Hexane-1,2,5,6-tetrol as a Versatile and Biobased Building Block for the Synthesis of Sustainable (Chiral) Crystalline Mesoporous Polyboronates

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Supporting Information

ABSTRACT: We report on the synthesis and characterization of novel mesoporous chiral polyboronates obtained by condensation of (RS)/(SS)-hexane-1,2,5,6-tetrol (HT) with simple aromatic diboronic acids (e.g., 1,3-benzenedi boronic acid) (BDB). HT is a cellulose-derived building block comprising two 1,2-diol structures linked by a flexible ethane bridge. It typically consists of two diastereomers one of which [(S,R)-HT] can be made chirally pure. Boronic acids are abundantly available due to their importance in Suzuki–Miyaura coupling reactions. They are generally considered nontoxic and easy to synthesize. Reactive dissolution of generally sparingly soluble HT with BDB, in only a small amount of solvent, yields the mesoporous HT/polyboronate materials by spontaneous precipitation from the reaction mixture. The 3D nature of HT/polyboronate materials results from the entanglement of individual 1D polymeric chains. The obtained BET surface areas (SAs) and pore volumes (PVs) depend strongly on HT’s diastereomeric excess and the meta/para orientation of the boronic acids on the phenyl ring. This suggests a strong influence of the curvature(s) of the 1D polymeric chains on the final materials’ properties. Maximum SA and PV values are respectively 90 m² g⁻¹ and 0.44 mL g⁻¹. Variably sized mesopores, spanning mainly the 5–50 nm range, are evidenced. The obtained pore volumes rival the ones of some covalent organic frameworks (COFs), yet they are obtained in a less expensive and more benign fashion. Moreover, currently no COFs have been reported with pore diameters in excess of 5 nm. In addition, chiral boron-based COFs have presently not been reported. Scanning electron microscopy reveals the presence of micrometer-sized particles, consisting of aggregates of plates, forming channels and cell-like structures. X-ray diffraction shows the crystalline nature of the material, which depends on the nature of the aromatic diboronic acids and, in the specific case of 1,4-benzenedi boronic acid, also on the applied diastereomeric excess in HT.

KEYWORDS: Boronates, Mesoporous, Chiral, Polymers, Cellulose

INTRODUCTION

Mesoporous materials have attracted significant attention as their wide pores (2–50 nm) allow for improved diffusion and accessibility, favoring applications as diverse as, controlled drug release, chromatography, adsorbents, electrodes, solar cells, and heterogeneous catalysis. Exemplary are (non)siliceous mesoporous oxides, periodic mesoporous organic silicas (PMOs), mesoporous carbons, hyperconjugated porous polymers, and some metal–organic frameworks. Often the introduction of mesopores in a material is linked to the use of (supra)molecular templates such as surfactants, but also other techniques are known like dealumination/desiliciation, nanoadamisteering, and local associations of helices, among others.

Borate ester polymers (BEP) and their tendency to form hierarchical supramolecular structures have been widely reported in the literature. However, few micro/mesoporous materials have been reported based on 1D polymeric boronate esters containing backbones not involving heteroatoms (e.g., nitrogen, sulfur). Exemplary are the flowerlike micro/mesoporous microparticles made by the condensation of 1,4-benzenedi boronic acid (1,4-BDB) with pentaerythritol (PE). These materials have a specific surface area of 184 m² g⁻¹.
interlocking of rings.17

achieved by the interpenetration of 2D/3D frameworks or weaving, entanglement in porous frameworks can also be one phenyl ring.19

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of a certain decrease in crystallinity.16 Most importantly, owing to the large flexibility of the 1D threads in demetalated COF-505, a 10-fold increase of its elasticity was observed.17 Besides weaving, entanglement in porous frameworks can also be achieved by the interpenetration of 2D/3D frameworks or interlocking of rings.17

In spite of the high surface areas and pore volumes, and the related plethora of potential and promising applications, the commercial use of mesoporous materials is still quite a challenge due to their chemical instability, expensive synthesis approaches, and nonstraightforward processability.18 The cost of the extended linkers, which is especially relevant for COFs and MOFs, is another major concern with, for instance, only two commercialized MOFs employing linkers with more than one phenyl ring.19

Recently, we have reported on the sustainable synthesis of hexane-1,2,5,6-tetroir (HT) in which two 1,2-diol moieties are joined by a flexible ethane bridge.20 HT is derived from cellulose in 3–4 steps many of which involve heterogeneous catalysts. In successive order these comprise (Figure 1) the following list:

- Conventional/microwave pyrolysis of (waste) cellulosic polysaccharides.21–23
- Hydrogenation of levoglucosanol to levoglucosanone over dihydrolevoglucosanone with heterogeneous Pd catalysts (one or two steps).24,25 It is noteworthy that presently a joint venture between the Australian Circa Company and the Norwegian multinational Norskenskog is running a fully operational 50 tonnes/year prototype plant in Tasmania.26
- Combined one pot hydrogenation/hydrolysis of levoglucosanone (one step).20

HT is typically obtained as a mixture of two diastereomers notably (S,R)-HT (mesomer) and (S,S)-HT. Boronic acids are abundantly available due to their importance in Suzuki–Miyaura coupling reactions.27,28 They are generally considered nontoxic and are easy to synthesize.27,29

Here we report on the synthesis of a novel porous organic polymer obtained by reactive dissolution of sparingly soluble HT with simple benzenediboronic acids (BDB) and concomitant precipitation of the mesoporous HT polyboronate. The reaction is conducted at RT and requires no special reaction conditions such as vacuum, inert atmosphere, or controlled water removal. It is shown that the properties of the resulting mesoporous polymers are largely determined by the chirality of HT and the meta/para orientation of the boronic acids on the phenyl ring. Thereby HT’s chirality is likely inducing coiling of the polymer chain, allowing for interweaving of the polymeric chains. These materials are crystalline and in spite of their lower surface areas (30–90 m² g⁻¹) display mesopore volumes up to 0.441 mL g⁻¹. Scheme 1A provides a visual of the potential polymeric chains indicative of their stereocenters and points of potential molecular rotation. Hereafter these materials are denoted as “HT/polyboronates” or more specifically using the HT/1,x-BDB/solvent formalism.

**RESULTS AND DISCUSSION**

Table 1 lists the obtained specific surface areas (SAs) and pore volumes (PVs) as a function of the type of diboronic acid used, the applied solvent, and HT’s diastereomeric excess. It is found that the meta orientation of the diboronic acids on the phenyl ring (1,3-BDB) gives higher SA and PV values compared to the para orientation of the diboronic acids (1,4-BDB) (Table 1,
**Scheme 1.** (A) Schematic Visualization of Two Different HT (hexane-1,2,5,6-tetrol)-BDB (1,4- or 1,3-Benzenediboronic Acid) Polymeric Chains with Indication of the Stereocenters and Points of Potential Rotation of the Molecular Chains, (B) Illustrative Variability of the Growing Polymeric Chain When Incorporating Chirally Pure (R,S)-HT [Shown as (R-S)], and (C) Unique Chiral Polymer When Constructing the Polymeric Chain from (S,S)-HT [Shown as (S-S)]

**Table 1.** Survey of the Specific Surface Areas and Specific Pore Volumes for a Range of HT (Hexane-1,2,5,6-tetrol) Containing Polyboronate Esters

| entry | component 1 | component 2 | spec surface area (m² g⁻¹ BET) | spec pore volume (mL g⁻¹) |
|-------|-------------|-------------|--------------------------------|--------------------------|
| a     | HT 38%      | 1,4-BDB     | 42                             | 0.227                    |
| b     | HT 38%      | 1,3-BDB     | 65.8                           | 0.408                    |
| c     | HT 38%      | 4,4'-BPDB   | 64.5                           | 0.359                    |
| d     | HT 38%      | 1,3-BDB     | 60.5                           | 0.315                    |
| e     | HT 38%      | 1,3-BDB     | 74.0                           | 0.397                    |
| f     | HT 38%      | 1,3-BDB     | 61.2                           | 0.290                    |
| g     | HT 38%      | 1,3-BDB     | 67.2                           | 0.441                    |
| h     | HT 38%      | 1,3-BDB     | 49.1                           | 0.289                    |
| i     | HT 38%      | 1,3-BDB     | 62.8                           | 0.407                    |
| j     | HT 38%      | 1,3-BDB     | 51.3                           | 0.271                    |
| k     | HT 98%      | 1,3-BDB     | 80.7                           | 0.412                    |
| l     | HT 98%      | 1,4-BDB     | 32.8                           | 0.148                    |
| m     | HT 98%      | 1,3-BDB     | 87.3                           | 0.441                    |

Entries a/b). This suggests for a distinct influence of the curvature of the polymeric chain. Interestingly, the use of 4,4'-biphenyldiboronic acid (4,4'-BPDB) led to quasi similar results as obtained with 1,3-BDB (Table 1, entry c). This can be rationalized by the nonplanarity of biphenyl units and the consequent absence of p–p stacking of the aromatic rings. The use of acetone or CH₂Cl₂ as the reaction solvents gives higher SAs and PVs than when THF or AcCN are used (Table 1 entries d–g). Additionally, the SA and PV values benefit also from higher reaction volumes (Table 1, entries d, h, and i).

Partial enrichment of (S,R)-HT over (S,S)-HT was described in a previous publication.²⁰ Specific crystallization of the (S,R)-HT diastereomer was achieved from hot 1,4-dioxane, allowing for the evaluation of HT’s chirality on the properties of the HT/polyboronates. Unfortunately, it has presently not been found possible to crystallize/purify (S,S)-HT. It is noteworthy that (S,R)-HT is a mesomer, making that it displays no net chirality on its own but does so effectively in a polymeric setting. Given that (S,R)-HT can build into the polymeric chain as (S,R) or (R,S), substantial variability in the curvature of the polymeric chain is likely present (Scheme 1B/C). Actual and distinct influences of the diastereomeric (S,R)/ (S,S) ratio in HT on the properties of the materials are observable:

(A) In using 38% d.e. (S,R)-HT, higher SA and PV are obtained then when 38% d.e. (S,S)-HT is used (Table 1, entries g vs j).

(B) Elevating the d.e. in (S,R)-HT from 38 to 98% is found to increase both the SA and PV by around 30% for HT/1,3-BDB/acetone, only an SA increase of 30% is observed with an already high PV (Table 1, entries m vs g).

(C) Using diastereomerically pure (S,R)-HT magnifies the effect of the meta/para orientation of BDB on the spread in SA and PV (Table 1, entries a, b, k, and l).

It is noteworthy that no cooperative stereochemical effect occurs as hydrolysis of the formed HT/polyboronates does not reveal a preferential incorporation of one of the HT diastereomers. Irrespective the nature of the HT/polyboronates, the obtained SA and PV values correlate to a certain degree (Figure S1_A).

In terms of yield, reactions between HT with 38% d.e. (1 > 2) and 1,4-BDB typically give between 65 and 75% yield of the solid HT/1,4-BDB polyboronate product. This is 73–85% of the theoretical yield which is 88.6% given the loss of two waters molecules during the condensation of HT and...
BDB. Reactions between HT with 38% d.e. (1 > 2 or 1 < 2) and 1,3-BDB typically give between 55 and 65% yield of the solid HT/1,3-BDB polyboronate product. This is 62–73% of the theoretical yield. Reactions using pure HT (i.e., > 98% d.e.) are with 26–30% much lower yielding, and this irrespective of the HT diastereomer and the BDB nature.

As shown in Figure 2A, the N₂ physisorption isotherms display a type IV isotherm and an H3 hysteresis. The latter is indicative of aggregates of platelike particles forming slitlike pores. This is reminiscent of the pentarythritol/1,4-BDB (PE/1,4-BDB) material reported by Fujiwara et al. and Matsushima et al. with the difference that the N₂ adsorption is fully reversible, strengthening our claim on some ordering in the pore wall structure. The nature of the irregular voids between the platelets, as channels or cell-like structures, are in the tens of nanometers and thus consistent with the N₂ physisorption results and the appearance of an H3 hysteresis in the N₂ physisorption experiments. In spite of the high melting temperatures of the HT/polyboronates (~600 K), TEM imaging was found impossible due to melting of the materials under the electron beam. XRD revealed a degree of crystallinity in all HT/polyboronates with the nature of the BDB (length, substitution pattern) affecting markedly the appearance of the different XRD patterns (Figure 4). Given the characteristic H3 hysteresis loop in the N₂ physisorption experiments and the absence of order in the SEM images (Figure 3), we assume that the observed crystallinity reflects some ordering in the pore wall structure. The nature of the used solvent has no effect on the crystallinity (Figure S2). The diastereoselectivity of HT affects the crystallinity only in the case of the 1,4-BDB linker (Figure 5).

The (S,R)-HT/1,4-BDB and (S,R)-HT/1,3-BDB polyboronates were characterized by ¹³C CP-MAS and ¹¹B spin echo MAS NMR, and the results are shown in Figure 6. Analytical simulations of the ¹¹B spectra confirmed the existence of only one type of boron species, and this is irrespective of the nature of the BDB unit. (Figure 6B/D, red curve). FT-IR spectroscopy further confirmed the presence of boronate esters with characteristic vibrational peaks in the 1320–1300 cm⁻¹ (B–O stretch) and 690–650 cm⁻¹ (characteristic boronate ester peak³⁵) spectral ranges (Figure S3_A). In addition, FT-IR spectroscopy revealed the clear absence of OH groups (3500–3000 cm⁻¹), characteristic for the residual presence of HT or incomplete boronate ester formation in the synthesized HT/polyboronates (Figures S3_B/C).

## CONCLUSIONS

A range of new crystalline (chiral) mesoporous materials were obtained by the facile and benign condensation of (chiral), biobased hexane-1,2,5,6-tetrol (HT) and simple aromatic diboronic acids. The so formed mesoporous HT/polyboronates are obtained by spontaneous precipitation from the reaction mixture, not requiring an extensive workup procedure. HT is easily accessible from cellulose and chiral pure (R,S)-HT is straightforward to crystallize. The applied simple diboronic acids are abundantly available due to their commercial use in Miyaura–Suzuki reactions, easy to synthesize, and generally

SEM analysis of 98% d.e. (R,S)-HT/1,4-BDB/THF (Figure 3A series), 98% d.e. (R,S)-HT/1,3-BDB/THF (Figure 3B series), and 98% d.e. (R,S)-HT/1,3-BDB/acetone (Figure 3C series) reveals the following:

- Micrometer-sized particles (Figure 3A1/B1/C1).
- Aggregates of plates for 98% d.e. (R,S)-HT/1,4-BDB/THF (Figure 3A2/A3).
- A more pronounced pore structure when 1,3-BDB is used over 1,4-BDB (Figure 3B1/B2). Here the plate aggregates form channels and cell-like structures. Interestingly, some of the observed plate-like structures seem to have delaminated which may have contributed positively to the observed enhanced porosity (Figure 3B3).
- Using acetone as the reaction solvent yields equally a highly porous structure yet with a substantially different fine-structure (on the micrometer level) than observed when THF is used (Figure 3B2/B3 vs Figure 3C2/C3).

Overall the dimensions of the irregular voids between the platelets, as channels or cell-like structures, are in the tens of nanometers and thus consistent with the N₂ physisorption results and the appearance of an H3 hysteresis in the N₂ physisorption experiments. In spite of the high melting temperatures of the HT/polyboronates (~600 K), TEM imaging was found impossible due to melting of the materials under the electron beam. XRD revealed a degree of crystallinity in all HT/polyboronates with the nature of the BDB (length, substitution pattern) affecting markedly the appearance of the different XRD patterns (Figure 4). Given the characteristic H3 hysteresis loop in the N₂ physisorption experiments and the absence of order in the SEM images (Figure 3), we assume that the observed crystallinity reflects some ordering in the pore wall structure. The nature of the used solvent has no effect on the crystallinity (Figure S2). The diastereoselectivity of HT affects the crystallinity only in the case of the 1,4-BDB linker (Figure 5).

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considered nontoxic.\textsuperscript{35–37} The low cost of these materials is ultimately considered to benefit potential commercial applications. The reported materials show an appreciable pore volume of up to 0.44 mL g$^{-1}$, rivalling the pore volumes of some reported COF materials yet obtained in a less expensive way. Importantly, presently no COF has been reported with pore diameters in excess of 5 nm. The (R$\,\!$,S$\,\!$)-HT/\polypolyboronate represents a first example of a chiral mesoporous polyboronate. The high pore volume and the larger pore sizes of 5–50 nm offer great potential to the incorporation of enzymes or as supports to immobilize catalysts to convert bulkier substrates. The meta/para orientation of the boronic acids on the phenyl ring was shown to influence the obtainable surface areas and pore volumes. Uniquely, a distinct influence of HT’s chirality (its d.e. value) on the materials’ properties was evidenced. Future work will focus on testing these new materials toward a wide range of different applications, thereby hopefully demonstrating its potential as a valuable material.

\section{ASSOCIATED CONTENT}

\subsection{Supporting Information}

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acssuschemeng.9b02772.

Figure 3. Scanning electron microscopy (SEM) images of 98% d.e. (R,S)-HT/1,4-BDB/THF (parts A1, A2, A3), 98% d.e. (R,S)-HT/1,3-BDB/THF (parts B1, B2, B3), and 98% d.e. (R,S)-HT/1,3-BDB/acetone (parts C1 C2 C3).

Figure 4. X-ray diffractograms (XRD) of (A) HT/1,4-BDB, (B) HT/1,3-BDB, and (C) HT/4,4$'\!$-BPDB; no background subtraction.

Figure 5. X-ray diffractograms (XRD) of HT/1,3-BDB (A 38% HT d.e.; B 98% HT d.e.) and HT/1,4-BDB (C 38% HT d.e.; D 98% HT d.e.); PHT = chirally pure HT.
Experimental procedures, additional X-ray diffractograms, and FT-IR spectra (PDF)

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The authors declare no competing financial interest.

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