic calculation of a simplified element species system shows that Na is preferably bonded in phosphate phases and K in silicates.

**Discussion**

**Relation between process reaction temperature and melting point of alkali additive**

Maximum $P_{NAC,rel}$ correlates very well with melting points of alkali sulfate in the phase diagram Na$_2$SO$_4$-K$_2$SO$_4$ by Rowe [26] (Table 4). The melting point of 3 Na$_2$SO$_4$ - K$_2$SO$_4$ (comparable to series CT-K25) is the lowest with 823 °C in this phase system. The melting points increased monotonously with increasing mass fractions of K$_2$SO$_4$. It is 910 °C for Na$_2$SO$_4$. K$_2$SO$_4$ (comparable to series CT-K50) and 1000 °C for Na$_2$SO$_4$. 3 K$_2$SO$_4$ (comparable to series CT-K75). These melting points are compared with reaction temperatures of the crucible trials series with high level of P$_{NAC,rel}$ ≥ 90%. The reaction temperature is minimal 30 °C (series CT-Na) and maximally 80 °C (series CT-K100) above the respective melting points showing that melting of the additive significantly supports the reaction between additive and crystalline P-phases. However, the increase of P$_{NAC,rel}$ already starts below the melting points [15] indicating that solid phase reactions between phosphate phases in SSA and additves (Na$_2$SO$_4$, K$_2$SO$_4$) take place. This is supported by thermogravimetric experiments suggesting that decomposition of Na$_2$SO$_4$ already starts at approx. 600 °C under reducing conditions producing sodium oxide that reacts with the crystalline P-phases [41].

**Preparation of mixtures of additive**

A further challenge was to ensure that the mixture of Na$_2$SO$_4$ and K$_2$SO$_4$ reacts as mixture with the respective melting points (Table 4) and not as individual phases. To ensure proper mixing and activating of the K$_2$SO$_4$ fraction, additive mixtures were milled and preheated at 600 °C for the laboratory-scale experiments. The results of the pilot-scale trials show that this preheating is not necessary. The samples L-5 to L-7 represent the mixture L-K50 and have comparable $P_{NAC,rel}$ as samples L-1 and L-2 representing mixture L-Na (Fig. 4). Thus, the K$_2$SO$_4$ fraction of the additive took part in the reaction with the phosphate phases presumably after forming an eutectic melt together with Na$_2$SO$_4$.

**Comparison of laboratory-scale and pilot-scale trial for phosphorus availability**

The lower values of $P_{NAC,rel}$ in pilot-scale trial (Fig. 4) compared to corresponding laboratory-scale experiments (Fig. 1) are based on the use of sulfates as additives. Comparable results with Na$_2$SO$_4$ as additive were observed in demonstration-scale trials that achieved a $P_{NAC,rel}$ of maximal 80% [10] and $P_{NAC,rel}$ of ~75% [15]. Phosphorus was nearly completely extractable with neutral ammonium citrate in case pilot-scale trials were conducted with Na$_2$CO$_3$ ($P_{NAC,rel}$ > 90%) instead of Na$_2$SO$_4$ [16, 17]. This implies that the choice of the type of additive has a strong influence on the success of thermochemical reaction in the rotary kiln (pilot-scale trials). Carbonates should be preferred.

Furthermore, $P_{NAC,rel}$ in the pilot-scale trials declined with increasing K content (L-8 to L-12) as it was observed after feeding the material mixture L-K75 (Fig. 4). We assume that the reaction temperature in the kiln was not high enough to force reaction between SSA and K-rich alkali sulfate mixtures. Laboratory-scale experiments (CT-K75) postulate that 1000 °C is sufficient to transform all phosphate phases to calcium alkali phosphate in case an additive containing 75 mol% K$_2$SO$_4$ and 25 mol% Na$_2$SO$_4$ is used.
(Fig. 1). The kiln set up temperature was 1000 °C but the measured temperature in the material bed was 950–970 °C. Thus, this material bed temperature was too low to promote reactions between SSA and K-rich sulfates in L-K75.

**Formation of (Ca,Mg)NaPO₄ or CaNaPO₄**

Crystalline phase analysis and $P_{\text{NAC,rel}}$ confirmed that low plant available phosphate phases in SSA were completely transformed to highly plant available (Ca,Mg)NaPO₄ or CaNaPO₄ in laboratory-scale experiments (Table 3). Previous studies postulate that CaNaPO₄ [14] and the associated phase 2CaNaPO₄·Ca₂SiO₄ [16] are main compounds after thermochemical treatment of SSA with Na₂SO₄ or Na₂CO₃. The presence of 2CaNaPO₄·Ca₂SiO₄ can be expected in case of high calcium mass fractions e.g. by dosing of CaCO₃ [16].

Steckenmesser et al. [19] identified also a magnesium containing calcium alkali phosphate (Ca,Mg)NaPO₄. This phase can be formed in case of low ratio of Ca to P [19]. In our study, a higher ratio of Ca to P (Table 1) compared to Steckenmesser et al. [19] is given indicating that the inclusion of magnesium in the phosphate phase was not forced by a shortage in Ca supply.

In our case, the process conditions and composition of additives presumably play a major role for the formation of (Ca,Mg)NaPO₄. The pilot-scale and corresponding laboratory-scale experiments resulted in different P-phases at similar chemical compositions. The product samples L-1 and L-2 of pilot-scale trial (Table S1) contain mainly (Ca,Mg)NaPO₄, whereas CaNaPO₄ is dominant in the corresponding laboratory-scale experiment sample CT-Na-1000 (Table 3).

Furthermore, the evaluation of crystalline phase composition in samples CT-K50-1000 (Table 3) indicates that (Ca,Mg)NaPO₄ is formed preferentially instead of CaNaPO₄ using K-containing additives (Fig. 2). Presumably, K is also incorporated in CaNaPO₄ and (Ca,Mg)NaPO₄ indicated by an extraction of K by neutral ammonium citrate solution (Table 2). CaNaPO₄ can incorporate up to 10 mol% K without crystal phase transition [42]. Presumably, (Ca,Mg)NaPO₄ can incorporation similar amounts.

Formation of (Ca,Mg)NaPO₄ is connected to incomplete phase transition from whitlockite to calcium alkali phosphate in pilot-scale trial and in case of using K-containing additives in laboratory-scale experiments. It can be assumed that (Ca,Mg)NaPO₄ is an intermediate phase between Mg-whitlockite (Ca,Mg)₃(PO₄)₄ and CaNaPO₄ [15]. Mg can substitute up to 15% of Ca in whitlockite [43].

**Incorporation of K preferred in silicates**

The additives Na₂SO₄ and K₂SO₄ react with silicates as well as with phosphates from SSA. More precisely, silicates are formed before calcium alkali phosphates [14]. In case only one alkali sulfate is used as additive, the order of formation is irrelevant. Addition of Na₂SO₄ or K₂SO₄ is resulting in the formation of CaNaPO₄ or CaKPO₄, respectively [10]. In case of using mixtures of both additives, the ratio of alkalis in additives and in the formed phosphates are different [44]. Calcium alkali phosphates contain less K as expected (Table 2). Samples CT-K25-1000 and CT-K50-1000 contains P-phases with low amounts of K (Table 2) despite of the addition of K₂SO₄ (25 mol% or 50 mol%). In contrast, K containing silicates ((Na,K)AlSiO₄, KAlSiO₄ and KAlSi₂O₆) are dominant in these samples (Table 3). Apparently, formation of K silicates is preferred compared to Na silicates. Thus, calcium alkali phosphates can first incorporate remaining K into the calcium alkali phosphates after finishing of formation of K containing silicates. This led to the formation of K containing calcium alkali phosphates only in case of addition of high amounts of K₂SO₄ (CT-K75-1050) (Table 2). Thus, CT-K75-1050 can be postulate as a promising PK-fertilizer which contains 88% of P and 35% of K extractable in neutral ammonium citrate (Figs. 1 and 6). The required reaction temperature of 1050 °C for production of this PK-fertilizers is feasible for an industrial-scale plant.

If Na or K is preferably incorporated in phosphates [Ca(Na,K)PO₄] could not be deduced by laboratory-scale experiments. Synthesis of K containing Ca(Na,K)PO₄ indicates that Na is preferably incorporated in Ca(Na,K)PO₄ [37]. This supports the preferred formation Na-rich Ca(Na,K)PO₄.

The thermodynamic calculation confirmed that K is preferably incorporated in silicates (Na,K)AlSiO₄ and predict that Na is preferably incorporated in Ca(Na,K)PO₄.
Nevertheless, the temporal progression (first silicate reaction, later phosphate reaction) could not be illustrated by thermodynamic calculation. For this reason, CaKPO₄ is available in thermodynamic calculation (Fig. S4 and Fig. 5) but is not present in corresponding laboratory-scale experiments CT-K25 and CT-K50 (Table 3).

The preferred incorporation of K in silicates could be a drawback to produce PK-fertilizers. Thermochemical products containing CaNaPO₄ are known to have high plant availability of P comparable to struvite and triple super phosphate [16, 17].

The plant availability of K cannot be certainly determined with neutral ammonium citrate solution since it is a non-standard extraction method for K-fertilizers. Other non-standard extraction methods (e.g., hydrochloric acid (0.5 mol/L) and HNO₃ (pH 5)) used for extraction of K-silicates without correlation to plant growth tests [45, 46] but were not tested for K containing phosphate fertilizers. Only, water solubility is a standard method for K which is certified for K-fertilizers containing KCl and K₂SO₄ [29].

Calcium alkali phosphates are a new category for PK-fertilizers which are not soluble in water [11]. Previous extraction and plant availability tests of K-containing calcium alkali phosphates are focus on phosphorus [23, 24] but also reported a complete extraction of K in in citric acid [23]. Data for extraction of K in neutral ammonium citrate are not reported previously.

Potassium bonded in Na,K)PO₄ is most likely plant available because K is extracted in neutral ammonium citrate (Table 2 and Fig. 6) and citric acid solution [23]. Alkali bonded in silicates are not extractable with neutral ammonium citrate solution and have a very low dissolution rate [45] and presumably less plant available. This is in line with finding by Santos et al. [47]. They measured a low extractability of K in citric acid for rocks containing KAlSiO₄. Franca et al. [48] postulate kalsilite as a slow release fertilizer but they investigated water containing kalsilite (KAlSiO₄*1.5 H₂O) and not the water free phase KAlSiO₄. Thus, we conclude the K bonded in silicates (KAlSiO₄, (Na,K)AlSiO₄, KAISi₂O₆) in our samples is poorly plant available and K bonded in calcium alkali phosphates are predicted to be plant available.

Conclusions

Our target was to show that K containing alkali sulfates are suitable to form highly plant available K and P products after thermochemical treatment of sewage sludge ash. The sufficient reaction temperature of thermochemical treatment is strongly connected to the melting point of used alkali sulfates. Thus, K₂SO₄ is unsuitable for thermochemical treatment due to its high melting point. If it is mixed with Na₂SO₄ the reaction temperature is decreasing according to the eutectic melting point. 25 mol% of Na₂SO₄ in additive mixture is enough to reduce the melting temperature to an economically suitable reaction temperature of 1050 °C.

Potassium is preferably incorporated in the silicate structure instead of calcium alkali phosphate phase. This could be deduced by laboratory-scale and pilot-scale experiments as well as by thermodynamic calculations. The formed highly plant available P phase (Ca,Mg)(Na,K)PO₄ contains less K as expected. The bonded K in silicates is assumed to be poorly plant available and will be investigated in ongoing studies. Thus, the produced PK-fertilizer is highly plant available for P as well as for K in phosphate phases.

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