Synthesis of 3D large pore germanosilicate zeolites using imidazolium-based long dications

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ABSTRACT: Introduction of small molar fractions of Ge in a synthesis system otherwise yielding the monodimensional large-pore zeolite MTW, produced three 3D large pore materials as the Ge fraction increased: *BEA, HPM-8 and HPM-7. HPM-8 is a zeolite consisting of an intergrowth of polymorphs D and E of the Beta family with a large predominance (>80%) of polymorph D, according to DIFFax simulations and High Resolution STEM analysis. After calcination, HPM-8 is remarkably stable upon contact with ambient air. HPM-7 likely possesses the POS topology, which is related to polymorph C. The density of double 4-ring units in the crystallized materials increases from *BEA to HPM-8 to HPM-7, i.e., as the Ge fraction increases. Interestingly, molecular mechanics show that in both HPM-7 and HPM-8 the organic dications prefer to sit with their imidazolium rings in close contact with the D4R units as a consequence of a stronger confinement in this position. Indeed, the structure-directing role of these long dications can be explained as a consequence of the appropriate distance match between adjacent D4R units in both zeolite frameworks (and hence the negative charges associated to the occluded fluoride) and the imidazolium rings of these organic dications (and hence the positive charges), which is dictated by the length of the alkyl spacers, thus driving the crystallization pathway towards these particular germanosilicate zeolites.

INTRODUCTION
The large industrial applicability of zeolites relies on their wide variability of properties, which may afford fine tuning for optimal performance in a particular application. This variability is, in turn, the result of the large structural and chemical diversity of this class of materials. In the search for new zeolite structures, scientists use “structure-directing” factors that may include the use of organic structure-directing agents (OSDA), use of fluoride, use of heteroatoms (i.e., atoms other than Si or Al that can enter tetrahedral sites of the framework) with a particular tendency to produce structures with specific characteristics (double four membered ring units (D4R), spiro-5 units...), use of concentrated conditions, etc. Frequently, an OSDA may fail in providing a strong structure-directing action, in which case a so-called “default structure” may crystallize, with the OSDA just serving as a space filler. A default structure is a zeolite that is pretty much stable and does not need a strong structure-direction to crystallize, and as a consequence they are obtained under a wide range of synthesis conditions. It goes without saying that default structures are, by definition, already known. When this
is the case, one can combine several structure-directing factors to force the crystallization to deviate from the default structure and produce another, hopefully new and interesting zeolite.\(^4\) Deionized water until no ammonia smell, then, the dibromide salts were dissolved in the resin-water mixture, which was kept under mild stirring overnight. The hydroxide solution was recovered by washing and filtration with deionized water and finally concentrated using rotoevaporation at 50 °C. The final concentration of the hydroxide solution was determined by titration using phenolphthalein as indicator.

**Synthesis of zeolites**

The starting gels for the synthesis of germanosilicates were prepared by adding GeO\(_2\) (Sigma-Aldrich, 99.998%) in the aqueous solution of the organic dication in a 50 ml plastic beaker, then tetraethylthiosilicate (TEOS, 98%, Sigma-Aldrich) was added. The mixture was magnetically stirred under room temperature overnight and then heated in a water bath until all the ethanol produced from the hydrolysis of TEOS plus some water (monitored by weight) were evaporated. In some cases, extra water was added in order to achieve the desired \(\text{H}_2\text{O}/\text{TO}_2\) ratio, where \(T\) stands for both Si and Ge. Hydrofluoric acid (HF, Sigma-Aldrich, 48%) was added and the mixture was homogenized with a spatula by hand for 15 min. The final gel composition was \((1-x)\text{SiO}_2\cdot x\text{GeO}_2\cdot 0.25\text{R(OH)}_2\cdot 0.5\text{HF}\cdot 5.0\text{H}_2\text{O}\), where \(0 \leq x \leq 0.4\).

The resulting thick gel was transferred into a Teflon liner, which was sealed into a stainless steel autoclave. The hydrothermal reaction was carried out by heating the autoclaves while tumbling (~60 rpm). The reaction temperature used was 150 °C, 175 °C or 190 °C. The autoclaves were taken out at different time intervals and the solid product was recovered, washed with deionized water, and dried at 100 °C. The encapsulated organics could be removed by calcination at 600 °C for 5 h in air.

**Characterization techniques**

The solid products were identified using Powder X-ray diffraction (PXRD) data obtained in a Bruker D8 Advance diffractometer using Cu K\(\alpha\) radiation (\(\lambda = 1.5418\) Å). The amount of organics occluded in the as-made zeolites was determined by CHN elemental analysis using a LECO CHNS-932 analyzer. Thermal gravimetric analysis was carried out in a SDT Q600 TA instrument under air flow (100 ml/min) heating from 25 °C to 1000 °C (with a heating rate of 10 °C/min). Routine field emission scanning electron micrographs (FE-SEM) were collected using Philips XL30 S-FEG. Multinuclear magic angle spinning (MAS) NMR spectroscopy of as-synthesized samples was obtained at room temperature on a Bruker AV-400-WB equipment and the details have been given elsewhere.\(^13\) Fourier Transform infrared spectra were obtained with a Bruker IFS 66/S spectrophotome-
ter using the KBr pellet technique. The $N_2$ adsorption/desorption isotherms were obtained at 77.35 K using a Micromeritics ASAP 2010 equipment.

**Computational details**

Molecular mechanics simulations were performed in order to find the most stable location of 8BDMI within HPM-7 (POS framework) and polymorph D of zeolite beta (as a representation of HPM-8). Simulations were carried out using the Forcite module implemented in Materials Studio software. The geometry of the zeolite structures were kept fixed during the calculations. Molecular structures and the interaction energies of the organic SDA cations with the frameworks were described with the CVFF forcefield. The atomic charge-distribution of the dications was obtained from DFT+D calculations, using the B3LYP hybrid functional and the ESP charge calculation method, setting the total net charge to +2. The positive charge of the organic dications was compensated by the framework by reducing the atomic charge of Si from 0.6 until neutrality. Framework oxygen charges were kept fixed to $-0.3$. For studies with the POS framework (taken from the database of zeolite structures), depending on the case 1$x$1$x$1 or 1$x$1$x$2 supercells have been used as the zeolite model, while for polymorph D of zeolite beta, 2$x$2$x$2 supercells were used. The organic dications were manually docked in the framework in the required position/orientation, and the most stable locations were obtained by simulated annealing. Relative energies are referred to the most stable case, and are given in kcal/mol.

**HRSTEM**

Transmission Electron Microscopy observations were performed in a double corrected JEOL GrandARM 300 which was operated at 300 kV. The microscope was equipped with two separated spherical aberration correctors for both TEM and STEM imaging conditions assuring in each case 0.7 Å point resolution. It was also equipped with a JEOL EDS detector and a Gatan Quantum Energy Filter (GIF) for spectroscopic analyses. For every experiment presented here, the microscope was operated in Scanning Transmission Electron Microscopy (STEM) mode, collecting the electrons simultaneously using an Annular Dark Field (ADF) detector and an Annular Bright field (ABF) detector. Prior to observation, the materials were deeply crushed using mortar and pestle for up to 5 minutes with the intention of breaking the sample into very thin crystals which allowed obtaining high resolution images. The finely divided powder was dispersed in ethanol and sonicated and few drops of the suspension were deposited onto holey carbon copper microgrids. Before any electron microscopy (EM) experiments, the corrector was aligned using amorphous carbon assuring that the maximum spatial resolution could be achieved. Due to the high sensitivity of zeolites under the electron beam, STEM low dose conditions were necessary in order to acquire high resolution images with atomic-resolution. For this purpose, the exposure time did not exceed 8 seconds for the high-resolution data, with an image size of 1024×1024 pixels.

High resolution scanning electron microscopy (SEM) was performed in a JEOL 7800 Prime. In order to achieve high-resolution images we use the GBSH (gentle beam super high) mode. In this mode, the gun voltage applied was 5.5 keV with a decelerating voltage -5 keV on the sample holder, giving a final landing energy of 0.5 keV on the samples.

**RESULTS AND DISCUSSION**

**Zeolite synthesis**

Table S1 summarizes the synthesis conditions and products obtained using 8BDMI and 10BDMI as SDAs under varying germanium molar fractions ($Ge_f = Ge/(Si+Ge)$). Additionally, the phase selectivity is schematically shown in Figure 1. The synthesis of zeolites under pure silica compositions was extensively reported recently for these dications of varying length: at high concentration 8BDMI produced zeolite *BEA that subsequently transformed into MTW, whereas 10BDMI yielded MWW that transformed also into MTW. Lower concentrations directly yielded MTW in both cases. When Ge was introduced into the gel using 8BDMI as SDA three 3D large pore zeolites, *BEA, HPM-8 and HPM-7, were obtained with the increase of $Ge_f$. The syntheses trials using 10BDMI with $Ge_f$ at 0.1 and 0.25 also produced HPM-8 and HPM-7, respectively.
As it is well known, *BEA is an intergrowth of two closely related polymorphs: polymorph A and polymorph B.\textsuperscript{16} As for HPM-8, it is an intergrowth of polymorphs D and E of the same family, which are the remaining two hypothetical polymorphs of zeolite beta predicted by Burton et al.\textsuperscript{17} and successfully synthesized later by Yu et al.\textsuperscript{14} as SU-78, an intergrowth of both polymorphs in a close to 1:1 ratio. All the polymorphs of the Beta family contain a 3D system of interconnected large 12MR pores. HPM-8 appears to be much closer to polymorph D than SU-78 (see below). We observed two types of XRD patterns for HPM-8 (Figure 2): The most frequent one presented somewhat broader peaks and a merge of the two central most intense peaks. This difference initially suggested HPM-8 could present different degrees of disorder, however this does not appear to be the case (see below). In the case of HPM-7, a phase with a much similar XRD pattern was recently reported and named PKU-16 (POS), which is structurally related to polymorph C of Beta by rotating half of the four-rings in double mtw units resulting in a unique three dimensional 11×11×12 channel system.\textsuperscript{18}

The synthesis temperature played a vital role for the crystallization of HPM-7 as demonstrated in Figure S3. Ill-crystallized HPM-7 with amorphous phase was obtained at 150 °C after 2 days, while products with well-resolved patterns were obtained at 175 °C and 190 °C under the same synthesis time. Prolonging the crystallization time rendered better-resolved patterns for all three synthesis batches. However, the reflections are still very broad for the products at 150 °C, suggesting that the crystallization of HPM-7 may have a relatively high activation energy.

![Figure 2](image)

Figure 2. The XRD patterns of as-made HPM-8 synthesized with 8BDMI and Ge\textsubscript{f} = 0.15 in comparison with the simulated pattern of SU-78B (bottom trace). Two HPM-8 samples with markedly differently resolved patterns are shown.

The structure-directing ability of 8BDMI and 10BDMI towards various beta-related structures at different Ge\textsubscript{f} suggests a significant altering role of Ge on the T-O-T connections. It has been long recognized that Ge prefers to locate in the D4R or D3R units especially in fluoride medium.\textsuperscript{6, 19-21} Indeed, in going from *BEA (no D4R units in polymorphs A or B) to HPM-8 (one D4R in a unit cell of 32 T atoms) and then to HPM-7 (four D4R per cell of 64 T atoms) the density of D4R units increases. Similar phenomenon of phase change along germanium fraction variation have been observed previously in other germanosilicate zeolites synthesis.\textsuperscript{3, 22} On the other hand, the selectivity of the long dications used in this work towards Beta-related zeolites contrasts with the tendency of shorter dications to produce STW, which must be clearly due to the size of 8BDMI and 10BDMI being too large to accommodate the imidazolium moieties into two consecutive stw cages.\textsuperscript{1}

Zeolite characterization

The XRD patterns of HPM-8 in comparison with the simulated pattern of polymorph D of the Beta family (i.e., SU-78B) are shown in Figure 2. The peak intensity increased after calcination for both HPM-8 samples, which is especially obvious at the low angle region, and also, some additional peaks appeared in the region between 12° and 16° (Figure 3). The as-made HPM-8 was indexed in the monoclinic system using DICVOL 6,\textsuperscript{23} with unit cell parameters a' = c = 13.9090 Å, b' = b = 12.6815 Å, c' = a = 12.5224 Å, β = 107.768° and cell volume of 2103.44 Å\textsuperscript{3}, which are close to those previously reported for SU-78. SU-78 was synthesized with two SDAs, namely N-ethyl-N-methyl-dicyclohexylammonium hydroxide and N,N-dimethyl-dicyclohexylammonium hydroxide, which are mono-cationic organics, in contrast with our imidazolium-based dications. Interestingly, HPM-8 shows much higher stability at room temperature against moisture than SU-78, which collapsed after being exposed in air for 1 h after calcination.\textsuperscript{14} The XRD patterns of calcined HPM-8 after exposure to ambient air for different periods of time demonstrates its improved stability (Figure 3), which could give HPM-8 a high potential for catalytic applicability. The stability difference between our HPM-8 and SU-78 may be due to the lower Ge content in HPM-8 (Ge\textsubscript{f} = 0.1-0.15 compared to 0.3). The chemical analysis of the HPM-8 material has been performed by means of X-ray Dispersive Spectroscopy using both Transmission and Scanning electron microscopes using at least 50 different crystallites. The obtained Si/Ge ratio was 7.3, resulting in a Ge\textsubscript{f} = 0.145 for a sample synthesized from a gel with Ge\textsubscript{f} = 0.1, confirming the much lower Ge value of HPM-8 compared to SU-78.
Our as-made HPM-7 exhibits a similar pattern to the simulated pattern of PKU-16 as shown in Figure 4. Different from PKU-16, synthesized using the relatively small neutral amine 4-methylaminopyridine (DMAP) as the SDA, HPM-7 was synthesized with long imidazolium-based dications. The corresponding calcined HPM-7 samples (Figure S4) show increased intensities at the low angle region, a normal phenomenon for the calcined zeolites indicating the release of the occluded organics. Additionally, new peaks showed up at about 13° and 15°. Unfortunately, the stability of HPM-7 upon calcination and subsequent exposure to ambient air is significantly lower than that of HPM-8, which we attribute to its increased Ge; while freshly calcined HPM-7 still presents a good crystallinity, exposure to ambient air significantly destroys the framework in a relatively short time (Figure S4).

Figure 3. The XRD patterns of calcined HPM-8 (from bottom to top): freshly calcined, stored in closed eppendorf for 3 months, after 3 h in open air and after 3 d in open air.
Table 1. Chemical analysis results of as-made zeolites.

<table>
<thead>
<tr>
<th>SDA</th>
<th>Phase</th>
<th>C(%)</th>
<th>H(%)</th>
<th>N(%)</th>
<th>C/N$^a$</th>
<th>H/N$^b$</th>
<th>TG$^c$</th>
<th>empirical composition$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>8BDMI</td>
<td>*BEA</td>
<td>11.8</td>
<td>1.97</td>
<td>2.78</td>
<td>4.95(4.5)</td>
<td>9.85(8.0)</td>
<td>80.47(82.8)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
<tr>
<td>8BDMI</td>
<td>*BEA</td>
<td>12.57</td>
<td>1.95</td>
<td>2.78</td>
<td>5.27(4.5)</td>
<td>9.75(8.0)</td>
<td>79.1(82.8)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
<tr>
<td>8BDMI</td>
<td>8HPM-8</td>
<td>12.31</td>
<td>1.91</td>
<td>2.90</td>
<td>4.95(4.5)</td>
<td>9.15(8.0)</td>
<td>80.0(82.8)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
<tr>
<td>8BDMI</td>
<td>HPM-8</td>
<td>11.36</td>
<td>1.81</td>
<td>2.85</td>
<td>4.65(4.5)</td>
<td>8.83(8.0)</td>
<td>81.2(82.5)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
<tr>
<td>8BDMI</td>
<td>HPM-7</td>
<td>11.04</td>
<td>1.76</td>
<td>2.43</td>
<td>5.0(4.5)</td>
<td>10.06(8.0)</td>
<td>82.6(84.9)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
<tr>
<td>10BDMI</td>
<td>8HPM-8</td>
<td>12.43</td>
<td>1.99</td>
<td>2.76</td>
<td>5.25(5.0)</td>
<td>10.02(9.0)</td>
<td>80.6(81.6)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
<tr>
<td>10BDMI</td>
<td>HPM-7</td>
<td>12.22</td>
<td>1.90</td>
<td>2.62</td>
<td>5.44(5.0)</td>
<td>10.08(9.0)</td>
<td>81.4(82.6)</td>
<td>[C$<em>{10}$H$</em>{12}$N$<em>{4}$F$</em>{6}$]$<em>{2.36}$[SiO$</em>{4}$]$<em>{4.08}$[H$</em>{2}$O]$_{1.67}$</td>
</tr>
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</table>

$^a$h Molar ratios, the theoretical values are given in parentheses. $^b$Solid residues after thermal analysis to 1000 °C (the amount in parentheses stands for the TO2 content in a unit cell given in the last column). $^c$SDA calculated from the N percent assuming the organics are intact, and are charge-balanced by F$^-$ ion, water calculated from the excess H and TO2 is the rest. $^d$The sample with the merged two central most intense peaks.

Table 1 lists the chemical composition details of the as-made SDA-zeolites, showing sample variability. For all samples synthesized, there are non-negligible C/N ratio deviations from the theoretical values, suggesting partial decomposition might have occurred during the crystallization. There are about 2.5 cationic-like species in one unit cell of the beta-related zeolites calculated based on the N weight percentage. In order to investigate the existing form of the SDAs occluded in the as-made zeolites, solid state $^{13}$C MAS NMR spectra were measured, as shown in Figure 5. The spectra show resonances close to the expected ranges for each C atom, suggesting the organics are essentially intact in all the samples. However, several resonances overlapped, like the ones centered around 122.6 ppm, the C atoms at the b and c positions on the imidazolium ring, and around 29.7 ppm, the C atoms at g, h, i and j positions on the methylene chain (see Figure 5 for the C labeling).

Figure 4. $^{13}$C$^{-1}$H CPMAS NMR spectra of as-made zeolites using 8BDMI and 10BDMI as SDAs (from bottom to top): 8BDMI-HPM-7-Ge0.25, 10BDMI-HPM-8-Ge0.10, 8BDMI-HPM-7-Ge0.25 and 8BDMI-HPM-8-Ge0.15.

Thermogravimetric analysis (TGA) combined with derivative thermogravimetry (DTG) of the samples are shown in Figure S5 and Figure S6. For the samples synthesized with 8BDMI, the amount of organics occluded in the zeolite showed a decreased trend with the increase of Ge, in the starting gel composition (Table 1 and Figure S5), which has also been observed in our previously reported work. The 10BDMI derived samples also followed the above trend despite the limited Ge range tested (Figure S6). This trend is likely due at least in part to the increased fraction of the heavier Ge.

Figure 6 (bottom three traces) shows the $^{19}$F MAS NMR spectra of several as-made HPM-7 zeolites, all showing a single resonance in the -7 to -8 ppm range, which shall be ascribed to F occluded in D4R units containing Ge pairs with no Ge having more than 2 Ge neighbors. The spectrum of HPM-7 sharply contrasts with that previously reported for PKU-16, in which F$^-$ not only shows a resonance at -7.6 ppm assigned to F occluded in the D4Rs but also presents signals at -7.2 ppm assigned to F in the lau units and at -15.4 ppm assigned to “dissociative F$^-$”. These large differences in the $^{19}$F spectra of HPM-7 and PKU-16 may reveal different interactions between the organic cation, the fluoride anion and the zeolite framework.
The $^{19}$F spectra of the HPM-8 samples are quite more complicated (Figure 6, top three traces). In addition to a main resonance at similar chemical shift regions, there are three resonances around -50, -61 and -72 ppm, which we attribute to fluoride occluded in other small cavities of the zeolite, such as [435] (similarly to the case of pure silica zeolite Beta, with resonances at -58, -65 and -70 ppm assigned to F in that cavity). These signals are more prominent for the sample with the lowest Ge fraction. An additional resonance around -20 ppm is assigned to F occluded in D4R containing Ge but without Ge-O-Ge pairs. This resonance is also more prominent in the sample with lower Ge content, for which still another resonance around -40ppm, assigned to F in pure silica D4R, is also noticeable. For all the samples of HPM-7 and HPM-8 there appears to be a very small resonance at a rather unusual low field (2.3 ppm) which we would tentatively assign to F occluded in defective D4R units, i.e., D4R with one T-O-T broken bond, see text).

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Structural Characterization of HPM-8

HRSTEM

Spherical aberration corrected (Cs-corrected) Scanning Transmission Electron Microscopy (STEM) using both Annular Dark Field and Annular Bright Field detectors were used to characterize the materials obtained. Low-magnification data of HPM-8 reveals well-defined faceted particles with two different morphologies clearly differentiated. Rhombohedral shaped crystals of 100 nm up to 1µm were identified, see Figure 7a. Such well-defined facets suggest a good crystallinity of the material. Figure 7b displays a closer observation where the pore arrangements showing multiple defects can be clearly appreciated. The Fourier Diffractogram (FD) obtained from two different regions marked as dashed A and B squares are presented. The FD coming from A shows sharp diffraction spots attributed to the single crystal nature of that region, meanwhile, the FD marked as B contains structural defects represented by the diffuse spots that are attributed to the presence of a faulted structure. Figure 7c displays the edge of the crystal in which the two distinct domains can be distinguished and attributed to polymorphs D and E. d) Atomic-resolution data, with the model of the polymorph D overlapped.
atoms are represented as blue and red spheres, respectively, corroborating the perfect matching between the model of polymorph D and the experimental data acquired. The two polymorphs can be described as two stacking sequences, in which the 12 MR of the polymorph E follows a zigzag trend along the c axis (see Figure 8a), where every other layer is shifted approximately 1/3 of a, 4 Å, (i.e., shifts are 0, 1/3, 0, 1/3...) whereas for polymorph D successive layers are always shifted by 1/3a (i.e., shifts are 0, 1/3, 2/3, 0, 1/3...), as seen in Figure 8b.

Despite the numerous defects that arise as a consequence of the coexistence of both polymorphs, when the crystals are observed perpendicular to this projection (this is along [100]) the materials exhibit a single crystal nature where the defects are not observable. This is due to the fact that both polymorphs D and E share a common projection along [100] (which is also common to polymorph C). In this case, the low-magnification micrograph reveals the morphology along this projection of HMP-8, elongated crystals with rectangular shapes which generally terminate with truncated ends corresponding to the (011) and (010) facets, see Figure 9a. High-resolution studies by means of C$_s$-corrected STEM-ADF were also performed along this zone axis. Figure 9b corresponds to the crystal edge that allows the observation of the pore distribution as well as extracting the FD diffractogram, Figure 9c. The FD can be indexed assuming Pcmn or P2/m obtaining a lattice constant b = 12.9 Å and c = 13.47 Å. The atomic resolution data is depicted in Figure 9d, where all T atoms can be identified corroborating that along this projection the framework is identical as BEC.

Thus, we can observe in HMP-8 defect-free projections that are shared by polymorphs C, D and E as well as faulted projections in which polymorphs A or E are intergrown with polymorphs B or D respectively, as expected for an intergrowth between polymorphs D and E. To unambiguously determine the HMP-8 structure three dimensional diffraction electron tomography (3D-EDT) was collected from several crystals obtaining in every case the same result. The typical data acquired for this material is presented in figure S7. Figure S7a and S7b depict two different orientations related by 90 ° of the same crystal which correspond to the [100] and the [010] projections. The shape of the crystals of these two images are in agreement with the images presented in figures 7 and 8 that correspond to HPM-8. The electron diffraction analysis by means of 3D-EDT, figures S7c and S7d shows two electron diffraction patterns extracted from the reconstructed three dimensional diffraction data along the a* and b* orientations. From these two ED patterns it is clear the absence of structural defects when the HPM-8 is along the [100] orientation, that, however appear when the crystal was tilted evidenced by the existence of the diffuse spots, as streaks, on the [010] orientation. Lattice cell parameters obtained by this method were a = 12.72Å; b = 12.89Å; c = 14.22Å; α = 89.89°; β = 107.02°; γ = 90.14° for a space group P2/m in good agreement with the predominant polymorph D.

Thus, we conclude that HMP-8 is a highly crystalline material formed by an intergrowth of polymorphs D and E where, after observations of over 15 different crystallites, polymorph D is the predominant phase as evidenced in Figure 7.

Powder XRD
In order to further characterize the structure of HPM-8, we simulated the powder XRD data of intergrowths between polymorphs D and E of the Beta family (the recently reported SU-78B and SU-78A, respectively) using DIFFaX. The layers were obtained from the reported SU-78A structure, but centering the so-called “helicopter” in the center of the cell. The length of the cell edges was deduced from the experimental pattern of calcined HPM-8 (reflections that are invariant when polymorphs D and E intergrow were used to refine the cell). We checked that only 2 layers (related by a 90° rotation) are necessary to build all the Beta family of polymorphs (A to E). Both layers need to strictly alternate, while they may be or not shifted by ±1/3 of the corresponding cell edge along x or y, depending on the polymorph. Along z a shift of 1/4 of the c edge needs to be applied to each successive layer. Figure 10 shows the effect of intergrowing polymorphs D and E on the XRD pattern and compares the patterns with the simulation for the pure polymorphs and the experimental pattern of HPM-8.

The most visible change in the pattern along this intergrowth refers to reflection (101) of the orthorhombic polymorph E (SU-78A), at around 7.8°, which splits into two broader reflections that are more separated as the prevalence of polymorph D (SU-78B) increases, finally becoming reflections (100) and (10-1) of this monoclinic polymorph (at around 7.4 and 8.3° in the pure polymorph). Comparison with the experimental pattern suggests that, contrarily to SU-78, HPM-8 is significantly enriched (around 80%) in polymorph D. Further comparison in the 76-86% percent range suggests HPM-8 is an intergrowth of polymorphs D and E in around an 82:18 ratio (Figure S8).

When we compare HPM-8 samples displaying less well resolved patterns with the simulations, the positions and general features of the peaks can be reproduced well with an intergrowth containing 84% D, see Figure S9. The cell in this case was a little bit smaller (a = 12.5645Å, b = 12.6787Å, c = 26.51984Å), Thus, the presence of a single central reflection rather than two sharper lines in HPM-8 turned out to be simply a consequence of the change in cell parameters and increase in peak broadness, which made the two reflections to merge into a broad and asymmetric one.

Figure 8. DIFFaX simulated powder XRD patterns of intergrowths of polymorphs E and D of the Beta family (color traces, with 10% increments). The bottom and top traces correspond to 99% pure polymorphs E and D, respectively, and are compared to a simulation of the corresponding polymorph obtained with the program VESTA. The experimental pattern for a HPM-8 sample is shown (noisy black trace) close to the 80% polymorph D simulation. The layer used in DIFFaX had a cell of 12.5465Å, 12.7086Å, 26.539Å, 90.0°. Lambda: 1.5418Å.

We note that all the reflections in the experimental pattern of HPM-8 are somehow broader than those in the simulations. Tweaking with peak broadening parameters in DIFFaX (U, V, W and Gaussian-Lorentzian mixing parameter) did not solve this difference: while all experimental peaks are broad, this is by far more apparent in the peaks that are affected by the intergrowth (while the U, V, W and mixing parameters affect all the peaks). Playing with the number of stacked layers or with the widths of the layers did not help either because the effect on the pattern was relatively small. This suggested that additional disorder may occur in HPM-8. Given that C-corrected STEM data shows the coexistence of projections that are compatible with polymorphs A, B and C (in addition to D and E), we introduced variable proportions of these polymorphs into the DE intergrowth simulation, which did not result in an improvement in the match (see Figures S10-S13). However, since an exhaustive simulation of the intergrowth of 3 or 4 polymorphs covering all range of possible relative amounts would be extremely time consuming, we cannot discard this possibility and, in fact, we have observed by HRSTEM a single occasion in which projections compatible with C (or D or E) intergrew with projections compatible with A and B (or D and E), see Figure S14. While this fact by itself does not prove or disprove the presence of additional polymorphs in HPM-8, we think there may be some additional disorder in this system that we can not specify. Anyway, we can safely conclude that HPM-8 is basically an intergrowth of polymorphs D and E much enriched in polymorph D (around 80%) compared to SU-78 (50%). Another difference is that HPM-8 is significantly richer in Si, as deduced not only from the lower Ge but centering the so-called “helicopter” in the center of the cell. The length of the cell edges were obtained from the reported SU-78A structure, but centering the so-called “helicopter” in the center of the cell. The length of the cell edges was deduced from the experimental pattern of calcined HPM-8 (reflections that are invariant when polymorphs D and E intergrow were used to refine the cell). We checked that only 2 layers (related by a 90° rotation) are necessary to build all the Beta family of polymorphs (A to E). Both layers need to strictly alternate, while they may be or not shifted by ±1/3 of the corresponding cell edge along x or y, depending on the polymorph. Along z a shift of 1/4 of the c edge needs to be applied to each successive layer. Figure 10 shows the effect of intergrowing polymorphs D and E on the XRD pattern and compares the patterns with the simulation for the pure polymorphs and the experimental pattern of HPM-8.

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Computational results

In order to investigate the structure-directing effect of these long dications, we finally analyzed by molecular mechanics simulations their fit with the zeolite void volume. The POS framework (corresponding to HPM-7) contains three channel systems, one 12MR channel running parallel to c, and two 11MR channels running perpendicular to c, and perpendicular to each other, with an elliptical shape and a slightly smaller cross-section. We
first introduced manually the 8BDMI dications in the two types of POS channel systems, the 12MR and 11MR channels (the two 11MR are topologically equivalent), and calculated the most stable location by simulated annealing; as we did before with STW, the OSDA dications were loaded in the two possible orientations of the imidazolium rings, with methyl groups of the two rings pointing to the same side (‘ss’) or to opposite sides (‘ops’). In order to have enough space to accommodate the long 8BDMI species along the 12MR channel, 1×1×2 supercell systems were used (‘a’ and ‘b’ directions are large enough as to host one dication while preventing interactions between image cells). Two possible locations for 8BDMI are possible within the 11MR channels, with the imidazolium rings in the intersections with the 12MR (labelled as ‘intersections’), or with the alkyl spacer sited in those intersections, and the imidazolium rings close to the D4Rs (labelled as ‘D4R’). The relative energies are given in Table S3, and images of some of the models (those with ‘ss’ orientation) are shown in Figure S16.

From the energy results, it is clear that the 8BDMI dication fits better within the 11MR channels, probably because of a stronger confinement and hence higher interaction with the channel walls due to their smaller cross-section; worth is noting that this type of dications typically structure-direct towards 10MR channels, like those in MFI or STW. On the other hand, the location of the dications with the imidazolium rings sited close to the D4Rs is rather more stable (Figure S16-bottom), again due to a stronger confinement with both D4R units surrounding the imidazolium rings than in the more open channel intersections (Figure S16-middle).

In the next model, two 8BDMI dications were included in the most stable location (11MR channels). As there are two of these channels per POS u.c. (perpendicular to each other), we loaded one dication on each 11MR channel, giving a packing of 2 SDAs per u.c. 8BDMI cations were loaded in the most stable site, with the imidazolium rings close to the D4Rs, and in the two possible conformations (‘ss’ and ‘ops’). In addition, there are two possible arrangements with the 2 dications in the 11MR channels: in the first (referred as ‘A’, Figure S17-left), one intersection with the 12MR channel is filled with the spacers of the 2 dications, while leaving free the other 12MR channel, while in the other (referred as ‘B’, Figure S17-right), the 8BDMI dications are shifted and hence the two 12MR channels are only partially filled with the spacers; the resulting relative energies are given in Table S4. The most stable arrangement corresponds to the ‘A’ configuration, where one of the 12MR channels is occupied by the spacers of the two dications, and the other remains empty (see Figure S17-left); the most stable orientation of the imidazolium rings in this case is in the ‘ops’ orientation (see Table S4), giving a rather symmetric arrangement of the dications which nicely fits with the POS topology. It should be remarked that this configuration enables the incorporation of further dications within the empty 12MR channel (see void volume surface in Figure S17-left). In the ‘B’ configuration, which is less stable (Table S4), each 12MR channel is partially filled with the organic spacers, and hence does not allow for additional dications loading (see Figure S17-right).

As previously mentioned, the ‘A’ organic arrangement allows for the inclusion of one additional 8BDMI dication along the empty 12MR channel. In this case, 1×1×2 supercell systems are required in order to accommodate the 8BDMI dication aligned with ‘c’ direction (12MR channel); hence, 5 dications were loaded in the 1×1×2 supercell as follows: 4 (2 per unit cell) in the 11MR channels in the A-ops configuration, and one additional dication aligned with the empty 12MR channel, both in ‘ss’ and ‘ops’ orientations. Again the most stable situation was found for the system where the 8BDMI dication in the 12MR channel was in the ‘ops’ orientation. Figure S18 shows the final arrangement of the 8BDMI dications, giving a packing value of 5/2 = 2.5 8BDMI/u.c., in close agreement with the value found experimentally.

Calculations with the longer 10BDMI dication within the POS framework (in the ‘A’ configuration) gave a similar location for this SDA (Figure S19). We note that in both cases (8BDMI and 10BDMI dications, Figures S17 and S19, respectively), the imidazolium rings always site very close to a D4R unit, which may be possible for both dications due to the inherent flexibility of the spacer.

We then studied the location of the 8BDMI dications within a hypothetical pure polymorph D of zeolite Beta (as a representation of HPM-8). In this case, one 8BDMI dication was loaded in a 2×2×2 supercell in three different orientations, with the molecular axis aligned with the [100], [010] or [001] channels, and again in ‘ss’ or ‘ops’ conformations of the imidazolium rings. Table S5 reports the final relative energies, and Figure S20 displays the final configuration for the ‘ops’ conformations. The most stable location for 8BDMI is when they site aligned with the [001] channels (Figure S20-bottom); the next more stable orientation is with 8BDMI aligned with the [010] channels, while the least stable is aligned with the [100] channels. We note that in the two most stable orientations (along [010] and [001], Table S5), the 8BDMI dications site with the two imidazolium rings very close to D4R units (Figure S20-middle and bottom), the same as occurred in the POS framework.
It is interesting to remark the tendency of these dications to locate their imidazolium rings close to D4R units in both frameworks (POS and beta polymorph D). We note here that this conclusion arises from a calculation without occluded fluoride so it must be topological in nature, and that it would be reinforced when considering electrostatic interactions between fluoride-containing D4R and the charged imidazolium rings. From a chemical point of view, fluoride and imidazolium rings are expected to be closely associated during the crystallization process. In fact, it seems that the efficient structure-directing effect of these long dications towards these particular frameworks is closely associated to the adequate distance match between adjacent D4R units in both frameworks and the two imidazolium rings in these long dications. Adjacent D4Rs in the POS framework are separated by 13.77 Å along the 11MR channels (the most stable location for 8BDMI, Table S4) and by 11.69 Å along the 12MR channels (the less stable position for 8BDMI), while those inter-D4R distances in polymorph D of zeolite Beta are 12.85 Å along the [010] channels and 13.98 Å along the [001] channels, the two most stable locations for 8BDMI (Table S5). In order to monitor such geometric match, we compared these inter-D4R distances with the intramolecular distance between the two imidazolium rings in the two dications. In order to have averaged values, we ran 500 ps of Molecular Dynamics of the dications in vacuum, and plot the Radial Distribution Functions of these inter-ring distances (Figure 11). We can observe that the range of distances between imidazolium rings falls close to the distance between adjacent D4R units in both frameworks, especially in the 11MR channel direction in POS (13.77 Å) and in the [001] channels in polymorph D (13.98 Å), for both cations. Indeed, this geometric relationship is even more appealing when considering the distance between N atoms (Figure S21), which should be an indication of the separation between the positive charges; two limits are defined, the lowest one is given by the distance between the internal N atoms (N<sub>int</sub>, N bonded to the alkyl spacer), and the highest one is given by the maximum separation of charges as represented by the distance between external N atoms (N<sub>ext</sub>, N bonded to methyl groups). Again we observe a perfect distance match between external N atoms of 8BDMI (solid blue lines in Figure S28) and the inter-D4R separation in the POS framework along the 11MR channels (and this can explain why these SDAs prefer to site along the 11MR channels), and also with the inter-D4R separation in beta polymorph D along the [001] channels (and also is not far from the separation along [010]). In the case of 10BDMI, the distance between external N atoms (solid red line) is longer (~16.7 Å) than the inter-D4R separation (13.8-14 Å), but in contrast the distance between the internal N atoms (dashed red line) falls right in the inter-D4R separation of both POS and beta polymorph D. In any case, we note that the positive charge of imidazolium rings in principle should be delocalized around the ring, but it is possible that the charge might localize on a particular N atom upon interaction with the D4R units and the occluded fluoride.

**CONCLUSION**

Introduction of small amounts of Ge in a synthesis using dimethylimidazolium-based long dications that would otherwise yield the well-known trite MTW produces interesting changes in the phase selectivity of the crystallization. As the Ge content increases, *BEA, HPM-8 and HPM-7 are obtained. The trend corresponds to an increase in the density of D4R. *BEA has no D4R (although it is difficult to discard the occurrence in this intergrown material of small proportions of polymorphs C, D and E, all containing D4R), while HPM-8 and HPM-7 contain 2 and 4 D4R, respectively, per 64 T atoms.

HPM-8 is an intergrowth of polymorphs D and E of the zeolite Beta family with a large predominance (over 80%) of polymorph D, as deduced from DIFFaX simulations and HRTEM. Compared to the recently reported SU-78, which contains an almost 1:1 proportion of both D and E polymorphs, HPM-8 is significantly more stable due to its much lower Ge content (Ge<sub>f</sub> = 0.1-0.15 compared to 0.3). On the other hand, HPM-7 is likely isostructural to POS in view of its PXRD data.

Computer simulations confirm that 8BDMI can accommodate in the POS channels in a high loading of up to 2.5 dications per unit cell and show that the most favorable host-guest interaction is within the 11MR channels (2 dications per cell), with the imidazolium moieties in between D4R units providing a strong confinement, and with an additional loading in the 12MR (up to 0.5 dications per cell) being possible. Interestingly, also in...
the framework of polymorph D of zeolite beta do 8BDMI dications site with imidazolium rings close to D4R units. In fact, our results suggest that the efficiency of these long organic dications to direct the crystallization of these frameworks is closely associated to the appropriate distance match between the positive charges of the imidazolium rings in 8BDMI and 10BDMI dications, determined by the length of the alkyl spacers, and adjacent D4R units in these germanosilicate zeolites.

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