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ARTICLE TYPE

High-throughput microwave-assisted discovery of new metal phosphonates Mark Feyand,^{*a*} Christopher F. Seidler^{*d*}, Carsten Deiter,^{*b*} Andre Rothkirch,^{*b*} Alexandra Lieb^{*c*}, Michael Wark^{*d*} and Norbert Stock^{*a*,*}

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A systematic study was carried out to investigate the influence of linker geometry, metal ionic radius as well as the nature of the counter ions on the structure formation of metal tetraphosphonates. Two tetraphosphonic acids *p*- and *m*-(H₂O₃PCH₂)₂N-CH₂-C₆H₄-CH₂-N(CH₂PO₃H₂)₂, six metal ions (Ca²⁺, Mn²⁺, Co²⁺, Ni²⁺, Zn²⁺, Cd²⁺) and two different counter ions (Cl⁻ and NO₃⁻) were employed using high

¹⁰ throughput methods. Microwave (MW)-assisted heating led to the discovery of ten new metalphosphonates which crystallize in three different crystal structures. The combination of direct methods and force field calculations allowed us to establish the crystal structures. The counter ion and the ionic radii of the metal ions have a profound influence on the crystalinity and the formed crystal structure. All compounds were characterized in detail by thermogravimetric analyses, IR spectroscopy and magnetic ¹⁵ susceptibility measurements. The proton conductivity on two selected compounds is also reported.

Introduction

Metal phosphonates are in the focus of recent research activities due to their interesting properties such as magnetism, porosity,^{1, 2} luminescence³ or catalysis.⁴ Metal phosphonates show a large

- ²⁰ structural variety depending on the linker molecule or the incorporated metal ion. The properties may change drastically upon variation of the constituents. The inorganic building units observed in metal phosphonates range from isolated clusters⁵ to chains^{2, 6-8} or layers.^{8, 9, 10}
- ²⁵ A suitable method to synthesize metal phosphonates are solvothermal reactions. Recently, microwave assisted heating was applied to synthesize new metal phosphonates.^{7, 11, 12} Microwave (MW) assisted reactions have proven to lead to higher purities, high yields and much shorter reaction times.¹³ However, the
- ³⁰ sealed reaction conditions make obtaining information about crystallization difficult. In addition, the outcome of such reactions, i.e. the product formation, is not predictable and large variable parameters such as temperature, pH or molar ration of reactants have to be studied explorative. A combinatorial and
- ³⁵ more systematical approach provides the high-throughput (HT) methods.^{14, 15} They allow to investigate efficiently and fast important reaction parameters such as reaction temperature,^{14, 16} concentration of reactants,¹⁷ ionic radii,¹⁸ the pH,^{10, 16, 17} or length of the organic moiety.¹⁰ Recently HT MW assisted reactions were
- ⁴⁰ employed to establish the fields of formation of new metalorganic framework compounds. However, these methods have never been employed to metal phosphonate chemistry before.¹⁹ We are interested in the synthesis of new metal phosphonates and for the present study we have chosen the tetraphosphonic acids *p*-
- ⁴⁵ and m-(H₂O₃PCH₂)₂N-CH₂-C₆H₄-CH₂-N(CH₂PO₃H₂)₂. They have previously been shown to lead to three new compounds containing Ca²⁺ and Cu²⁺ ions.^{20, 21}

Here we present the results of the systematic HT investigation of

the synthesis of metal tetraphosphonates using MW-assisted 50 heating and varying the linker geometry as well as the metal and counter ion. In addition, the crystal structures are determined from X-ray powder diffraction data and the detailed characterisation of the products is shown.

Experimental section

55 Materials and methods.

All starting materials were obtained from Aldrich or ABCR and were used without further purification. The two tetraphosphonic acids were synthesized using a slightly modified Mannich-type procedure.22 XRPD measurements were carried out on a Stoe 60 Stadi P diffractometer in transmission geometry with Cu-Kal radiation, equipped with an image plate detector system or a linear PSD detector system for high resolution data. The MIR spectra were recorded on a Bruker ALPHA-P FT-IR spectrometer in the spectral range 4000-400 cm⁻¹. For the thermogravimetric 65 analyses under air a NETSCH STA 409 CD analyzer was used with a heating rate of 4 K/min and an air flow rate of 75 ml/min. High resolution powder XRD patterns were measured at beamline G3 (light source DORIS)²³ and beamline P08 (light source PETRA)²⁴ both at HASYLAB, DESY. Scale up syntheses under 70 conventional heating and were carried out under solvothermal conditions using DURAN culture tubes D50 GL 14 M.KAP, SCHOTT. An Anton Paar Synthos 3000 microwave reactor was used for the high-throughput syntheses. Temperature dependent X-ray diffraction patterns were recorded on a Panalytical 75 Empyrean diffractometer equipped with a PIXcell detector using a MRI TC radiation chamber with Pt heater and Al₂O₃ crucibles.

Synthesis of $p-(H_2O_3PCH_2)_2NCH_2-C_6H_4-CH_2N(CH_2PO_3H_2)_2$ ($p-H_8L$)

⁸⁰ 5.00 g (36.7 mmol) *p*-xylenediamine und 12.0 g (147 mmol)

phosphorous acid were added to 60 mL half concentrated hydrochloric acid.²² The mixture was heated to reflux and 13 mL (161 mmol) 37 % formalin solution were slowly added. The reaction was then heated for additional 14 h and cooled to room

- ⁵ temperature. The white precipitate was filtered and washed with deionized water. The product was identified as *p*-H8L by ¹H and ³¹P-NMR spectroscopy (yield 45.17 g 60 % based on *p*-xylenediamine) (see supporting information).
- 10 Synthesis of m-(H2O3PCH2)2NCH2-C6H4-CH2N(CH2PO3H2)2 (m-H8L)
- 10.00 g (73.4 mmol) *m*-xylenediamine und 24.0 g (294 mmol) phosphorous acid were added to 60 mL half concentrated hydrochloric acid.²² The mixture was heated to reflux and 26 mL ¹⁵ (322 mmol) 37 % formalin solution were slowly added. The reaction was then heated for additional 14 h and cooled to room temperature. The white precipitate was filtered and washed with
- deionized water. The product was identified as m-H₈L by ¹H and ³¹P-NMR spectroscopy (yield 67.76 g 45 % based on m-
- 20 xylenediamine) (see supporting information).

HT Microwave Assisted Reactions

The HT syntheses were carried out in 4 ml microwave glass vials in a 24 reactor setup using an Aton Parr HT reactor system. The 25 reactions were carried out at 150 °C for 5 h using a power of 400

- W. The molar ratio H_8L : M^{2+} was kept at 1:1 and the concentration was fixed by using 25 mg of H_8L per reaction. To investigate the influence of the linker geometry 12 reactions per linker were carried out. To investigate different metal ion radii
- ³⁰ we studied Mn²⁺, Ca²⁺, Co²⁺, Ni²⁺, Zn²⁺ and Cd²⁺ using Cl⁻ and NO₃⁻ as counter ions. The obtained products were identified by X-ray powder diffraction. The results of the high-throughput investigation are summarized in Figure 2. The exact reaction conditions are given in the supporting information (Table S1).

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Scale-up synthesis of $Cd[p-(HO_3PCH_2)_2N(H)-CH_2-C_6H_4-CH_2-N(H)(CH_2PO_3H)_2]$ (1)

40 mg p-H₈L (78mmol), 117 μ L (234 mmol) of a 2 M CdCl₂ solution and 3 ml deionized water were mixed in a DURAN ⁴⁰ culture tubes, sealed and heated up to 150 °C for 12 h. The reaction mixture was filtered and washed with water. Using pH paper pH = 2 was determined for the initial and final reaction solution. The yield was 33 mg, 68 % based on p-H₈L.

45 Scale-up syntheses of M[p-(HO3PCH2)2N(H)-CH2-C6H4-CH2-N(H)(CH2PO3H)2(H2O)] (M= Mn, Co, Ni, Zn) (**2**)

- 40 mg p-H₈L (78 mmol), 39 μ L (78 mmol) of a 2 M M(NO₃)₂ solution and 3 ml deionized water were mixed in a DURAN culture tubes, sealed and heated up to 150 °C for 12 h. The
- ⁵⁰ reaction mixture was filtered and washed with water. Using pH paper pH = 2 was determined for the initial and final reaction solution. The yields were 34, 13, 23, 26 mg, 70, 25, 45, 50 % based on p-H₈L, for M= Mn, Co, Ni, Zn, respectively.

55 Scale-up syntheses of M[m-(HO₃PCH₂)₂N(H)-CH₂-C₆H₄-CH₂-N(H)(CH₂PO₃H)₂] · H₂O (M=Ca, Mn, Co, Ni, Zn) (3) 40 mg m H₂ (78 mmg) 20 mJ (78 mmg)) of a 2 M MCla

40 mg *m*-H₈L (78 mmol), 39 μ L (78 mmol) of a 2 M MCl₂ solution. 20 μ L (39 mmol) of a 2 M NaOH solution and 3 ml deionized water were mixed in a DURAN culture tubes, sealed ⁶⁰ and heated up to 150 °C for 12 h. The reaction mixture was

filtered and washed with water. Using pH paper pH = 2 was determined for the initial and final reaction solution. The yield was 38 mg, 75 % based on m-H₈L.

Crystal Structure Determination

⁶⁵ All compounds were obtained as microcrystalline powders. Thus, the structures had to be determined from X-ray powder diffraction data. The important crystallographic parameters are given in Table 1.

Table 1. Crystallographic data of compound **1**, **2** and **3**. (SH = spherical ⁷⁰ harmonics series)

	compound 1	compound 2	compound 3
formula sum	C10H18N2CdO	$C_{10}H_{20}N_2ZnO$	C10H20N2NiO1
	${}_{12}P_4$	${}_{13}P_4$	$_{3}P_{4}$
radiation source	Synchrotron	Synchrotron	Laboratory
wavelengths / Å	1.54296	0.824516	$CuK\alpha_1$
a / Å	12.1198(4)	10.5960(3)	22.2633(4)
b/Å	15.1091(4)	7.5569(2)	8.5904(2)
c∕Å	5.5632(1)	6.9120(2)	9.8834(2)
lpha / °	90	106.42(2)	90
β / °	97.28(2)	98.50(3)	90
γ/°	90	86.79(1)	90
space group	$P2_{1}/a$	<i>P</i> -1	Pmcn
\mathbf{R}_{wp}	2.80	3.60	5.77
R _{Bragg} (without SH)	1.03(3.25)	0.79(4.25)	1.86(8.74)
GoF	1.42	1.97	1.81
number of restrains	6	8	10
number of	80	85	78
parameters			
volume / Å ³	1010.62(5)	524.99(4)	1890.22(6)
Z	2	1	4

Crystal Structure Determination of compound 1

High resolution XRPD data of compound 1 (Cd) was collected at beamline G3, HASYLAB, DESY, Hamburg using a wavelengths 75 of 1.54296(2) Å selected by a double silicon single crystal monochromator and determined with a LaB₆ standard. The diffracted beam was detected by a scintillation counter mounted behind a soller slit system. The sample was measured in a capillary (0.7 mm diameter). The powder pattern was successful ⁸⁰ index using Topas Academic and the possible space group $P2_1/a$ was suggested.¹¹ The crystal structure was solved in $P2_1/a$ using direct methods implement in EXPO2009. The heavy atom positions and the phosphonate groups could be located directly from the Fourier map. The structure model was completed using 85 Materials Studio 5.3 applying force field calculations.²⁵ An universal force field was used as implemented in Materials Studio. The obtained model was further refined by Rietveld methods in the 2θ range of 9 to 80° using Topas Academic. The final Rietveld refinement included six background points, three 90 Thompson-Cox-Hastings profile parameters, four lattice parameters, one scaling factor, a simple axial model, a fourth order spherical harmonics series, 45 atomic coordinates and three temperature factors (Cd, P, and C/N/O). The P-O and P-C distances were soft restrained. The final Rietveld plot is given in 95 Figure 3.

Crystal Structure Determination of compound 2

A high resolution XRPD pattern of compound 2 (Zn) was

recorded at beamline P08, PETRA, DESY, Hamburg with a wavelengths of 0.824516 Å. The powder pattern was successfully indexed using Topas Academic. Due to the small cell volume of 525(1) Å³ only one twofold deprotonated linker ion and one Zn²⁺

- ⁵ ion are present in the unit cell. A structure model was derived in the space group *P*-1 by molecular modeling using Materials Studio 5.3. Details are given in the supporting information (Fig. S3). The starting model was refined by Rietveld methods in the range of 4-40° 2θ using Topas Academic. The final Rietveld
- ¹⁰ refinement included six background points, three Thompson-Cox-Hastings profile parameters, six lattice parameters, one scaling factor, a simple axial model, a fourth order spherical harmonics series, 48 atomic coordinates and three (Zn, P, and C/N) temperature factors. The P-O and P-C distances were soft
- 15 restrained. The final Rietveld plot is given in Figure 5.

Crystal Structure Determination of compound 3

The powder pattern of **3** (Ni) was measured on a Stoe Stadi P diffractometer equipped with a linear PSD detector. The structure

- ²⁰ was solved using direct methods implemented in EXPO2009. The inorganic building unit was directly located from the Fourier map and completed by force field calculations using Materials Studio 5.3. The structural model was refined by Rietveld methods in the range of 9 to 75° 2θ using Topas Academic. The final Rietveld
- ²⁵ refinement subsumed six background points, three Thompson-Cox-Hastings profile parameters, four lattice parameters, one scaling factor, a simple axial model, a fourth order spherical harmonics series, 45 atomic coordinates and three (Ni, P, and C/N/O) temperature factors. The P-O, P-C and some C-C ³⁰ distances were soft restrained. The final Rietveld plot is given in
- Figure 7.

Proton Conductivity

- Proton conductivity was determined by impedance spectroscopy ³⁵ (IS).^{26, 27} An oscillating voltage of 100 mV was applied over a frequency range from 1 to 10⁶ Hz using a Zahner Zennium electrochemical workstation. The powder samples were pressed into small pellets (diameter 8 mm, thickness 0.5–1.5 mm) and incorporated into a stack comprising the sample sandwiched
- ⁴⁰ between two graphitic slices. The stack was placed in a PTFE sample holder, and two sintered metal electrodes were used to make electrical contact to the stack. The sample holder was placed in a gastight, temperature-controlled stainless steel chamber with an attached water reservoir.²⁸ The relative humidity
- ⁴⁵ (RH) in the cell was determined by the Clausius-Clapeyron relation and controlled by the heating of the water reservoir and the cell. Preceding each series of measurements, samples were equilibrated for 24 hours at 313K and at 50% RH. The sample was equilibrated for 1 hour at desired temperature and RH before
- ⁵⁰ measuring each data point. Each point was measured three times to determine reproducibility.

Results and discussion

To investigate the influence of the linker geometries the two ⁵⁵ tetraphosphonic acids $p-(H_2O_3PCH_2)_2N-CH_2C_6H_4CH_2 N(CH_2PO_3H_2)_2$ ($p-H_8L$) and $m-(H_2O_3PCH_2)_2N-CH_2C_6H_4CH_2-$ $N(CH_2PO_3H_2)_2$ (*m*-H₈L) were synthesized (Figure 1).

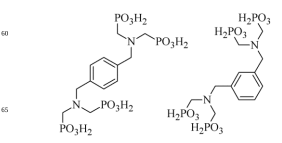


Fig. 1 Linker molecules used in this study. Left: $p-(H_2O_3PCH_2)_2N-CH_2C_6H_4CH_2-N(CH_2PO_3H_2)_2$ ($p-H_8L$), right: $m-(H_2O_3PCH_2)_2N-70$ CH₂C₆H₄CH₂-N(CH₂PO₃H₂)₂ ($m-H_8L$).

High Throughput Investigation

The influence of the geometry of tetraphosphonic acid on the formation of metal phosphonates was investigated using high-throughput methods.¹⁵ Thus six different metal ions with ionic ⁷⁵ radii between 0.74 and 1.14 Å (Mn²⁺, Ca²⁺, Co²⁺, Ni²⁺, Zn²⁺ and Cd²⁺) and different counter ions (NO3⁻ and Cl⁻) were used. 24 reactions were carried out under microwave assisted heating at 150°C for 5 h and a molar ratio of H₈L : M²⁺ = 1 : 1 (details are given in Table S1). The results of the study are presented in ⁸⁰ Figure 2.

Using p-H₈L and metal chlorides CdCl₂ leads to a the highly crystalline compound of composition [Cd(HO₃PCH₂)₂NH- $CH_2C_6H_4CH_2-NH(CH_2PO_3H)_2$] (1). The use of metal nitrates instead of metal chlorides leads to the isostructural compounds 85 [M(HO₃PCH₂)₂NH-CH₂C₆H₄CH₂-NH(CH₂PO₃H)₂(H₂O)] (2)with $M = Mn^{2+}$, Co^{2+} , Ni^{2+} and Zn^{2+} , respectively. Using Cd^{2+} as the cation only X-ray amorphous powders were obtained. Both Ca^{2+} salts leads to the literature known compound [$Ca(p-H_6L)$]]·2H₂O in much shorter reaction times of 5 h compared to 48 h $_{90}$ using conventional heating. In contrast, the reactions of $m-H_{8L}$ with metal nitrates only lead to X-ray amorphous compounds while reactions with the metal chlorides result in crystalline [M(HO₃PCH₂)₂NH-CH₂-C₆H₄-CH₂compounds $NH(CH_2PO_3H)_2](H_2O)$ (3) for $M = Mn^{2+}$, Ca^{2+} , Co^{2+} , Ni^{2+} and 95 Zn^{2+} and an X-ray amorphous product for $M = Cd^{2+}$.

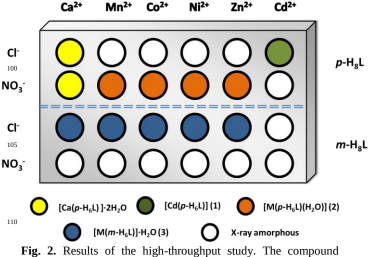


Fig. 2. Results of the high-throughput study. The compound $[Ca(p-H_6L)] \cdot 2H_2O$ has already been reported before.²¹

Crystal Structure of $[Cd(HO_3PCH_2)_2NH-CH_2C_6H_4CH_2-NH(CH_2PO_3H)_2]$ (1)

Using high resolution powder diffraction data the structure of **1** was determined by a combination of direct methods and ⁵ molecular modeling. The final results of the structure refinement are given in Figure 3.

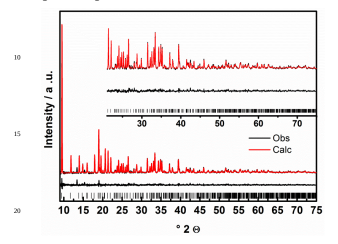


Fig. 3 Final Rietveld plot of **1**. The observed powder pattern is shown in black, the calculated powder pattern as an overlay in red and the ²⁵ difference (observed-calculated) of both is given by the lower black line. The allowed positions of the Bragg peaks are given as tics. The inset shows the higher 2θ range.

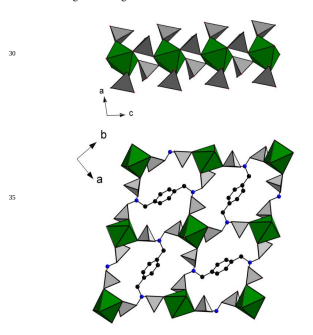


 Fig. 4. Crystal structure of compound 1. Chains of corner-sharing CdO₆
 ⁴⁰ and O₃PC polyhedra along the *c*-axis (top) and their connection to a threedimensional network (bottom). MO₆ polyhedra are shown in green, O₃PC polyhedra in grey, nitrogen and carbon atoms in blue and black, respectively.

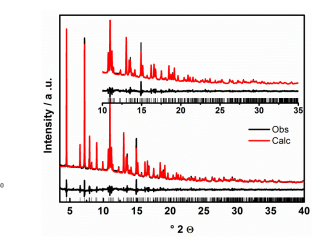


Fig. 5 Final Rietveld plot of 2 (Zn). The observed powder pattern is shown in black, the calculated powder pattern as an overlay in red and the difference (observed-calculated) of both is given by the lower black line.
⁵⁵ The allowed positions of the Bragg peaks are given as tics. The inset shows the higher 2θ range.

The crystal structure of **1** is shown in Figure 4 (and S4). Each Cd^{2+} ion is coordinated by one oxygen atom of six different phosphonate groups. These polyhedra are connected by the ⁶⁰ phosphonate groups to one-dimensional chains along the *c*-axis. These chains are interconnected by rest of the linker molecules to form a three dimensional network. The non-coordinating phosphonates groups are involved in hydrogen bonds which are shown in the supporting information (Fig. S6).

Crystal Structure of $[Zn(HO_3PCH_2)_2NH$ - $CH_2C_6H_4CH_2$ - $NH(CH_2PO_3H)_2(H_2O)]$ (2 (Zn))

- Taking into account the unit cell dimension, the space group symmetry and the unit cell content a structural model for **2** was ⁷⁰ derived by molecular modeling. The model was successfully refined using high resolution X-ray diffraction data (Figure 5). The coordination environment of the Zn²⁺ ions and the interconnection of the metal ions by the linker molecules are shown in Figure S7, a section of the crystal structure in Figure 6. ⁷⁵ The Zn²⁺ ions are sixfold coordinated by oxygen atoms of four different phosphonate groups and two water molecules. These ZnO₆ polyhedra are connected by the organic linker molecules along [010] to sheets (Fig. 6, middle), which are connected via two different kinds of hydrogen bonds (PO-H…OP and PO-⁸⁰ H…OH) as indicated by the donor acceptor distances (Fig. S8)
- between HO₃PC-groups phosphonate dimers.

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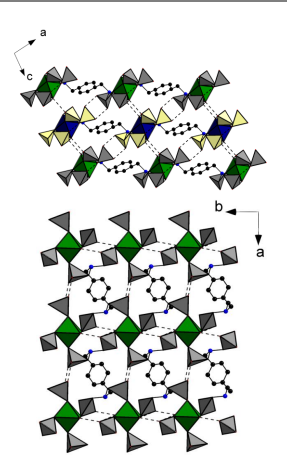


Fig. 6 Crystal structure of compound **2** (Zn). Connection of the Zn²⁺ ions by p-H₆L²⁻ ions in the *a,b*-plane (bottom) and possible interconnection of ¹⁵ the layers by postulated hydrogen bonds (top). MO₆ polyhedra are shown in green, O₃PC polyhedra in grey, nitrogen and carbon atoms in blue and black, respectively. One layer is shown by blue MO₆ and yellow CPO₃ polyhedra

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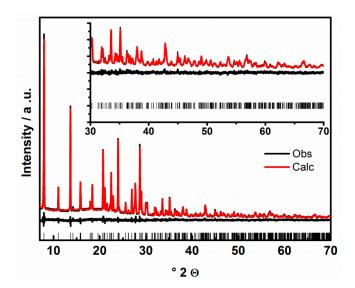


Fig. 7. Final Rietveld plot of **3** (Ni). The observed powder pattern is shown in black, the calculated powder pattern as an overlay in red and the difference (observed-calculated) of both is given by the lower black line. The inset shows the higher 2θ range.

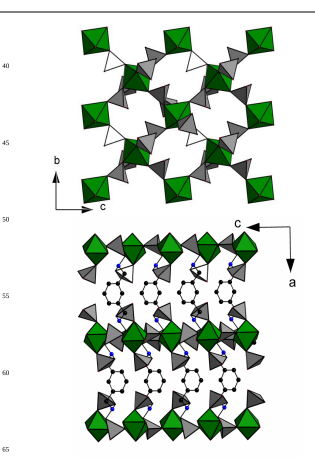


Fig. 8. Crystal structure of **3**. View along the *a*-axis (top) onto the layers and view along the layers connected by the organic linker molecules along the *b*-axis (bottom). Water molecules were omitted for clarity.

70 Crystal Structure of [Ni(HO3PCH2)2NH-CH2C6H4CH2-NH(CH2PO3H)2]·H2O (**3** (Ni))

In house XRPD data was used for the structure determination of **3** (Ni). Direct methods in combination with difference Fourier calculations and molecular modeling lead to a structural model

- ⁷⁵ that was successfully refined by Rietveld methods (Fig. 7). The coordination environment of the Ni²⁺ ions and the interconnection of the metal ions by the linker molecules are shown in Figure S7 and the crystal structure is shown in Figure 8. Each Ni²⁺ ion is six fold coordinated by oxygen atoms of six different phosphonate
- ⁸⁰ groups and layers in the *b*,*c*-plane are formed. These layers are interconnected to a three dimensional network by the organic linker molecules. Based on P-O···O-P and P-O···OH₂ distances between 2.50(2) and 3.06(2) Å hydrogen bonds can be postulated that further stabilize the structure (Fig. S9).

Structural Trends

As a result from the crystal structures we can conclude that the ionic radius has crucible influence on the crystal structure formation. Using *p*-H₈L with Ca²⁺ ions with the largest ionic ⁹⁰ radius (r_{ionic} = 1.14 Å) leads to the literature known compound [Ca(*p*-H₆L)]·2H₂O²¹ whereas the much smaller ions Mn²⁺, Ni²⁺, Co²⁺, and Zn²⁺ (r_{ionic} = 0.81, 0.79, 0.83, 0.88 Å, respectively) form an isostructural compound. Cd²⁺ ions with an intermediate ionic radius of 1.09 Å crystallizes in a different crystal structure ⁹⁵ (compound 1). All structures are pseudopolymorphic with 1

containing no, **2** one and $[Ca(p-H_6L)] \cdot 2H_2O$ two water molecules per formula unit. Thus, the Cd²⁺ ions in **1** seem to have the right size to allow for the formation of a three dimensional coordination polymer. In contrast, layers are formed which are

⁵ connected by hydrogen bonds for structure of compound **2**. This leads to a higher flexibility of the network and can adjust to the different ionic radii. In contrast the use of m-H₈L leads to one isostructural compound for Mn^{2+} , Ni^{2+} , Co^{2+} , Ca^{2+} and Zn^{2+} .

10 IR Spectroscopy

To learn more about the properties of the compounds we carried out IR spectroscopic and thermogravimetric measurements. The IR spectra of compound 1 (Cd), 2 (Zn) and 3 (Ni) are shown in Figure 9. All other IR spectra are given in the supporting 15 information (Fig. S9 and S11).

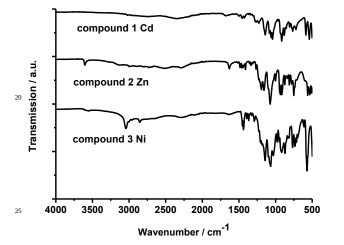


Fig. 9. IR-spectra of compound 1, 2 and 3.

- ³⁰ All three compounds show a very similar IR-spectrum. In the range between 1250 and 980 cm⁻¹ the typical CPO₃- bands are observed. The aromatic and aliphatic C-H stretching vibration are observed in the region between 3098-2970 cm⁻¹ and broad P-OH bands can be observed between 2840 and 2200 cm⁻¹. In ³⁵ accordance to the crystallographic data a broad band around 3600
- cm^{-1} from the coordinated and uncoordinated water molecules of compound 2 and 3 are present.

Thermal Stability

- ⁴⁰ The thermal stability was investigated by thermogravimetric analyses (TG). The results of all TG measurements are given in the supporting information (Fig. S12 - S13). All compounds show different thermal stabilities. Compound **1** (Cd) is stable up to 300 °C and decomposes in a multiple step decomposition. As
- ⁴⁵ expected, compound 2 (Zn) and 3 (Ni) show in addition to the decomposition in the range between 100 and 200 °C one step of weight loss which is due to the coordinated and uncoordinated water molecules. The removal of the uncoordinated water molecule in 3 (Ni) was further investigated by temperature
- ⁵⁰ dependent XRPD in the range of 40-400 °C (Figure 10).

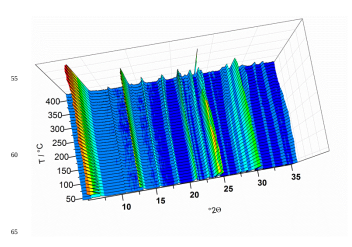


Fig. 10. Results of the temperature dependent XRPD measurements of compound 3 (Ni). The Bragg peak at ~ 23.2 °2 θ is due to the sample holder.

The sample is stable up to at least 400°C and shows slightly shifts ⁷⁰ of the Bragg peaks and changes in the intensity above 200°C. This can be explained by the loss of the water molecule. The lattice parameters of each pattern were extracted using parametric Pawley refinement methods as implemented in Topas academic¹¹ using the program Powder3D parametric.²⁹ The results are shown ⁷⁵ in Figure 11. Up to 200 °C all lattice parameters and the cell volume are slighly increasing as expected due to the thermal expansion of the lattice. Above 200°C the lattice parameters *a*and *c*-axis as well as the cell volume are rapidly decreasing due to the loss auf the water molecule while the *b*-axis lengths is ⁸⁰ increasing further. The cell volume is decreasing above 200°C.

These lattice parameter changes with temperature can be explained by inspecting the crystal structure of **3** (Ni). Along [100] and [001] PO···OP and PO···Ow···OP hydrogen bonds are observed (Fig. S9). Removal of the water molecule (Ow) st therefore results in a decrease of the *a*- and *c*-parameter. In contrast along [010] no hydrogen bonds are observed which is reflected in the increase of the *b*-parameter with temperature.

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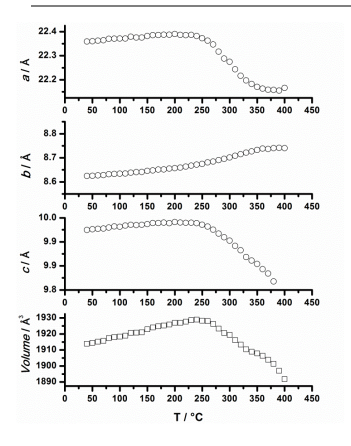


Fig. 11. Evolution of lattice parameters of compound 2. Parameters were obtained by parametric Pawley refinement.

Magnetic Properties of 3 (Mn, Co and Ni)

- ¹⁵ Since compounds 2 (Mn, Co, Ni) decompose upon evacuation of the samples only the compounds 3 (Mn, Co, Ni) were investigated for their magnetic properties. Thus, temperature and field dependent magnetic susceptibility measurements were performed.
- 20

Magnetic properties of [Mn(HO₃PCH₂)₂NH-CH₂C₆H₄CH₂-NH(CH₂PO₃H)₂]· H₂O

The temperature dependent magnetic susceptibility was measured in the temperature range of 300-2 K applying an external field of

- ²⁵ 0.1 T. The χ vs T and $1/\chi$ vs T plot for **3** (Mn) is given in Figure 12. The compound shows typical Curie-Weiss behavior in the temperature range of 300-9 K. Below this temperature regime the compound shows antiferromagnetic coupling. A least-square fit using the Curie-Weiss law leads to C = 4.48 K cm³ mol⁻¹, and θ =
- $_{30}$ -8.15 K, both in agreement with antiferromagnetic coupling. We observe a μ_{eff} of 5.98 μ_b which is in good agreement with the theoretical value of 5.92 μ_b for a high spin Mn^{2+} ion.



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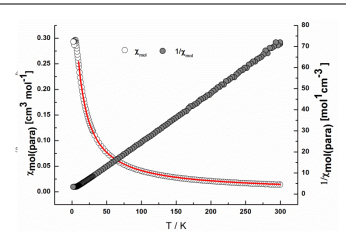


Fig. 12. Temperature dependent magnetic susceptibility of [Mn(HO₃PCH₂)₂NH-CH₂C₆H₄CH₂-NH(CH₂PO₃H)₂]·(H₂O). The molar ⁵⁵ magnetic susceptibility is shown as white dots, the red line shows the Curie-Weiss fit and the reciprocal magnetic susceptibility is shown as grey dots.

Magnetic properties of [Ni(HO₃PCH₂)₂NH-CH₂C₆H₄CH₂-NH(CH₂PO₃H)₂]·H₂O

⁶⁰ The results of the temperature dependent magnetic susceptibility measurement of compound **3** (Ni) are shown in Figure 13. The measurement was carried out in the temperature range of 300-2 K. The compound shows typical paramagnetic behavior between 300 and 50 K. Below 13.3 K ferromagnetic coupling is observed. ⁶⁵ Linear regression of the 1/χ vs T plot leads to values of C = 1.10 K cm³ mol⁻¹ and θ = 13.26 K which indicates with ferromagnetic coupling. We observe a μ_{eff} of 2.98 μ_b which is in good agreement with the expected value for a Ni²⁺ ion of 2.83 μ_b. To learn more about the ferromagnetic coupling a field dependent ⁷⁰ measurement at 4.2 K in the range of -9 T to 9 T was carried out. The obtained hysteresis loop is shown in Figure 14. From that loop a magnetic reminiscence of +M_r= 0.013 Nβ mol⁻¹, -M_r= -0.013 Nβ mol⁻¹ and H_c = 0.160 T, -H_c = -0.159 T was determined.

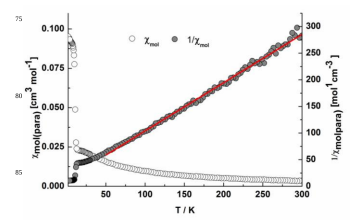


Fig. 13. Temperature depend magnetic susceptibility of $[Ni(HO_3PCH_2)_2NH-CH_2C_6H_4CH_2-NH(CH_2PO_3H)_2]\cdotH_2O$. The molar ⁹⁰ magnetic susceptibility is shown as white dots, the reciprocal magnetic susceptibility is shown as grey dots and the red line shows the linear fit of $1/\chi$ vs T.

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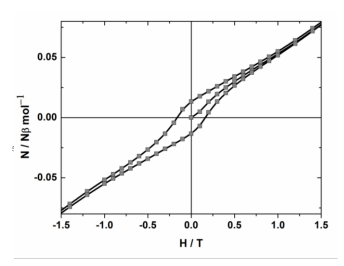


Fig. 14. Field dependent magnetic susceptibility measurement of 3 (Ni) at 10 4.2 K.

Magnetic properties of [Co(HO₃PCH₂)₂NH-CH₂C₆H₄CH₂-NH(CH₂PO₃H)₂]·H₂O

The temperature dependent magnetic susceptibility of compound 3 (Co) is shown in Figure 15. The measurements were carried out 15 in the temperature range of 300-2 K with zero field cooling (ZFC). The magnetic behavior is similar to the canted antiferromagnetic coupling observed in

Co[HO₂C(CH₂)₃NH(CH₂PO₃H)₂]₂ which is build up from the same inorganic layered motif.³⁰ The temperature dependent ²⁰ measurement shows paramagnetic behavior in the temperature range between 300-13 K and antiferromagnetic coupling below 13 K. Nevertheless the susceptibility splits upon ZFC. A fit of the χ vs T curve with the Curie-Weiss law leads to C = 3.03 K cm³ mol⁻¹ and θ = -29.50 K. We observe a μ_{eff} of 5.02 μ_b which is

²⁵ much higher than the expected spin-only value of $\mu_{eff} = 3.87 \ \mu_b$. However, this is in good agreement with the literature value of the canted antiferromagnet ($\mu_{eff} = 5.23 \ \mu_b$) which can be explained by the spin orbital interactions of the Co²⁺ ions.

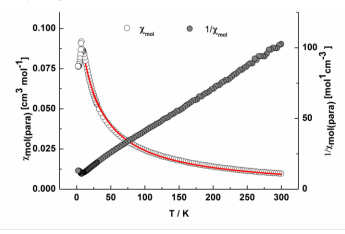


Fig. 15. Temperature dependent magnetic susceptibility of [Co(HO₃PCH₂)₂NH-CH₂C₆H₄CH₂-NH(CH₂PO₃H)₂]·H₂O. The molar magnetic susceptibility is shown as white dots, the red line shows the ⁴⁰ Curie-Weiss Fit and the reciprocal magnetic susceptibility is shown as grey dots.

To understand the magnetic behaviour a field dependent initial curve and AC measurements were carried out (Fig. 16). The initial curve shows no step in the range of 0 to 9 T. Thus, a ⁴⁵ metamagnet can be excluded. The AC measurements show no shift in the maximum susceptibility. These results exclude the presence of spin glas or a ferromagnet. Thus, all results are in agreement with the presence of a canted antiferromagnet.

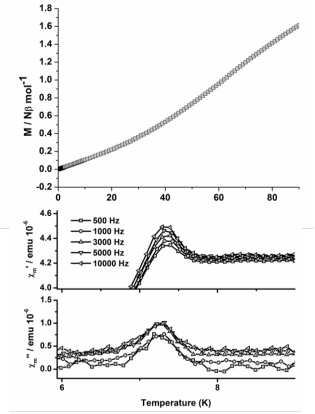


Fig. 16. Initial curve of the field dependent measurement of **3** (Co) at 4.2 K (top) and AC measurements of the susceptibility (bottom).

Proton conductivity

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⁶⁵ The compounds 2 (Zn) and 3 (Mn) were investigated for their proton conductivity properties because of their expected low toxicity and good crystallinity. While in 2 (Zn) one water molecule per formula unit is coordinated to the zinc ion in 3 (Mn) it is close to the –PO₃H⁻ groups and involved in hydrogen ⁷⁰ bonding. The proton conductivity was determined using equation (1).²⁶

$$\sigma = \frac{1 \Box L}{R \Box A} \tag{1}$$

Z: impedance, R: ohmic resistance, σ : proton conductivity,

L: sample thickness, A: sample cross-section area

C

The ohmic resistance was taken from the Bode phase plot of the impedance, which shows the impedance phase shift versus frequency. The impedance corresponding to the phase shift so closest to zero is approximately equal to the ohmic resistance of

the sample.^{26, 27} Each impedance value was measured three times, and the arithmetic mean was used in the analyses.

Both metal phosphonates exhibit proton conductivity (Fig.17),

- ⁵ however the proton conductivity at 413K and 100% RH of compound **3** (Mn) (1,53 \cdot 10⁻³ S/cm) is not only 35 times higher than that of **2** (Zn) (4,26 \cdot 10⁻⁵ S/cm), but also much more reproducible (rep.). The proton conductivity of **3** (Mn) is in the same order of magnitude as observed for the compound
- ¹⁰ [Mg((HO₃PCH₂)₂NH-C₈H₁₆-NH(CH₂PO₃H)₂]·6H₂O, which exhibits a value of 1.6×10^{-3} S cm⁻¹ at 298 K.³¹ As expected, relative humidity has a strong influence on the proton conductivity of the investigated phosphonates: for example, a 10% decrease of RH at 413 K results in a 5-fold decrease in the ¹⁵ proton conductivity of **3** (Mn).

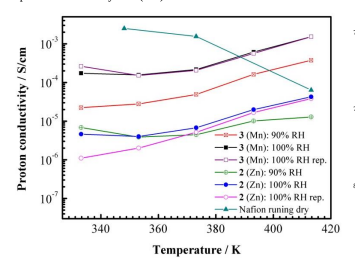


Fig. 17. Proton conductivity of the two phosphonates $([Zn(HO_3PCH_2)_2NH-CH_2C_6H_4CH_2-NH(CH_2PO_3H)_2(H_2O)] - 2$ (Zn) and $[Mn(HO_3PCH_2)_2NH-CH_2C_6H_4CH_2-NH(CH_2PO_3H)_2]\cdot H_2O - 3$ (Mn)) at 35 different temperatures and relative humidities (90% RH and 100% RH), compared to the polymer Nafion[®] at 100% RH.³²

At temperatures above 393 K, the proton conductivity of **3** (Mn) exceeds that of Nafion[®], which is the most widely used polymer as membrane in Proton Exchange Membrane Fuel Cells

⁴⁰ (PEMFC). In contrast to Nafion[®] for which the loss in conductivity at these high temperatures is well-documented due to dehydration of the membrane,³³ the metal ions and the phosphonate groups seem to interact well enough with the water molecules at the measurement conditions to keep water ⁴⁵ molecules in the structures.

Thus, the metal phosphonates are comparable to oxide nanoparticles or functionalized ordered mesoporous oxides being widely discussed as additives for PEMFCs in order to avoid the shortcomings of Nafion[®] at high temperatures and low relative

⁵⁰ humidities.³⁴ Since especially sulfonic acid functionalized variants of MCM-41, obtained by co-condensation exhibit very high proton conductivities up to 0.2 S/cm,³² new metal sulfonates with defined porosity might even be further prospective.

⁵⁵ XRD measurements reveal that during the impedance spectroscopy measurement (Fig S14-S15) **2** (Zn) is in part transformed into a new crystalline product whose structure is unknown. The structural change during causes the conductivity losses and corresponding irreproducibility observed in the **2** (Zn) ⁶⁰ impedance spectra. In contrast, the second metal phosphonate, **3**

(Mn), shows no significant changes in the XRD pattern after IS, and thus features reproducible impedance data.

Conclusion

High-throughput methods employing microwave-assisted heating ⁶⁵ were shown to be an excellent tool for the rapid discovery of metal phosphonates. Although the counter ions of the metal salts employed are not incorporated in the structure they have a distinct influence on the crystallinity of the resulting compound. Thus the systematic investigation using different counter ions of ⁷⁰ metal salts is important.

Since all compounds in this study are obtained as microcrystalline powders, the structure determination had to be carried out from X-ray powder diffraction data. Here, the combined approach of direct methods and force field calculations 75 to establish a structural model has been proven to be very valuable and subsequent Rietveld refinements could be carried

out to get the crystal structures. Here the results observed for reactions with p-H₈L are especially interesting. Depending on the ionic radii of the metal ions three different crystal structures are

⁸⁰ formed. This suggests that systematic studies on the influence of ionic radii are important for the discovery of new structures. Finally the isostructural compounds **3** (Mn, Ni and Co) show magnetic properties ranging from antiferromagnetic, to ferromagnetic and canted antiferromagnetic interactions.

85 Notes and references

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- 95 † Electronic Supplementary Information (ESI) available: Assymetric Units, detaislabout the structure determination, thermogravimetric measurements See DOI: 10.1039/b000000x/

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contains the supplementary crystallographic data for this paper. These 100 data can be obtained free of charge via the Internet at ww.ccdc.cam.ac.uk/conts/retrieving.html (or from the CCDC, 12 Union Road, Cambridge CB2 1EZ, U.K; fax, +44 1223 36033; e-mail, deposit@ccdc.ac.uk)

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