Water-Stable, Nonsiliceous Hybrid Materials with Tunable Porosity and Functionality: Bridged Titania-Bisphosphonates
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ABSTRACT: Combining the properties of organic and inorganic moieties with high surface areas and pore volumes offers endless possibilities to design materials adapted to a wide range of advanced applications. The vast majority of mesoporous hybrid materials are siliceous materials, and developing low-cost synthetic methodologies leading to water stable non-siliceous hybrid materials with controlled texture and functionality is essential. We report here an original strategy for the synthesis of mesoporous bridged titania-bisphosphonate hybrids based on a one-step, templateless non-hydrolytic sol–gel route. The reaction of Ti(OiPr)4 and a rigid bisphosphonate ester in the presence of Ac2O leads to the formation of TiO2 anatase nanorods cross-linked by fully condensed bisphosphonate groups. The porosity can be readily adjusted over a wide range by changing the reaction conditions, and very high specific surface areas (up to 720 m² g⁻¹) and pore volumes (up to 1.85 cm³ g⁻¹) can be reached. The texture is stable in aqueous media between pH 1 and pH 12. Furthermore, accessible functional organic groups can be easily incorporated using either functional bisphosphonates or easily available monophosphonate compounds. The accessibility of bipyridyl organic groups was checked by Cu²⁺ adsorption from aqueous solutions. The unique combination of texture, functionality, and stability displayed by bridged titania-bisphosphonates makes these promising materials complementary of other hybrid materials such as organosilicas, MOFs, or mesoporous metal phosphonates.

1. INTRODUCTION

Over the last 30 years, mesoporous hybrid materials have received much attention because combining the properties of organic and inorganic moieties with high surface areas and pore volumes offers unprecedented possibilities to design materials adapted to a wide range of advanced applications.¹,²

The vast majority of mesoporous hybrid materials with strong ionic-covalent bonds between organic and inorganic species (Class II hybrid materials)³ are siliceous materials based on organotrialkoxysilane coupling molecules. These compounds have been employed to prepare a large variety of hybrid materials with disordered or ordered mesoporosity, including surface modified silicas, (bridged) silsesquioxane gels and cogels, and periodic mesoporous organosilicas.⁴–⁸

Research efforts are increasingly shifting toward the development of non-siliceous mesoporous hybrid materials. These materials are mostly based on carboxylate or phosphonate coupling molecules.

Carboxylate-based metal–organic frameworks (MOFs) have known an exponential growth during the past decade.⁹,¹⁰ They are renowned for their high crystallinity, extremely large surface areas, and well-defined micropores. However, designing mesoporous MOFs requires elaborated strategies,¹⁰,¹¹ and the limited water stability of MOFs may be an issue.¹²

Phosphonates, which bind strongly to a wide range of metal atoms, are considered the best coupling molecules candidates for the design of water-stable, non-siliceous class II hybrids.¹³–¹⁵ However, metal phosphonates classically yield dense layered structures with low intrinsic porosity.¹⁶

Several approaches have been used to obtain intrinsically porous metal phosphonates of titanium, zirconium, or aluminum, notably mixing bridging bisphosphonates and short monophosphonates¹⁵,¹⁷ to generate microporosity or using a bulky multiphosphonic acid to hinder the formation of layered structures.¹⁸

In addition, metal phosphonates with external mesoporosity have been prepared by sol–gel and/or hydrothermal processing using bis- or multiphosphonic acids.¹⁴,¹⁹ In these materials, the walls are built by amorphous or semicrystalline metal phosphate particles with high phosphorus to metal ratios (P/M), ranging from approximately 1.3 to 2.²⁰–²⁶ In
most cases, the use of a template is required to reach specific surface areas and total pore volumes above \( \sim 300 \text{ m}^2 \text{ g}^{-1} \) and \( \sim 0.30 \text{ cm}^3 \text{ g}^{-1} \), respectively.\(^{14}\) For instance, an amorphous titanium phosphonate with hierarchical porosity and superior textural properties (surface area and pore volume of 448 m\(^2\) g\(^{-1}\) and 1.06 cm\(^3\) g\(^{-1}\), respectively) has been obtained through a hydrothermal process by reacting TiCl\(_4\) and 1-hydroxyethylene-1,1-diphosphonic acid in the presence of both a cationic surfactant and an anionic polymeric template.\(^{27}\)

However, the range of functional organic groups incorporated in metal phosphonate hybrid materials remains limited, and the cost of templates is a major drawback that hinders the industrial application of ordered mesoporous materials. Actually, the main advantages of periodic mesoporous materials lie in their high surface area and their large pore size, which minimizes diffusion limitations and improves accessibility for large molecules. For most applications, notably heterogeneous catalysis, the ordering of the porosity and even the narrow pore size distribution are not expected to bring any advantage,\(^{28-31}\) and an aperiodic, three-dimensional mesoporous network (as in aerogels) can provide better accessibility and thus higher rates in applications involving mass-transfer.\(^{32}\)

Accordingly, developing a template-free and general methodology for the synthesis of nonsiliceous hybrids with tunable mesoporosity and adjustable functionality remains highly desirable.

Metal oxide-phosphonates, built of metal oxide nanodomains modified by phosphonate groups, form another family of nonsiliceous hybrid materials. In this case, the P/M ratio is much lower than in metal phosphonates (0.02 < P/M < 1). There have been very few attempts to prepare metal oxide-phosphonates by sol–gel methods and up to now this approach has met with little success.\(^{33-35}\) Actually, the sol–gel synthesis of porous phosphonate-based hybrids is challenging, because of the very high reaction rates between precursors (phosphonic acids and metal alkoxides or chlorides).\(^{14,36}\)

Nonhydrolytic sol–gel, which offers much more manageable rates, can provide an elegant solution to this problem.\(^{37}\) Recently, we used a nonhydrolytic route involving the reaction of titanium isopropoxide, acetophenone and diethyl octylphosphonate to prepare mesoporous titania-phosphonate hybrids, but the octylphosphonate groups prevented the growth of titania domains; as a result, the pore volume was moderate and decreased rapidly when increasing P/Ti ratio.\(^{38}\)

Here we show that the key to high pore volumes is to combine nonhydrolytic sol–gel with rigid bis-diethylphosphonate precursors. In this case, the growth of TiO\(_2\) is not prevented and the final material is built of anatase nanorods cross-linked by fully condensed bridging bisphosphonates. This very original structure leads, in the absence of a template, to outstanding textural properties (specific surface areas up to 830
m² g⁻¹, pore volumes up to 1.85 cm³ g⁻¹) and very high hydrolytic stability. Furthermore, the pore size can be easily tuned by changing the reaction conditions, and high amounts of accessible functional groups can be introduced using either functional bisphosphonate or monophosphonate groups. This low-cost method is general and can be extended to different bisphosphonate groups and metal precursors, giving access to a wide family of new mesoporous nonsiliceous hybrid materials.

2. RESULTS AND DISCUSSION

2.1. Synthesis and Characterization of bBzP-Ti. Recently, we reported that the reaction at 200 °C between Ti(OiPr)₄ and acetic anhydride afforded mesoporous anatase TiO₂ with hierarchical porosity and high specific surface area. Here this nonhydrolytic route is extended to the synthesis of titania-bisphosphonates, using phosphonic acid diethyl esters as phosphonate precursors. Thus, a bridged titania bisphosphonate monolithic gel (denoted as bBzP-Ti) was prepared by reaction at 200 °C between Ti(OiPr)₄ and 0.1 equiv of 4,4′-bis(diethylphosphonomethyl)biphenyl in the presence of Ac₂O (Scheme 1). The P/Ti ratio is equal to 0.2, which is 10 times lower than in a layered titanium phosphonate. For comparison, a sample with nonbridging groups (mBzP-Ti) was obtained under the same conditions but using 0.2 equiv of diethyl benzylphosphonate to keep the P/Ti ratio constant. In both cases, the P/Ti ratios measured by MEB-EDX are, within experimental error, identical to the expected ones, showing that all of the precursors are incorporated in the gel network and that the materials are homogeneous at the micrometer scale (see Table S1).

Nitrogen physisorption (Figure 1a) shows that bBzP-Ti is mainly mesoporous with a specific surface area of 170 m² g⁻¹ and a pore volume of 0.36 cm³ g⁻¹. Accordingly, the presence of phosphonate groups bridges favors the formation of a high specific surface area, but bisphosphonate bridges lead to a highly mesoporous material, whereas terminal phosphonates lead to a mostly microporous one.

In both cases, solid-state ¹³C MAS NMR (Figure 1b) indicates that the bridging organic group was not damaged during the synthesis. Signals at 22 and 180 ppm point to the presence of residual acetate groups, while the absence of resonances around 60 ppm (OCH₂CH₃ sites) and 75 ppm (OCH(CH₃)₂ sites) indicates complete condensation of P-OEt and Ti(OiPr) groups. The ³¹P MAS NMR spectra (Figure 1c) display in both cases a single broad resonance centered at 20 ppm, ascribed to tridentate CP(OTi)₃ sites. FTIR spectroscopy (Figure S1a) confirms in both cases the presence of residual acetate groups (bands at 1450 and 1540 cm⁻¹) and the predominantly tridentate bonding mode of phosphonate groups (absence of P-OC stretching vibration at 950 cm⁻¹ and very low intensity of the P=O stretching band at 1250 cm⁻¹). The wide-angle powder XRD pattern of bBzP-Ti (Figure 1d) shows the presence of anisotropic TiO₂ anatase nanoparticles elongated along the c-axis, as shown by the higher intensity and narrower (004) reflection compared to the (200) reflection. Such anatase nanorods have recently attracted attention as cathode materials for aluminum-ion batteries.

The nanorods in bBzP-Ti are quite similar to those found for pure TiO₂ samples prepared by the same nonhydrolytic route in toluene were purely mesoporous, with a specific surface area of 170 m² g⁻¹ and a pore volume of 0.36 cm³ g⁻¹. Accordingly, the presence of phosphonate groups bridges favors the formation of a high specific surface area, but bisphosphonate bridges lead to a highly mesoporous material, whereas terminal phosphonates lead to a mostly microporous one.

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route in toluene. The average crystallite size obtained by the Scherrer equation from the (101) and (200) peaks were 15 and 12 nm, respectively. On the other hand, mBzP-Ti appears amorphous to XRD, indicating that monophosphonate groups drastically hinder the growth of TiO2 domains, probably by capping the surface of these domains. In both cases, low angle XRD showed no evidence for a layered titanium (bis)-phosphonate phase, which is the other stable phase in this system. As such a phase is usually non porous, its formation would be detrimental to the texture and to the organic groups accessibility. SEM and HR-TEM images (Figure 2 and Figure S2) confirm the presence in bBzP-Ti of anatase nanorods about 15 nm wide (in good agreement with the size determined by XRD) and up to 60 nm long. These nanorods are aggregated to form a three-dimensional porous network. On the other hand, mBzP-Ti appears built of compact aggregates of very small (~3 nm) amorphous nanoparticles, with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm). TEM-EDX images of bBzP-Ti (Figure S3) show that P atoms are homogeneously dispersed with small pores (<4 nm).

This nonhydrolytic route is particularly well-suited to the preparation of hybrid materials: the diethylphosphonate groups are compatible with most functional organic groups, and the byproducts formed are stable, volatile compounds, which facilitates their removal and avoids secondary reactions.

2.3. Shaping. Shaping porous materials is important for applications such as separation and catalysis. It is remarkable that in the case of bBzP-Ti a gentle drying at room temperature (without any solvent exchange) afforded a monolithic, crack-free xerogel (Figure 3). Again, this has to be

![Figure 3](https://dx.doi.org/10.1021/acs.chemmater.9b05095)

Figure 3. Monolithic bBzP-Ti gel (left) and bBzP-Ti xerogel (right) obtained by drying the gel at room temperature for 15 h under ambient pressure then for 2 h under vacuum.

The very different textures of bBzP-Ti and mBzP-Ti can be rationalized based on XRD and TEM results. Bisphosphonate groups do not hinder the formation of TiO2 nanorods, moreover the strong, covalent network formed by the nanorods linked by the fully condensed bisphosphonate groups prevents the collapse of pores during the drying step and accounts for the high mesopore volume.

On the contrary, terminal monophosphonate groups severely hinder the growth of the TiO2 domains and cannot link the resulting very small nanoparticles, which form compact, microporous aggregates.

2.2. Reactions Involved. In order to identify the reactions involved in this nonhydrolytic sol–gel route, we analyzed the reaction byproducts in the synthesis of bBzP-Ti by 1H and 13C liquid-state NMR (Figure S4). The only byproducts formed were isopropyl acetate and ethyl acetate (>98%) in a 9:1 ratio. This is consistent with a clean, 2-step nonhydrolytic mechanism, involving first acetoxylation (eq 1) then condensation with ester elimination leading to the simultaneous formation of Ti–O–Ti bonds (eq 2) and Ti–O–P bonds (eq 3).

\[
\begin{align*}
\text{Ti-O}^\circ\text{Pr(CH}_2\text{CO})_2\text{O} & \rightarrow \text{Ti-OCOCH}_3 + \text{CH}_3\text{COO}^\circ\text{Pr} \\
\text{Ti-O}^\circ\text{Pr} + \text{Ti-OCOCH}_3 & \rightarrow \text{Ti-O-Ti} + \text{CH}_3\text{COO}^\circ\text{Pr} \\
\text{RP-OEt} + \text{Ti-OCOCH}_3 & \rightarrow \text{RP-O-Ti} + \text{CH}_3\text{COOEt}
\end{align*}
\]

(eq 1) and (eq 2)

Therefore, the ideal reaction scheme for the synthesis of bBzP-Ti is as depicted in eq 4.

\[
\begin{align*}
\text{Ti-(O}^\circ\text{Pr)}_4 + 0.1(\text{EtO})_2\text{PO-bBz-PO(OEt)}_2 + 2.2\text{Ac}_2\text{O} & \rightarrow \text{Ti(O)}_{1.8}(\text{O}_5\text{P-bBz-PO}_3)_{0.1} + 4\text{AcO}^\circ\text{Pr} + 0.4\text{AcOEt}
\end{align*}
\]

where -bBz- is the bridging biphenylene(methylene) group.

The amount of residual acetoxy groups can be evaluated from the thermogravimetric analysis in air of bBzP-Ti (Figure S5). The 22.5% weight loss between 120 and 600 °C would correspond to the conversion of \(\text{Ti(O)}_{1.68}(\text{OAc})_{0.20}(\text{O}_5\text{P-bBz-PO}_3)_{0.1}\) into 0.895 TiO2 + 0.105 TiP2O7. The corresponding degree of condensation around Ti (given by (1.68 + 2.0.105)/ 2-100%) is very high, 94.5%.

![Figure 4](https://dx.doi.org/10.1021/acs.chemmater.9b05095)

Figure 4. Evaluation of the water stability of bBzP-Ti under acidic and basic conditions: N2 physisorption isotherms at 77 K and corresponding BJH pore size distribution of bBzP-Ti (black), bBzP-Ti treated at pH 1 (blue), and pH 12 (brown) for 24 h at 30 °C in aqueous solutions at different pH, washed, then dried at 120 °C under vacuum. As shown in Figure 4, for pH values of 1 and 12 the physisorption isotherm and pore size distribution were only marginally impacted, and the specific surface area and total pore volume remained above 430 m² g⁻¹ and 1.0 cm³ g⁻¹, respectively. Conversely, treatment at pH 0 or pH 13 reduced significantly the specific surface area and, in the case of pH 13, the total pore volume. In addition, the mesoporous texture of bBzP-Ti is maintained after treatment
in neutral water at 100 °C for 2 h (see Table S2 and Figure S6).

The remarkable water stability of bBzP-Ti can be explained by its structure: both Ti−O−Ti and Ti−O−P bonds are known for their high hydrolytic stability, as shown by the very low aqueous solubility of crystalline TiO2 and of M(IV) phosphonates.

2.5. Tuning the Texture of bBzP-Ti. As demonstrated in Figure 5 and Figure S7, playing on the reactions kinetics offers simple and efficient ways to tune the texture of bridged titania-bisphosphonates, from micromesoporous to meso-macroporous. Starting from the "standard" synthesis of bBzP-Ti, performed with 8 mL of toluene (for 5.25 mmol Ti(OiPr)4) at 200 °C for 18 h, different bBzP-Ti samples were prepared by changing either the reaction temperature, the solvent volume, or the reaction time. Depending on these reaction conditions, the specific surface area varies between 525 and 720 m^2 g^-1 and the pore volume from 0.43 to 1.85 cm^3 g^-1 (Table 1). As shown on the BJH pore size distributions (Figure S5), the maximum pore size varies from 200 nm to less than 5 nm. In all cases, the ^31P MAS NMR spectra are similar, pointing to a complete condensation of phosphonate precursors. On the other hand, the crystallinity of the samples depends on the reaction conditions: the size of the anatase crystallites varies from 8 to 15 nm, and the sample obtained at 180 °C is practically amorphous. As a rule, a higher reaction temperature, a higher concentration, or a longer reaction time lead to a higher mesopore volume, a lower micropore volume and larger micropores (Figure S8), and a higher crystallite size.

2.6. Introducing Organic Functionality. A major advantage of hybrid materials is the possibility to tailor the functional organic groups depending on the targeted applications. We explored several approaches for preparing bridged titania bisphosphonate materials with functional organic groups. The first approach consists in using functional bisphosphonate bridges. As the number of commercially available bridging bisphosphonates is limited, an interesting alternative consists in modifying a simple (nonfunctional) bridged bisphosphonate with functional monophosphonate groups, either by co-condensation or by postmodification (Scheme 2).

In order to demonstrate the first approach, we prepared a titania bisphosphonate sample with bridging bipyridyl groups (bPyP-Ti) by exactly the same synthesis as bBzP-Ti but using tetraethyl 2,2'-bipyridine-5,5'-bisphosphonate (bPyP) instead of bBzP. As shown in Figure 6, this functional material is mesoporous, with high specific surface area (520 m^2 g^-1) and pore volume (0.93 cm^3 g^-1) (Table 2).

**Table 1. Influence of Reaction Temperature (T_R), Volume of Toluene (V_s, for 5.25 mmol Ti(OiPr)_4), and Reaction Time (t_R) on the Texture of bBzP-Ti Samples**

<table>
<thead>
<tr>
<th>sample</th>
<th>T_R(°C)</th>
<th>V_s(mL)</th>
<th>t_R(h)</th>
<th>S_BET(m^2 g^-1)</th>
<th>V_p(cm^3 g^-1)</th>
<th>D_p(nm)</th>
<th>d_cryst(nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>bBzP-Ti</td>
<td>200</td>
<td>8</td>
<td>18</td>
<td>525</td>
<td>1.20</td>
<td>10.4</td>
<td>12</td>
</tr>
<tr>
<td>bBzP-Ti-180 °C</td>
<td>180</td>
<td>8</td>
<td>18</td>
<td>620</td>
<td>0.43</td>
<td>0.19</td>
<td>3.4 am.</td>
</tr>
<tr>
<td>bBzP-Ti-220 °C</td>
<td>220</td>
<td>8</td>
<td>18</td>
<td>575</td>
<td>1.75</td>
<td>0.07</td>
<td>13.4</td>
</tr>
<tr>
<td>bBzP-Ti-4 mL</td>
<td>200</td>
<td>4</td>
<td>18</td>
<td>625</td>
<td>1.69</td>
<td>0.05</td>
<td>18.2</td>
</tr>
<tr>
<td>bBzP-Ti-16 mL</td>
<td>200</td>
<td>16</td>
<td>18</td>
<td>720</td>
<td>0.89</td>
<td>0.13</td>
<td>5.7</td>
</tr>
<tr>
<td>bBzP-Ti-9h</td>
<td>200</td>
<td>8</td>
<td>9</td>
<td>570</td>
<td>0.73</td>
<td>0.15</td>
<td>7.0</td>
</tr>
<tr>
<td>bBzP-Ti-36h</td>
<td>200</td>
<td>8</td>
<td>36</td>
<td>715</td>
<td>1.85</td>
<td>&lt;0.01</td>
<td>11.4</td>
</tr>
</tbody>
</table>

^BET specific surface area. ^Total pore volume at P/P_0 = 0.99. ^Volume of micropores estimated by DFT analysis. ^BJH average pore diameter in the 2–100 nm range calculated from the desorption branch. ^Average crystallite size estimated by the Scherrer equation from the width of the (101) reflection.
13C NMR indicates that the bipyridyl groups are not damaged during the synthesis. As in the case of bBzP-Ti, XRD indicates the formation of anatase nanorods and 31P NMR shows the presence of fully condensed bisphosphonate groups (Figure S9). Electron microscopy images (Figure 6d and Figure S10) show that these nanorods assemble to form a three-dimensional mesoporous network, as in the case of bBzP-Ti, demonstrating the generality of our synthetic route.

The adsorption of Cu2+ ions was investigated to determine the adsorption capacity of bPyP-Ti and evaluate the accessibility of the bridging bipyridyl groups (see Figure S11 and Table S3). The maximum adsorption capacity was 0.87 mmol Cu g\(^{-1}\) for bPyP-Ti, and only 0.13 mmol Cu g\(^{-1}\) for bBzP-Ti. Accordingly, Cu\(^{2+}\) adsorption by bPyP-Ti is mostly due to the presence of bipyridyl groups and at least 86% of these groups are accessible, as calculated from the bipyridyne content (0.86 mmol g\(^{-1}\), see Table S1) after subtraction of the value found for bBzP-Ti, and assuming the formation of a 1:1 complex. The maximum Cu\(^{2+}\) adsorption capacity of bPyP-Ti is similar to values reported for diamine-functionalized organosilica sorbents,\(^48\) such as PMOS cogels,\(^49\) hexagonal silica cogels \(^50\) or postmodified SBA15.\(^51\) Conversely, the maximum adsorption capacity reported for periodic mesoporous phosphonate hybrid materials prepared from ethyl-

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Table 2. Textural Data for Functional Bridged Titania Bisphosphonate Samples

<table>
<thead>
<tr>
<th>sample</th>
<th>(S_{\text{BET}}) (m(^2) g(^{-1}))</th>
<th>(V_{\text{p}}) (cm(^3) g(^{-1}))</th>
<th>(V_{\text{m}}) (cm(^3) g(^{-1}))</th>
<th>(D_{\text{p}}) (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>bPyP-Ti</td>
<td>520</td>
<td>0.91</td>
<td>0.06</td>
<td>6.5</td>
</tr>
<tr>
<td>bBzP-PyP-Ti</td>
<td>830</td>
<td>1.37</td>
<td>0.17</td>
<td>9.0</td>
</tr>
<tr>
<td>BAPA/bBzP-Ti</td>
<td>314</td>
<td>0.75</td>
<td>0.04</td>
<td>11.4</td>
</tr>
</tbody>
</table>

\(^{a}\)BET specific surface area. \(^{b}\)Total pore volume at \(P/P_0 = 0.99\), \(^{c}\)Volume of micropores estimated by DFT analysis, \(^{d}\)BJH average pore diameter in the 2 to 80 nm range calculated from the desorption branch.
enediamine tetra(methylene phosphonic acid) and TiCl₄ was much lower (36.5 μmol/g), indicating a very low accessibility of the diamine groups in these materials. Two samples were prepared to illustrate the functionalization with monophosphonate groups. The first one, bBzP-PyP-Ti, was obtained by cogelation using 0.1 equiv of both diethylpyridin-3-ylphosphonate and bBzP, while the second sample, BPPA/bBzP-Ti was prepared by surface modification of bBzP-Ti with 3-bromopropyl phosphonic acid (Scheme 2). Functionalization with a bromoalkyl group opens many possibilities for postfunctionalization, notably for the grafting of polymer chains by atom transfer radical polymerization (ATRP).

Both bBzP-PyP-Ti and BPPA/bBzP-Ti samples exhibit a mesoporous texture (Figure 7a). ³¹P MAS NMR (Figure 7b) and ³¹C MAS NMR (Figure S12) confirm the incorporation of the functional groups. The functional group contents are quite high, between 0.80 and 1.03 mmol g⁻¹ (Table S1). The specific surface area (830 m² g⁻¹) and pore volume (1.37 cm³ g⁻¹) of the Cogel bBzP-PyP-Ti are even higher than those of bBzP-PyP-Ti. The specific surface area and pore volume (Table 2); however, this method could be useful in the case of thermally sensitive functional groups.

3. CONCLUSIONS

In summary, we demonstrate a simple and original non-hydrolytic sol–gel route giving access to a promising family of porous nonsiliceous hybrid materials, bridged titania bisphosphonates. The final materials can be described as titanium oxide nanodomains assembled by rigid, completely condensed bisphosphonate groups. The porosity of these materials can be adjusted over a wide range simply by playing on reaction parameters. Furthermore, functional bridged titania bisphosphonates with high contents of functional groups can be obtained by several methods involving either functional bisphosphonates or easily available monophosphonate esters or monophosphonic acids. The stability over a wide pH range of these original hybrid materials combined to their tunable texture and functionalnality makes them complementary of other mesoporous hybrid materials such as organosilicas or metal phosphonates and opens exciting opportunities notably in the fields of heterogeneous catalysis, drug delivery, ion-exchange, or adsorption/separation.

4. EXPERIMENTAL SECTION

4.1. Chemicals. Titanium(IV) isopropoxide (Ti(OiPr)₄, 97%) and acetic anhydride (Ac₂O, 99%) were purchased from Sigma-Aldrich. 4,4′-Bis(diethylphosphonomethyl)biphenyl (bBzP, 98%) and 3-Bromomethylphosphonic acid (BPPA) were purchased from Tokyo Chemical Industry. Diethylpyridin-3-ylphosphonate (PyP) and tetraethyl 2,2′-bipyridine-5,5′-bisphosphate (bPyP) were purchased from SiKEMIA in Montpellier. Copper(II) nitrate hemi-pentahydrate was purchased from Sigma-Aldrich. All these reactants were used without further purification. Toluene (Sigma-Aldrich, 99.7%) was dried over a PureSolve MD5 solvent purification system (H₂O < 10 ppm, controlled with a Karl Fischer coulometer).

4.2. Synthesis. To avoid the presence of water, the addition of reagents and sealing of the autoclaves were carried out in a glovebox under an argon atmosphere (<10 ppm of water).

4.2.1. bBzP-Ti. To a solution of titanium(IV) isopropoxide (1.50 g, 5.25 mmol) in toluene (8 mL), 4, 4′-bis(diethylphosphonomethyl)-benzyl, bBzP, (0.24 g, 0.53 mmol) then acetic anhydride (1.18 g, 11.55 mmol) were added slowly. After stirring for 5 min, the obtained solution was transferred to a stainless steel autoclave (Parr Instruments) with a PTFE lining (23 mL), which was then sealed. The autoclave was heated in an oven at 200 °C for 18 h under autogenous pressure. After reaction, the resulting monolith gel was thoroughly washed with ether (5 × 30 mL) and acetone (5 × 30 mL). The gel was dried under vacuum (100 Pa) at 120 °C and finally ground into a fine powder for characterization. %C 13.9; P/Ti 0.22.

4.2.2. bPyP-Ti. To a solution of titanium(IV) isopropoxide (1.50 g, 5.25 mmol) in toluene (8 mL), 4, 4′-bis(diethylphosphonomethyl)biphenyl, bBzP, (0.24 g, 0.53 mmol) then acetic anhydride (1.18 g, 11.55 mmol) were added slowly. After stirring for 5 min, the obtained solution was transferred to a stainless steel autoclave (Parr Instruments) with a PTFE lining (23 mL), which was then sealed. The autoclave was heated in an oven at 200 °C for 18 h under autogenous pressure. After reaction, the resulting monolith gel was thoroughly washed with ether (5 × 30 mL) and acetone (5 × 30 mL). The gel was dried under vacuum (100 Pa) at 120 °C and finally ground into a fine powder for characterization. %C 13.6; %N 2.3; P/Ti 0.31.

4.2.3. bBzP-PyP-Ti. To a solution of titanium(IV) isopropoxide (1.50 g, 5.25 mmol) in toluene (8 mL) were slowly added 4, 4′-bis(diethylphosphonomethyl)biphenyl, bBzP, (0.24 g, 0.53 mmol), diethylpyridin-3-ylphosphonate, PyP, (0.11 g, 0.53 mmol), and then acetic anhydride (1.23 g, 12.08 mmol). After being stirred for 5 min, the obtained solution was transferred to a stainless steel digestion vessel (Parr Instruments) with a PTFE lining (23 mL). After sealing, the vessel was heated in an oven at 200 °C for 18 h under autogenous pressure. After reaction, the resulting monolith was thoroughly washed with ether (5 × 30 mL) and acetone (5 × 30 mL). The gel was dried under vacuum (100 Pa) at 120 °C and finally ground into a fine powder for characterization. %C 13.6; %N 2.3; P/Ti 0.31.

4.2.4. BPPA/bBzP-Ti. Three hundred milligrams of bBzP-Ti powder (2.72 mmol) was added to a solution of (3-bromopropyl)-phosphonic acid, BPPA, (10 mmol) in MeOH (100 mL). The mixture was stirred at room temperature for 2 days. After washing
with MeOH (5 × 30 mL) and acetone (5 × 30 mL), the obtained solid was dried under reduced pressure at 120 °C. P/Ti 0.35; Br/Ti 0.14.

4.3. Stability Tests. To investigate the hydrolytic stability of bBzP-Ti, we stirred the powder (0.20 g) for 24 h at 30 °C in 400 mL of aqueous solutions of HCl (pH0 and pH1) or NaOH (pH12 and pH13). The hydrothermal stability was tested by refluxing the powder in neutral water at 100 °C for 2 h. After washing with water and acetone, then drying under reduced pressure at 120 °C, the N2 sorption measurements were repeated.

4.4. Testing of Cu²⁺ Adsorption by bPyP-Ti and bBzP-Ti. One-hundred eighty milligrams (1.67 mmol) of bPyP-Ti or 185 mg (1.67 mmol) of bBzP-Ti were first washed with water several times to eliminate the acetate groups. They were then separately added to 30 mL of a copper nitrate standard solution (10 mM). After stirring at room temperature for 24 h, the solid was separated by centrifugation and the Cu²⁺ content in the supernatant was analyzed by UV–vis spectroscopy. The amount of adsorbed Cu²⁺ was calculated from the change in Cu²⁺ concentration of the solutions.

4.5. Characterization. Fourier-transform infrared (FTIR) spectra were collected in ATR mode on a Spectrum II PerkinElmer spectrometer. UV/vis absorption spectra were obtained with a JASCO V-760 UV–visible spectrophotometer in 10 mm quartz cells (Hellma). Spectra were recorded over the 900–3000 nm range with a bandwidth of 1.0 nm at a rate of 200 nm min⁻¹. The powder X-ray diffraction (XRD) patterns were collected with a PANalytical X-Pert Pro MPD diffractometer (CuKα1 = 0.1540598 nm). The scanning electron microscopy (SEM) images were obtained with a Hitachi S-4800 electron microscope. Energy-dispersive X-ray spectroscopy (SEM-EDX) was done on an Oxford Instruments X-Ma SDD. The high-resolution transmission electron microscopy (TEM) analyses were acquired with a JEOL CENTURIO TEM-EDX apparatus; the specific area was measured at 77 K with a Micrometrics Triac. FTIR spectra, DFT pore size distributions, SEM and TEM images, solution and solid-state NMR spectra, detailed data on stability assessment and on Cu²⁺ adsorption (PDF)

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.chemmater.9b05095.

Composition of the samples, FTIR spectra, DFT pore size distributions, SEM and TEM images, solution and solid-state NMR spectra, detailed data on stability assessment and on Cu²⁺ adsorption (PDF)

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Author Contributions

Y.W. performed the experiments. J.A. and P.M. supervised the experiments. F.M. wrote the manuscript with the help of Y.W. and J.A.

Notes

The authors declare no competing financial interest.

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