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8262Thermally processed Ni- and Co-struvites as
functional materials for proton conductivity†Stephanos Karafiludis,^{a,*} Biswajit Bhattacharya,^b Ana Guilherme Buzanich,^a
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Here, we describe how to synthesise proton-conductive transition metal phosphates (TMPs) by direct thermal processing of precursor M-struvites, $\text{NH}_4\text{MPO}_4 \cdot 6\text{H}_2\text{O}$, with $\text{M} = \text{Ni}^{2+}, \text{Co}^{2+}$. In the as-derived TMP phases their thermal history and bulk proton conductivity were linked with the structural information about the metal coordination, phosphate groups, and volatile compounds. These aspects were investigated with vibrational and synchrotron-based spectroscopic methods (FT-IR, FT-RS, XAS). We elucidated the structures of amorphous and crystalline Ni- and Co phosphate phases in association with different coordination changes and distortion degrees of the metal polyhedra as they developed upon heating. Ni-struvite transformed to a stable amorphous phase over a broad range of temperatures ($90^\circ\text{C} < T < 600^\circ\text{C}$), in which it remained in an octahedral coordination environment, but the degree of distortion changed with T . In contrast, heating of Co-struvite led to several successive crystalline phases with only unstable transitional and short-lived amorphous components. Among the as-occurring phases, a highly functional layered M-dittmarite $\text{NH}_4\text{MPO}_4 \cdot \text{H}_2\text{O}$ obtained at low temperatures ($T < 200^\circ\text{C}$) demonstrated high proton conductivity values of $4.2 \times 10^{-5} \text{ S cm}^{-1}$ for Ni-dittmarite and Co-dittmarite $> 10^{-4} \text{ S cm}^{-1}$ at room temperature. Even at low humidity, these values are comparable with those found for Nafion, MOFs, some perovskites or composite materials. Coprecipitation of phosphates and transition metal cations in the form of struvite is potentially a viable method to extract these elements from wastewater. Thus, we propose that recycled M-struvites could be potentially further directly upcycled into crystalline and amorphous TMPs useful for electrochemical applications.

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Introduction

Transition metal phosphates (TMPs) have attracted considerable attention due to their favourable properties for electro-, photo- or organic catalysis and proton exchange membranes.^{1–7} For example, TMPs are known as cost-effective cathode materials in lithium/sodium battery technologies,^{8–11} for their adequate catalytical performance in the oxygen evolution

reaction^{12–14} or in organic reactions^{15–17} such as the selective oxidation of methane.¹⁸

TMPs are highly suitable for this broad range of applications due to their ability to form thermally stable diverse crystalline and amorphous structures, including transition metals at various oxidation states and in several coordination environments. Each of these structures can exhibit beneficial properties for specific applications. For instance, the distorted phosphate tetrahedrons in TMPs promote the adsorption or incorporation of water molecules,^{19,20} advantageous for proton conductivity. TMPs are known to exhibit relatively high proton conductivity of the order of $>10^{-2}$ – $10^{-8} \text{ S cm}^{-1}$ from 25°C to temperatures as high as 400°C .^{21–24} For instance, Zr- and Sn-phosphates demonstrate a broad range of applications due to their sufficiently high proton conductivities at ambient conditions and high temperature.^{22,25–27}

Importantly for mass applications, phosphates are readily inexpensive compounds providing potentially high mechanical strength,²⁸ high thermal stability of the associated stable hydrogen-bonded network, and chemical compatibility with fuel cell device components (electrodes, catalysts, seals, etc.).

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† Electronic supplementary information (ESI) available: X-ray diffraction (XRD), scanning electron microscopy (SEM), infrared spectroscopy (IR), Raman spectroscopy (RS), thermal gravimetric analysis with differential scanning calorimetry (TGA/DSC), X-ray absorption spectroscopy (XAS), proton conductivity (PC) and dynamic vapor sorption (DVS) measurements, Figs. S1–S12; Tables S1–S23. See DOI: <https://doi.org/10.1039/d3dt00839h>

In addition, TMPs with tuneable mesoporous frameworks can be synthesized either by using template-based^{3,14,20} or template-free methods.²⁹ Thus, these materials can be obtained in many form factors such as bulk ceramics, thin films, or membrane materials.^{30–32} Due to the different condensation degrees of the phosphate units and variable exchangeable metal cations, TMPs provide a wide range of chemical compositions.

Moreover, TMPs can be made through a variety of synthesis procedures such as hydrothermal, mechanochemical, molten salt, or precipitation routes, resulting in highly crystalline to purely amorphous materials, depending on specific requirements.^{19,20} Even for near-identical chemical composition, crystalline and amorphous compounds could exhibit completely different physicochemical properties such as solubility, conductivity, or adsorption.^{33,34} Nevertheless, due to the lack of long-range order, the structure–property relationships in amorphous solid phases of TMPs are still relatively poorly explored in contrast to their crystalline counterparts.

The necessary challenge in the research on the application use of TMPs in all their varieties is an ability to obtain, elucidate and optimise different compositions and structures. This goal can be achieved by processing a universal TMP precursor with a single yet flexible crystal structure. Such potential TMP compound is M-struvite $\text{NH}_4\text{MPO}_4 \cdot 6\text{H}_2\text{O}$, where M^{2+} is usually Mg^{2+} , but other divalent cations such as Ni^{2+} , Co^{2+} , Zn^{2+} , Cu^{2+} readily substitute Mg^{2+} in the struvite structure due to similar ionic radii.^{35–38} In this work, we focus on Ni- and Co-containing phosphate materials as these are the most promising for electrochemistry.^{3,14,39} In our previous study, we showed how to precipitate Co- and Ni-struvites of various morphologies and sizes from aqueous solutions at concentrations

similar to those found in wastewaters.⁴⁰ Building on this previous work, we present how these recycled M-struvite materials could be potentially further upcycled for electrochemical applications by undergoing a direct thermal treatment (Fig. 1). Consequently, to upcycle and modify these materials for electrochemical applications, a complete understanding of the phase transition sequence during thermal heating and resolving of the complex structures is required. Amorphous and crystalline phases are formed upon the thermal decomposition of Ni- and Co-struvites. We show that some of such as-derived from M-struvite phases, namely M-dittmarites, exhibit high proton conductivity.

Synthesis

$(\text{NH}_4)_2\text{HPO}_4$ (DAP) (ChemSolute, 99%), $\text{NiSO}_4 \cdot 6\text{H}_2\text{O}$ (ChemSolute, 99%) and $\text{CoSO}_4 \cdot 7\text{H}_2\text{O}$ (Alfa Aesar, 98%) were used to synthesise precursor M-struvites ($\text{M} = \text{Ni}$ or Co). A detailed description of the synthesis of the precursor transition metal struvites can be found elsewhere.⁴⁰ For further analysis, the precursor M-struvites were thermally treated in a temperature series (T -series = 25 °C, 90 °C, 120 °C, 150 °C, 180 °C, 210 °C, 240 °C, 270 °C, 300 °C, 400 °C, 500 °C, 600 °C, 700 °C and 800 °C) with a constant dwell time of $t = 24$ h to evaluate the thermal decomposition products and related crystalline or amorphous phase transitions. The long dwell time was used to assure equilibration. The following abbreviations are used for the occurring phases: nickel struvite $\text{NH}_4\text{NiPO}_4 \cdot \text{H}_2\text{O}$ (NIS), amorphous phase (am), nickel pyrophosphate $\text{Ni}_2\text{P}_2\text{O}_7$ (NPY), cobalt struvite $\text{NH}_4\text{CoPO}_4 \cdot 6\text{H}_2\text{O}$ (COS), cobalt dittmarite $\text{NH}_4\text{CoPO}_4 \cdot \text{H}_2\text{O}$ (COD), hydrogen cobalt(II)

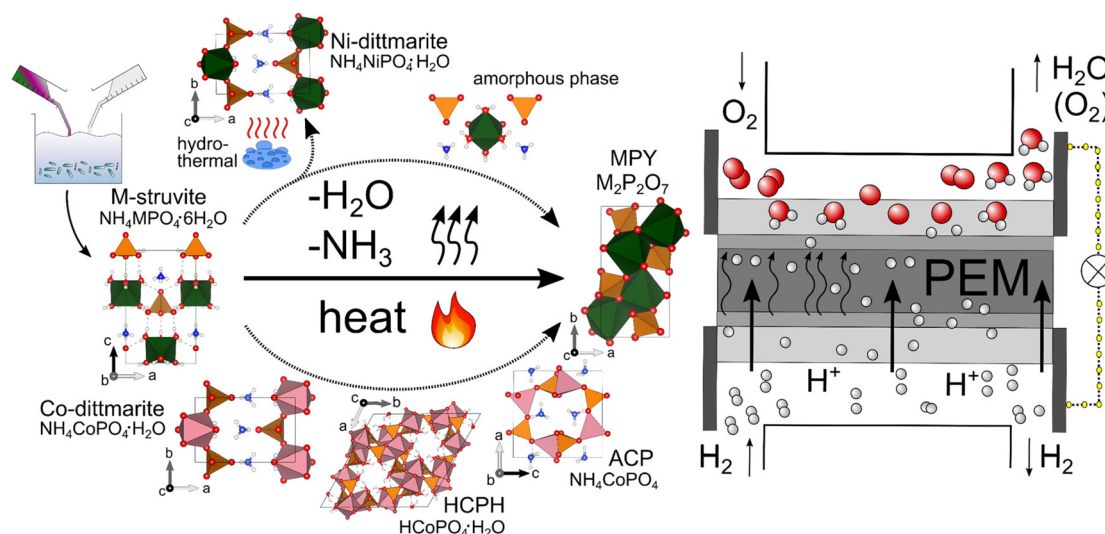
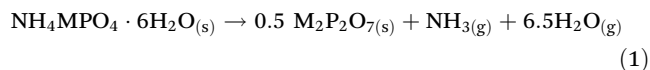


Fig. 1 The precursor M-struvites are obtained by a simple one-pot synthesis and further thermally treated, which results in degassing of $\text{H}_2\text{O} + \text{NH}_3$ and the associated occurrence of many phases, e.g. Ni- and Co-dittmarite, amorphous phases, HCPH, ACP and finally $\text{M}_2\text{P}_2\text{O}_7$ (MPY) with $\text{M} = \text{Ni}$, Co . Arrows indicate thermal history, lower arrow (Ni), upper arrow (Co). The synthesis leads to TMP materials potentially suitable for proton exchange membranes (PEMs) in fuel cells. A balance between high thermal stability and significant proton conductivity is the key to their promising application use.



phosphate monohydrate $\text{HCoPO}_4 \cdot \text{H}_2\text{O}$ (HCPH), ammonium cobalt phosphate NH_4CoPO_4 (ACP), cobalt pyrophosphate $\text{Co}_2\text{P}_2\text{O}_7$ (CPY) or M-pyrophosphate (MPY) with $\text{M} = \text{Ni}$ or Co .



Upon the heat treatment, M-struvites converted sequentially to amorphous and crystalline phases through many phase transitions on a pathway to a final crystalline M-pyrophosphate, which formed at high temperatures, $>500^\circ\text{C}$.

The thermal treatment of Co-struvite resulted in the formation of a crystalline Co-dittmarite phase, while Ni-struvite converted completely to an amorphous phase. The Ni-dittmarite analogue was not found directly in the heating experiments. Therefore, in an additional series of experiments, the ~ 5 g of Ni-struvite precursor powder was put into a hydrothermal autoclave reactor with a small amount of water (<50 ml) and heated up to $T = 90^\circ\text{C}$ to obtain directly crystalline Ni-dittmarite, allowing for a comparison of its properties with those of Co-dittmarite.

Complete technical descriptions of the used characterisation methods, including X-ray diffraction (XRD) for identification of the phase composition, scanning electron microscopy (SEM), combined thermogravimetric analysis, and differential scanning calorimetry (TGA/DSC) for imaging and tracking phase transitions, infrared spectroscopy (IR), Raman spectroscopy (RS), X-ray absorption spectroscopy (XAS) for determination of the coordination environment, proton conductivity (PC) and dynamic vapour sorption (DVS) measurements for evaluation of the applicational use can be found in the ESI (Methods†).

Results and discussion

Composition and morphology of crystalline and amorphous phases

For a clear presentation of the occurring complex phase transitions, we summarise the most important aspects of our synthesis and phases in Fig. 1. Highly crystalline, phase-pure, transition metal M-struvite powders with $\text{M} = \text{Ni}$ and Co were obtained as precursor materials through a simple one-pot synthesis (Fig. 1),⁴⁰ as was evidenced by the X-ray diffraction (Fig. 2A–C).

After treating the precursor Ni- and Co-struvites at different temperatures ($T = 90$ – 800°C , T -series) in air with a dwell time of $t = 24$ h, chemical and phase composition changes occurred and involved transformations to amorphous and crystalline compounds. The changes were also associated with colour changes of the powders. In the T -series of Ni-struvite, the crystal decomposed to an amorphous phase already at 90°C (Fig. 2A). Simultaneously, the colour turned from an emerald green (Ni-struvite precursor) to yellow-green at 90°C to brown at 120°C . Up to 600°C , the amorphous phase remained stable. At temperatures $>600^\circ\text{C}$ $\text{Ni}_2\text{P}_2\text{O}_7$ began to crystallise,

accompanied by a colour change of the powders from brown to pure yellow (Fig. 2A).

In contrast to Ni-struvite, Co-struvite $\text{NH}_4\text{CoPO}_4 \cdot 6\text{H}_2\text{O}$ transformed to crystalline Co-dittmarite $\text{NH}_4\text{CoPO}_4 \cdot \text{H}_2\text{O}$ already after the first heating step at 90°C (Fig. 2B and C). Up to 210°C Co-dittmarite remained present until it converted to a mixture of hydrogen cobalt(II)phosphate hydrate $\text{HCoPO}_4 \cdot \text{H}_2\text{O}$, ammonium cobalt(II)phosphate NH_4CoPO_4 , and an amorphous phase (Fig. 1). The amount of NH_4CoPO_4 increased with higher temperatures up to 300°C while the reflections of $\text{HCoPO}_4 \cdot \text{H}_2\text{O}$ disappeared. At $T = 300^\circ\text{C}$, a binary mixture of majorly NH_4CoPO_4 and minorly the amorphous phase(s) was present (Fig. 2B and C). At $T = 400^\circ\text{C}$ $\text{Co}_2\text{P}_2\text{O}_7$ crystallised, although the amorphous phase was still present. Heating to $T \geq 500^\circ\text{C}$, resulted in higher intensities of peaks in the diffractogram. At the same time, the proportion of the amorphous phase decreased successively, meaning that the crystalline phase evolved at the expense of the amorphous component.

Thus, Ni- and Co-struvite systems differ significantly in their thermal decomposition behaviour (Fig. 1). The TGA/DSC analysis was carried out to elucidate the mentioned thermally induced phase transitions, and SE images of the heated samples were collected (ESI: Fig. S2–S4†). A detailed description of the TGA/DSC data can be found in the ESI (Supplementary Data Analysis 1 and Fig. S4†). The thermally induced phase transitions and resulting mass losses due to degassing of volatile compounds are in agreement with the phases detected by XRD measurements. Based on the mass losses and DSC signals, the range of possible chemical compositions of the amorphous phase at given temperatures could be significantly narrowed down (ESI: Supplementary Data Analysis 1†).

The SE images show the presence of a “relict” Ni-struvite crystal morphology in the thermally treated amorphous Ni samples, overall indicating a gradual crystalline-amorphous phase transition and an associated pseudomorphism of the amorphous phases after the crystalline Ni-struvite (Fig. 2D and ESI: Fig. S2†). Even at higher temperatures $T \geq 600^\circ\text{C}$, the relict Ni-struvite morphology can still be observed (Fig. 2D, ESI: Fig. S2†). In this case, the pseudomorphism merely signifies a formation of the amorphous phase(s) without any crystalline intermediate step (Fig. 1A). Crystalline-crystalline pseudomorphism involving usually low volume changes is possible,⁴¹ but is unlikely to occur for these transformations, as the decomposition of M-struvite demonstrates significant volume dilatation due to loss of water and ammonia. Notably, the amorphous inhabited Ni-struvite structure is mechanically weak and can be easily distorted during sample manipulation (ESI: Fig. S2†).

In contrast, the transformation of Co-struvite (Fig. 2E and ESI: Fig. S3†) to other Co-phosphate phases is directly visible as changes in the morphology of the crystals. Here, partial crystalline-crystalline pseudomorphism also occurs *i.e.* of Co-dittmarite after Co-struvite, although the formation of other crystalline phases quickly extinguishes these structures (ESI:



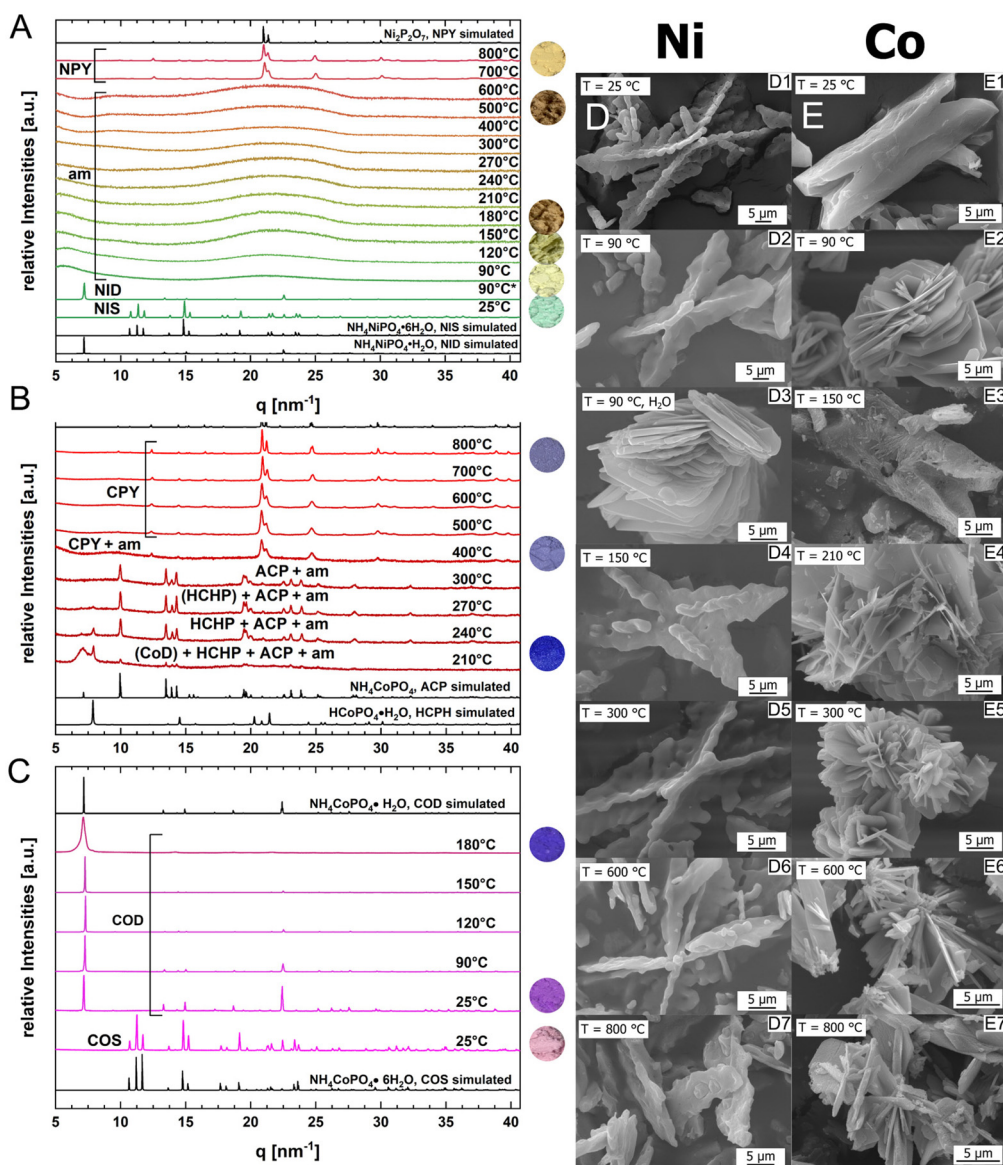


Fig. 2 Diffraction patterns of *T*-series of (A) Ni-struvite and (B and C) Co-struvite from 25 °C to 800 °C; 90 °C* = hydrothermal synthesis at RH \approx 100% at 90 °C; simulated XRD patterns were always added at the beginning and end of the diffraction pattern; The colour of the powder in the circles are placed next to the diagram; selected SE images of (D) Ni- and (E) Co-phosphates heated at different temperatures from 25 to 800 °C. SE images of the complete *T*-series of Ni and Co can be found in the ESI (Fig. S2 and S3†). Ni-struvite (NIS) reference database ICSD 403058; Ni-dittmarite (NID) reference database ICSD 424553; α -Ni₃P₂O₇ (NPY) reference ICSD 403058; Co-struvite reference ICSD 170042; Co-dittmarite (COD) with COD reference database 2008122; α -NH₄CoPO₄ (ACP) reference PDF 00-018-0402; Hydrogen cobalt(II)phosphate hydrate (HCPH) reference database ICSD 33828; α -Co₂P₂O₇ (CPY) reference database ICSD 280959.

Fig. S3†). At $T > 90$ °C, Co-dittmarite replaces Co-struvite as a pseudomorph at progressing temperatures (Fig. 2E and ESI: Fig. S3†). This is remarkably well-visible as the habit of the former original X-shaped Co-struvite crystals can still be identified (Fig. 2E and ESI: Fig. S3†), although Co-dittmarite covers the whole surface. The crystals' thin 2D sheet morphology indicates a crystalline-crystalline phase transition from Co-dittmarite to mainly HCoPO₄·H₂O and NH₄CoPO₄ (Fig. 1, 2E and ESI: Fig. S3†). Above 270 °C, the NH₄CoPO₄ exhibits its final crystal habit as two-dimensional plates similar to Co-dittmar-

ite. When Co-pyrophosphates begin to crystallise at 400 °C (Fig. 2E and ESI: Fig. S3†), the crystal habit changes again to an elongated prism or a tabular habit. The single crystals form massive aggregates of tens of microns.

Tracking the removal of volatile components

As the phase composition and structure changed due to degassing (see eqn (1) and Fig. 1), we wanted to track the evaporation of the volatile components, mainly NH₃/H₂O, and the condensation of the phosphate environment, by performing



infrared (IR) and Raman spectroscopy (ESI: Supplementary Note 2 and Fig. S5†).

The IR and Raman bands correlate well with the phase compositions and the degrees of crystallinity of the Ni and Co samples^{42,43}. The degree of crystallinity decreases in the *T*-series up to 300–400 °C due to the decomposition of the highly crystalline precursor *M*-struvite. It increases up to 800 °C during the crystallisation of Ni- and Co-pyrophosphates (Fig. 1). This trend is reflected in the progressive broadening and later narrowing/crystal field splitting of the phosphate stretching bands. Surprisingly, ammonium and water bands are still present at elevated temperatures of *T* = 400 °C, implying that these two species remain incorporated in the compounds.

The thermal effect on the coordination environment in the amorphous and crystalline Ni- and Co-phases

We further investigated the structures of amorphous and crystalline phases by following the changes/distortion in the metal coordination environment using X-ray absorption spectroscopy (XAS). Since XAS probes local coordination, it is particularly

well-suited for establishing any potential differences among the structures of amorphous phases. Especially, the signal in the pre-peak region (X-ray absorption near-edge structure, XANES) contains information about the 1s-3d orbital transitions and expresses the degree of centrosymmetry of the first metal coordination sphere. Near-ideal centrosymmetric coordination shows very low pre-peak intensities, while more distorted geometries exhibit higher intensities. As only 3d-4p hybridisation enables parity-allowed transitions, the extent of those is strongly correlated with the coordination geometry. Due to an inversion centre, ideal octahedral coordination sites show lower pre-peak intensities than tetrahedral symmetries ($A_{pp} = \text{area of the pre-peak}$, $A_{pp}^{\text{tetrahedral}} > A_{pp}^{\text{octahedral}}$) (Fig. 3).

The signal beyond the absorption edge in the EXAFS region (extended X-ray absorption fine structure) depicts the excited photoelectron's interactions with the compound's next neighbour atoms. Here, the degree of crystallinity directly influences the signal's intensity and width. Since amorphous phases do not have an established long-range order, they show broader, and less intense peaks in the R-space mainly in the low

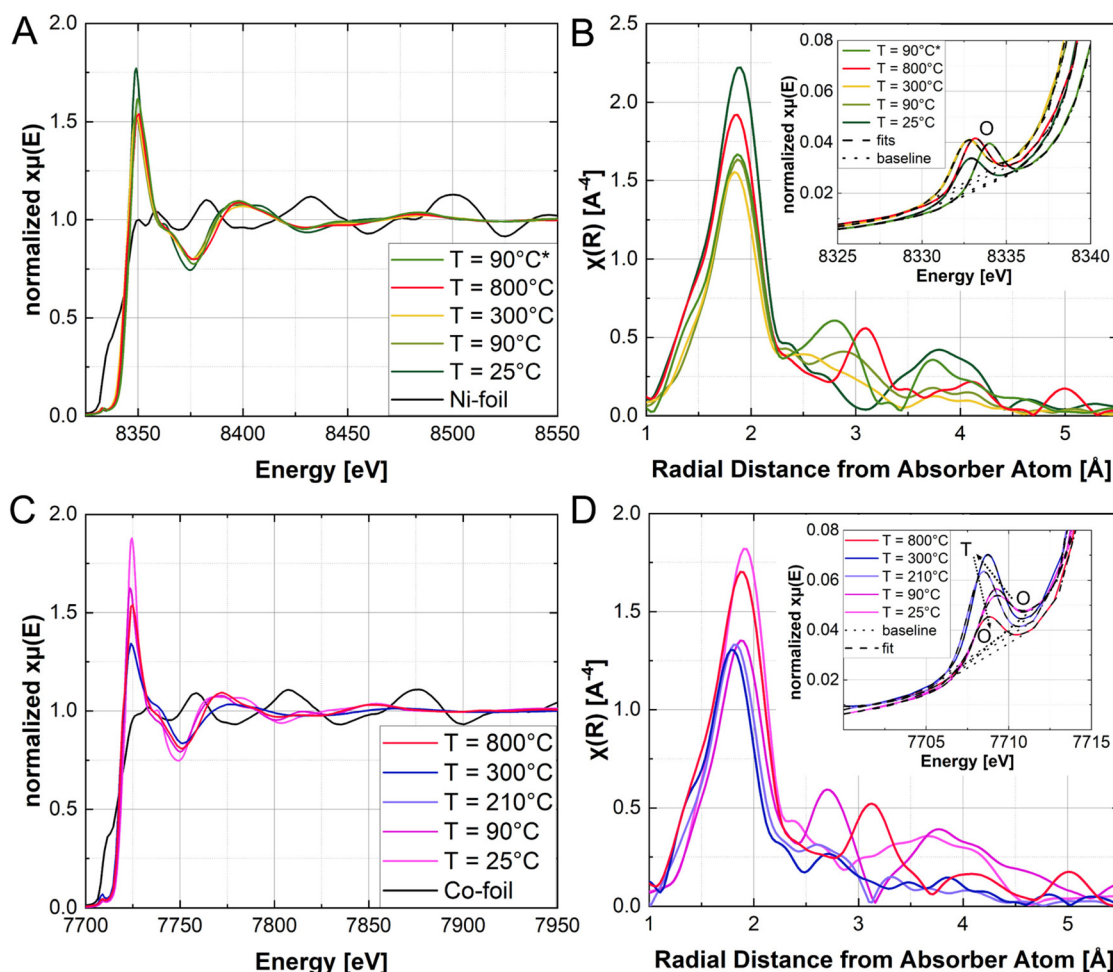


Fig. 3 XAS spectra of the thermally processed TMPs plotted in the real space and the R-space of (A and B) Ni-phosphates and (C and D) Co-phosphates; 90°C^* = hydrothermal synthesis at RH \approx 100% at 90°C . Inset in (B and D): pre-peak region and integration from Gaussian fits, O = octahedron, T = tetrahedron.



R-region due to an irregular arrangement of the lattice ions. The XAS spectra (Fig. 4) prove that Co and Ni are present exclusively as divalent M^{2+} species in all the samples (due to absorption edge at around 7730 and 8350 eV), as it is evidenced by comparing them with several M^{2+} standards (only metal foils are shown for the sake of clarity). Quantitative pre-peak integration results were calculated to compare the distortion in the first coordination sphere (ESI: Table S3†). The thermally induced change in the pre-peak area, indicative of a higher degree of distortion, differs significantly between the Ni- and Co-phosphate systems (Fig. 1). Up to 600 °C, crystalline Ni-struvite decomposes to an amorphous phase with a higher degree of distortion in remanent octahedral coordination [Ni: $A_{pp}^{25^\circ C} = 0.028(2)$ (precursor), $A_{pp}^{300^\circ C} = 0.034(2)$ and $A_{pp}^{600^\circ C} = 0.041(2)$]. In contrast, at low $T < 200$ °C the Co-system comprises crystalline octahedrally coordinated Co-phases, exhibiting a higher degree of distortion than the precursor Co-struvite [Co: $A_{pp}^{25^\circ C} = 0.026(1)$, $A_{pp}^{90^\circ C} = 0.032(1)$ and $A_{pp}^{150^\circ C} = 0.041(2)$]. In the further heating process due to the occurrence of a tetrahedrally coordinated Co-phase (NH_4CoPO_4) at $T > 180$ °C, the pre-peak intensity rises significantly [$A_{pp}^{210^\circ C} = 0.071(1)$ and $A_{pp}^{300^\circ C} = 0.075(3)$]. When the crystallisation to $M_2P_2O_7$ ($M = Ni, Co$) occurs at $T > 400$ °C (for Ni at $T = 600$ °C, for Co at 400 °C), the pre-peak intensities in both systems decrease again. Here, the metal octahedra are still more distorted than those in the near-ideal octahedrally coordinated precursor M-struvites, which we associate with the presence of twisted and linked pyrophosphate chains ($A_{pp}^{800^\circ C}$ (Ni and Co) = 0.033(1)).

Based on the fitting of the spectra in the R-space, radial distances from the absorber metal atom and average coordination numbers of the first coordination sphere could be calculated (ESI: Fig. S6–S8 and Tables S4–S20†). In the Ni-system, the calculated average coordination number, CN (*i.e.* degeneracy of oxygen), changes slightly from CN = 6.7(7) in the crystalline precursor Ni-struvite ($T = 25$ °C) to CN = 6.3(5) – 5.3(2) in the amorphous phase(s) ($T = 90$ °C–500 °C). Finally, Ni exhibits average coordination numbers of CN = 5.6(3) in the crystallised pyrophosphates, proving a remanent octahedral coordination with a slight degree of distortion in all occurring phases. The occurrence of several amorphous phases (polyamorphism) cannot be entirely ruled out. If this were the case, they all would have to exhibit similar local structures, as we do not observe any peak splitting in the Fourier-transformed spectra in R-space. In the Co-system the calculated average coordination number varies strongly from 6.4(5) in the precursor Co-struvite, to CN = 6.0(4) in Co-dittmarite heated at $T = 90$ °C to CN = 5.1(3) in the phase mixture of NH_4CoPO_4 , $HCoPO_4 \cdot H_2O$, an amorphous phase and residues of Co-dittmarite heated at 210 °C. Finally, at 300 °C, the average coordination number of Co decreases to a value of CN = 3.6(2), indicative for a tetrahedral coordination. When the crystallisation of $Co_2P_2O_7$ begins at ≥ 400 °C, the coordination number increases again to a value near 6 (CN = 5.3(4) at 400 °C and CN = 5.6(6) at 800 °C), which suggests octahedral coordination. In this way, the anticipated phase-dependent coordination transitions from octahedral ($NH_4CoPO_4 \cdot 6H_2O$, $NH_4CoPO_4 \cdot H_2O$,

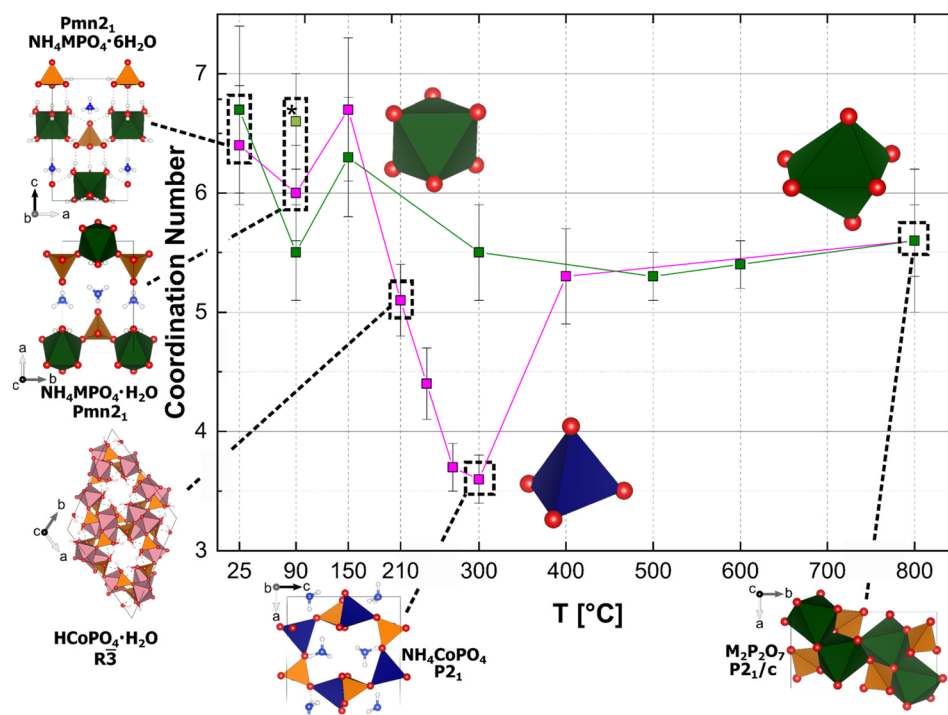


Fig. 4 Calculated coordination numbers through the thermal history of Ni (green squares)- and Co (pink squares)-phosphate compounds; Occurring phases with their crystal structure in tetrahedral and octahedral coordination are depicted. The asterisk marks the hydrothermal synthesised Ni-dittmarite compound.



HCoPO₄·H₂O) to tetrahedral (NH₄CoPO₄), and back to octahedral (Co₂P₂O₇) could be successfully derived by fitting EXAFS spectra in the R-space.

Our EXAFS characterisation and observations from other methods paint a coherent image of changes in metal coordination, correlating with the phase compositions. These results are summarised in Fig. 4. Here the degeneracy of oxygen, *i.e.* the distinct metal's calculated coordination number (CN), is plotted against the processed temperature. A change in the coordination geometry causes the intense colour change of Co in the heated samples from an octahedral one in [CoO₆] (NH₄CoPO₄·6H₂O, NH₄CoPO₄·H₂O, HCoPO₄·H₂O, amorphous phase, see also Fig. 1) to a tetrahedral one in [CoO₄] (NH₄CoPO₄) and again to an octahedral coordination environment (Co₂P₂O₇). Due to an increasing Jahn–Teller effect as Co²⁺ is no longer coordinated by six O^{2−} of H₂O molecules but by five O^{2−} of PO₄ and one remaining O^{2−} from H₂O, a color change can be observed from pink (Co-struvite) to purple (Co-dittmarite). When heated at 210 °C, Co-dittmarite transforms into HCoPO₄·H₂O and secondly into NH₄CoPO₄ accompanied by an intense colour change from purple to “cobalt blue”. The analogous colour change caused by an octahedral-tetrahedral coordination change was also observed in the related potassium compound of KCoPO₄·H₂O and KCoPO₄.⁴⁴ Here, the degree of distortion increases even more indicated by an increase in pre-peak area due to a Co²⁺ coordination change from an octahedron to a tetrahedron. The associated change in d-orbital splitting with a change in coordination significantly affects the colour of the Co-compounds. When the coordination switches again to an octahedral configuration accompanied by the phase transformation of NH₄CoPO₄ to Co₂P₂O₇, the CoO₆ octahedron in Co-pyrophosphate exhibits a lower degree of distortion than the CoO₄ tetrahedron in NH₄CoPO₄. Therefore, the pre-peak area decreases while the intensity of the white line increases again (Fig. 4). Based on the calculations on the degeneracy, Co²⁺ has a near octahedral coordination in the amorphous compound. Due to their presence in the *T*-range of 210–500 °C, a low amount of Co is always bound in octahedral coordination.

From the crystal field stabilisation energy (CFSE) perspective, Ni²⁺ as a d⁸ ion strongly prefers octahedral coordination. The difference in CFSE between tetrahedral and octahedral geometries is significantly higher ($|\Delta\text{CFSE}_{\text{O-T}}| = 0.84$ with $\text{CFSE}_{\text{O}} = -1.2 \Delta_{\text{O}}$, $\text{CFSE}_{\text{T}} \approx -0.36 \Delta_{\text{O}}$ assuming $\Delta_{\text{T}} = 4/9 \Delta_{\text{O}}$ ⁴⁵) than the difference between both geometries in the Co-system. Co²⁺ as a d⁷ ion exhibits only a slight difference between the CFSE_O and CFSE_T in high spin configurations ($|\Delta\text{CFSE}_{\text{O-T}}| = 0.27$ with $\text{CFSE}_{\text{O}} = -0.8 \Delta_{\text{O}}$ and $\text{CFSE}_{\text{T}} \approx -0.53 \Delta_{\text{O}}$). This difference in energy/stability in comparison with Ni²⁺ promotes an additional occurrence of tetrahedral coordination in the Co-system.^{46,47}

Summarised order of phase transitions with increasing temperature

Based on the combined multi-modal characterisation, we established the order of phase transitions in our M-struvite systems. At first, Ni-struvite decomposes to an amorphous

phase while Co-struvite transforms through a crystalline-crystalline phase transition to Co-dittmarite at *T* = 90 °C (Fig. 1). The pseudomorphic phase transition of the amorphous Ni-PO₄ phase formed from Ni-struvite, leads to a relict “Ni-struvite crystal skeleton” visible in the SE images (ESI: Fig. S2†). Although pseudomorphism indicates a weak degree of densification and structural reconfiguration (otherwise precursor crystal framework would collapse), the relict crystal morphology is not in every case preserved as in the 500 °C sample (ESI: Fig. S2K†). Looking at the degassing events of heated Co-struvite based on XRD and DSC, an amorphous phase formed in a second and third thermal event at 140 °C and 250 °C, associated with the partial decomposition of Co-dittmarite and HCoPO₄·H₂O (Fig. 1). The amorphous phase is proposed to be akin to HCoPO₄ based on the TGA/DSC, as is also suggested by other studies in the literature^{48,49} Surprisingly, ammonia is still present in the amorphous phases up to 400 °C in both metal systems, which is important for potential proton carrier molecules. Based on the EXAFS fits, the spectra of the amorphous phases *T* > 400 °C resemble mostly those of crystalline Ni-pyrophosphate with less pronounced long-range order above *R* distances > 4 Å. This fact indicates a beginning of pyrophosphatisation already at 400 °C in agreement with FT-IR and FT-RS. Further heating leads to the actual crystallisation of M-pyrophosphate. At 400 °C Co₂P₂O₇ forms, although a major proportion of amorphous phases is present. In contrast, the amorphous nickel phosphate phase remains stable up to 600 °C until it converts to Ni₂P₂O₇. The humidity and heating time/rate can significantly influence the phase stability and crystallinity, especially in the low *T*-region below 400 °C. This ability of M-struvite to sequentially convert into various amorphous and crystalline transition metal phosphate phases opens the space for possible crystal and phase engineering and upcycling of the waste-derived struvite^{29,40,50,51} into tailored TMPs, as we indeed explore below.

Proton conducting properties of selected Ni- and Co-phosphates

The structural results presented above point to the potential presence of extensive hydrogen-bonded frameworks involving the coordination of water H₂O and ammonia NH₃ molecules in some of the amorphous or crystalline TMP phases of Ni and Co. Such frameworks indicate a potential for good proton conductive properties of these compounds. To characterise our processed TMP phases as functional proton conductive materials, proton conductivity (PC) measurements were performed on the pressed powder pellets at different relative humidity values and temperatures (Fig. 5, see Methods Proton conductivity measurements for more details, Fig. S9–S11†). Before and after each PC measurement, a diffraction pattern was recorded to demonstrate the phase stability (Fig. S12†). The impedance spectra were fitted with a minimal circuit model consisting of a parallel constant phase element (CPE) and resistor (R) in series with two parallel Warburg elements (*W*_s and *W*_o, inset: Fig. 5B).



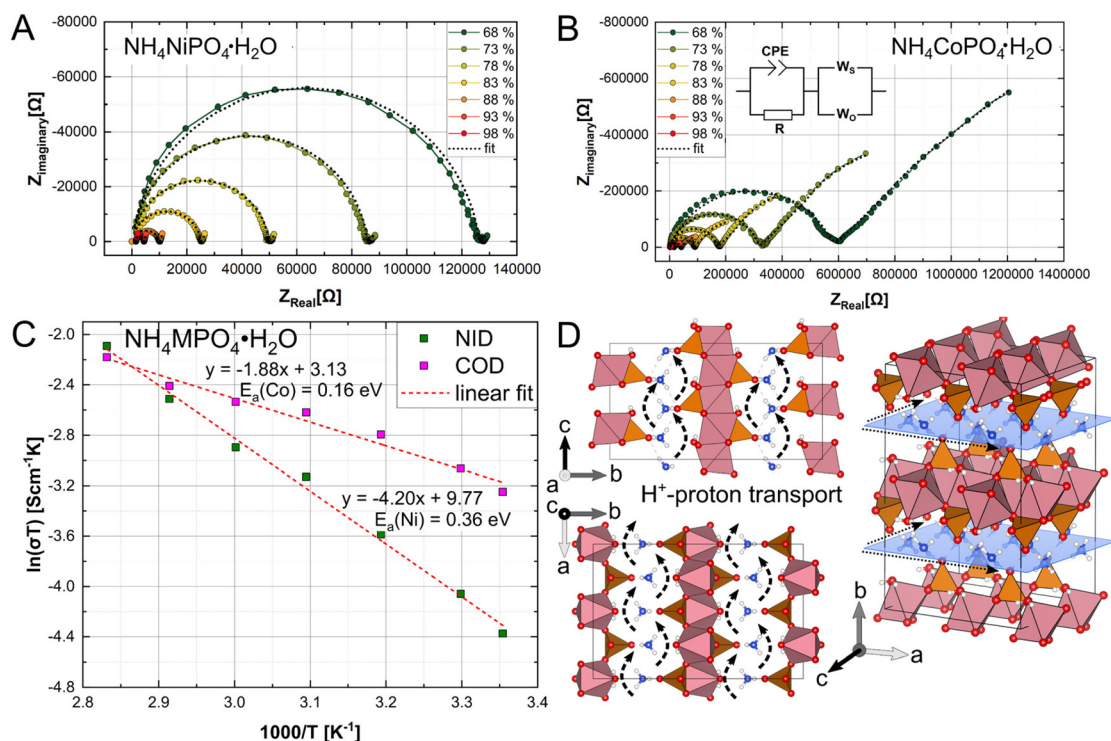


Fig. 5 Nyquist plots with real and imaginary impedance of (A) Ni- and (B) Co-dittmarite, $\text{NH}_4\text{MPO}_4 \cdot \text{H}_2\text{O}$ with $\text{M} = \text{Ni}, \text{Co}$ at $\text{RH} = 68\%–98\%$ at $T = 25^\circ\text{C}$; inset in B: equivalent circuit of the fitting model; (C) Arrhenius plots of Ni- and Co-dittmarite with $\ln(\sigma T)$ vs. $1000/T [\text{K}^{-1}]$ ($T = 25^\circ\text{C}–80^\circ\text{C}$), red dashed line = fit of the data; inset: DVS isotherms of Ni- and Co-dittmarite; (D) view on the $(0k0)$ plane of the crystal structure of M-dittmarite $\text{Pmn}2_1$ from different perspectives; dashed black arrows and blue marked $(0k0)$ planes indicate enhanced proton migration pathways.

The high-frequency R-CPE circuit represents the electronic resistivity of the metal phosphate phase. In our model, the low-frequency $W_s\text{--}W_o$ circuit models the effects of the proton transport through grain boundaries. The Warburg short element expresses the transmissive, while the Warburg open element points to the reflective character of the grain boundary. No singular Warburg element (either a general, short or open W), fitted the impedance spectra sufficiently well. Including two Warburg elements resulted in much more accurate fitting results. From the intersection of the fits and the abscissa axis (real part of impedance Z'), the proton conductivity was calculated (more details in ESI: Methods proton conductivity measurements†). As the current is measured perpendicularly to the pellet base, the proton conductivity is interpreted as bulk proton conductivity under the assumption of randomly orientated grain boundaries.⁵² The activation energy (E_A) of proton migration is the amount of energy required for a proton to jump from one ionic domain to another, an important value to distinguish between Grotthuss and vehicle mechanism.

Selected samples of the Ni- and Co T-series (Fig. 1) were analysed. For Co: Co-dittmarite processed at $T = 90^\circ\text{C}$, NH_4CoPO_4 at $T = 300^\circ\text{C}$ and Co-pyrophosphate at $T = 800^\circ\text{C}$. For Ni: Ni-dittmarite at $T = 90^\circ\text{C}$ in $100\% \text{H}_2\text{O}$, amorphous NiPO_4 at $T = 500^\circ\text{C}$ and Ni-pyrophosphate at $T = 800^\circ\text{C}$. All data and fits associated with the proton conductivity are summarised in the ESI (Fig. S9–S11 and Tables S22, S23†). The electrochemical impedance data were plotted in Nyquist diagrams

with the imaginary part of the impedance on the y -axis $Z_{\text{imaginary}}$ and the real part of the impedance on the x -axis Z_{real} . All Nyquist plots demonstrate semicircles in the high-frequency region and an elongated tail in the low-frequency region. The M-dittmarite compounds of Ni and Co, heated at $T = 90^\circ\text{C}$ and stable up to 200°C , exhibit the highest proton conductivities of all analysed phases at $1.4 \times 10^{-4} \text{S cm}^{-1}$ and $4.2 \times 10^{-5} \text{S cm}^{-1}$ for Co- and Ni-dittmarite, respectively, at $\text{RH} = 98\%$ and $T = 25^\circ\text{C}$. Considering the low-frequency region, the Warburg coefficients from the fits show a slight increase of the W_s/W_o ratio with higher RH, visible in both M-dittmarite components from nearly near zero to 2.5(1) for Ni and 6.5(9) for Co (ESI: Table S23†). This could indicate a more transmissive nature of the grain boundaries at high RH. From a crystallographic point of view, protons can “jump” in the hydrogen-bond network mainly in the $(0k0)$ plane in two directions, where ammonium NH_4 and the coordinating water of the metal octahedron are located.

Since protons are most likely transported through 2D layers, build up by the ammonium sites (Fig. 5) in the M-dittmarite structure $\text{Pmn}2_1$, the compound probably demonstrates a significant anisotropic proton conductivity in directions in the (010) plane. Perpendicularly to this direction, the proton conductivity is most likely decreasing. As we measured the bulk proton conductivity of a pressed pellet of the material, grain boundaries, microstructure and the orientation of the different grains effect the overall bulk properties. Due to these



anisotropic features, the overall bulk proton conductivity could be influenced. Furthermore, to evaluate the water affinity and adsorption, a major factor influencing the proton conductivity, dynamic vapour sorption (DVS) measurements were carried out (inset in Fig. 6C and ESI: Fig. S11†). Co-dittmarite displays a final water uptake of $0.0263 \text{ cm}^3 \text{ g}^{-1}$ at RH = 98%, corresponding to the sorption of 0.27 H_2O molecules per formula unit. Ni-dittmarite exhibits lower values of $0.0164 \text{ cm}^3 \text{ H}_2\text{O/g}$ at 98% RH equating to 0.17 molecules of H_2O per formula unit. However, for all the RH values and $T = 25^\circ\text{C}$, the Co compound exhibits proton conductivities around one order of magnitude higher than those found for the Ni analogue. This discrepancy disappears with higher temperatures up to 80°C (Fig. 6C). Although both compounds share the same structure, Co^{2+} exhibits higher proton conductivity at $T = 25^\circ\text{C}$ in Co-dittmarite, which can be correlated with slight structural differences compared to Ni-dittmarite, as indicated by our XAS measurements. The differences in bond lengths, especially of the first coordination sphere Co–O and Ni–O could cause different local polarities resulting in a changed proton mobility (ESI: Tables S11 and S13†). The adsorbed amount of water could only promote an increase of proton conductivity at high RH > 80%, but not at lower RH $\leq 80\%$. The dependency of the proton conductivity on the water partial pressure (associated with the RH) is similar for both dittmarite phases, because their $\log(\sigma)$ vs. $\log(\text{pH}_2\text{O})$ slope values are relatively close to each other (16 ± 2 (Ni) vs. 13 ± 1 (Co), Table S22†). When examining other processed phosphate phases, NH_4CoPO_4 shows a significant proton conductivity of $6.42 \times 10^{-5} \text{ S cm}^{-1}$ at RH = 98% and $T = 80^\circ\text{C}$. The other phases, $\text{Ni}_2\text{P}_2\text{O}_7$, $\text{Co}_2\text{P}_2\text{O}_7$ and the amorphous Ni- PO_4 phase of heated at 500°C , demonstrate low to medium proton conductivities in the range of 10^{-6} – $10^{-7} \text{ S cm}^{-1}$ (ESI: Fig. S8 and S9†). The M-dittmarite and $\text{M}_2\text{P}_2\text{O}_7$ phases did not change its phase composition after the PC measurement, while the amorphous Ni-phase and NH_4CoPO_4 crystallised. In addition, the water partial pressure-dependent proton conductivity behaviour of the $\text{M}_2\text{P}_2\text{O}_7$ compounds is significantly different from that of Ni-, Co-dittmarite, and NH_4CoPO_4 . Due to a highly condensed structure, the adsorbed water in the pyrophosphates, even at high RH (>85%), does not provide a significant jump in proton conductivity (ESI: Fig. S10 and Table S22†).

This work's goal was to obtain a TMP material with high thermal stability and significant proton conductivity optimal for their potential applicational use, e.g., proton exchange membranes in fuel cells (Fig. 1). For this purpose, the mechanism of ionic conductivity/proton migration in the material had to be investigated by determining its activation energy. In general, proton conductivity is assumed to rely on two major transport mechanisms, the Grotthuss mechanism and the vehicle mechanism.⁵³ The Grotthuss mechanism *i.e.* proton jumping, points to the migration processes of protons through translation from an $\text{H}_{2n+1}\text{O}_n^+$ ion to a water molecule by tunnelling in a hydrogen bond with subsequent molecule rotation. This diffusing proton in the newly formed $\text{H}_{2n+1}\text{O}_n^+$ ion can attach to the next neighbour water molecules resulting

in a successive motion of H^+ over small molecular distances through the compound.^{53,54} Recent O–K-edge XAS results could prove the dominant occurrence of an H_7O_3^+ proton complex in contrast to the conventional explanation of Zundel or Eigen ions, H_5O_2^+ and H_9O_4^+ .⁵⁵ In the vehicle mechanism, molecules such as water H_2O , ammonia NH_3 , or alkali metals Na, K, *etc.* act as proton acceptors/Brønsted bases and transport the protons as carriers through the compound over large molecular distances.⁵³ The activation energy of proton migration E_A , calculated from the Arrhenius plots, is a key parameter to distinguish between those mechanisms.^{53,56} A value between 0.1–0.4 eV per proton indicates proton jumping, meaning the transfer of protons is accomplished between the proton donor and acceptor. A value greater than 0.4 eV per particle implies a vehicle mechanism where a proton transport proceeds *via* the diffusion of protonated carriers.^{52,53} A low content of ammonia and water in the compound resulting from heating at elevated temperatures would decrease the proton conductivity performance as a weaker hydrogen-bond network and a lower number of potential carriers of protons are available. The crystal structure determines the possibilities and pathways for preferential proton transport *e.g.* through 1D channels, 2D layers, or 3D frameworks, and the binding strength of the hydrogen network (*e.g.* “free” crystal water vs. bound hydroxyl groups). Therefore, the absolute number of protons and carrier molecules in the structure and the internal structure significantly influence the overall PC performance. The dittmarite structure $\text{Pmn}2_1$ seems to provide a beneficial structure for enhanced proton conduction as protons can migrate through the weakly bound ammonium layers in two dimensions. The increase of the Warburg ratio W_s (transmissive boundary)/ W_o (reflective boundary) with increasing RH could represent a more transmissive behaviour of the grain boundaries as interpreted by other studies.^{57–59} More water units are adsorbed at the interfaces providing an enhanced proton transfer due to a higher density of potential transporting molecules. The calculated values for the activation energy E_A infer a Grotthuss mechanism (<0.4 eV) in most compounds except for the Ni- and Co-pyrophosphates. In these compounds, no structural water or ammonia are present, and carrier molecules only accomplish proton conduction, indicated by the high activation energy E_A and the slopes of $\log(\sigma)$ vs. $\log(\text{pH}_2\text{O})$ (ESI: Table S22†). The similar increase of proton conductivity of Ni- and Co-dittmarite at higher RH/partial pressures of water majorly agrees with the uptake of water visible in the DVS isotherms and the slopes of $\log(\sigma)$ vs. $\log(\text{pH}_2\text{O})$ (Fig. 6 and ESI: Fig. S9–S11, Table S22†). The activation energies for proton migration of 0.16 eV for Co-dittmarite and 0.36 eV for Ni-dittmarite are relatively low compared to other oxides and phosphate compounds (Table 1 and ESI: Table S23†).

In addition, based on recent research,⁶⁰ the dittmarite structure shows high suitability as a framework crystal for high-entropy phosphate materials (HEMPs). HEMPs are known to be electrochemically more active than their pure endmembers as the high configurational entropy of mixing



Table 1 Summarized proton conductivity data of most promising metal phosphate compounds. o.w. = our work

Material	Chemical composition	T [°C]/RH [%]	σ [S cm ⁻¹]	E_a (eV)	Ref.
Ionomer	NAFION®	25/98	7.80×10^{-2}	—	63
Na-Al-PO ₄	Na ₆ [(AlPO ₄) ₈ (OH) ₆].9H ₂ O	20/98	3.60×10^{-3}	0.21	64
Doped Sn ₂ P ₂ O ₇	In _{0.1} Sn _{0.9} P ₂ O ₇	300/—	0.195	—	65
Sn-PO ₄	Sn(HPO ₄) ₂ ·3H ₂ O	100/—	1.00×10^{-2}	—	27
ZrP ₂ O ₇	Zr(P ₂ O ₇) _{0.81} (O ₃ POH) _{0.38}	20/98	1.30×10^{-3}	—	25
TiP ₂ O ₇	TiP ₂ O ₇	100/—	4.40×10^{-3}	0.14	66
Ti-PO ₄	Ti ₂ (HPO ₄) ₄	20/95	1.20×10^{-3}	0.13	67
org./inorg. Mn-PO ₄ -comp.	(C ₂ H ₁₀ N ₂) [Mn ₂ (HPO ₄) ₃](H ₂ O)	20/99	1.60×10^{-3}	0.22	68
org./inorg. Fe-PO ₄ -comp.	(C ₄ H ₁₂ N ₂) _{1.5} [Fe ₂ (OH)(H ₂ PO ₄)(HPO ₄) ₂ (PO ₄)] 0.5(H ₂ O)	40/98	5.10×10^{-4}	—	69
org./inorg. Co-PO ₄ -comp.	(C ₂ N ₂ H ₁₀) _{0.5} CoPO ₄	56/98	2.00×10^{-3}	1.01	70
org./inorg. Zn-PO ₄ -comp.	NMe ₄ Zn[HPO ₄][H ₂ PO ₄]	30/98	1.30×10^{-2}	0.92	71
K-V-PO ₄	K ₂ [(VO) ₂ (HPO ₄) ₂ (C ₂ O ₄)]	40/95	1.15×10^{-2}	—	72
NH ₄ -Ni-PO ₄	NH ₄ NiPO ₄ ·H ₂ O	25/98	4.20×10^{-5}	0.36	o.w.
NH ₄ -Co-PO ₄	NH ₄ NiPO ₄ ·H ₂ O	25/98	1.40×10^{-4}	0.16	o.w.

promotes their reactivity.^{61,62} In such a way, a high-entropy transition metal dittmarite compound derived from a high-entropy struvite could show even better performance than Ni- or Co in proton conductivity, but this aspect has to be further evaluated.

Finally, we showed in our previous study,⁴⁰ that M-struvite could be precipitated from industrial wastewater. Here we demonstrate that through a simple thermal treatment, these reclaimed TMP materials could be upcycled for applicational purposes (Fig. 1). The processed M-dittmarite compounds of Ni- and Co demonstrate high to superprotonic conductive behaviour comparable to ionomers, *i.e.* Nafion 117,^{63,73,76} MOFs,⁷⁴ perovskites,⁷⁵ or other oxide/phosphate compounds.

A summary of selected high-performing proton conductive transition metal phosphate compounds can be found in Table 1 in further detail in the ESI (Table S23†). While most of the reported compounds require a complex multistep synthesis, our as-obtained materials could be precipitated in a facile one pot way directly out of wastewater. For example, ceramic oxide proton conductors, often perovskite-type, require in most cases an energy intensive, and complex solid state or solvothermal syntheses. Even if the absolute bulk proton conductivity values of the as-obtained Ni- and Co-dittmarites are not among the highest measured, they are still sufficiently conductive considering that we are discussing recycled waste materials. Due to their high thermal and chemical stability up to 210 °C, the M-dittmarite compounds could endure the optimal operating temperatures in conventional fuel cells. These compounds may be upcycled directly or could be reused indirectly by processing them even further to an advanced proton conductive material. In combination with their high compatibility with other cell components (electrodes, electrolytes, seals) and low cost, M-dittmarites are promising proton-conducting materials.^{77,78}

Conclusions

In this study, we followed the thermal history of Ni- and Co-struvite decomposition all the way to their crystallisation into

M-pyrophosphates. Ni-struvite forms an amorphous phase during the thermal treatment, which preserves octahedral metal coordination with slight distortion at all temperatures. The complex amorphous structure increasingly resembles a local structure of a Ni-pyrophosphate with progressive temperatures. In contrast, heated Co-struvite exhibits several crystalline-crystalline phase transitions with an associated change of metal coordination from octahedral to tetrahedral, and back to octahedral. In both systems, volatile compounds such as H₂O or NH₃ are still present in the solids up to 400 °C, potentially making transition metal phosphates promising high-temperature water-retaining materials. The metal and phosphate coordination environment associated with the amount of bound water and ammonia in the thermally treated TMPs could be directly linked to their proton conductivity performance. Indeed, with values of $>10^{-4}$ S cm⁻¹ at room temperature, a low activation energy of 0.16 eV, and high thermal stability above 200 °C, the layered Ni- and Co-dittmarites are the most promising candidates as proton exchange material, among all the obtained phases. Importantly, we showed that such functional materials could be directly made by thermally processing M-struvites, which might be precipitated from various wastewater streams.

Conflicts of interest

There are no conflicts to declare.

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