

Guest Molecule-Responsive Functional Calcium Phosphonate Frameworks for Tuned Proton Conductivity

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Supporting Information



ABSTRACT: We report the synthesis, structural characterization, and functionality (framework interconversions together with proton conductivity) of an open-framework hybrid that combines Ca^{2+} ions and the rigid polyfunctional ligand 5-(dihydroxyphosphoryl)isophthalic acid (PiPhtA). $Ca_2[(HO_3PC_6H_3COOH)_2]_2[(HO_3PC_6H_3(COO)_2H)(H_2O)_2] \cdot SH_2O$ (Ca-PiPhtA-I) is obtained by slow crystallization at ambient conditions from acidic (pH \approx 3) aqueous solutions. It possesses a high water content (both Ca coordinated and in the lattice), and importantly, it exhibits water-filled 1D channels. At 75 °C, Ca-PiPhtA-I is partially dehydrated and exhibits a crystalline diffraction pattern that can be indexed in a monoclinic cell with parameters close to the pristine phase. Rietveld refinement was carried out for the sample heated at 75 °C, Ca-PiPhtA-II, using synchrotron powder X-ray diffraction data, which revealed the molecular formula $Ca_2[(HO_3PC_6H_3COOH)_2]_2[(HO_3PC_6H_3(COO)_3H)(H_2O)_2]$. All connectivity modes of the "parent" Ca-PiPhtA-I framework are retained in Ca-PiPhtA-II. Upon Ca-PiPhtA-I exposure to ammonia vapors (28% aqueous NH₃) a new derivative is obtained (Ca-PiPhtA-NH₃) containing 7 NH₃ and 16 H₂O molecules according to elemental and thermal analyses. Ca-PiPhtA-NH₃ exhibits a complex X-ray diffraction pattern with peaks at 15.3 and 13.0 Å that suggest partial breaking and transformation of the parent pillared structure. Although detailed structural identification of Ca-PiPhtA-NH₃ was not possible, due in part to nonequilibrium adsorption conditions and the lack of crystallinity, FT-IR spectra and DTA-TG analysis indicate profound structural changes compared to the pristine Ca-PiPhtA-I. At 98% RH and T = 24 °C, proton conductivity, σ , for Ca-PiPhtA-I is 5.7 × 10^{-4} S·cm⁻¹. It increases to 1.3×10^{-3} S·cm⁻¹ upon activation by preheating the sample at 40 °C for 2 h followed by water equilibration at room temperature under controlled conditions. Ca-PiPhtA-NH₃ exhibits the highest proton conductivity, 6.6×10^{-3} S cm⁻¹, measured at 98% RH and T = 24 °C. Activation energies (E_a) for proton transfer in the above-mentioned frameworks range between 0.23 and 0.4 eV, typical of a Grothuss mechanism of proton conduction. These results underline the importance of internal H-bonding networks that, in turn, determine conductivity properties of hybrid materials. It is highlighted that new proton transfer pathways may be created by means of cavity "derivatization" with selected guest molecules resulting in improved proton conductivity.

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■ INTRODUCTION

Metal—organic framework (MOF) materials are now at the epicenter of research efforts of many groups around the globe. An obvious exegesis for the "explosion" in the field is two-fold: (a) new structural motifs are constantly being sought,^{1a} and (b) functionality in these materials becomes increasingly important.^{1b–d} More specifically, potential applications are investigated in fields such as gas adsorption, storage and separation,² magnetism,³ catalysis,⁴ photoluminescence, sensors,⁵ controlled release of pharmaceuticals, medicinal and dental materials,⁶ chiral separations,⁷ and ion exchange,⁸ just to mention a few.

A plethora of organic linkers/ligands have been widely utilized to construct these hybrid MOF materials, with the majority being carboxylate, imidazolate, or polypyridyl based. On the other hand, multichelating phosphonate ligands are "alternative" organic linkers (to the widespread polycarboxylates, imidazoles, and various derivatives) and provide synthetic access to a number of thermally and chemically stable MOFs.⁹

of thermally and chemically stable MOFs.⁹ Recently, phosphonates,¹⁰ phosphocarboxylates,¹¹ and phosphonosulfonates¹² that possess an aromatic entity (usually one or more phenyl rings) have been exploited in synthetic efforts, yielding multidimensional (mostly 2D and 3D) platforms.

Proton conductivity has long been investigated as a desirable property in a variety of diverse materials, whether organic,¹³ inorganic,¹⁴ or hybrid.¹⁵ The field of MOFs and more specifically metal phosphonates has provided an opportune arena for elegant efforts for accessing suitable and highly conducting materials.^{15f,16} Furthermore, postsynthetic modification^{1d} of the internal surfaces of MOFs offers opportunities for enhancing desired properties of crystalline solids.^{16c} The main requirement for a MOF material to be proton conductive is the presence of acidic protons within an appropriately structured metal-organic framework that facilitates charge transport. In the case of metal phosphonate materials the phosphonic acid group (or partially protonated phosphonate) assumes the role of the acidic group. Such well-ordered structural motifs, with highly ordered phosphonate groups located in close proximity to lattice water molecules, often present appropriate platforms that are candidates for high proton conductivity.¹² For instance, proton conductivity higher than 10^{-2} S cm⁻¹ has been reported for PCMOF21/2,^{15f} a mixed sulfonate—phosphonate MOF, and for derivatized MIL-101 with H₂SO₄ and/or H₃PO₄.^{15g} Other notable examples are metal derivatives containing the *N,N*-amino-bis(methylenephosphonate) moiety, such as ZrChDTMP,¹⁷ MgH₆ODTMP,¹⁸ and LaH₅DTMP,¹⁹ phases, which exhibit conductivity values from 1×10^{-4} to 8×10^{-3} S cm⁻¹. The high flexibility imparted by the ligand is considered to exert a key role in determining internal proton transfer pathways.

In the present work, we report the synthesis, structural characterization, and proton conductivity of an open-framework hybrid that combines Ca²⁺ ions and the rigid polyfunctional ligand 5-(dihydroxyphosphoryl)isophthalic acid (**PiPhtA**). This material was obtained by slow crystallization at ambient conditions from acidic (pH \approx 3) aqueous solutions. It possesses a high water content (both Ca coordinated and in the lattice), and importantly, it exhibits water-filled 1D channels. Both characteristics are, in principle, considered to be remarkable attributes to evaluate the proton conductivity of this hybrid compound. The effects of the guest species (water and NH₃) on the conductivity properties will be highlighted.

EXPERIMENTAL SECTION

Materials and Common Instrumentation. The starting salt CaCl₂·2H₂O was from EM Science Merck and used as received. Deionized (DI) water from an in-house laboratory cation-exchange column was used for all syntheses. Stock solutions of HCl and NaOH were used for pH adjustments. The pH meter used was a wTw pH315i setup, equipped with a SeTix 41 electrode. Elemental analyses (C, H, N) were measured on a Perkin-Elmer 240 analyzer. Infrared spectra were collected in the 4000–400 cm⁻¹ range at 4 cm⁻¹ resolution, and 50 scans were accumulated. The ¹H, ¹³C{¹H}, and ³¹P{¹H} NMR (see below; operating frequency, solvent) spectra in solution were recorded on a Bruker Avance DRX300 instrument. Chemical shifts (δ) are given in ppm.

Syntheses. The ligand 5-(dihydroxyphosphoryl)isophthalic acid $[5-(H_2O_3P)C_6H_3-1,3-(COOH)_2]$ (**PiPhtA, 1**) was synthesized through a number of intermediates, shown in Figure 1, as follows.



Figure 1. Protocol for the synthesis of 5-(dihydroxyphosphoryl)isophthalic acid (**PiPhtA**, **1**): (i) NaNO₃, HCl, KI; (ii) SOCl₂, CH₃OH; (iii) HP(O)(OR)₂, (i-C₃H₇)₂NC₂H₅, Pd(OOCCH₃)₂, dppf, CH₃CN; (iv) aqueous HCl, H₂O; (R = C₂H₅).

5-lodoisophthalic Acid (2). This compound was prepared from commercially available 5-aminoisophthalic acid according to a literature procedure.²⁰ Specifically, 5-aminoisophthalic acid (5.00 g, 0.0276 mol) was dissolved in a mixture of concentrated hydrochloric acid (49 cm^3) and water (49 cm^3). This solution was cooled in an ice—water bath, and a solution of sodium nitrite (2.05 g, 0.0295 mol) in water (9.6 cm^3) was added with such a rate that the temperature of the mixture did not exceed 4 °C. Next, a solution of potassium iodide (15.0 g, 0.0903 mol) in water (38 cm^3) was slowly added. Evolution of nitrogen was observed, and the mixture turned a yellow-reddish color. Afterwards the mixture was refluxed for 1 h. After cooling the precipitate was filtered off and air dried to yield 5-iodoisophthalic acid: 5.76 g (72%), mp 284–285 °C (lit. 288–289 °C²⁰).

Dimethyl 5-lodoisophthalate (3). 5-Iodoisophatlic acid (2, 14.83 g, 0.050 mol) was heated under reflux with thionyl chloride (65 cm³, 0.89 mol) for 3.5 h. Next, the excess of thionyl chloride was distilled off. Methanol (50 cm³) was added to the residue carefully due to an exothermic effect and evolution of hydrogen chloride, and the mixture was heated to reflux for a few hours. A precipitate formed during the heating process and was filtered off and recrystallized from methanol to obtain dimethyl 5-iodoisophthalate (3): 13.32 g (82%), mp 103–105 °C (lit. 104 °C²¹).

Dimethyl 5-(Diisopropyloxyphosphoryl)isophthalate (4a). A 50 cm³ round-bottom flask was charged with dimethyl 5-iodoisophthalate (3, 2.00 g, 6.25 mmol), acetonitrile (20 mL), diisopropyl phosphite (1.246 g, 7.5 mmol), and diisopropylethylamine (1.05 g, 8.125 mmol). The flask was flushed with argon for 15 min, after which palladium acetate (0.014 g, 0.0625 mmol) and 1,1'-bis(diphenylphosphine)ferrocene (dppf, 0.038 g, 0.0687 mmol) were added. The reaction mixture was stirred and heated in an oil bath at 90 °C for 24 h under constant flow of argon. After cooling to room temperature the mixture was purified by column

chromatography on silica gel using ethyl acetate as eluent to yield dimethyl 5-(diisopropyloxyphosphoryl)isophthalate (4a): 1.434 g (64%), $R_{\rm f}$ = 0.40 (silica gel on TLC-PET foil, ethyl acetate, UV light 254 nm). ¹H NMR (300 MHz, CDCl₃) δ : 8.82 (m, 1H, CH arom.), 8.63 (dd, $J^{\rm HH}$ = 1.68 Hz, $J^{\rm HP}$ = 16.35 Hz, 2H, CH arom.), 4.74 (m, 2H, CH), 3.96 (s, 3H, CH₃), 1.39 (d, $J^{\rm HH}$ = 6.18 Hz, CH₃), 1.24 (d, $J^{\rm HH}$ = 6.18 Hz). ¹³C NMR (75 MHz, CDCl₃) δ : 165.43, 136.64 (J = 10.94 Hz), 133.88, 131.88 (J = 192.01 Hz), 130.97 (J = 14.94 Hz), 71.51 (J = 5.59 Hz), 52.60, 24.02 (J = 3.81 Hz), 23.86 (J = 4.58 Hz). ³¹P{¹H} NMR (121 MHz, CDCl₃) δ : 14.16 (s).

Dimethyl 5-(Diethoxyphosphoryl)isophthalate (4b). Compound 4b was synthesized using the same procedure as 4a, starting from 1.173 g of dimethyl 5-iodoisophthalate, although diethyl phosphite was used. Yield of the ester 4b: 42%, $R_f = 0.30$ (silica gel on TLC-PET foil, ethyl acetate, UV light 254 nm). ¹H NMR (300 MHz, CDCl₃) δ : 8.83 (m, 1H, CH arom.), 8.63 (dd, $J^{\text{HH}} = 1.68 \text{ Hz}$, $J^{\text{HP}} = 13.32 \text{ Hz}$, 2H CH arom.), 4.16 (m, 4H, CH₂), 3.96 (s, 6H, CH₃), 1.34 (t, J = 7.06 Hz, 6H, CH₃). ³¹P{¹H} NMR (121 MHz, CDCl₃) δ : 16.33 (s).

5-(Dihydroxyphosphoryl)isophthalic Acid (**PiPhtA**, **1**). A 25 cm³ round-bottom flask was charged with dimethyl 5-(diisopropyloxyphosphoryl)isophthalate (**4a**, 1.072 g, 3.00 mmol), water (4 cm³), and concentrated hydrochloric acid (4 cm³). The mixture was refluxed for 20 h. After cooling down to room temperature it was evaporated to dryness in vacuo to yield the product **1** as a white solid: 0.734 g (99%). ¹H NMR (300 MHz, D₂O) δ : 8.29 (d, *J* = 3.82 Hz, 1H, CH arom.), 8.21 (dd, *J*^{HP} = 13.36 Hz, *J*^{HH} = 2.62 Hz, 2H, CH arom.);. ³¹P{¹H} NMR (121 MHz, D₂O) δ : 12.63 (s).

 $Ca_2[(HO_3P-C_6H_3-COOH)_2]_2[(HO_3P-C_6H_3-(COO)_2H)-(H_2O)_2])$. 5 H_2O (*Ca-PiPhtA-I*). A quantity of the ligand 5-(dihydroxyphosphoryl)isophthalic acid (1, 0.061 g, 0.250 mmol) was dissolved in 10 cm³ of DI water. A small quantity of a 0.1 M NaOH stock solution was added dropwise until the ligand dissolved completely and a clear, colorless solution was obtained. A quantity of CaCl₂·2H₂O (0.037 g, 0.250 mmol) was separately dissolved in 5 cm³ of DI water. The two solutions were subsequently mixed, and the final solution pH was adjusted to 3.00 using the 0.1 M NaOH stock solution. The transparent colorless solution was left undisturbed for 1 week, after which the product appeared as a white crystalline material. It was isolated by filtration, washed with DI water, and air dried. Yield: 92%. Anal. Calcd for Ca₂[(HO₃P-C₆H₃-COOH)₂]₂[(HO₃P-C₆H₃-(COO)₂H)(H₂O)₂]·5H₂O, fw 940.56, C₂₄H₃₁Ca₂O₂₈P₃: C, 30.81; H, 2.80. Found: C, 29.27; H, 3.36.

 $Ca_2[(HO_3P-C_6H_3-COOH)_2]_2[(HO_3P-C_6H_3-(COO)_2H)-(H_2O)_2]$ (*Ca-PiPhtA-II*). This compound was obtained by thermal treatment of solid **Ca-PiPhtA-I** at 75 °C for 2 h.

Absorption of NH₃ Vapors (Aqueous Solution). A 150 mg amount of Ca-PiPhtA-I was placed in contact with vapors from a commercial solution of 28 wt % of NH₃ in water for 2 h in a closed container. Subsequently, the product was air dried for 15 min and kept at 98% relative humidity at 10 °C for 18 h following the same procedure described below to measure the proton conductivity. The obtained solid, $Ca_2[(HO_3P-C_6H_3-COOH)_2]_2[(HO_3P-C_6H_3-(COO)_2H)]-(NH_3)_7(H_2O)_{16}$ (Ca-PiPhtA-NH₃), has the following elemental analysis. Anal. Calcd: C, 23.69; H, 5.38; N, 8.06. Found: C, 23.86; H, 4.75; N, 7.65. TG -analysis: weight loss up to 400 °C, found 32.67%, calcd 33.36%.

Thermal Analysis. Differential thermal analysis (DTA) and thermogravimetric analysis (TGA) data were recorded on an SDT-Q600 analyzer from TA Instruments. The temperature varied from room temperature to 900 °C at a heating rate of 10 °C·min⁻¹. Measurements were carried out on samples in open platinum crucibles under a flow of air.

X-ray Crystallography. Laboratory X-ray powder diffraction (XRPD) patterns were collected on a PANanalytical X'Pert Pro automated diffractometer. Powder patterns were recorded in Bragg–Brentano reflection configuration using a Ge(111) primary monochromator (Cu K α_1) and the X'Celerator detector. Thermodiffractometric studies for Ca-PiPhtA-I and Ca-PiPhtA-NH₃ were carried out on samples loaded in an Anton Paar HTK 1200N chamber under static air. Data were collected between room temperature and 240 °C. All measurements were performed with a heating rate of 10 °C·min⁻¹ and a

delay time of 5 min to ensure thermal stabilization. Data acquisition was carried out between 4° and 40° (2 θ) due to "less-than-ideal" crystallinity of the samples. A step size of 0.017° and an equivalent counting time of 150 s step⁻¹ were used.

Crystal Structure Determination. Suitable-size crystals of **Ca-PiPhtA-I** were kept under inert conditions and immersed in perfluoropolyether as protecting oil for manipulation. A suitable crystal was mounted on MiTeGen Micromounts, and this sample was used for data collection. Data were collected with a Bruker SMART APEX diffractometer. Data were processed with APEX2²² suite and corrected for absorption using SADABS.²³ The structure was solved by direct methods,²⁴ which revealed the position of all non-hydrogen atoms. These atoms were refined on F^2 by a full-matrix least-squares procedure using anisotropic displacement parameters.²⁴ All hydrogen atoms were located in difference Fourier maps, except those corresponding to C–H groups, which were placed geometrically and included as fixed contributions riding on attached atoms with isotropic thermal displacement parameters 1.2 (C–H) or 1.5 (O–H) times those of the respective bonded atom.

Synchrotron X-ray powder diffraction data (SXRDP) for Ca-PiPhtA-II were collected in Debye–Scherrer (transmission) mode using the X-ray powder diffraction station of ALBA, the Spanish Synchrotron Radiation Facility (Barcelona, Spain).²⁵ The wavelength, 0.62020(1) Å, was selected with a double-crystal Si (111) monochromator and determined from a Si640d NIST standard (a = 5.43123 Å) measurement. The capillary was rotated during data collection to improve diffracting particle statistics. The detector system used was the MYTHEN, which allows fast collection of good-resolution patterns. The data acquisition time was ~20 min per pattern to attain very good signal-to-noise ratio over the angular range $1-41^{\circ}$ (2θ).

The X-ray powder pattern was autoindexed using the DICVOL06 program.²⁶ The crystal structure of **Ca-PiPhtA-II** compound was determined by the Rietveld method²⁷ using the structure of **Ca-PiPhtA-I** available from single-crystal data as starting model. Rietveld refinement was carried out using the GSAS package²⁸ with soft constraints to maintain chemically reasonable geometries for the phosphonate, aromatic ring, and carboxylic groups. The soft constraints were as follows: /PO₃C tetrahedron/P–O, 1.53(1) Å; P–C, 1.80(1) Å; O···O, 2.55(2) Å; O···C, 2.73(2) Å; /aryl/aromatic ring/C–C [1.40(1) Å], C_{ring}···C_{ring} [2.40(1) Å], C_{cring}···C_{ring} [2.78(1) Å], /COO group/, C_{aryl}···O_{carb}, 2.36(2) Å. Two isotropic atomic displacement parameters were refined, one for calcium and phosphorus atom and a second one for carbon and oxygen atoms. Hydrogen atoms were not included in the refinements. Crystallographic data and CCDC reference codes are given in Table 1.

Proton Conductivity Studies. Impedance data were collected on cylindrical pellets (~10 mm diameter; ~0.9–1.1 mm thickness) obtained by pressing ~0.15 g of sample at 500 MPa for 2 min between porous C electrodes (Sigracet, GDL 10 BB, no Pt). Impedance data were collected using a HP4284A impedance analyzer over the frequency range from 20 Hz to 1 MHz with an applied voltage of 1 V. All measurements were electronically controlled by the winDETA package of programs.²⁹

Electrical measurements were taken in a double-walled, temperaturecontrolled glass chamber with a gas inlet and outlet. The temperature of the chamber was controlled by a Julabo F32-MA refrigerated/heating circulator from 10 to 24 °C every 2 °C with a heating rate of 0.4 °C ·min⁻¹ using EasyTEMP software. Samples were equilibrated for 15 min after each step in temperature, measured in closest vicinity to the sample. The relative humidity (RH) was obtained by a continuous flow of water-saturated nitrogen at different temperatures through the cell. Pellets were equilibrated at 98% RH for 18 h to ensure a fixed water content of the sample and stable conductivity values. RH is approximately constant in the measured temperature range from 10 to 24 °C.

RESULTS AND DISCUSSION

Synthesis. The product Ca-PiPhtA-I is formed at a relatively low reaction pH (\sim 3.0). At this pH the carboxylic acid groups are

Table 1. Crystallographic Data for Ca-PiPhtA-I and Ca-PiPhtA-II

	Ca-PiPhtA-I	Ca-PiPhtA-II
chemical formula	$Ca_2C_{24}H_{31}O_{28}P_3$	Ca2C24H21O23P3
formula mass	940.56	850.49
cryst syst	orthorhombic	monoclinic
cryst size (mm) color/ shape	$0.10 \times 0.09 \times 0.08$ colorless/ block	
$\lambda/\text{\AA}$	0.71073	0.62020(1)
a/Å	23.112(4)	23.0652(7)
b/Å	6.9534(13)	6.9731(1)
c/Å	22.638(4)	22.4165(7)
$\alpha/{ m deg}$	90.0	90
β/\deg	90.0	91.540(4)
γ/deg	90.0	90
unit cell volume/Å ³	3638.1(12)	3604.1(2)
temp/°C	-173	25
space group	$Pca2_1$	Pa
Ζ	4	4
R _{int}	0.0832	
no. of independent reflns	4603	3455
data (reflns)/restraints/ params	6716/1/522	5500/299/357
<i>R</i> factor $[I > 2\sigma(I)]$	$R1^{a} = 0.0715$; w $R2^{a} = 0.1610$	
R factor (all data)	$R1^{a} = 0.1075; wR2^{a} = 0.1834$	
R _{WP}		0.0654
$R_{\rm P}$		0.0464
GoF/R_F	1.032	0.075
CCDC number	968321	968322
^a R1(F) = $\Sigma F_o - F_c / \Sigma F_o $; wR2(F ²) = $[\Sigma w (F_o^2 - F_c^2)^2 / \Sigma F^4]^{1/2}$.		

expected to be deprotonated, whereas the phosphonic acid group is monodeprotonated. This would render the ligand trianionic, and a possible 1:1 adduct with Ca^{2+} would be anionic. However, based on elemental analyses and the structure solved, a solid with a Ca:ligand ratio of 2:3 is instead obtained, indicating that the protonation state of the ligand is complex. As suggested by the empirical formula, the ligand is found in different protonation states, which are adjusted depending on the various modes of coordinating to Ca^{2+} (Figure 2).

Structural Study. Ca-PiPhtA-I shows a complex pillared layered structure with a unit cell containing two calcium atoms, three **PiPhtA** ligands, and seven water molecules (two are Ca coordinated and the remaining five are situated within the lattice). The coordination geometry of both calcium centers is pentagonal bipyramidal (see Figure 3).

In this geometry, four of the equatorial positions are occupied by oxygen atoms of three different phosphonate groups, one of them chelating the metal ion in a bidentate fashion. The fifth equatorial coordination position is occupied by an oxygen atom of a carboxylate group. All of these equatorial oxygen atoms belong to two ligands, P2 and P3, which configure edge-sharing calcium polyhedral chains and connect them into the layers, while leaving alternately arranged free carboxylic groups inside the interlayer space (Figure 4). The axial positions of the calcium polyhedron are occupied by a water molecule and a phosphonate or carboxylate oxygen of the third organic ligand, P1, which is pillaring adjacent layers.

Within the layers, each phosphonate moiety uses only two oxygen atoms to coordinate calcium in such a way that one oxygen atom is bridging two metal centers, thus leaving a free POH pointing to the interlayer space. One of the carboxylate moieties of both dicarboxyphosphonate ligands, P2 and P3, is



Figure 2. Three deprotonation states of the ligand PiPhtA in the structure of Ca-PiPhtA-I.



Figure 3. Coordination environment of the two Ca centers in the structure of Ca-PiPhtA-I.

coordinated to calcium atoms through one of the oxygens, while the other one is protonated. Besides, the second carboxylic group of each these ligands is pointing toward the interlayer space and faced up to each other interacting strongly by a H-bond. This disposition creates H-bond rings contributing to the stabilization of the pillared layered structure. The third dicarboxyphosphonate ligand, P1, is linking adjacent layers only by one phosphonate oxygen, by one side, and one carboxylate oxygen atom, from the other side. This connectivity mode leaves a POH and a free carboxylic group inside the interlayer space (see Figure 4).

The overall structure architecture reveals interlayer hydrophilic 1-D channels running parallel to the b axis, filled with five lattice waters per unit cell forming zigzag chains inside the cavities (Figure 5).

The parallel zigzag chains acquire a left-handed helix and a right-handed helix along the c axis. The water-to-water interactions between these lattice waters create an extended network of H bonds along the 1-D channels and therefore a potential proton transfer pathway, which may play an important role in the proton conductivity exhibited by the **Ca-PiPhtA** materials (see Table S1, Supporting Information).

The title compound shares some similarities with compounds $Ca_2(H_2O)[H(OOC)_2C_6H_3PO_3H]_2$ and $[Sm_2(H_2O)_4(H_2O)_2C_6H_3PO_3)_2]\cdot 2H_2O$ obtained by hydrothermal treatment.³⁰ In the former, each ligand is 2-fold deprotonated resulting



Figure 4. (Top) 3D structure of **Ca-PiPhtA-I** along the *b* axis, showing the relative arrangement of the water-filled 1D channels and the moieties protruding from the channel wall to the interior. (Bottom) Longitudinal view (*b*-axis horizontal) of the 1D water columns within the pillared scaffold.



Figure 5. (Top) Two neighboring water channels. Upper channel (in blue) represents a left-handed helix, whereas the lower (in red) represents a right-handed helix. (Bottom) Close contacts between the O atoms within a water channel.

in three distinctive coordinating groups, one carboxylate, one carboxylic, and one hydrogen phosphonate group. Ca^{2+} ions are also coordinated by seven oxygen atoms, and Ca polyhedrons are found sharing edges. However, only discrete Ca_2O_{12} dimers are present in the structure of the reported Ca derivative instead of lineal chains. In addition, the connectivity of the three groups is also modified, the title compound showing a higher connectivity of the hydrogen phosphonate groups relative to the carboxylate/ carboxylic groups. Regarding the samarium derivative, the main structural feature common with the title compound is the presence of edge-sharing Sm polyhedron chains. However,

differently, Sm³⁺ is nine coordinated and, additionally, side arms of edge-sharing polyhedra of Sm³⁺ are attached to the main chain. Thus, the chains are interconnected by phosphonate and carboxylate groups coordinated to Sm³⁺ ions, giving hybrid layers interconnected by hydrogen bonding between two carboxylic acid groups.

FT-IR Spectroscopy. Comparison of the FT-IR spectrum of the "free" ligand PiPhtA with that of Ca-PiPhtA-I is revealing. The ligand shows antisymmetric ν (C=O) stretches at 1705, 1665, and 1599 cm⁻¹ and symmetric ν (C–O) stretches at 1454, 1437, and 1395 cm⁻¹. These are shifted in the spectrum of Ca-**PiPhtA-I**, with the antisymmetric ν (C=O) stretches appearing at 1692, 1605, and 1572 $\rm cm^{-1}$ and the symmetric $\bar{\nu}(\rm C{-}\rm O\bar{)}$ stretches located at 1434, 1406, and 1381 cm⁻¹. Such shifts upon ligand coordination in bands associated with the carboxylate group are well established in the literature.³¹ There are also notable shifts in the vibrations related to the phosphonate group. There is a set of three strong intensity bands in the ligand spectrum at 1109, 1135, and 1242 cm⁻¹, which are appropriately shifted to 1132, 1255, and 1278 cm⁻¹ in the compound spectrum. These are assigned to the $-PO_3$ antisymmetric stretch. Lastly, a band at 932 cm⁻¹ in the ligand spectrum assigned to the P–O stretch shows up at 913 cm^{-1} in the compound spectrum (Figure S1, Supporting Information). All phosphonate-related bands appear in the same region as that reported for several other metal phosphonate materials.

Thermal Behavior. The thermal stability of Ca-PiPhtA-I compound was studied by TG-DTA analysis and X-ray thermodiffraction. The TG and heat flow curves, Figure 6,



Figure 6. TGA (green) and DTA (blue) curves for Ca-PiPhtA-I (dashed line) and after adsorption of NH_3 , Ca-PiPhtA-NH₃ (solid lines).

show that dehydration of the solid takes place in three overlapping steps between 50 and 220 °C. The two first steps are centered at 75 and 130 °C and correspond to removal of five lattice waters (observed weight loss 9.63%, calculated 9.62%). The third step, centered at 180 °C, is due to elimination of the Ca-bound water. The dehydration process ends at 220 °C with an associated weight loss of 13.24%, which is consistent with the loss of seven water molecules (calculated 13.5%) and the results from the crystallographic study. Above this temperature, a small but progressive mass loss is observed that suggests partial decomposition of the solid before combustion of the organic moieties at 380 °C.

Dehydration of **Ca-PiPhtA-I** was also followed by thermodiffraction (Figure 7). Although no water is lost between room temperature and 50 $^{\circ}$ C, a slight change in the symmetry of the



Figure 7. Thermodiffractometric study for Ca-PiPhtA-I at selected temperatures.

compound is already apparent in this temperature range, as indicated by the splitting of some of the diffraction peaks. At 75 °C, the sample is partially dehydrated and exhibits a diffraction pattern that can be indexed in a monoclinic cell with parameters very close to the "as-synthesized sample". This framework remains stable upon heating up to 135 °C, as confirmed by the corresponding LeBail fits. The appearance of additional diffraction peaks in the range 135-220 °C, which cannot be indexed in the monoclinic cell, is indicative of further structural changes taking place concurrently with removal of the coordinated water and a progressive loss of crystallinity. However, the crystallinity is restored when the sample preheated at 220 °C is exposed to air or put in contact with a controlled humidity atmosphere (saturated K₂SO₄ solution) at room temperature for 4 days (Figure 7). The resulting diffraction pattern agrees well with a mixture of the orthorhombic and monoclinic phases. Thermogravimetric analysis also reveals that this sample is fully rehydrated (13.39% of weight loss at 250 °C).

Rietveld refinement was carried out for the sample heated at 75 °C, **Ca-PiPhtA-II**, in order to gain better insight on the structural identity of the monoclinic phase. For refinement, the crystal structure of **Ca-PiPhtA-I** was used as a starting model after reducing symmetry to the monoclinic space subgroup *Pa*. A total of 104 atoms in the asymmetric unit were refined using synchrotron powder X-ray diffraction data. The final Rietveld plot is given as Supporting Information (Figure S2). The water content for this sample was determined to be 9.70% by thermogravimetric analysis, at a time scale similar to that of the thermodiffractometric study. The weight loss agrees well with complete removal of the lattice water.

Removal of lattice waters provokes small structural changes, according to the refined unit cell parameters (see Table 1), which are more significant along the pillaring direction (*c* axis). The resulting monoclinic phase, **Ca-PiPhtA-II**, has 4 Ca atoms, 6 ligands, and 4 water molecules in the asymmetric unit. All connectivity modes of the "parent" **Ca-PiPhtA-I** framework are retained in **Ca-PiPhtA-II**. However, the higher complexity of the unit cell results in higher distortion of the calcium polyhedra and hence formation of an alternate interlayer region with slightly different cavity dimensions (see Figure 8), the alteration of the axial angles O–Ca–O along the *c* axis being the most significant structural change in the monoclinic cell. Hence, new O_{5b}–C_{a1b}– O_{w6b} = 153(3)° and O_{1b}–Ca₂–O_{w76} = 153(3)° angles are observed, together with O_{ax}–Ca–O_{ax} angles close to 180° characteristic of the orthorhombic cell. A consequence of these



Figure 8. *b*-axis view (*a*-axis horizontal) of the crystal structures for **Ca-PiPhtA-I** (left) and **Ca-PiPhtA-II** (right) showing the slightly different conformation of the 1D channels along the *c* axis.

changes in the interlayer region of the monoclinic cell is the appearance of asymmetric H-bond rings between facing-up carboxylic groups, with interaction distances $-COO-H\cdotsO-C-OH-$ of 2.1 and 2.8 Å. The existence of different 1-D channels may affect the H-bond strength in the monoclinic cell upon rehydration.

Absorption of NH₃ Vapors. Ca-PiPhtA-I was exposed to vapors from a concentrated aqueous ammonia solution (28%). A new derivative was obtained (Ca-PiPhtA-NH₃) containing 7 NH₃ and 16 H₂O molecules according to elemental and thermal analyses. This material exhibits a complex X-ray diffraction pattern with peaks at 15.3 and 13.0 Å that suggest partial breaking of the pillared structure (Figure S3, Supporting Information). This hypothesis is strongly supported by the inability of the original narrow 1-D channels to accommodate the high number (23) of intercalated guest species. Although a detailed structural analysis of Ca-PiPhtA-NH₃ has not yet been possible, due in part to the lack of crystallinity, inspection of the IR spectra and DTA-TG analysis indicate profound structural changes compared to the pristine Ca-PiPhtA-I. Hence, in addition to the characteristic IR bands corresponding to metal carboxyphosphonates observed for Ca-PiPhtA-I and in agreement with the crystallographic data,³² the IR spectrum of Ca-PiPhtA-NH₃ (Figure S1, Supporting Information) shows a shift to lower wave numbers in the water stretching vibrations region, which is indicative of the existence of strong hydrogen bonds between intercalated water and ammonia guest molecules. In addition, the characteristic bands corresponding to the carboxylic³¹ (\sim 1700 cm⁻¹) and hydrogen phosphonate³³ (915–930 cm⁻¹) groups are practically absent in the vibrational spectrum of Ca-PiPhtA-NH₃ as a consequence of the reaction of these acidic groups with NH₃. In addition, the DTA-TG curves of the ammonia derivative Ca-PiPhtA-NH₃ (Figure 6) show a higher guest molecule content between room temperature and 400 °C. Importantly, a different weight loss profile is noted, revealing a new endothermic peak at 210 °C, indicative of strongly retained guest species. Thermodiffraction of Ca-PiPhtA-NH₃ shows a progressive amorphization of the sample with removal of the guest species, although the peak at 15.3 Å (corresponding to the new intercalated phase) persists up to 125 °C. This is a good indication of the high stability of this phase.

In the absence of precise structural data, we speculate that **Ca-PiPhtA-NH**₃ is a multiphase compound with peaks at 15.3 and 13 Å representing the $d_{(oo2)}$ basal spacing of two new layered phases originated by rupture of the interlayer phosphonate/Ca²⁺ or carboxylate/Ca²⁺ bonds (see Figure 9). The similar bond lengths for the distances Ca-O_{carboxylate} [2.378(9) Å] and



Figure 9. Schematic representation for the rupture of the 3D framework to give new 2D intercalated derivatives.

Ca– $O_{phosphonate}$ [2.394(9) Å] may support both breaking mechanisms. Other phases at lower *d* spacing, if existing, are more difficult to distinguish due to overlapping with other reflections of the crystal. The reactivity of **Ca-PiPhtA-I** toward a concentrated solution of ammonia is in clear contrast to the almost inexistent adsorption by running a typical NH₃-TPD experiment on a sample previously dehydrated at 200 °C. This suggests that ammonia and water adsorption occurs through a cooperative mechanism of hydrogen-bond interaction.

Proton Conductivity Study. The **Ca-PiPhtA-I** framework possesses 1D channels hosting zigzag chains of water molecules that simultaneously interact with the pendant–POH and –COOH groups. These structural features prompted us to undertake a study of its proton conductivity. Effective pathways for proton transfer could be created through the H-bond interactions among the acidic groups and both lattice and Ca-bound water molecules. Thermogravimetric analysis revealed that the starting samples exposed to 98% RH were fully hydrated, as the measured weight losses were very close to the observed value of 13% at 250 °C for the as-synthesized material **Ca-PiPhtA-I**. All conductivity measurements were carried out after equilibration for 18 h to ensure that the samples exhibit stable conductivity values and were fully reproducible after several cycles of measurements (Figure S4, Supporting Information).

The impedance plots, Figure 10, show similar features with spikes inclined to the Z' axis by $\sim 70^{\circ}$, indicating a partialblocking electrode response that allows limited diffusion and, therefore, confirms the conducting species must be ionic, i.e., H^+ ions. The total pellet resistances, R_T , were obtained from the intercept of the spikes and/or the arcs (low frequency end) on the Z' axis. At 98% RH and T = 24 °C, the conductivity values, $\sigma_{\rm r}$ are 5.7 × 10⁻⁴ and 3.6 × 10⁻⁴ S·cm⁻¹ for Ca-PiPhtA-I and **Ca-PiPhtA-II**, respectively. These conductivities rise up to $1.3 \times$ 10^{-3} S·cm⁻¹ upon preheating the sample at 40 °C for 2 h and subsequently water-equilibrated (Ca-PiPhtA-III). The X-ray diffraction pattern for Ca-PiPhtA-III is similar to that of Ca-PiPhtA-I, and its water content remains unchanged as confirmed by thermal analysis. We attribute the increase in conductivity to a thermal activation mechanism implying small but effective structural and/or textural changes (surface effect) leading to an enhancement of the proton conduction properties.

The conductivity value obtained for the **Ca-PiPhtA-I** sample after thermal activation is of the same magnitude as for other functionalized metal phosphonates (from 1.6×10^{-3} to 8×10^{-3} S·cm⁻¹)^{18,19} and other MOF materials, some of them reaching 10^{-2} S·cm⁻¹.^{15g,f} However, the proton conductivity of **Ca-PiPhtA**



Figure 10. Plot of the complex impedance plane for Ca-PiPhtA compounds at 24 $^{\circ}\text{C}$ and 98% RH.

materials is higher than those reported for other Ca-based MOFs, the latter exhibiting values in the range from 10^{-5} to 4×10^{-4} S·cm⁻¹ under similar experimental conditions.^{34,35}

As observed in Figure 11, the proton conductivity of Ca-PiPhtA-III remains practically constant up to \sim 40 °C,



Figure 11. Plot of log σ vs 1000/*T* for Ca-PiPhtA-III from 24 to 75 °C.

notwithstanding the relative humidity is progressively decreasing. Higher temperatures cause a profound drop in conductivity, most likely due to dehydration, according to the TG curve. This behavior is characteristic of water-mediated proton conductors when a drastic reduction in proton carrier number occurs.^{15d}

As the nature and size of the internal cavities may strongly influence the proton conductivity behavior, it was decided to measure the conductivity of the NH₃ derivative, which can host a considerably high number of proton carriers and thus possesses impressive swelling properties. As shown in Figure 10, **Ca-PiPhtA-NH**₃ exhibits the highest proton conductivity in the selected temperature range. The maximum conductivity value measured at 98% RH and T = 24 °C was 6.6×10^{-3} S·cm⁻¹. These results underline the importance of internal H-bonding networks that, in turn, determine conductivity "derivatization" with selected guest molecules may enhance these properties by creating new proton transfer pathways.

The activation energies (E_a) for the proton transfer in **Ca-PiPhtA-I**, **Ca-PiPhtA-III**, and **Ca-PiPhtA-NH**₃ range between 0.23 and 0.4 eV (Figure 12). Preheating the sample at



Figure 12. Arrhenius plots vs T^{-1} from 10 to 24 °C.

40 °C seems to induce small structural distortions favorable for proton transfer through a pathway of sites separated by a smaller energy barrier.^{15f,16} The activation energy of Ca-PiPhtA-NH₃ although increased with respect to the two other Ca-PiPhtA compounds is still within the range of values consistent with the Grotthuss H⁺ transfer mechanism.³⁶

CONCLUSION

We reported the synthesis, structural characterization, and some important properties (framework interconversions and proton conductivity) of an open-framework hybrid material containing 1D channels filled with hydrogen-bonded water molecules, namely, $Ca_2[(HO_3PC_6H_3COOH)_2]_2[(HO_3PC_6H_3(COO)_2H) (H_2O)_2$]·5H₂O (Ca-PiPhtA-I). Its partial dehydration at 75 °C produces Ca-PiPhtA-II, which exhibits lower water content but otherwise retains the framework structural features of its parent Ca-PiPhtA-I. Structure solution using synchrotron powder X-ray diffraction data revealed its molecular formula $Ca_{2}[(HO_{3}PC_{6}H_{3}COOH)_{2}]_{2}[(HO_{3}PC_{6}H_{3}(COO)_{2}H)(H_{2}O)_{2}].$ Upon Ca-PiPhtA-I exposure to ammonia vapors (28% aqueous NH₃) a new derivative is obtained (Ca-PiPhtA-NH₃) containing 7 NH₃ and 16 H₂O molecules. The above-mentioned materials were studied for proton conductivity. At 98% RH and T = 24 °C, proton conductivity for Ca-PiPhtA-I is 5.7×10^{-4} S·cm⁻¹. The proton conductivity properties of this phase can be enhanced up to 1.3×10^{-3} S·cm⁻¹ by activation upon preheating the sample at 40 °C for 2 h followed by water equilibration. Ca-PiPhtA-NH₃ exhibits the highest proton conductivity, 6.6×10^{-3} S·cm⁻¹ measured at 98% RH and $T = 24 \,^{\circ}$ C. Activation energies (E_a) for proton transfer in the above-mentioned frameworks range between 0.23 and 0.4 eV, typical of a Grothuss mechanism of proton conduction. These results emphasize the significance of internal H-bonding networks and their profound effect on conductivity properties, at least in the case of Ca-PiPhtA-I. Finally, cavity "derivatization" with selected guest molecules may

be an important mechanism to create new proton transfer pathways that improve the proton conductivity.

ASSOCIATED CONTENT

Supporting Information

ATR-IR spectra for Ca-PiPhtA-I and Ca-PiPhtA-NH₃; Rietveld refinement powder XRD profile of Ca-PiPhtA-II; powder X-ray diffraction patterns for (a) Ca-PiPhtA-I and (b) Ca-PiPhtA-NH₃; plot of log σ vs time for Ca-PiPhtA-I and Ca-PiPhtA-IIII for two cycles; table of bond lengths and H-bond distances for Ca-PiPhtA-I; CIF files for Ca-PiPhtA-I and Ca-PiPhtA-II. This material is available free of charge via the Internet at http://pubs. acs.org.

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Notes

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