Chiral Silica Zeolite

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HPM-1: A Pure Silica Chiral Polymorph with Helical Pores**

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Since the discovery in the late 1980's of the complex structure of zeolite Beta,^[1] comprising a disordered intergrowth of several polymorphs one of them being chiral, the synthesis of zeolite materials with a chiral framework has been a very challenging goal:^[2] It can be argued that, when proper spatial conditions are met, a chiral zeolite could transfer its chirality to the product of a chemical or physical process occurring within its pores. The importance of stereochemical control arise from the fact that chiral molecules typically provoke a different response in living beings depending on their chirality, a phenomenon already recognized by Pasteur over 150 years ago.^[3] Despite an early report claiming a minor enantiomeric excess (ee \sim 5%) in a reaction catalyzed by a zeolite Beta synthesized using a chiral structure directing agent (SDA),^[4] only in the last few years there have been significant advances. The chiral nature of several zeolites had passed unnoticed until very recently, and calorimetric measurements on a natural zeolite indeed suggested a potential for use in processes entailing enantiomorphic molecular discrimination.^[5] However, those chiral zeolites contain non helical pores, and it can be expected that will impose less severe sterical restrictions. Also recently a germanosilicate zeolite,^[6] structure type code STW, with a helical channel was reported but its stability is severely limited by its very high germanium content (Ge/Si>1): amorphization occurs above 673K, and it is likely that even at room temperature ambient humidity could degrade the structure in a large extend.^[7] This is also the case of a mesoporous chiral zeolite-like material, with an -ITV interrupted framework, which can not withstand ambient humidity at room temperature without fast structural collapse.^[8] None of the known chiral zeolites has been synthesized in a pure homochiral form yet. The synthesis of a chiral silica-based, as opposed to germania-based, zeolite with helical channels would be thus highly desirable. Here we present HPM-1,^[9] the first pure silica chiral zeolite with a helical pore, with STW structure type and displaying the high stability of silica-based zeolites.

The new zeolite HPM-1 has been synthesized using achiral 2ethyl-1,3,4-trimethylimidazolium cations as new SDA. Under the synthesis conditions reported for SU-32 using diisopropylamine as SDA and with Ge/Si ratios exceeding 1, there is a strong

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competition for crystallization between SU-32 and SU-15 (an achiral germanosilicate, framework type **SOF**, currently unknown in pure silica composition) and frequently crystals of both phases, plus unreacted GeO₂, are found in the final solid product.^[6] By contrast, no **SOF** or amorphous silica are found under our synthesis conditions.

Calcination of HPM-1 at 973K affords removal of fluoride anions and organic cations, without structural degradation. Furthermore, HPM-1 withstands calcination up to at least 1173K, and it is not affected by ambient humidity. This outstanding stability is due to the SiO₂ composition and the lack of connectivity deffects of HPM-1: after calcination, ²⁹Si MAS NMR reveals four resonances with aproximate 1:2:1:1 relative intensities in the Q⁴ region, with no hints of Si-OH or Si-O⁻ defects in the region around -102 ppm (Figure 1). This essentially perfect SiO₂ network has a micropore volume of 0.22 cm³g⁻¹.

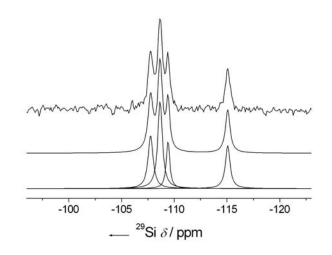


Figure 1. ²⁹Si MAS NMR spectrum of calcined pure silica HPM-1 (top to bottom: experimental, simulated and deconvoluted components). The four $\underline{Si}(OSi)_4$ resonances are assigned to (left to right) Si2, Si1+Si3, Si4 and Si5.

Rietveld refinement of a $P6_122$ model based on the **STW** structure type went smoothly with no restrains on bond lengths or angles in the final stages (see Supporting Information). The refined structure contains Si-O bonds and O-Si-O angles (1.578-1.630Å, average 1.599Å and 106.6-114.0°, average 109.5°, respectively) in the usual ranges for silica zeolites, while the Si-O-Si angles are more spread and tend to be slightly sharp, although still within typical ranges (139.7-170.7°, average 150.6°). The small Si-O-Si angles of calcined HPM-1 are confirmed by inspection of its ²⁹Si MAS NMR spectrum (Figure 1): three of the four distinct resonances observed (4/5 of the total Si) appear at quite low fields (-107.7/-109.4ppm), and are assigned to Si in double four ring units, D4R. The remaining resonance appears at a chemical shift of -115.1ppm, which is in a more typical range for Si(OSi₄) species in tectosilicates and is assigned to the only crystallographic site not

belonging to D4R units. A calculation of the Si-O-Si average angle of each cristallographic Si site using the empirical equation of Thomas et al. and the chemical shifts observed in Figure 1,^[10] is in good agreement with the refinement results although the angles are always sharper when calculated from the ²⁹Si spectrum (Supporting Information).

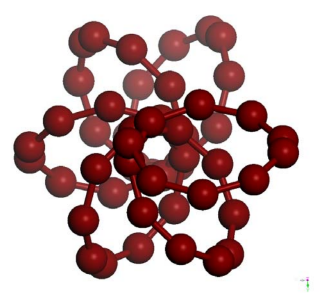


Figure 2. The helical arrangement of the rings of 10 Si-O pairs limiting diffusion through the main pores of HPM-1 in space group $P6_{1}22$ (viewed along [001], O omited for clarity).

The main pore in HPM-1 consists of an helical arrangement of $[4^{6}5^{8}8^{2}10^{2}]$ cavities with access through windows of 10 tetrahedra, 10MR, with minimum and maximum crystallographic free diameters of 5.36 and 5.80 Å, respectively. Figure 2 shows the helical arrangement of 10MR windows in the main pore. Two smaller 8MR pores (3.29-4.56 Å) run perpendicular to the screw axis. From powder XRD data it is not possible to discern the relative proportion of P6₁22 and P6₅22 crystals. Although no chiral reagent was used in the synthesis spontaneous resolution can not be ruled out: Spontaneous chiral symmetry breaking during the crystallization of chiral crystals from achiral units has been reported,^[11] and, according to recent findings, may be more frequent than thought.^[12] This may provide the basis for synthetic strategies toward a purely homochiral STW zeolite.

We would like to suggest one point for reflection regarding the feasibility of silica zeolites. Silica-STW has no "flexibility window",^[13] and its feasibility has been explicitly ruled out on the basis of molecular mechanics calculations.^[14] The existence of a flexibility window has been very convincingly proposed as a useful discriminating parameter for the real existence of hypothetical frameworks,^[15] and it was recently proposed that the existence of a broad flexibility window would increase the likelihood of a good matching between an organic structure-directing agent (SDA) and the framework, thus increasing the probability of a favourable stabilization energy.^[16] Conversely, the lack of such a window would translate into a low probability for finding favourable hostguest interactions unless a particularly suitable SDA could adequately fit into the voids of the rigid framework. On the other hand, molecular mechanics calculations have concluded that, due to the existence of TOT angles that are stressed for a pure silica composition (T=Si) pure silica STW is an unfeasible material (see Table 3 in reference [14]). Here we want to stress that criteria or calculations applied on a calcined zeolite may eventually fail because the synthesized zeolite is in fact a host-guest complex. It has been recently demonstrated that fluoride occluded in pure silica D4R affords a flexibilization of the whole framework by polarization of the Si-O bond.^[17] Thus, rigid-tetrahedra models may fall short in their predictions if host-guest interactions provide an enhanced flexibility relaxing an otherwise strained framework.

In summary, we have synthesized a chiral SiO₂ zeolite, HPM-1, with helical channels that may offer the possibility to test asymmetric processes on zeolites if the next step, obtaining a homochiral solid or selectively blocking homochiral pores,^[18] is realized.

Experimental Section

In a typical synthesis, tetraethylorthosilicate was hydrolyzed in a hydroxide solution of 2-ethyl-1,3,4-trimethylimidazolium (ROH), and the mixture was stirred while allowing ethanol and some water to evaporate. Then, HF (48% aq., Aldrich) was added. The mixture with composition SiO₂:0.5ROH:0.5HF:4.5H₂O was heated to 175°C for 11 days in teflon vessels inside stainless steel bombs tumbled at 60 rpm. The structure of the pure SiO₂ material was refined using laboratory powder diffraction data (Cu K α_1 radiation) and details are given in the Supporting Information.

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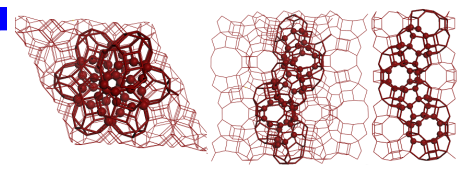
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HPM-1 is a new crystalline microporous polymorph of SiO₂. Its chiral structure ($P6_{1}22$ or $P6_{5}22$, STW zeolite framework type) contains helical pores. The defect-free pure SiO₂ composition, which has been previously considered unfeasible for this structure type, bestow a high thermal and hydrothermal stability upon this new material.