Enantiomerically enriched, polycrystalline molecular sieves

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Zeolite and zeolite-like molecular sieves are being used in a large number of applications such as adsorption and catalysis. Achievement of the long-standing goal of creating a chiral, polycrystalline molecular sieve with bulk enantiomeric enrichment would enable these materials to perform enantioselective functions. Here, we report the synthesis of enantiomerically enriched samples of a molecular sieve. Enantipure organic structure directing agents are designed with the assistance of computational methods and used to synthesize enantioenriched, polycrystalline molecular sieve samples of either enantiomer. Computational results correctly predict which enantiomer is obtained, and enantiomeric enrichment is proven by high-resolution transmission electron microscopy. The enantioenriched and racemic samples of the molecular sieves are tested as adsorbents and heterogeneous catalysts. The enantioenriched molecular sieves show enantioselectivity for the ring opening reaction of epoxides and enantioselective adsorption of 2-butanol (the R enantiomer of the molecular sieve shows opposite and approximately equal enantioselectivity compared with the S enantiomer of the molecular sieve, whereas the racemic sample of the molecular sieve shows no enantioselectivity).

Chirality | zeolite | asymmetric catalysis | chiral adsorption

The synthesis of zeolites and zeolite-like molecular sieves has been accomplished using organic structure-directing agents (OSDAs) (1–4). This synthetic method uses an organic molecule (OSDA) to interact with and influence the assembly pathway of the inorganic components to create a crystalline, organic-inorganic composite material. Upon removal of the OSDA, the microporous void space that is created can be exploited in a variety of applications (i.e., catalysis, separations, ion exchange, and adsorption) (5). These microporous, polycrystalline materials are 3D networks of oxide tetrahedra [zeolites contain only silicon and aluminum, whereas zeolite-like molecular sieves can have a broader range of elements (1–5)] that create highly ordered, hydrothermally stable framework structures with pores of sizes less than 2 nm. Molecular sieves provide shape-selective properties and, coupled with the inclusion of catalytic active sites, are capable of innumerable highly selective chemical reactions. Despite the abundance of chirality in nature, the discovery of a zeolite or zeolite-like microporous material with enantioselective properties has remained elusive. Enantioenriched, chiral zeolitic materials are of particular interest for their potential to provide robust, new, enantiospecific, shape-selective catalytic pathways and separation processes (6–13).

Several inherently chiral molecular sieves have been synthesized to date (although the bulk, polycrystalline samples are racemic), including *BEA, CZP, GOO, -ITV, JRY, LTI, OSO, SFS, and STW [molecular sieve framework types are designated by three-letter codes that define the unique connectivity of the oxide tetrahedral (14)]. In particular, several studies have reported polymorph A (which possesses a chiral helical pore) enriched *BEA. However, zeolite *BEA is limited in that the material crystallizes as highly faulted intergrowths of a racemic mixture of polymorph A and polymorph B (achiral). In 1992, Davis and Lobo (15) discussed the concept of synthesizing a chiral *BEA molecular sieve via the use of a chiral OSDA and reported low enantioenrichment (ee) for both a chemical reaction and an adsorption experiment. Recently, Tong et al. (16) have reported a high-fluoride method of synthesizing polymorph A enriched *BEA using achiral OSDAs (although it is difficult to understand the origin of the proposed enantioenrichment) and reported low ee’s from a chemical reaction. Other authors have reported samples enriched in polymorph A with similar ambiguity in enantioenrichment (17–20). These studies suggest the viability of synthesizing an enantioenriched, chiral molecular sieve. However, the inherent difficulty in controlling the synthesis of polymorphic domains, and the enantiomeric domains of only polymorph A in *BEA, has made it very problematic to conclusively prove that a bulk sample of a molecular sieve does in fact have an enantioenriched framework content. Davis and Lobo (15) discussed this issue and suggested that a preferred approach would be to design syntheses that target chiral molecular sieve structures where individual crystals are single enantiomers (microporous analogs to quartz). The STW framework is an example of this type of molecular sieve; recent evidence shows that each individual crystal is a single enantiomorph (i.e., there are no polymer or opposite enantiomeric domains within a single crystal) (21). A molecular sieve with the STW framework was initially synthesized as a germanosilicate (denoted SU-32) in 2008 (22) and then as a hydrothermally stable pure-silica material in 2012 (23).
The STW structure contains 10-membered rings (MRs, the number of oxygen or tetrahedral atoms that make up the ring) that form chiral helical pores (22). Past synthesis protocols that used achiral OSDAs resulted in bulk, polycrystalline samples that were racemic mixtures of the two structural enantiomers. STW, therefore, provides a good model for designing the synthesis of bulk, polycrystalline samples of either the “R” or “S” enantiomers of the framework. Here, we use computational methods to aid in our design of chiral OSDAs for the synthesis of enantiopure STW and show that a bulk sample of a polycrystalline, molecular sieve can be synthesized with significant enantiomeric enrichment. Our design (shown in Fig. 1) provides for the synthesis of either the R or the S enantiomers of STW, thus yielding materials that enable appropriate control experiments when elucidating structures and functions.

Results and Discussion

Directed Computational Design of a Chiral OSDA for STW. Schmidt et al. (24) reported on the synthesis of STW using a computationally predicted OSDA. This work demonstrated the feasibility of a priori predicting chemically synthesizable monoquaternary, imidazolium OSDAs to create a specified, fluoride-mediated, pure-silica framework. Additionally, STW has been reported to form using diquaternary imidazolium-based OSDAs that are of sufficient size to conform to 10MR channel structure, implying more rigid, chiral analogs may be included in the framework of STW, and may potentially impart structural chirality (24–28). Here, we implemented the previously published computational method implemented by Schmidt et al. (24) and Pophale et al. (29) and used the molecular design constraints suggested by Davis and Lobo (15). The computational method was modified such that a given enantiomer of each potential OSDA molecule was simulated in both enantiomers of STW, with successful candidates producing a strong stabilization in only a single enantiomer. Based on our previous work with STW, we believed that a computed stabilization energy larger than −15 kJ (mol Si)\(^{-1}\) would be needed to form STW. Ultimately, a single OSDA candidate was selected (Table 1). Relative to the other predicted OSDAs for STW, 2 has the largest energy difference between enantiomers, making it the most suitable target for experimental evaluation. Coupled with energy predictions, this molecule was also selected after ensuring that the both enantiomers were synthetically attainable (24). A full description of the computational method is provided in SI Appendix.

Synthesis and Characterization of Enantiopure OSDA 2. Detailed procedures for the chiral resolution and reaction schemes used to synthesize OSDA 2 are given in SI Appendix, Figs. S1 and S2. Chiral resolution of the starting compound (trans-2-phenylcyclopropane-1-carboxylic acid, 1) was performed by successive crystallizations with either dehydroabietylamine or quinine as chiral derivatization agents to obtain the R (1a) or S (1a') forms of the starting material (SI Appendix, Fig. S1). Polarimetry measurements were taken for the salt and free-acid forms of 1a and 1a' to confirm enantiopurity, after the separation was complete. Further details of the synthesis of 2 are provided in SI Appendix (note: enantiomers of 2 will be denoted R-2 and S-2). \((\Lambda,R)^{-}\)BINPHAT tetrabutylammonium salt was used as a chiral shift reagent to detect the enantiopurity of 2 after executing the reactions outlined in SI Appendix, Fig. S2. As shown in Fig. 2, the neat product \(^1\)H NMR resonances at 8 4.80 and 4.04 ppm are singular and distinct multiplets. Addition of BINPHAT to a racemic sample of 2 results in a twofold spectral change: (i) a distinct shift upfield of only the aforementioned peaks from their initial position and (ii) an observed peak split. Integration of the split peaks indicates that they maintain a 1:1 ratio, expected for a racemic compound. However, the two enantiomers of 2 show only a single peak. This method confirms the enantiopurity of 2 and demonstrates that no racemization occurs throughout the organic synthesis scheme.

Synthesis and Characterization of Enantiopure STW. Syntheses of STW were conducted with enantiopure samples of the R and S enantiomers of 2, as well as the racemic mixture of the two. These syntheses were performed at temperatures and H\(_2\)O/SiO\(_2\) ratios that are typical for silica-enriched, fluoride-mediated syntheses. Because STW is composed of ~80% double four-tetrahedral atom rings that are known to be stabilized by inclusion of germanium, addition of varying quantities of germanium to the synthesis gels was explored (30, 31). Following the work of Schmidt et al. (24), reagent molar ratios that led to the synthesis of STW in the shortest times were 1 SiO\(_2\)-x GeO\(_2\):5 H\(_2\)O:0.5 HF:0.5 2 at 160 °C (where x varies from 0.05 to 0.5) using 10% (wt/wt) seeds produced from an achiral diquaternary OSDA with crystal sizes on the order of 1 μm. Aluminum-containing samples were also synthesized (in the presence of Ge) using aluminum isopropoxide. In these syntheses, initial gel Si/Al ratios were maintained above 50; otherwise, RTH impurities were observed in the resultant products (25). Aged the synthesis gels over the course of 24 h was found to reduce crystallization times. A complete summary of the synthesis results is provided (SI Appendix, Tables S1–S3) along with representative powder X-ray diffraction (PXRD) patterns (SI Appendix, Fig. S3). Additionally, \(^{13}\)C, \(^{29}\)Si, and \(^{27}\)Al solid-state NMR (SS NMR) spectra, a representative thermogravimetric analysis, and a selection of scanning electron micrographs are listed as supporting

![Image](https://example.com/image.png)

Fig. 1. Illustration of the synthesis of enantioenriched STW samples using enantiopure, chiral OSDAs.
characterizations (SI Appendix, Figs. S4–S8). In general, the concentrations of germanium and aluminum were found to increase in the product materials relative to the synthesis gel. Representative results from energy-dispersive X-ray spectroscopy are given in SI Appendix, Table S4 for products from germanosilicate and aluminogerminosilicate synthesis gels. In addition to STW, numerous other phases were produced without significant variation in synthesis conditions, including LTA, RTH, IWV, CSV, as well as several phases that could not be identified or consisted of layered organosilicate materials. Synthesis of these microporous materials is not unexpected, because they have all been shown to be derived from similarly shaped imidazole-based OSDAs under crystallization conditions similar to those reported here (25–28, 32).

As-made STW crystals obtained using 2 were analyzed by $^{13}$C cross-polarization magic-angle spinning SS NMR to evaluate whether the OSDA remains intact and directs the formation of STW. Fig. 3 compares the neat OSDA liquid $^{13}$C NMR spectrum (Fig. 3A) with those obtained from analyzing the occluded organic (from SS $^{13}$C NMR) and the organic recovered by dissolution of the framework structure (liquid $^{13}$C NMR). Agreement between all spectra indicates that 2 is occluded intact within the STW framework and is the structure-directing agent. As such, STW is not formed from an organic decomposition product or as a consequence of spontaneous, directed crystallization from the seed crystals. Efforts to perform chiral shift experiments on recovered OSDA 2 collected from dissolution of the STW framework by hydrofluoric acid were not successful due to solubility issues (2, chiral shift reagent, and solvent combinations).

Although it has been speculated that chiral OSDAs (when directed toward a chiral structure) should necessarily lead to an enrichment in the chirality of framework, this concept has never been conclusively proven. To do so, the chirality of the organic occluded within the STW structure must be known, as must the chirality of the framework structure for a bulk, polycrystalline sample. CD provides an effective method of accomplishing the former for as-made STW samples. The results for R-, S-, and racemic STW samples are given in Fig. 4. As shown by the data provided in Fig. 4, the molecules occluded within R- and S-STW absorb light at 242 nm with equal and opposite polarities. No measured rotation of light is observed for the racemic sample because, statistically, any polarization effects are negated for a sample containing equivalent quantities of chiral crystals. These data collectively demonstrate that 2 remains chiral and enantiopure within the STW framework. Although it may be possible to analyze the structural chirality of a calcined STW structure, no adsorption in the 200- to 300-nm range was detected from our germanosilicate samples. Incorporation of UV-active heteroatoms beyond those that have been used in this work may provide a pathway for future investigation via CD studies.

In previous reports involving the enrichment of polymorph A in *BEA (which does not guarantee enantioenrichment), characterization of any true structural enrichment of polymorph A has been limited to transmission electron microscopy (TEM) analysis (PXRD analyses are fraught with problems) of the polymorph domains and demonstration of enantioenrichment by measuring some function (e.g., catalysis or adsorption) (15, 16). A distinct issue with this methodology, however, is that the synthesis of these materials lacks proper controls with regard to the enantiomorphs obtained in the material products, and therefore it is not possible to perform appropriate analytical and experimental controls to confirm chirality. The methods we have developed and used here are not subject to these problems. We are able to control the synthesis of a chiral OSDA (and obtain the R-, S-, and racemic forms thereof) to yield bulk, polycrystalline STW samples composed of individual crystals of a single chirality. As such, characterization and functional results that are capable of probing the bulk chirality of the two enantioenriched samples of opposing direction must demonstrate enantiomeric excess (ee) that is approximately equal and opposite, with the racemic preparation revealing no ee. The CD experiments for the R-, S-, and racemic STW samples provide an illustration of how these types of controls can be exploited to derive meaningful conclusions.

Characterization of the structural chirality of STW is complicated. An examination of the literature yields few methods for effectively determining this property for siliceous microporous materials beyond the demonstration of enantiomeric excesses derived from reactions or adsorption experiments. Analytical difficulty partially stems from the small crystal sizes (as is the case for STW synthesized with OSDAs used here) that prohibit the use of single-crystal XRD. Rojas et al. (31) point toward the possibility of examining the optical activity to quantify enantiomeric excess within a bulk, polycrystalline sample. SEM and high-resolution TEM (HRTEM) have proven to be highly advantageous methods for analyzing crystallite morphologies. HRTEM has been used previously to analyze the polymorph domains in *BEA (16, 20). However, specific chiral space groups (i.e., structural enantiomers) within the detected polymorph A domains were not able to be determined. Fundamentally, the difficulty in unequivocally distinguishing between the space groups of a chiral material using traditional TEM techniques arises from properties inherent to both structural chirality and experimental limitations. Specifically, on a given 2D plane (as is observed in a typical TEM experiment) the structures of the two enantiomers are superimposable and indistinguishable. A method was therefore developed to perform 3D HRTEM experiments by Ma et al. (21) to effectively characterize chiral space groups. The projection of the STW framework shifts after rotation along the
screw axis as schematically illustrated in Fig. 5A. The rotation direction can be either clockwise (to the right) or counterclockwise (to the left). A zeolite crystal [that has been deposited with gold nanoparticles (diameter ∼5 nm) that serve as reference points with the microporous structure] is first aligned to [2110] zone axis and a through-focus series of HRTEM images are taken from a thin area. The crystal is then tilted continuously to the right or to the left by 30° about the screw axis. Upon rotation, the [1100] zone axis is observed if the tilting is clockwise. Based on simulated results, the crystal is right-handed if the shift direction is downward (Fig. 5B and C). Upon rotation, the [1100] or [1010] zone axes are observed if the tilting is to the right or left, respectively. A series of through-focus images are then taken again along either zone axis. Two images along [2110] and [1100] are obtained by structure projection reconstructions using the through-focus series of HRTEM images and are aligned based on the positions of gold nanoparticles (Fig. 5D–G and SI Appendix, Figs. S9–S11). By comparing the aligned images from the two zone axes there is an observable shift between the two projections from which the space group of the crystal can be assigned. Additional images are provided in SI Appendix.

The results collected from using this method to analyze STW crystals selected from bulk, polycrystalline samples synthesized using the R-, S-, and racemic versions of 2 are given in Table 2. Out of the six crystals analyzed for both the R- and S-OSDA-derived samples, five were determined to possess the P6$_2$22 and P6$_3$22 space groups, respectively, demonstrating (within this dataset) notable enantioenrichment. Moreover, these experimental results are consistent with computational predictions whereby R-OSDA 2 is expected to yield crystals with P6$_2$22 space group and, similarly, S-OSDA 2 is anticipated to result in crystals with the P6$_3$22 space group (Table 1). Analysis of the racemic sample resulted in an equal number of crystals from each space group, as expected.

Ideally, for a given sample synthesized using a chiral OSDA, HRTEM analysis would subsequently demonstrate only crystals from the expected space group. However, as shown in Table 2, a crystal from both chiral OSDA-derived samples was determined to be from the unexpected space group (e.g., P6$_3$22 from the R-OSDA). Possible reasons for incomplete purity include that (i) the crystal of the opposite space group was a seed crystal, (ii) there was OSDA degradation that occurred during that yielded a racemic portion of the polycrystalline sample, and (iii) the inherent nature of the crystallization process does not synthesize an enantiopure, but only an enantioenriched, polycrystalline sample. Although currently the number of crystals that have been analyzed is not sufficient to draw a statistically conclusive picture of the bulk chirality of the samples, these HRTEM data (coupled with the distinct differences detected between R- and S-OSDA synthesized materials) are useful to demonstrate enantioenrichment of a bulk, polycrystalline sample of a molecular sieve.

CD and HRTEM characterizations demonstrate that a given enantiomer of OSDA 2 maintains its respective chirality in the occluded state and is capable of producing an enantioenriched, molecular sieve framework. Although there may exist crystallization processes wherein achiral molecules lead to enantioenrichment of product solids, such as by spontaneous chiral symmetry breaking by self-catalyzed crystallization (33), such systems would not allow for the directed synthesis of specific enantiomers of crystals, as we are able to demonstrate in this work. Thus, it follows that the initial hypothesis by Davis and Lobo (15) that any bulk, polycrystalline sample having enantioenrichment (not obtained solely from chiral

![Fig. 4](image-url) CD spectra for as-made samples of STW using R-, S-, and racemic OSDA 2.

![Fig. 5](image-url) (A) Schematic representation of a sixfold rotation of STW frameworks with different handedness. (B and C) Simulated HRTEM images of the zeolite with right- and left-handedness, respectively. The atomic structure models, where blue and red balls represent Si and O atoms, were overlaid on top of the simulated images. The shift between two images for right-handed and left-handed STW frameworks has the same length but reverse directions. (D and E) Comparison of two HRTEM images with gold nanoparticles as markers. A crystal was tilted from [2110] (d) to [1010] (e) and a shift down was observed, which indicates a space group of P6$_3$22. (F and G) The processed images of D and E after Fourier filtering that only includes spatial frequencies within a particular range to enhance the contrasts of gold nanoparticles. (Scale bars, 5 nm.)
seeds) must be derived from a chiral OSDA with a particular set of properties seems valid. Additionally, this synthesis methodology ensures that all forms (R-, S-, and racemic) are attainable, thus allowing for appropriate controls to be performed.

Adsorption and Catalysis with Enantioenriched STW. The chirality in STW is defined over the specific distance within the helical STW pore structure (Fig. 1) (34). As a consequence, adsorption and catalysis that show ee’s will involve molecules that are of sufficient size to effectively experience the chirality of the structure, yet still be able to pass through the limited size of the pores. As such, we expect that any measured ee’s will be greatly dependent on the selection of molecules used to test for a function. Moreover, the external surfaces of the crystallites may behave nonspecifically, resulting in diminishing ee’s.

To examine whether the enantioenriched STW samples are capable of performing enantioselective functions, catalysis and adsorption experiments were conducted. Epoxide ring-opening reactions on 1,2-epoxyalkanes were selected for catalysis, because they have been shown to take place within molecular sieves. Under typical silica-enriched, fluoride-mediated inorganic reaction conditions the chiral OSDA directs toward the formation of the STW framework. Solid-state NMR and CD characterizations demonstrate that the occluded OSDA remains intact within the STW pores and retains enantipurity. Newly developed 3D HRTEM techniques are applied to determine the chirality of individual crystals. These data show that the samples synthesized using a chiral OSDA are enantioenriched with the chirality of the STW framework. Solid-state NMR and CD characterizations demonstrate that the occluded OSDA remains intact within the STW pores and retains enantipurity. Newly developed 3D HRTEM techniques are applied to determine the chirality of individual crystals. These data show that the samples synthesized using a chiral OSDA are enantioenriched with the chirality of individual crystals. These data show that the samples synthesized using a chiral OSDA are enantioenriched with the chirality of the STW framework.

Summary
We have computationally designed large, rigid, and stable chiral OSDAs to synthesize the two polycrystalline, enantiomorphs of STW (enantioenriched) following the synthesis heuristics specified by Davis and Lobo (15) for chiral molecular sieves. Under typical silica-enriched, fluoride-mediated inorganic reaction conditions the chiral OSDA directs toward the formation of the STW framework. Solid-state NMR and CD characterizations demonstrate that the occluded OSDA remains intact within the STW pores and retains enantipurity. Newly developed 3D HRTEM techniques are applied to determine the chirality of individual crystals. These data show that the samples synthesized using a chiral OSDA are enantioenriched with the chirality of the STW framework. Solid-state NMR and CD characterizations demonstrate that the occluded OSDA remains intact within the STW pores and retains enantipurity. Newly developed 3D HRTEM techniques are applied to determine the chirality of individual crystals. These data show that the samples synthesized using a chiral OSDA are enantioenriched with the chirality of the STW framework.

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<tr>
<th>Substrate</th>
<th>Enantiomeric excess of products using enantioenriched STW</th>
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<tbody>
<tr>
<td></td>
<td>R-STW</td>
</tr>
<tr>
<td>1,2-epoxybutane</td>
<td>a, % 0.14</td>
</tr>
<tr>
<td>1,2-epoxyhexane</td>
<td>a, % 1.29</td>
</tr>
<tr>
<td>1,2-epoxyoctane</td>
<td>a, % 4.13</td>
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Product a is the less-substituted 1-methoxyalkan-2-ol product, and product b is the more-substituted 2-methoxyalkan-1-ol product. The reaction solutions were analyzed after 48 h.
molecular sieve asymmetric catalysis may follow a similar pathway.

Materials and Methods
Reagents, OSDA Synthesis, and Characterization.
A full list of chemicals used in this study and their sources can be found in SI Appendix. The full synthesis procedure for the chiral SDA used in this study as well as the respective 1H and 13C NMR characterization data are described in detail in SI Appendix.

SWT Synthesis. The hydrothermal synthesis procedures are given in SI Appendix, along with a comprehensive list of starting material composition ranges and the resultant products.