

Cite this: *Dalton Trans.*, 2023, **52**, 15697

## 2-Isopropyl-1,3-dimethylimidazolium as a versatile structure-directing agent in the synthesis of zeolites†

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An organic cation lacking specificity in its structure-directing action offers the possibility, through the screening of other structure-directing parameters, to synthesize a variety of zeolites. In this work we show that the organic structure-directing agent 2-isopropyl-1,3-dimethylimidazolium (2iPr13DMI) can produce up to seven different zeolite phases depending on water concentration, the presence of inorganic impurities, crystallization temperature and time, and germanium molar fraction. The obtained phases are very different in terms of pore system, connectivity of the zeolite structure and structural units. At the pure SiO<sub>2</sub> side, ZSM-12 and SSZ-35 dominate, with ZSM-12 being favored by the presence of potassium impurities and by less concentrated conditions. The introduction of Ge at low levels favors SSZ-35 over ZSM-12 and as the Ge fraction increases it successively affords **CSV**, **-CLO** and two distinct **UOS** zeolites, HPM-11 and HPM-6. These two zeolites have the same topology but distinct chemical compositions and display powder X-ray diffraction patterns that are much different from each other and from that of as-synthesized IM-16 (**UOS** reference material). They also show different symmetry at 96 K. Rietveld refinements of the three as-made **UOS** materials mentioned are provided. HPM-6 and HPM-11 are produced in distinct, non-adjacent crystallization fields. The frequent cocrystallization of the chiral **STW** zeolite, however, did not afford its synthesis as a pure phase. Molecular mechanics simulations of the location of the organic cation and host–guest interactions fail to explain the observed trends, but also considering the intrinsic stability of the zeolites and the effect of germanium help to rationalize the results. The study is completed by DFT calculations of the NMR chemical shifts of <sup>13</sup>C in **UOS** (helping to understand splittings in the spectrum) and <sup>19</sup>F in **CSV** (supporting the location of fluoride inside the new [4<sup>4</sup>5<sup>2</sup>], which is an incomplete double 4-ring).

Received 28th July 2023,  
Accepted 17th September 2023

DOI: 10.1039/d3dt02414h

rsc.li/dalton

## Introduction

Zeolites are microporous crystalline materials with interesting commercial applications that critically depend on their structures and compositions.<sup>5</sup> The crystalline structure enables

control of access in and out of the intrazeolitic pore space and, in this respect, pores ranging from small,<sup>6</sup> to medium,<sup>7</sup> to large<sup>8</sup> and extralarge<sup>9</sup> sizes all find interesting applications. On the other hand, the zeolite composition determines properties such as chemical, thermal and hydrothermal stability,

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† Electronic supplementary information (ESI) available: 3D ED and supplementary PXRD patterns, Rietveld refinement plots, characterization data, supplementary calculation figures, detailed synthesis results, and crystallographic tables. Structures for as-synthesized IM-16, HPM-6 (Ge<sub>f</sub> = 1), HPM-6 (Ge<sub>f</sub> = 0.88) and HPM-11 (Ge<sub>f</sub> = 0.58) were previously published as CCDC Communications (ref. 1–4). See DOI: <https://doi.org/10.1039/d3dt02414h>

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cation exchange,<sup>10</sup> acidity,<sup>11</sup> redox activity<sup>12</sup> and hydrophilicity vs. organophilicity characteristics,<sup>13</sup> which may prove valuable in different applications.<sup>14</sup> The wide variability in structures and compositions of zeolites offers opportunities to select and fine tune materials to maximize the performance in a given process. This has led to huge research efforts to control the structure, composition and textural properties of zeolites. Organic cations, usually referred to as organic structure-directing agents (OSDAs), are among the most critical factors allowing the synthesis of a particular zeolite. However, they may exert a structuring/crystallizing role with varying degrees of specificity, from a close to key-and-lock templating effect all the way down to mere pore filling.<sup>15</sup> The latter typically produces “default structures”,<sup>16</sup> *i.e.*, zeolites that are rather stable by themselves and that, thus, are not very in need of additional stabilization by a host–guest interaction with the OSDA. As a consequence, default materials can be obtained using a very large number of different OSDAs. By contrast, OSDAs that truly act as templates may afford unique materials.

When an OSDA is unspecific in its structure-directing action, other synthesis parameters may come into play to control the output of crystallization.<sup>17</sup> Among them, the use of fluoride anions,<sup>18</sup> the concentration of the synthesis mixture,<sup>19</sup> the presence of heteroatoms<sup>20</sup> and temperature<sup>21</sup> can all make a large difference. As we show here, the presence of inorganic impurities should be added to this list. With regard to the role of heteroatoms, it is interesting to realize that, on occasion, a given heteroatom (for instance Ge) can have absolutely no structure-directing effect, producing instead a solid solution over the whole range of  $Ge_f = Ge/(Ge + Si) = 0-1$  molar fractions,<sup>22</sup> while in other cases a structure-directing effect definitely exists.<sup>20,23</sup> Thus, for unspecific OSDAs, a screening of synthesis conditions by varying the above factors may be of interest and provide opportunities for discovery.<sup>24</sup> Here we show that 2-isopropyl-1,3-dimethylimidazolium (2iPr13DMI) is a rather unspecific OSDA that can produce up to seven different zeolite phases. Interestingly, two of these phases possess the same UOS topology but distinctly different compositions and are produced in significantly different, non-adjacent crystallization fields [bold three letter codes refer to zeolite framework type codes approved by the Structure Commission of the International Zeolite Association and, when preceded by a dash, they correspond to “interrupted frameworks”].<sup>25</sup> Due to different symmetries and unit cell parameters, the PXRD diagrams of these zeolites differ from that of IM-16 zeolite, the reference UOS material, the monoclinic structure of which in the as-synthesized form is also reported here for the first time.<sup>26</sup>

## Experimental section

### Synthesis of the OSDA

**Method A.** The first synthesis attempts were carried out through single step nucleophilic substitution reactions, following a methodology similar to the ones we previously used for the synthesis of other imidazolium cations. This consisted of

full methylation at the N positions of imidazole compounds using methyl iodide in the presence of potassium carbonate sesquihydrate as a proton scavenger. However, in this case, the organic salt obtained (2iPr13DMI<sup>+</sup>I<sup>-</sup>) showed low solubility in chloroform (as well as in other organic solvents) so separation of the solid resulting from the use of K<sub>2</sub>CO<sub>3</sub>·1.5H<sub>2</sub>O was very difficult. To overcome this problem, after filtering the reaction mixture, the solid residues were washed with methanol and the salt was obtained by rotary evaporation. However, it turned out that methanol eventually also dissolved some potassium compound (coming from the potassium carbonate) and, while its amount was very small according to CHN analysis (which suggested that the particular OSDA analyzed was close to 100% pure, see Table S1†), it very significantly affected the subsequent zeolite synthesis experiments (see below).

**Method B.** We then changed the synthesis to a two-step methylation that avoided the solubility problem. First 0.075 mol of 2-isopropylimidazole (Aldrich or Alfa Aesar, 98%) was dissolved in around 100 mL chloroform (Aldrich, 99%). Then, 0.0825 mol of iodomethane (Aldrich, 99%) was added to the solution in the presence of 0.206 mol of finely ground K<sub>2</sub>CO<sub>3</sub>·1.5H<sub>2</sub>O (Aldrich, 99%). After 3 days under magnetic stirring at RT, the solution was filtered and the solid was washed with chloroform. Then, an excess of iodomethane was added to the chloroform solution, and the reaction was left for a further three days under magnetic stirring. The solvent was removed by rotary evaporation and the organic salt was washed with acetone. The overall yield was generally close to 80% and the nature and purity of the cation were confirmed by <sup>1</sup>H and <sup>13</sup>C NMR and chemical analysis (see below).

For both synthesis methods, the obtained iodide salt was converted into the organic hydroxide form by anion exchange using Dowex Monosphere 550A (OH) anion exchange resin (Sigma Aldrich). The hydroxide solution, 2iPr13DMIOH (aq), was concentrated by rotary evaporation to achieve a final concentration of around 1.2 mol kg<sup>-1</sup>. The concentration before and after rotary evaporation was determined by titration with HCl 0.1 N (Aldrich) using phenolphthalein as an indicator.

### Zeolite synthesis

Tetraethylorthosilicate (TEOS 98%, Aldrich) and GeO<sub>2</sub> (Aldrich, 99+%) were used as the silicon and germanium sources, respectively. Appropriate amounts of these reagents were hydrolyzed at room temperature under stirring in the aqueous 2iPr13DMIOH solution. Evaporation of all the ethanol produced and as much water as needed was allowed and monitored by weight. Finally, hydrofluoric acid (HF 48%, Aldrich) was added (caution: work under a fume hood), followed by manual stirring using a spatula for approximately 15 minutes. The highly viscous gel so obtained was placed inside the inner Teflon vessels of stainless steel autoclaves. For crystallization, temperatures of 423, 448, 458 and 468 K were used, under tumbling (rotation speed *ca.* 54 rpm). The final composition of the gel was (1 - *x*)SiO<sub>2</sub>:*x*GeO<sub>2</sub>:0.5OSDAF: *w*H<sub>2</sub>O, where  $x = Ge_f = Ge/(Si + Ge)$  was the molar fraction of germanium to germanium and silicon and was varied from 0



to 1 and  $w$  was varied in the 2.9 to 6.7 range for the pure SiO<sub>2</sub> syntheses and kept fixed at 4 for the syntheses containing Ge. After predefined times, the autoclaves were removed from the oven, cooled with water, then the final pH was checked and the solids were washed with deionized water and then dried in an oven at 373 K. The reference material IM-16 was prepared hydrothermally at 443 K for 14 days according to the procedure published by Lorgouilloux *et al.* with 3-ethyl-1-methyl-3*H*-imidazol-1-ium as the OSDA.<sup>26</sup>

### Characterization

Powder X-ray diffraction (PXRD) was performed using a Bruker D8 Advance diffractometer, with Cu K $\alpha$  radiation. C, N, H analyses were performed with a LECO CHNS-932 instrument. <sup>19</sup>F, <sup>29</sup>Si and <sup>13</sup>C MAS NMR experiments were recorded on a Bruker AV 400WB spectrometer, as described elsewhere.<sup>27</sup> Field emission scanning electron microscopy (FE-SEM) images were obtained on an FEI NOVA NANOSEM 230 instrument without metal coating. Thermogravimetric (TG) analyses were achieved with an SDT Q600 TA Instruments device at a heating rate of 10 K min<sup>-1</sup> under an air flow of 100 mL min<sup>-1</sup>. Synchrotron PXRD (SPXRD) data for HPM-6 and HPM-11 were collected at the bl04 MSPD beamline of the Spanish synchrotron light source (ALBA, Cerdanyola del Vallès, Barcelona), in capillary mode (0.6–0.8 mm diameter). The PXRD pattern of as-synthesized IM-16 was collected on a PANalytical MPD X'Pert Pro diffractometer in the Debye–Scherrer geometry equipped with a capillary sample holder, a hybrid mirror monochromator ( $\lambda = 1.5406$  Å), which gave the mono-chromatic parallel beam geometry, and an X'Celerator real-time multiple strip detector (active length = 2.122°,  $2\theta$ ).

### Electron diffraction data collection, structure solution and Rietveld refinement

Electron diffraction data on as-made HPM-6 and HPM-11 were collected at 96 K using the continuous rotation method (cRED) on a JEOL JEM2100 TEM (LaB<sub>6</sub> filament) instrument operating at 200 kV and equipped with a quad hybrid pixel detector (Timepix). The reciprocal space reconstruction (Figs. S1 and S2†) was carried out using the program REDp,<sup>28</sup> and the reflection intensity extraction was conducted by the program XDS.<sup>29</sup> *Ab initio* structure solution was performed using ShelXT,<sup>30</sup> then refined with atomic scattering factors for electrons by Olex 2.<sup>31</sup> The final structures of as-made HPM-6 and HPM-11 were refined against the SPXRD data using TOPAS-V6.<sup>32</sup> The model from cRED data was used as the initial model, although for HPM-11 we then had to change symmetry (see below). The positions of OSDAs were determined by using a combination of simulated annealing (global optimization) and Rietveld refinement using a previously described procedure.<sup>33</sup> The structure of as-synthesized IM-16 was refined using the GSASII suite of programs.<sup>34</sup> The initial model of the framework was generated from the structure of the orthorhombic unit cell (space group *Cmcm*)<sup>26</sup> of the calcined sample transformed into the monoclinic subunit cell (space group *P2<sub>1</sub>/m*) using the program PowderCell.<sup>35</sup> Two OSDA molecules were placed on a mirror

plane, as revealed by difference Fourier maps. The structure was minimized with the Forcite module of Materials Studio<sup>36</sup> before Rietveld refinement.

### Computational details

Molecular mechanics simulations were performed in order to find the most stable location of 2iPr13DMI cations within the different frameworks; simulations were carried out using the Forcite module implemented in Materials Studio,<sup>36</sup> using the Dreiding forcefield.<sup>37</sup> Suitable supercells were built for each system in pure silica composition. 2iPr13DMI cations were manually docked in the framework in different positions/orientations, and the most stable location was obtained by simulated annealing (200 cycles). Due to the distinct porosity and framework density of different zeolite structures (and hence of the different organic contents per Si) as well as the different intrinsic stabilities of the frameworks as a function of network composition, direct comparison of the total interaction energies is meaningless. Hence, in order to analyze the intrinsic fitting of the cation in the different zeolite frameworks, the interaction energies were normalized to the free volume available, and these values (which were referred to as the normalized interaction energy, N.I.E.) were used to estimate the structure-directing ability of the organic cation for the different frameworks. Free volumes were calculated with a probe radius of 1.0 Å. Lattice energies were calculated using the GULP code,<sup>38</sup> with the potential developed by Catlow<sup>39</sup> and Gale and Henson, and taking that of SiO<sub>2</sub>-quartz from ref. 40. In this way, the final stability of the host–guest zeolite systems was assessed by comparing both the relative framework stability through the lattice energy and the structure-directing ability of the organic cation through the normalized interaction energy.

The DFT relative framework stability as well as the theoretical <sup>13</sup>C and <sup>19</sup>F NMR chemical shifts were studied by full geometry optimization of zeolite models with DFT+D methods, which were performed with the CASTEP code,<sup>41</sup> using plane waves (with an energy cut-off of 571.4 eV) and the PBE functional (including the Grimme dispersion term).<sup>42</sup> In the case of the theoretical NMR calculations, F anions were located in the *4d*r units (for UOS) and in the [4<sup>4</sup>5<sup>2</sup>] cages (for CSV). Calculation of the <sup>13</sup>C and <sup>19</sup>F NMR chemical shieldings was carried out with the gauge-including projector augmented-wave method (GIPAW) developed by Pickard and Mauri,<sup>43</sup> as implemented in the CASTEP code. The chemical shift for a nucleus at a given position ( $\delta(r)$ ) is defined as  $\delta(r) = \sigma_{\text{ref}} - \sigma(r)$ , where  $\sigma(r)$  is the isotropic shielding obtained in the calculations. For comparison with experimental chemical shifts,  $\sigma_{\text{ref}}$  values of 174 ppm (for <sup>13</sup>C) and 91.6 ppm (for <sup>19</sup>F) were used, so that the experimental and theoretical values roughly coincided.

## Results and discussion

### Synthesis results

The results for the synthesis of pure silica zeolites using 2iPr13DMI obtained by the two different methodologies under



varying  $\text{H}_2\text{O}/\text{SiO}_2$  ratios and temperatures are listed in the first eighteen rows of Table S2.† Using the OSDA obtained by the first methodology at 423 K, the only zeolite that crystallized was the default phase ZSM-12 (structure code **MTW**) for any  $\text{H}_2\text{O}/\text{SiO}_2$  ratios tried. We also occasionally observed the coprecipitation of a  $\text{K}_2\text{SiF}_6$  impurity (Fig. S3†), which revealed contamination by  $\text{K}^+$  for the OSDA prepared by method A. Thus, further experiments were done using only the OSDA prepared by method B.

When the OSDA obtained by method B was used at low  $\text{H}_2\text{O}/\text{SiO}_2$  ratios and 423 or 448 K, it resulted in the crystallization of pure silica SSZ-35 (structure code **STF**), another zeolite without high demand for specificity. This phase was also obtained in a pure silica composition using 2iPr13DMI by Schmidt *et al.*<sup>44</sup> SSZ-35 tends to be replaced by ZSM-12 over time (suggesting ZSM-12 is thermodynamically more stable than SSZ-35), which is further favored as the concentration decreases. These observations are in accordance with Villaescusa's rule (the experimental observation that less dense zeolites tend to appear at higher concentrations),<sup>19</sup> since **STF** is less dense than **MTW** ( $\text{FD}_{\text{Si}} = 16.9$  vs.  $18.2$  T per  $1000 \text{ \AA}^3$ ).<sup>45</sup>

Thus, under the reported conditions, 2iPr13DMI does not appear to be a very interesting OSDA, merely yielding two different default structures. It was however interesting that the presence of minute amounts of potassium affected the selectivity of crystallization, since, when the OSDA synthesized by method A was used, only ZSM-12 was obtained, even under conditions in which SSZ-35 should have prevailed. Possibly, either  $\text{K}^+$  plays a structure directing role towards the crystallization of ZSM-12 zeolite in a short crystallization time or the crystallization of ZSM-12 is caused by fluoride depletion in solution resulting from the precipitation of highly insoluble  $\text{K}_2\text{SiF}_6$ . We reckon that the second possibility is more likely because the low solubility of potassium hexafluorosilicate would largely impede potassium from acting in solution. Also, the amount of this salt cannot be very large so it is unlikely that it may somehow promote the nucleation of ZSM-12 on its

surface. Upon the introduction of germanium into the synthesis gel, the scenario changes drastically (see Fig. 1). First, very small Ge fractions (around 0.01) stabilize the SSZ-35 structure with regard to its transformation into ZSM-12, suggesting that the accommodation of Ge within the **MTW** topology is more difficult. Slightly increasing the Ge content to just around 0.1 at 448 K consistently produces CIT-7, a recently discovered zeolite to which the **CSV** code has been assigned.<sup>46</sup> With a further increase in Ge, the system enters a region of very low specificity where three additional zeolites, to which we assigned lab codes HPM-10, HPM-6 and HPM-11, appear, most frequently as mixtures and with HPM-1 (**STW**) in minor proportions. The first one, HPM-10, the zeolite with lowest framework density obtained in this system, tends to appear at low  $\text{Ge}_f$  (0.2) in this region and transforms into CIT-7, which in turn starts to transform into HPM-1 during long runs. As  $\text{Ge}_f$  increases further (0.4), CIT-7 is no longer observed and HPM-10 starts to transform over time into HPM-1 and, with more Ge (0.6), into HPM-6. With a further increase in  $\text{Ge}_f$  to 0.7, HPM-10 disappears from the field and HPM-6 dominates until  $\text{Ge}_f = 1$ . An increase in temperature to 458 K at  $\text{Ge}_f = 0.4$  favors the transformation of initially crystallized HPM-10 into another phase, to which we assign the lab code HPM-11, and then its mixtures with **CSV**, while pure HPM-11 can be obtained with a further increase in temperature to 468 K, although small amounts of  $\text{GeO}_2$  argutite tend to precipitate over long crystallization times. All the phase transformations observed are towards denser phases. The structural features of all the zeolites obtained are listed in Table 1. The group of zeolites synthesized in this work using 2iPr13DMI is very much heterogeneous from any point of view, as it contains a very low-density interrupted framework (**-CLO**) and five true, fully connected zeolites of varying density, pore apertures and building units. It is true, however, that the materials requiring a higher  $\text{Ge}_f$  ( $>0.1$ ) all contain double four membered ring (*d4r*) units in their structures. The observed **-CLO** and **UOS** phases received an HPM lab code because their true nature remained unknown for some time (see below).

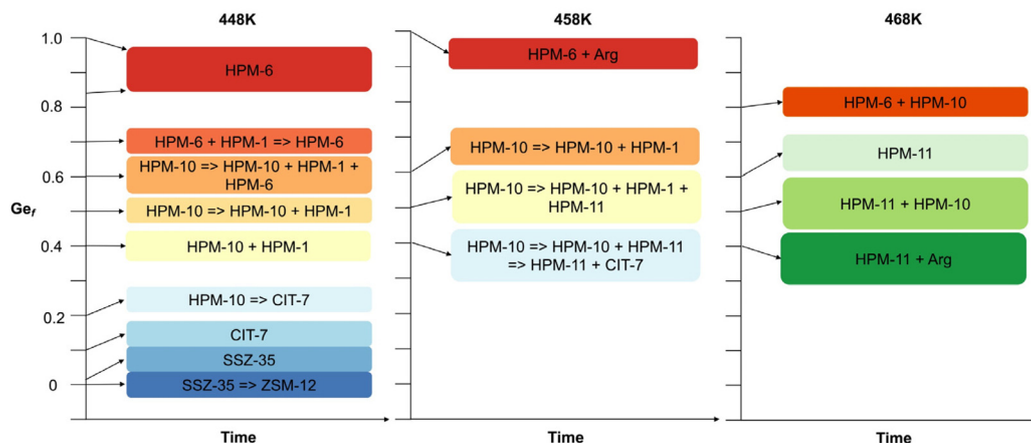


Fig. 1 Schematic representation of the synthesis results at  $\text{H}_2\text{O}/\text{SiO}_2 = 4$  and 448, 458 and 468 K (see Table S2†).



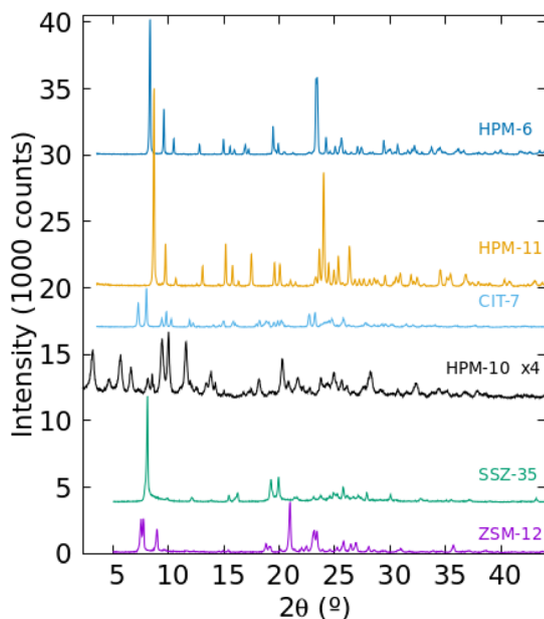
**Table 1** Structural characteristics of the materials synthesized in this work using 2iPr13DMI<sup>45</sup>

Zeolite	ZFT <sup>a</sup>	Channel system <sup>b</sup>	FD <sub>Si</sub> <sup>c</sup>	CBU <sup>d</sup>
ZSM-12	<b>MTW</b>	1D 12 MR	18.2	<i>jbw, cas, bik, mtw</i>
SSZ-35	<b>STF</b>	1D 10 MR	16.9	<i>stf, cas</i>
CIT-7	<b>CSV</b>	2D 10 × 8 MR	16.2	<i>sti, cas, mtw</i>
HPM-10	<b>-CLO</b>	3D 20 × 20 × 20 + 8 × 8 × 8 MR	11.1	<i>d4r, clo, lta</i>
HPM-1	<b>STW</b>	3D 10 × 8 × 8 MR	16.4	<i>d4r</i>
HPM-6, HPM-11	<b>UOS</b>	3D 10 × 8 × 8 MR	17.6	<i>d4r, mtw</i>

<sup>a</sup> Zeolite framework types. <sup>b</sup> Dimensionality and number of tetrahedra in the window limiting diffusion along the pore. <sup>c</sup> Framework density (FD, tetrahedra per 1000 Å<sup>3</sup>). <sup>d</sup> Composite building units.

### Powder X-ray diffraction

The PXRD patterns of the six zeolitic phases prepared in pure form in this work are presented in Fig. 2. They could be assigned to the **MTW**, **STF**, **-CLO**, **CSV** and **UOS** zeolite structure types but it was not immediately clear in the case of **-CLO** (HPM-10) and **UOS** (HPM-6 and HPM-11). In Figs. S4 and S5† we compare the PXRD patterns of these three materials with the patterns simulated with the program VESTA for **-CLO** and **UOS**,<sup>47</sup> respectively, using the cif files provided in the IZA's database of zeolite structures and a single average Cu K $\alpha$  wavelength,  $\lambda = 1.5418$  Å.<sup>45</sup> In the case of HPM-10, the very low intensity of its reflections, its most frequent appearance in mixtures (with HPM-1, HPM-6 or CIT-7, see Table S2†) and the omission of the first reflection around  $2\theta = 3.2^\circ$  under our then standard PXRD measurements prevented its identification for months. Fig. S4† clearly demonstrates, however, that



**Fig. 2** Powder XRD of the six zeolitic phases synthesized as pure phases in this work as Ge<sub>f</sub> in the gel increases (from bottom to top).

HPM-10 has the **-CLO** framework type and is thus similar to the recently reported germanosilicate PKU-12.<sup>48</sup> The comparatively broad and weak reflections are probably a consequence of its nanocrystalline nature (see below). In the case of HPM-6, its high Ge content (Ge<sub>f</sub> ≥ 0.7 for any pure solid in Table S2† that we have analyzed, compared to 0.43 in the original IM-16 (**UOS**) report<sup>26</sup> and up to 0.8 claimed in the patent<sup>49</sup>) together with a change in symmetry (see below and Fig. S5†) changed the PXRD pattern greatly enough so as to prevent identification of this phase. Then, its structure was solved in space group *Pbma* (no. 57) using EXPO2009 and conventional PXRD data for a phase with Ge<sub>f</sub> = 1.<sup>50</sup> First a partial solution lacking some Ge atoms in *d4r* was obtained and these units were then completed manually to yield the fully connected zeolite. Only then was HPM-6 identified as belonging to the **UOS** type through the use of KRIBER with an up-to-date version of the coseq file.<sup>51</sup> As far as we know, this is the first report of a pure germanate with **UOS** topology. Finally, HPM-11, synthesized at a higher temperature but with a lower Ge<sub>f</sub> (around 0.4 in the gel), appeared to be related but distinctly different from HPM-6 and from the reported IM-16. For instance, IM-16 has 5 peaks below 12° while HPM-11 has only 3 with much different relative intensities (Fig. S5†). This is attributed to its slightly larger Ge<sub>f</sub> (0.58 compared to 0.42, although the Ge<sub>f</sub> in the gel was lower for HPM-11 than for IM-16) and a change in symmetry from *Cmcm* in the reported calcined IM-16 to *Pbcm* in as-made HPM-11. In Fig. S5†, we have included the simulated PXRD pattern of ECNU-16 (**EOS**), a very recently reported zeolite, to show that these two zeolites (reported as having different but closely related zeolite framework types)<sup>52</sup> display patterns (simulated from the corresponding cif files) that are very similar to each other but greatly differ from those of HPM-11 and HPM-6. The sharp differences in PXRD patterns between HPM-11 and HPM-6 contrast with the fact that data for materials identified as HPM-6 but synthesized from quite different Ge<sub>f</sub> (0.7, 0.85, 1.0) hardly change (Fig. S6†). This is likely due, in part, to the fact that the measured Ge<sub>f</sub> in the HPM-6 solid is in a narrower range (0.88, 0.96 and 1.0) than in the gel. We have confirmed beyond doubt by cRED that both HPM-11 and HPM-6 samples indeed have **UOS** topology. Interestingly, there is a difference in symmetry between them at the temperature of the electron diffraction experiments (96 K). cRED data for eleven crystals of HPM-11 indicated *Pmcm* symmetry (Fig. S2†), but the PXRD pattern at room temperature showed several reflections that could not be indexed to that symmetry. The whole pattern could be indexed instead to a *Pbcm* cell with a doubling of the *b* parameter. In the case of HPM-6, both cRED (Fig. S1†) and PRXD agreed on *Pbcm* symmetry with double *b*. Thus, there must be a phase transition in HPM-11 between RT and 96 K that does not occur in the higher Ge<sub>f</sub> analog HPM-6. The Rietveld refinements of HPM-11 and HPM-6 against RT synchrotron PXRD data were performed under *Pbcm* symmetry with double *b* (see results in Figs. S7 and S8 and Tables S3–S8†). For completeness, the Rietveld refinement results for as-made IM-16 are shown in Fig. S9 and Tables S9–S11.†



Table 2 lists crystallographic data for several phases prepared in this work and Fig. S10† shows that the volume per tetrahedral atom in the topologically related materials we discuss here (IM-16, HPM-6, HPM-11 and ECNU-16) roughly increases as  $Ge_f$  increases. This tendency is clearly more linear for the samples synthesized with the same OSDA (2iPr13DMI in HPM-6 and HPM-11). In summary, the distinct patterns of HPM-6 and HPM-11, compared to UOS, which are particularly obvious in the  $2\theta = 20\text{--}30^\circ$  range, are a consequence of their different symmetries and chemical compositions, which for HPM-6 and HPM-11 arise from different synthesis conditions using the same OSDA.

**Elemental and thermal analyses.** The CHN analysis of the relevant zeolites in this study are listed in Table 3. The C/N and H/N molar ratios are close to, but generally a bit higher and lower, respectively, than the theoretical and experimental ratios for the pristine OSDA cation (Table S1†). The total organic content (based on N analysis and assuming the charge is balanced by fluoride) indicates that close to 2 OSDAs per unit cell are occluded in CIT-7. In HPM-6 and HPM-11, there are four OSDAs in the *Pbcm* double unit cell. In HPM-10, there are 24 per cell, due not only to the much larger cell of -CLO (192 T atoms compared to 20 for CSV and 24 for UOS), but also to its significantly more open character ( $FD_{Si} = 11.1$  T per  $1000 \text{ \AA}^3$  for -CLO compared to 16.2 and 17.6 for CSV and UOS, respectively).<sup>45</sup> The calculated amount of OSDAs in all these phases, if balanced by fluoride in *d4r* cages, suggests a total

occupancy of *d4r* cages in -CLO (24 *d4r*/u.c.) and UOS (4 *d4r*/u.c. in *Pbcm* symmetry) and also a total occupancy of the two [ $4^45^2$ ] cages per cell in CSV (see below).

The thermogravimetric (TG) and differential thermal analyses (DTA) of the five germanosilicate phases are shown in Fig. S11,† where in addition to the large weight loss presented by HPM-10, the most noticeable feature is observed in the thermograms of the phases synthesized with a higher  $Ge_f$  (HPM-6 with  $Ge_f = 0.7$  and 1), where two weight gaining stages are observed, starting at around 660–680 and 900–940 °C, respectively (see arrows in Fig. S11†). The temperatures of both weight gaining events are close to those we observe in the case of pure  $GeO_2$ -AST phases,<sup>53</sup> despite the large difference in pore apertures of the 6MR windows in AST and the 10 + 8 MR pores in UOS. This is due to the fact that the gaining steps are not related to diffusion along the pores but to reoxidation of previously reduced framework  $Ge(IV)$ .<sup>53</sup> Also, in HPM-11, a much smaller but clear weight gaining event is observed by the end of the thermal trace.

The noted reduction itself cannot be directly observed, but it has an influence on the corresponding DTA trace: instead of the large exothermic DTA peak observed in the remaining cases, due to OSDA combustion, the materials that present large weight gaining steps show minor and endothermic DTA variations associated with the weight loss steps and a small exothermic peak at the onset of the weight gaining step, very similarly to our previous observations in the  $GeO_2$ -AST

Table 2 Crystallographic data of materials relevant to this work

Zeolite (T/K)	ZFT	$Ge_f^c$	Space group	$a/\text{\AA}$	$b/\text{\AA}$	$c/\text{\AA}$	$\alpha/^\circ$	$\beta/^\circ$	$\gamma/^\circ$	Vol $\text{\AA}^3/T$
CIT-7 <sup>a</sup> (448)	CSV	0.22(0.2)	<i>P</i> $\bar{1}$	12.949	11.233	9.388	92.89	107.14	103.04	63.06
HPM-10 <sup>a</sup> (458)	-CLO	0.51(0.6)	<i>Pm</i> 3 <i>m</i>	26.179	26.179	26.179	90	90	90	93.4
IM-16 (443)	UOS	0.43(0.5)	<i>P2</i> <sub>1</sub> / <i>m</i>	11.82694	19.95425	11.71845	90	99.71	90	56.79
ECNU-16 <sup>b</sup> (443)	EOS	0.48(0.5)	<i>C2</i> / <i>m</i>	17.83400	15.12600	10.67100	90	108.56	90	56.85
HPM-11 (468)	UOS	0.58(0.4)	<i>Pbcm</i>	7.6212	18.1181	20.2859	90	90	90	58.36
HPM-11 <sup>a</sup> (468)	UOS	0.66(0.6)	<i>Pbcm</i>	7.644	18.114	20.410	90	90	90	58.88
HPM-6 (448)	UOS	0.88(0.7)	<i>Pbcm</i>	7.66042	18.07804	20.72105	90	90	90	59.78
HPM-6 (448) <sup>a</sup>	UOS	0.96(0.85)	<i>Pbcm</i>	7.677	18.095	20.822	90	90	90	60.26
HPM-6 (448)	UOS	1(1)	<i>Pbcm</i>	7.68166	18.10273	20.89530	90	90	90	60.54

<sup>a</sup> By Le Bail fitting (others by Rietveld refinement). <sup>b</sup> From ref. 52. <sup>c</sup>  $Ge_f$  in zeolite, determined by EDS (the value for the gel is listed inside parentheses).

Table 3 Elemental and thermal analysis data of relevant as-made zeolites prepared in this work

Sample	ZFT	$Ge_f$	C/%	H/%	N/%	C/N <sup>a</sup>	H/N <sup>a</sup>	Residue <sup>b</sup> /%	Empirical formula per cell <sup>c</sup>
ZSM-12	MTW	0	5.50	1.23	1.57	4.1	11.1	88.6 (87.7)	$[\text{SiO}_2]_{28}(\text{C}_8\text{H}_{15}\text{N}_2\text{F})_{1.1}(\text{H}_2\text{O})_{3.7}$
SSZ-35	STF	0	8.16	1.14	2.39	4.0	6.7	84.8 (86.5)	$[\text{SiO}_2]_{32}(\text{C}_8\text{H}_{15}\text{N}_2\text{F})_{1.9}$
CIT-7	CSV	0.22	11.01	1.66	3.16	4.1	7.3	79.9 (82.1)	$[\text{Ge}_{0.22}\text{Si}_{0.78}\text{O}_2]_{20}(\text{C}_8\text{H}_{15}\text{N}_2\text{F})_{1.9}$
HPM-10	-CLO	0.42	12.88	1.91	3.53	4.3	7.5	75.2 (77.6)	$[\text{Ge}_{0.42}\text{Si}_{0.58}(\text{OH})_{0.12}\text{O}_{1.94}\text{F}_{0.12}]_{192}(\text{C}_8\text{H}_{15}\text{N}_2)_{24.6}(\text{H}_2\text{O})_{27.1}$
HPM-11	UOS	0.58	8.53	1.31	2.45	4.1	7.4	86.0 (86.2)	$[\text{Ge}_{0.58}\text{Si}_{0.42}\text{O}_2]_{48}(\text{C}_8\text{H}_{15}\text{N}_2\text{F})_{4.2}$
HPM-6	UOS	0.88	7.39	0.99	2.04	4.2	6.7	86.4 (88.5)	$[\text{Ge}_{0.88}\text{Si}_{0.12}\text{O}_2]_{48}(\text{C}_8\text{H}_{15}\text{N}_2\text{F})_{3.9}$
HPM-6	UOS	1.0	6.81	0.92	1.93	4.1	6.7	88.9 (89.1)	$[\text{GeO}_2]_{48}(\text{C}_8\text{H}_{15}\text{N}_2\text{F})_{3.9}$

<sup>a</sup> Theoretical C/N = 4.0 and H/N = 7.5. <sup>b</sup> TG residue at 1000 °C. In parentheses, the  $\text{TO}_2$  weight percent in the empirical formula (note that the residue will be altered if  $\text{GeO}$  sublimation occurs during thermal analysis). <sup>c</sup> Calculated from the N content and assuming the OSDA is balanced by F anions. In the case of -CLO, a 2.5% weight lost at low T is accounted for by including 27  $\text{H}_2\text{O}$  molecules per cell.



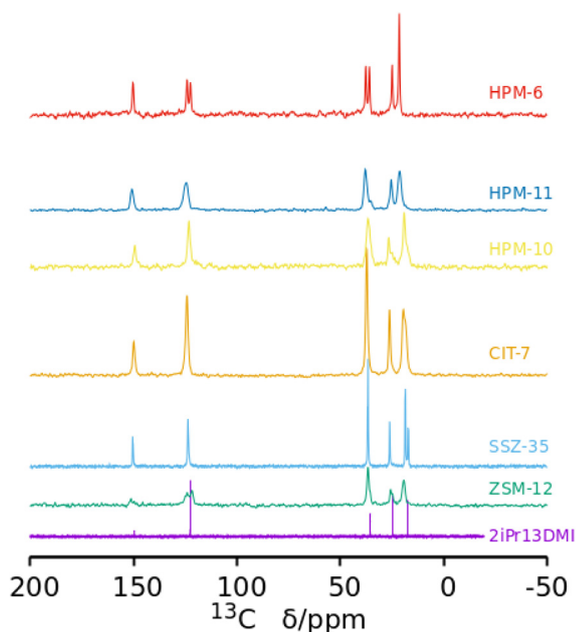
system.<sup>53</sup> This kind of DTA feature is likely due to the fact that there is no OSDA combustion with oxygen but, at least partly, an organothermal reduction of the framework  $\text{GeO}_2$ . We warn that, even when weight gaining is not clearly seen by TG/DTA (as in the case of CSV, HPM-10 and HPM-11), reduction and re-oxidation of Ge, and even sublimation of  $\text{GeO}$ , may still occur and pass unnoticed.<sup>53</sup> This is because germanium monoxide, which may be produced by the reaction of metallic Ge with  $\text{GeO}_2$ , sublimates at temperatures above 480–490 °C,<sup>54,55</sup> *i.e.* at a temperature below the first weight loss in all five germanosilicates. Given that -CLO has two independent channel systems and one of them is just a small 8MR pore, it is not surprising that thermal reduction also occurs in these small pores. This emphasizes our previous warning that much caution is needed in the interpretation of thermal events for germanosilicate zeolites.<sup>53</sup>

### Multinuclear NMR

The  $^{13}\text{C}$  CPMAS NMR spectra of the six pure phases (Fig. 3) show, in every case, signals close to the expected positions corresponding to the five unique C atoms in 2iPr13DMI, strongly suggesting the cations are occluded intact in the zeolites. This happens even when phase transformations occur, showing that during the transformations, which are likely driven by the increased density/stability of the framework, the OSDA continues to act as a pore filler (under the conditions of this study, it is highly unlikely that a zeolite could crystallize without an organic compound helping to stabilize the void

space). However, in several zeolites, some signal splittings are evident. For SSZ-35, the signal corresponding to the methyl groups of the isopropyl moiety are split, suggesting that both methyl groups are not equivalent in the occluded cation. In the case of HPM-6, the signals that can be assigned to  $\text{CH}$  groups of the imidazolium ring (around 123 ppm) and to  $\text{NCH}_3$  (around 37 ppm) are both split into two signals of about equal intensity. These splittings indicate the zeolites impose unequal environments for C atoms that would otherwise be, in principle, equivalent. In the case of isostructural HPM-11, the significantly broadened resonances, which can result from the heterogeneity of Si–Ge occupancies of sites, show no splitting except for the resonance around 37 ppm, which splits into two resonances of unequal intensities. For the remaining as-made zeolites, the NMR signals are significantly broader but there are, however, clear indications of signal splittings at least in ZSM-12 and HPM-10. The  $^{29}\text{Si}$  MAS NMR spectra of the samples containing silica are shown in the ESI (Fig. S12†). The only remarkable features appear in the pure silica SSZ-35 material, which shows very high resolution of crystallographic sites (not less than 10, possibly 14 tetrahedrally coordinated sites) plus a doublet at –147 ppm characteristic of pentacoordinated silicon ( $\text{SiO}_{4/2}\text{F}^-$  units) with a  $J$  coupling of around 175 Hz.<sup>58</sup> This agrees with previous results obtained for SSZ-35 zeolites prepared using unrelated OSDAs,<sup>59–61</sup> although the resolution and evidence of  $\text{SiO}_{4/2}\text{F}^-$  units are not always as clear in the  $^{29}\text{Si}$  MAS NMR spectra of as-made SSZ-35 zeolites as they are here.<sup>62</sup> In contrast, as-made pure silica ZSM-12 displays a poorly resolved spectrum, as commonly observed for this zeolite structure type.<sup>63</sup> On the other hand, the poorly resolved spectra of the Ge-containing materials is not surprising, since they are not pure  $\text{SiO}_2$  materials and, hence,  $\text{Si}(\text{OT})_4$  is expected to be spread over a wide chemical shift range.

Of most interest are the  $^{19}\text{F}$  MAS NMR spectra (Fig. 4). Fluoride occluded in small cages of zeolites displays  $^{19}\text{F}$  resonances with chemical shifts in ranges that typically depend on the fluoride–framework interaction, which in turn is a function of the composition and type of cage in which it is occluded. In pure silica zeolites, the typical chemical shifts can span from –35/–42 ppm in the small  $[4^6]$  cages ( $d4r$ ) to over –80 ppm in larger cages.<sup>64</sup> The pure silica ZSM-12 and SSZ-35 phases display resonances in the typical range for fluoride occluded in cages of silica zeolites: –75 and –78 ppm in ZSM-12 and –80 ppm in SSZ-35. ZSM-12 also shows sharp resonances at higher fields (–119, –126 and –127 ppm) that are ascribed to penta- and/or hexa(oxofluoro)silicate impurities. Several of the other spectra in Fig. 4 also present broad resonances in this region, although with much lower intensity. The germanium containing zeolites show, in contrast, resonances at significantly lower fields, which typically indicate strong F–Ge interactions and which depend on the Ge content and distribution in addition to the enclosing cage. Both Ge and fluoride typically favor the formation of  $d4r$  in zeolites.<sup>18,19,22,23</sup> The sharp and single resonance at –13.5 ppm in pure Ge HPM-6 has to be ascribed to fluoride occluded in  $d4r$  made of germanium



**Fig. 3**  $^{13}\text{C}$  NMR spectrum of 2iPr13DMI iodide in  $\text{D}_2\text{O}$  solution and  $^{13}\text{C}$  MAS NMR spectra of six as-made phases prepared in this work as Ge, increases (from bottom). In the bottom spectrum, the C at position 2 of the aromatic ring (between both N) is barely visible at around 150 ppm. That carbon typically displays a very small relative intensity in the proton-decoupled  $^{13}\text{C}$  NMR spectra of this kind of compound.<sup>56,57</sup>



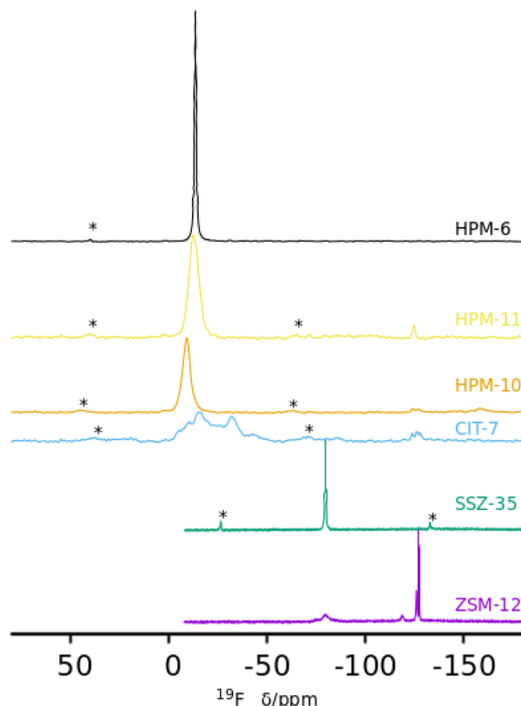


Fig. 4  $^{19}\text{F}$  MAS NMR spectra of six as-made phases prepared in this work: ZSM-12, SSZ-35, CIT-7, HPM-10, HPM-11 and HPM-6 ( $\text{Ge}_f$  0.88 and 1.0) (from bottom).

( $\text{GeO}_2$ - $d4r$ ), which agrees with previous reports on chemical shifts of  $\text{F@GeO}_2$ - $d4r$  whether as discrete units,<sup>65</sup> or in zeolites.<sup>22</sup> Interestingly, the isostructural HPM-11 synthesized with a significantly lower Ge content ( $\text{Ge}_f = 0.4$  in gel, 0.58 in the zeolite) has a single resonance in the close chemical shift of  $-12.7$  ppm. This resonance, however, cannot be assigned in this case to fluoride occluded in  $d4r$  made only of germanium because this would require a much higher Ge content even if there were an absolute preference for Ge to occupy  $d4r$  sites (16 out of the 24 tetrahedral sites in a unit cell, or  $\text{Ge}_f = 0.67$ ). For  $\text{Ge}_f = 0.58$  (13.9 Ge per unit cell) that signal needs to be assigned to fluoride occluded in  $d4r$  units containing Ge in closed clusters, *i.e.*, with at least one Ge having three Ge as next nearest neighbors in the  $d4r$  unit. For this a minimum of 4 Ge per unit cell are needed; this is indeed achievable for this Ge content. This agrees with our proposition that the main factor determining the chemical shift of  $^{19}\text{F}$  in Ge-containing  $d4r$  is not the total Ge content in the  $d4r$  but the number of Ge–O–Ge pairs resulting in four categories of resonances: with no Ge (I), non-paired Ge (II), Ge pairs without larger clusters (III) and with larger clusters (IV).<sup>22</sup> For HPM-10, a single resonance at  $-9.2$  ppm is assigned to fluoride occluded in  $d4r$  units of the -CLO structure with between 2 and 6 Ge atoms forming pairs but no larger clusters (*i.e.*, resonance type III where no Ge has three Ge neighbors in the  $d4r$  unit). This chemical shift contrasts with the one reported for PKU-12,  $-4.3$  ppm.<sup>48</sup> It is interesting that for both HPM-10 and PKU-12 there is a single and symmetrical  $^{19}\text{F}$  resonance despite the

fact that there are two different types of  $d4r$  in the -CLO structure: one is a “normal” unit in which each T atom is oxygen-bridged to 4 T atoms; the other contains two “dangling” T–O bonds where the structure is interrupted. In both PKU-12 and HPM-10, there is likely to be full occupancy of both types of  $d4r$ . For HPM-10, this is deduced from the CHN analysis and charge balance considerations (see above).

The reported  $^{19}\text{F}$  spectrum of as-made silica CIT-7 displays an asymmetric resonance around  $-45$  ppm, which is not specifically assigned.<sup>46</sup> CSV is the first zeolite containing the  $[4^45^2]$  cage, which is composed of an *sti* unit with an additional T atom inserted between the open corners. However, similar tiling containing the *sti* unit exists in other fluoride-containing materials: the truly open and interrupted “[4<sup>4</sup>6]” cage, which is not properly a tile but half the  $[4^88^2]$  tile, in the layered germanosilicate PKU-22, with no T atom between the open corners,<sup>66</sup> and the  $[4^46^2]$  cage with 2 T atoms inserted between the open corners in the interrupted germanosilicate PKU-26,<sup>67</sup> see Fig. 5. Reported  $^{19}\text{F}$  resonances at 1.0 and 0.3 ppm in PKU-22 and PKU-26, respectively, have been assigned to F occluded in the *sti* unit with a direct very short F–Ge bond (2.206 and 2.20 Å, respectively) to a specific Ge site. Interestingly, the  $^{19}\text{F}$  spectrum of our CSV germanosilicate contains at least six overlapped resonances in the 0 to  $-45$  ppm range, which we attribute to fluoride occluded in the  $[4^45^2]$  units of the structure because of its similarity to the  $d4r$  ( $[4^6]$ ), within which  $^{19}\text{F}$  spans a similar chemical shift range depending on the Ge content and distribution. Also, the range in our germanosilicate CSV is between that of pure silica CIT-7 and those of Ge-containing PKU-22 and PKU-26 with short F–Ge bonds in the *sti* units. Thus, we propose that the multiple resonances that can be discerned in the spectrum correspond to different Ge contents and distributions in the  $[4^45^2]$  cage and, possibly, to fluoride interacting with different crystallographic sites within that cage (see calculations below).

### FE-SEM

Fig. S13† shows FE-SEM micrographs of the germanosilicate phases. HPM-10 is nanocrystalline in nature (crystal size below

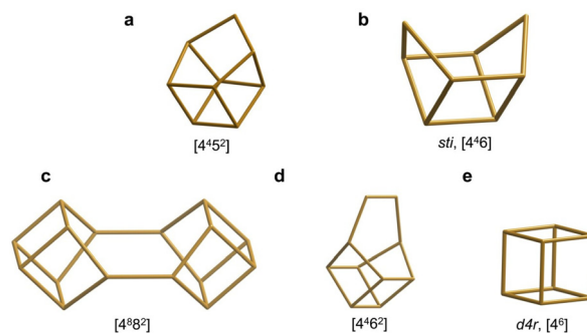


Fig. 5 Several cages that can be formally considered as being derived from the  $d4r$  cage by modification at one corner and which may provide similar environments for fluoride at the opposite side. They appear in CSV (a), STI (b), PKU-22 (c), PKU-26 (d) and, among others, AST (e).<sup>68</sup>



50 nm), while all the other phases are microcrystalline with sizes ranging from up to 3  $\mu\text{m}$  for CIT-7, up to 15  $\mu\text{m}$  for HPM-11 and up to 40  $\mu\text{m}$  and 80  $\mu\text{m}$  for HPM-6 synthesized at  $G_{\text{e}_f} = 0.7$  and 1, respectively. In the last sample we observed a considerable amount of smaller particles in the shape of circular wafers of around 0.8  $\mu\text{m}$ . This impurity phase was not apparent in the powder XRD data, which was fully indexed.

### Computational results

**Pore-filling and interaction energies.** We then analyzed the most stable location of the 2iPr13DMI cations within the different frameworks by molecular mechanics calculations in order to understand their structure-directing ability for the different zeolite structures. Due to the complexity of the -CLO host-guest system, with 24 cations per unit cell with a high degree of freedom, which might result in non-reliable results associated with local minima, this system was excluded from the calculations. Two 2iPr13DMI cations per unit cell (1 per cavity) were loaded in the STF framework (Fig. 6, top-left), with the organic cations sited in the center of the SSZ-35 cavities, showing good geometric host-guest complementarity between its molecular shape and that of the cavity. Indeed, 99.7% of the void pore volume is filled upon occlusion of this cation, showing efficient pore filling, with a N.I.E. of  $-1.08 \text{ kJ mol}^{-1} \text{ \AA}^{-3}$  (Table 4).

In the case of the ZSM-12 structure (MTW), two 2iPr13DMI cations per  $\text{Si}_{56}\text{O}_{112}$  unit cell were loaded; higher loadings were also tried, but these led to unstable situations. As can be seen in Fig. 6 (top-right), 2iPr13DMI cations are located within the 12MR channels of the ZSM-12 framework, with the imidazolium ring aligned with the channel direction, and the C3 methyl groups bonded to N sited in the lateral side pockets of the framework, showing once again a good geometric host-guest fit. However, in this case, a lower pore-filling efficiency is observed (92.3%) because one of each of the two side-pockets remains free, and this results in a relatively low N.I.E. of  $-1.04 \text{ kJ mol}^{-1} \text{ \AA}^{-3}$ .

Two 2iPr13DMI cations per unit cell were loaded in the CSV framework, corresponding to two cations per CSV cavity. The most stable location of the cations (Fig. 6, bottom-left) shows that the CSV cavity has the appropriate dimensions to host two OSDA cations parallel to each other, siting the bulky isopropyl groups of the two cations on opposite sides to avoid steric repulsion interactions. In this case, the high pore-filling efficiency of 2iPr13DMI in this framework (100%) and the good host-guest geometric fit brings an improvement in the N.I.E. to  $-1.22 \text{ kJ mol}^{-1} \text{ \AA}^{-3}$ , suggesting a good structure-directing efficiency of the cation towards this cavity-based framework. We then analyzed the location of the cation within the chiral STW framework, despite it not being obtained as a

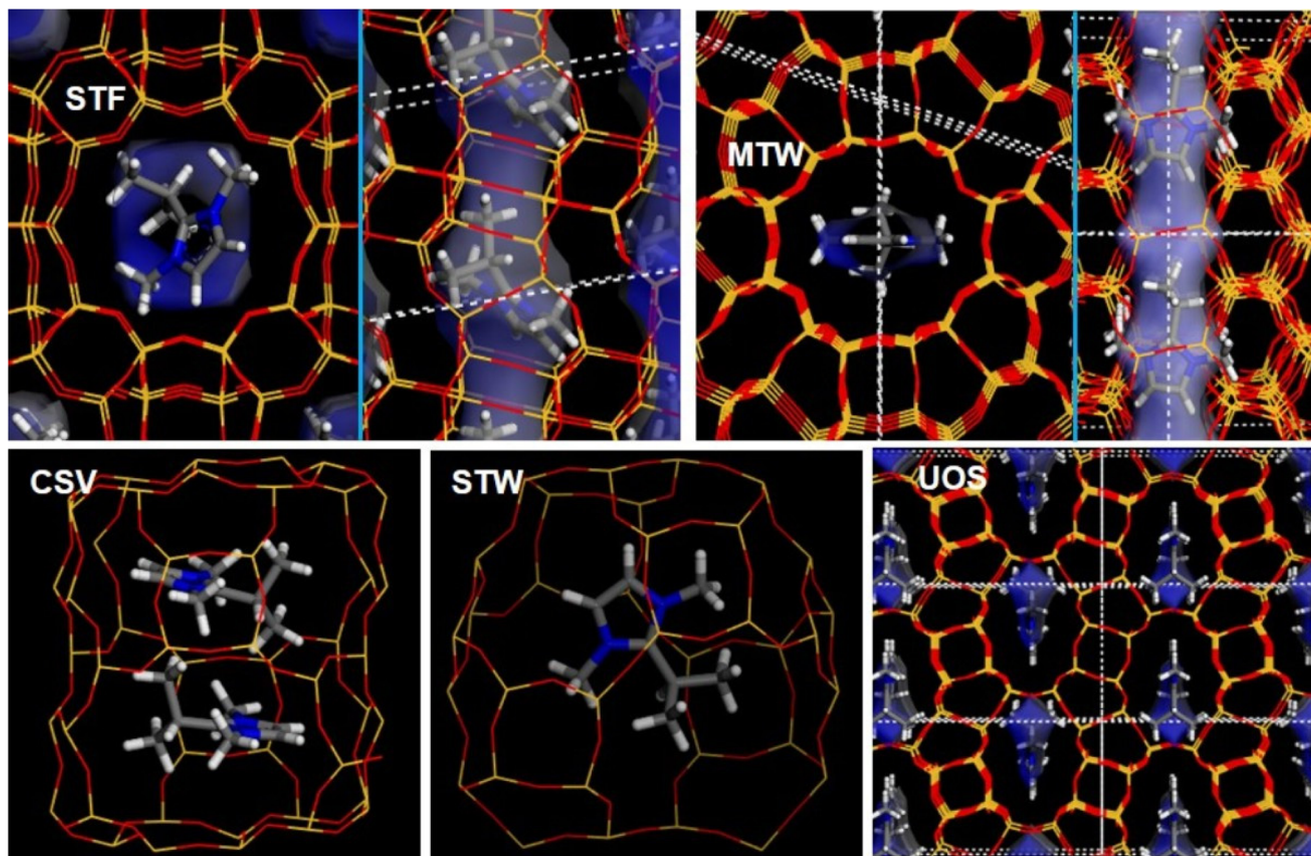


Fig. 6 Location of 2iPr13DMI within the different frameworks studied. Void volume is shown as a transparent white/blue surface to facilitate pore visualization.



Table 4 Different parameters studied for the 2iPr13DMI cations in different frameworks

ZFT (u.c.)	SDA per		Free volume (Å <sup>3</sup> /Si)	Pore filling (%)	I.E. (kJ mol <sup>-1</sup> Si)	N.I.E. (kJ mol <sup>-1</sup> Å <sup>-3</sup> of free V)	GULP (kJ mol <sup>-1</sup> Si)	DFT
	u.c.	Si						
<b>STF</b> (Si <sub>32</sub> O <sub>64</sub> )	2	0.0625	9.756	99.7	-10.51	-1.08	13.17	11.29
<b>MTW</b> (Si <sub>56</sub> O <sub>112</sub> )	2	0.0357	5.186	92.3	-5.37	-1.04	8.13	7.32
<b>CSV</b> (Si <sub>20</sub> O <sub>40</sub> )	2	0.1000	12.544	100	-15.34	-1.22	16.63	12.03
<b>STW</b> (Si <sub>60</sub> O <sub>120</sub> )	6	0.1000	9.867	99.9	-15.50	-1.57	19.94	14.06
<b>UOS</b> (Si <sub>24</sub> O <sub>48</sub> )	2	0.0833	6.056	100	-12.52	-2.07	15.48	11.41

pure phase experimentally. Each cavity can host one cation, resulting in a high N.I.E. of  $-1.57 \text{ kJ mol}^{-1} \text{ \AA}^{-3}$ , indicating a good geometric host-guest match for this system (99.9% of pore-filling efficiency), as can be seen in Fig. 6 (bottom-middle).

Finally, two 2iPr13DMI cations were loaded per **UOS** unit cell, giving 100% pore-filling efficiency. In this case, a clear geometrical match between the molecular cross-like shape of the imidazolium cation and the void space of the **UOS** framework is also apparent (Fig. 6, bottom-right). This framework shows the best host-guest fit for the 2iPr13DMI cations, giving a N.I.E. of  $-2.07 \text{ kJ mol}^{-1} \text{ \AA}^{-3}$  (Table 4), the highest value found among all the frameworks, suggesting a high specificity towards the **UOS** framework, independently of the inherent structural framework stability of the distinct zeolites.

Thus, our results suggest that the 2iPr13DMI cation has a molecular shape that can fit within all these zeolite frameworks, possibly favored by the conformational freedom of the isopropyl group, although this must be significantly limited by steric hindrance. However, by themselves, the N.I.E. results fail to explain the observed experimental trends in structure direction since **STW**, which displays the second-highest N.I.E. value (after **UOS**), should appear as a frequent framework, but it is only marginally observed as a competing phase at high Ge loadings. The lack of strong specificity of this cation implies that other factors come into play to determine the phase selectivity of crystallization. The best calculated fit (in terms of pore-filling efficiency) is **UOS** > **CSV** > **STW** > **STF** > **MTW** while the NIE is in the order **UOS** > **STW** > **CSV** > **STF** > **MTW**. Both orders are in pretty good agreement, except for **CSV** and **STW**. The fact that our theoretical results cannot be straightforwardly correlated with the experimental results is attributed to the different intrinsic stabilities of the different topologies and the strong effect of Ge on them. Although most zeolites typically exhibit a clear correlation between stability and density,<sup>69</sup> this is not the case for the phases involved in this work because of the existence of *d4r* units in several of them. The *d4r* units are strained in pure silica zeolites,<sup>70</sup> and require the presence of F and/or Ge for flexibilization.<sup>71</sup> In fact, the density (FD<sub>Si</sub>) of the phases involved decrease in the order **MTW** > **UOS** > **STF** > **STW** > **CSV**, while, based on previous reports, we can consider **MTW** and **STF** as default structures but **UOS** is clearly not (there has been, so far, only a single reported synthesis of **UOS**, and it required the presence of Ge).<sup>26</sup> The density of **STW** is also close to that of **STF**, but,

while it cannot be considered a very stable zeolite, it was once considered unfeasible.<sup>72</sup> Thus, the discussion must take into account the energies of the SiO<sub>2</sub> phases plus the likely effect of F and Ge on the synthesis. To account for the intrinsic stability of the different topologies, we have calculated their energies as pure silica polymorphs relative to quartz using both interatomic potentials (with GULP code) and DFT methodologies and the results are included in Table 4. Both GULP and DFT provide the same order of relative stabilities (**MTW** > **STF** > **UOS** > **CSV** > **STW**), which agree with our previous considerations (**MTW** and **STF** as default structures and **STW** as the least stable phase among those considered here). In the absence of Ge, **STF** is the phase that first crystallizes, which agrees with a fairly good host-guest match and with its known stability (default structure). It tends to transform into **MTW**, which is denser and, hence, this agrees with the expected stability trend. The predominance of **MTW** when the OSDA is prepared by method A suggests that, under these conditions (the presence of K, precipitation of K<sub>2</sub>SiF<sub>6</sub>), the phase with higher thermodynamic stability prevails, possibly because fluoride depletion is a catalyst to promote thermodynamic over kinetic control.

When Ge is introduced into the synthesis, the phase selectivity landscape changes dramatically. For Ge<sub>f</sub> equal to or above 0.4, **UOS** is clearly the dominant phase, which agrees with its not too low intrinsic stability as a pure SiO<sub>2</sub> zeolite, the stabilization effect of F and Ge on *d4r* units and the excellent fit and strongest host-guest interaction with the OSDA used. This does not happen for **STW**, which is never obtained in this system as a pure phase; this is likely due to its lower host-guest stabilization energy and, especially, to its lower intrinsic stability, which may require a relatively high Ge<sub>f</sub>, for which it competes with **UOS**, which has a stronger interaction energy. The appearance of **CSV** and **-CLO** at fairly low Ge<sub>f</sub> suggests they are not as highly unstable as **STW** and **UOS**, which, in the case of **CSV** may be easily understood, since it does not contain *d4r* but the more open and presumably more flexible *sti* cages.

Comparison with other OSDA systems may be illustrative. In the synthesis of HPM-1 (pure silica **STW**) using 2-ethyl-1,3,4-trimethylimidazolium, 2E134TMI, the OSDA was able to stabilize the pure silica material and, since **STW** contained *d4r*, the introduction of Ge did not change the phase selectivity and the whole solid solution series from Ge<sub>f</sub> 0 to 1 could be synthesized. Other examples from the literature show how



**Table 5** Predicted  $^{13}\text{C}$  NMR shifts as a function of the Ge content, and unit cell parameters and volume of the optimized unit cell

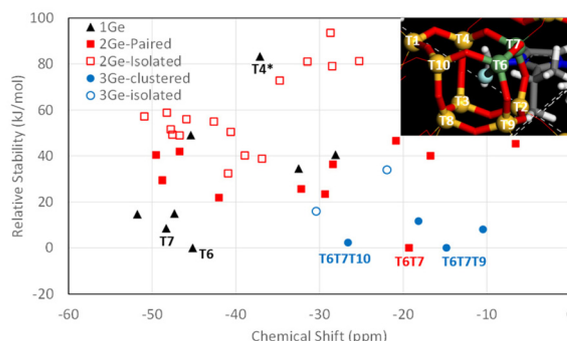
$\text{Ge}_f$	$\text{N}_2\text{CC}$	$\text{CH}$	$\text{NCH}_3$	$\text{CC}_3$	$\text{CCH}_3$	$V(\text{\AA}^3)$	$a$	$b$	$c$			
0	151.0	127.6	129.1	41.4	40.6	28.5	20.7	20.7	1378.08	9.114	19.994	7.563
0.33	151.7	127.2	128.8	40.7	40.2	28.3	21.2	20.5	1430.83	9.19	20.154	7.725
0.66	150.7	127.5	129.4	40.8	37.9	28.0	22.2	22.2	1456.41	9.169	20.344	7.809
1	151.6	127.7	128.6	40.9	37.2	28.8	22.8	22.8	1536.13	9.273	21.076	7.867

this is not always the case and totally different phases are obtained as  $\text{Ge}_f$  is systematically varied.<sup>73</sup> Thus, an interesting point to consider is that, when no single structure-directing effect dominates, the opportunities for discovery arise. This is the case for HPM-8, an intergrown zeolite of the Beta family largely enriched in polymorph E, which has a niche crystallization field with a low  $\text{Ge}_f$  (around 0.1 or 0.15).<sup>74</sup>

**Theoretical prediction of  $^{13}\text{C}$  and  $^{19}\text{F}$  NMR.** We then wanted to gain some insights into the splitting of the  $^{13}\text{C}$  NMR signals observed for the unsubstituted C of the imidazolium ring (CH) and the methyl groups attached to N ( $\text{NCH}_3$ ) as a function of Ge content in the UOS framework. Starting from the location of the 2iPr13DMI cations found previously by molecular mechanics, four different 2iPr13DMI/UOS systems were built: (i) with all T atoms being Si ( $\text{Ge}_f = 0$ ), (ii) with 8 Ge and 16 Si atoms per unit cell, siting 4 Ge in each  $d4r$ , having one Ge surrounded by 3 Ge (Ge clusters) ( $\text{Ge}_f = 0.33$ ), (iii) with 16 Ge and 8 Si atoms per unit cell, with all  $d4r$  atoms being Ge ( $\text{Ge}_f = 0.67$ ), and (iv) with all T atoms being Ge ( $\text{Ge}_f = 1$ ). All these systems were DFT geometry optimized with variable unit cell parameters, and then the  $^{13}\text{C}$  NMR shieldings were calculated by the GIPAW method (Table 5).

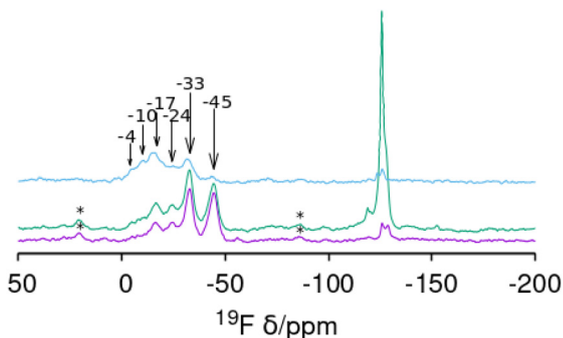
Interestingly, the results show that an increase of the Ge fraction involves a larger splitting of the  $\text{NCH}_3$  signals: they have very similar shifts for Ge-poor systems (with chemical shift differences of 0.8 and 0.5 ppm for  $\text{Ge}_f = 0$  and 0.33, respectively), while showing larger differences for Ge-rich systems (2.9 and 3.7 ppm for  $\text{Ge}_f = 0.67$  and 1, respectively) because of a high field displacement of one of the  $\text{NCH}_3$  chemical shifts from  $\sim 40$  to  $\sim 37$  ppm. This change is probably associated with the change in the unit cell upon incorporation of Ge, where the unit cell volume and specifically the  $c$  parameter, are notably increased, and the pores become slightly larger as well. Indeed, such a distortion of the unit cell might explain the splitting of the  $\text{NCH}_3$  NMR signal, since when  $\text{Ge}_f$  is low (0.33), the environment of both  $\text{NCH}_3$  atoms seems more similar (Fig. S14, left†) than when  $\text{Ge}_f$  is high (1) (Fig. S14, right†), where the distortion of the unit cell slightly changes the orientation of the organic cation, possibly involving a different environment for both  $\text{NCH}_3$ . In contrast, we do not observe a clear trend as a function of  $\text{Ge}_f$  for the two CH signals, which are always slightly split (0.9 to 1.9 ppm, Table 5). However, we need to take into account that these are just simplified systems with regular Ge distributions, while in the real case a more heterogeneous Ge distribution must be present, in particular for HPM-11, which may explain its broader  $^{13}\text{C}$  resonances.

On the other hand, since the  $^{19}\text{F}$  NMR spectrum of  $\text{F}^-$  occluded in the  $[4^45^2]$  cages of the CSV framework was reported only recently, and no specific assignments have been provided yet, we calculated the  $^{19}\text{F}$  NMR chemical shift of the system with different Ge configurations in order to test our above assignment. Systems with 1, 2 or 3 Ge per cage (giving overall  $\text{Ge}_f$  of 0.1, 0.2 and 0.3, respectively) were studied by DFT geometry optimization, and the theoretical  $^{19}\text{F}$  chemical shifts were calculated. For  $\text{Ge}_f = 0.1$ , each  $[4^45^2]$  cage contained 1 Ge and all the different configurations were tried.  $\text{F}^-$  was initially located in the center of the cage, and the systems were geometry optimized (keeping  $P\bar{1}$  symmetry, *i.e.* both  $[4^45^2]$  cages are symmetry-equivalent); in all cases, F bonded to Ge instead of Si upon geometry optimization (see Fig. 7 caption), evidencing higher stability of the Ge–F bonds (the only exception was T4, marked with \* in Fig. 7, where F–Ge4 was very unstable, and instead bonded to Si7, also resulting in a very unstable system). Results are shown in Fig. 7, where the relative stability (calculated independently for systems with the same  $\text{Ge}_f$ ) is plotted against the theoretical  $^{19}\text{F}$  NMR chemical shift. Energy results show that the most stable position for 1 Ge is T6 (see inset), followed by T7 (8.6 kJ mol $^{-1}$  u.c.); calculations predict a  $^{19}\text{F}$  NMR signal at around  $-45$  ppm for cages with 1 Ge in T6; indeed, all the more stable 1 Ge systems gave shifts close to this ( $-45$  to  $-52$  ppm). We then included a second Ge in the cage (red squares) ( $\text{Ge}_f = 0.2$ ); in this case, a notably higher stability for systems with Ge in adjacent positions (Ge-paired, solid red squares) was found (compared with



**Fig. 7** Relative stability vs.  $^{19}\text{F}$  NMR chemical shift for  $\text{F}^-$  in  $[4^45^2]$  cages with 1 Ge (black triangles), 2 Ge (red squares) in paired (solid) or isolated (empty) positions and 3 Ge (blue circles) in clustered (solid) or non-clustered (empty) positions. Relative stabilities are calculated independently for 1Ge, 2Ge and 3Ge systems. The inset shows the labels for the T atoms in the  $[4^45^2]$  cage, with 2Ge in T6 and T7 positions.





**Fig. 8**  $^{19}\text{F}$  MAS NMR spectra of CSV samples synthesized at  $\text{Ge}_f$  equal to (from bottom) 0.09, 0.1 and 0.2. The resonances around  $-115$  to  $-130$  ppm are assigned to impurities. Spinning side bands are marked with \*. Measured chemical shifts are signaled by arrows.

isolated Ge, empty red squares), clearly showing a trend to form Ge–O–Ge bonds in these silicogermanate systems. The most stable system corresponds to a paired Ge–O–Ge position with Ge in contiguous T6 and T7 positions with F bonded to T6 (Fig. 7, inset), which correspond to the most stable T sites found for 1 Ge systems. In this case, a broad dispersion of chemical shifts is observed, with signals ranging from  $-6$  to  $-50$  ppm, and with the most stable system (T6T7) giving a chemical shift of  $-19$  ppm. We finally studied systems with 3 Ge in the cage ( $\text{Ge}_f = 0.3$ ), with two of them in the more stable and contiguous T6 and T7 positions, varying the location of the third Ge. Once again, energy results evidence the tendency of Ge to cluster since the most stable systems correspond to Ge bonded to T6 in the T6T7T9 and T6T7T10 positions, both forming GeOGeOGe clusters, and giving theoretical  $^{19}\text{F}$  NMR chemical shifts at  $-15$  and  $-27$  ppm, respectively. These results agree well with the experimental chemical shifts found for CSV materials with different  $\text{Ge}_f$  (Fig. 8).

When compared with the experimental  $^{19}\text{F}$  NMR spectra observed for the CSV samples, the overall trends observed in our computational results can provide a tentative explanation for the bands observed (although the exact chemical shifts are not the same, trends can be appreciated). At low  $\text{Ge}_f$  (0.09 and 0.1), two more intense signals are observed at  $-33$  and  $-45$  ppm, which can be assigned to F bonded to Ge in cages with 1 Ge at positions T6 and T7, respectively. When  $\text{Ge}_f$  is increased to 0.2, the former bands decrease their intensity, and a higher intensity in the low-field region is observed, with a band at around  $-20$  ppm now displaying the highest intensity, which according to our computational results should be ascribed to F bonded to Ge in cages with 2 paired Ge at positions T6–T7, which increases its intensity because of the higher Ge content.

## Conclusions

This work shows how the complex interplay between different factors determines the phase selectivity of zeolite crystalliza-

tion when the structure direction exerted by the OSDA is not highly specific and strong enough. When using 2iPr13DMI as the OSDA, several phases might be stabilized but they differ in their intrinsic stability and in the effect of Ge (and F) on that stability. Here, only two default, intrinsically stable, zeolites (ZSM-12 and SSZ-35) were obtained under pure silica conditions. When Ge is added to the synthesis mixtures, up to four additional zeolites can be obtained: HPM-10 (-CLO), HPM-1 (STW, although we never got it as a pure phase), CIT-7 (CSV) and HPM-11 and HPM-6 (intermediate and high  $\text{Ge}_f$  UOS, respectively). These two UOS materials have PXRD patterns that are markedly different from each other and from that of as-synthesized IM-16 (reference UOS material with lower  $\text{Ge}_f$  and symmetries). Calculations suggest 2iPr13DMI may rather comfortably fit in all the obtained phases, providing some moderate stabilization, likely helped by the flexibility, though limited, provided by the isopropyl group.

Calculations also shine some light on  $^{13}\text{C}$  MAS NMR resonances that split in UOS when the  $\text{Ge}_f$  increases, although this is more clearly observed for the methyl groups attached to N than the imidazolium C–H. Finally, multiple overlapped  $^{19}\text{F}$  resonances in the  $-5$  to  $-45$  ppm range in germanosilicate CSV are ascribed to F occluded in the  $[4^{45^2}]$  cage interacting with different crystallographic sites and with different contents and distributions of Ge atoms. This is supported by comparison with similar systems and by DFT calculations.

## Author contributions

H. Y., A. R., Z. R. G. and L. A. V.: synthesis and physico-chemical characterization; L. G. H.: calculations; J. L.: structural characterization; J.-L. P.: synthesis, structural characterization; M. A. C.: supervision, physicochemical characterization, writing-original draft. All the authors discussed the results and revised the manuscript.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors thank the Spanish Research Agency, MCIN/AEI/10.13039/501100011033, and the European Union for funding (Projects PID2019-105479RB-I00, PID2019-107968RB-I00, and PID2021-128141OB-C22, FEDER and TED2021-131223B-I00, “NextGenerationEU”/PRTR). H. Y. is grateful to the China Scholarship Council (CSC) for a PhD fellowship. L. A. V. also thanks the Generalitat Valenciana (PROMETEO CIPROM/2021/007).



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