Ge-Based Hybrid Composites from Ge-Rich Zeolites as Highly Conductive and Stable Electronic Materials

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ABSTRACT: Ge-containing zeolites were used as precursors for the synthesis of highly conductive and stable hybrid electronic materials by postsynthetic thermal treatment of the crystals in the presence of an olefin. Treating the as-prepared Ge zeolites in 1-butene at 700 °C formed a graphitic matrix within and outside the crystals due to the thermal degradation of the organic structure-directing agent inside the pores and the polymerization of the olefin. Importantly, these conditions forced Ge out of the framework, leading to the collapse of the crystalline structure and subsequent formation of metallic Ge nanoparticles distributed either as small, well-dispersed nanoparticles within the silica matrix or larger carbon-coated core–shell Ge@C nanoparticles on the external surface of the carbon–silica composite. Varying the zeolite topology influenced the size of the Ge@C nanoparticles, with those obtained using the multipore zeolite ITQ-22 (IWW, 12 × 10 × 8 rings) featuring smaller sizes (30–60 nm) than those obtained with the large-pore zeolite ITQ-33 (ITT, 18 × 10 × 10 rings) (80–120 nm), where the lack of diffusional limitations increased metal sintering rates. The zeolite topology also influenced the final carbon content and dispersion of the Ge nanoparticles. The best performing Ge-based hybrid material was obtained by thermal treatment of ITQ-22 (Si/Ge = 4) at 700 °C in 1-butene. Unlike Ge-free hybrid controls, an ITQ-22 (Si/Ge = 4) sample treated at 700 °C in 1-butene showed a conductivity value of ~2 S/m (measured at 1 V), which is in the range of a commercially available graphene. The simple methodology presented here is an alternative route for the efficient preparation of highly stable Ge-based hybrid composites with excellent conductivity for potential use as high-capacity electrodes.

1. INTRODUCTION

In light of the electronics revolution that our society has experienced in the last 20 years with a continuous development of information and communication technologies, many research efforts have been devoted to the design of more efficient, durable, and inexpensive electronic materials.1,2 The most studied electronic materials are, in general, based on metals, polymers, ceramics, graphenes, and/or composites.2

In particular, germanium-based electronic materials have received significant attention because Ge presents outstanding electron and hole mobilities, exemplified, for instance, by an electrical conductivity that is 10⁴ times larger compared to that of silicon.3,4 However, the use of metallic Ge in electronic devices has several drawbacks, including high raw material costs and low stability caused by facile oxidation.5,6 To circumvent these limitations, efficient synthesis methods have to be developed not only to maximize Ge utilization but also to adequately protect its external surface to impede undesired oxidation and/or metal sintering.

State-of-the-art strategies to address these challenges include nanostructuring Ge crystals into nanoparticles7,8 or nanowires (NWs),9 their dispersion on large surface supports,10−13 and their encapsulation within porous matrices.14,15 The controlled synthesis of Ge nanoparticles with sizes below 50 nm is important to enhance their electronic performance in different applications. For example, when used as electrodes in lithium-ion batteries, small Ge particles can alleviate the physical strains associated with the Li uptake/release.7,8,16 Unfortunately, the agglomeration of very small Ge nanoparticles under operation conditions is a serious problem that severely compromises their overall electronic capacity, thus hindering their performance in these devices.13,15

The synthesis of Ge@C core–shell particles and the encapsulation of Ge nanoparticles into three-dimensional interconnected graphene networks have been proposed as efficient approaches to prevent undesired agglomeration.13,15 Notably, the presence of aromatic C species in the hybrid Ge@C composites can enhance chemical properties, reduce metal
sintering, and offer superior electron transport and electrical contact between the metal nanoparticles.2,8 The synthesis of these materials typically involves the reduction of a previously deposited Ge precursor with hydrogen and/or inert gases (i.e., N2 or Ar) at high temperatures (above 600 °C),8,15,19 which does not allow the proper control of particle size or degree of encapsulation/dispersion in the final Ge@C composite.

Here, we demonstrate that crystalline Ge-containing zeolites are ideal precursors to synthesize Ge-based hybrid composites with outstanding conductivity and stability. The use of Ge-rich zeolites as precursors has many advantages. First, the wide range of accessible Si/Ge ratios coupled with the labile nature of the tetrahedrally coordinated Ge atoms in the zeolite framework21–23 allows the controlled release of targeted Ge amounts by simple postsynthetic treatments. Second, the microporous topology of the zeolite25 automatically confines these extra-framework Ge species without the need of additional encapsulation protocols. Third, the presence of organic molecules used as organic structure-directing agents (OSDAs) in the pores of the as-prepared Ge-rich zeolites26 can be exploited to prepare homogeneous inorganic–organic conductive matrix composites containing well-dispersed polycyclic aromatic carbons that can be further complemented by postsynthetic thermal treatments in the presence of light olefins.25 Accordingly, the preselection of zeolite pore topology and initial Ge content can be used to control the distribution of extra-framework Ge particles, both in terms of encapsulation and particle size.

We investigated the electrical properties of Ge-based hybrid composites obtained from two Ge-rich zeolites, namely, ITQ-22 (IWW, 12 × 10 × 8 rings)26 and ITQ-33 (ITT, 18 × 10 × 10 rings)27,28 with Si/Ge contents ranging from 2 to 4, that were subjected to different postsynthetic thermal treatments. The hybrid composite obtained from IWW zeolite with a Si/Ge ratio of ∼4 showed a dual distribution of Ge nanoparticles of ∼5 nm encapsulated within the silica matrix and core−shell Ge@C nanoparticles of ∼30–60 nm located on the external surface of the composite. It featured outstanding conductivity compared to that of commercially available graphenes and related C-free and/or Ge-free hybrid composites. Our simple methodology combined with the excellent electrical and chemical properties observed for these Ge-containing composites paves the way for their evaluation as durable, high-capacity electrode materials in diverse devices, including lithium-ion batteries.

2. EXPERIMENTAL SECTION

2.1. Zeolite Synthesis. 2.1.1. ITQ-22 with Si/Ge ∼2 and 4 (IWW-2 and IWW-4, Respectively). The ITQ-22 zeolites were synthesized according to synthesis procedures described in the literature using hexamethonium as OSDA.26 The theoretical molar ratios of the gels are the following: Si/Ge = 2 and 4; B/(Si + Ge) = 0.05; OH/(Si + Ge) = 0.1; OSDA/(Si + Ge) = 0.25; H2O/(Si + Ge) = 20. The final gels were heated at 175 °C in Teflon-lined stainless steel autoclaves for 7 days, and the resultant solids were filtered, washed, and dried at 100 °C.

2.1.2. ITQ-33 with Si/Ge ∼2 (ITT-2). The ITQ-33 zeolite was synthesized according to the synthesis procedure described in the literature using hexamethonium as OSDA.26 The theoretical molar ratios of the gel are the following: Si/Ge = 2; Al/(Si + Ge) = 0.05; OSDA/(Si + Ge) = 0.25; F/(Si + Ge) = 0.3; H2O/(Si + Ge) = 2. The final gel was heated at 175 °C in a Teflon-lined stainless steel autoclave for 1 day, and the resultant solids were filtered, washed, and dried at 100 °C.

2.1.3. Silicate (MFI). The synthesis of the silicate zeolite (MFI) was carried out as follows: 2 g of tetraethylorthosilicate (TEOS) was mixed with 2.3 g of aqueous solution of tetrapropylammonium hydroxide (TPAOH, 25 wt %, Across) and 5 mL of water. The mixture was stirred overnight to favor the hydrolysis of TEOS and, afterward, the evaporation of ethanol and water was controlled until reaching the desired gel concentration. The final gel composition is 1 SiO2:0.3 TPAOH:15 H2O. The gel was loaded into a Teflon-lined stainless steel autoclave, aged for 12 h at 80 °C, and followed by hydrothermal synthesis for 72 h at 175 °C. The resultant solids were filtered, washed, and dried at 100 °C.

2.1.4. Ge-Containing Graphene Materials. For comparison purposes, a Ge/graphene and a Ge/SiO2/graphene have been prepared. For the preparation of Ge/graphene, 49.2 mg of GeO2 (Sigma-Aldrich) has been physically mixed with 180.8 mg of graphene (Strem Chemicals), followed by a reduction with H2 at 700 °C for 2 h. For the preparation of Ge-SiO2/graphene, 74.8 mg of GeO2 (Sigma-Aldrich) has been physically mixed with 229.9 mg of SiO2 (Aerosil) and 45.3 mg of graphene (Strem Chemicals), followed by a reduction with H2 at 700 °C for 2 h.

2.1.5. Postsynthetic Treatments. 2.1.5.1. Treatments with N2. The as-prepared zeolite (0.3 g) was heated at 2 °C/min to the desired temperature (580 or 700 °C) under a N2 flow of ∼100 mL/min and maintained for 6 h. The sample was cooled down under a flow of N2. The as-prepared zeolite (0.3 g) was first heated at 2 °C/min to 80 °C under a N2 flow of ∼100 mL/min and maintained at this temperature for 6 h. Then, the temperature was increased to 150 °C in N2 at 2 °C/min and maintained for 2 h. Next, the temperature was increased to the desired value (580 or 700 °C) in N2 at 2 °C/min, and once this temperature was reached, the N2 flow was kept for 2 h. Finally, a gas mixture formed by 160 mL/min N2 and 4 mL/min butene was passed through the sample at the desired temperature (580 or 700 °C) for 4 h. The sample was cooled down under a flow of N2.

2.2. Characterization. Powder X-ray diffraction (PXRD) measurements were performed with a multisample Philips X-Pert diffractometer equipped with a graphite monochromator, operating at 40 kV and 35 mA and using Cu Kα radiation (λ = 0.1542 nm).

Chemical analyses were carried out by a Varian 715-ES ICP optical emission spectrometer, after solid dissolution in HNO3/HCl/HF aqueous solution. Elemental analyses were performed by combustion analysis using a Eurovector EA 3000 CHNS analyzer.

The morphology of the samples was studied by field-emission scanning electron microscopy (FE-SEM) using a ZEISS Ultra-55 microscope and by field-emission transmission electron microscopy (TEM) using a JEM 2100F microscope.

Raman spectra were recorded at ambient temperature with 514 nm excitation on a Renishaw In Via Raman spectrometer equipped with a CCD detector.

2.3. Conductivity Measurements. All of the conductivity measurements were performed in a homemade system using a Keithley 2601 sourcemeter connected to a PC using LabTracer software that controls the scan rate and also provides data storage capability. The sourcemeter was previously calibrated by using a copper foil as a standard. Briefly, 15 mg of each powdered sample was placed in a sample holder of 13 mm diameter between two polished steel pellets that allow pressurizing during the measurements. The samples were pressed at 8 Ton.cm−2 with a pneumatic press piston, and the electrical conductivity was measured between the top and bottom metallic surface of the press holder at 25 °C. The thickness of the pressed samples was measured by using a Mitutoyo instrument with an accuracy of ±0.0001 in 2 μm.

The electrical resistivity of the materials is calculated using eq 1 and using the resistance (R) calculated at 1 V with the dimensions of the pressed pellet

$$\rho = R \times \text{a/l}$$  (1)

where R is the resistance, l the thickness, and a the area of the circular pellet (13 mm diameter).
Since the electrical conductivity is the reciprocal of the resistivity, it is calculated by taking the inverse of resistivity ($\sigma = 1/\rho$).

For comparison purposes, a commercially available graphene was employed (graphene nanoplatelets aggregates, submicron particles, surface area 300 m$^2$/g, Strem Chemicals, product 06-0225). The mass employed for the electrical measurements both for the controls and zeolite-based materials was 15 mg.

2.3.1. Cyclic Voltammetry Experiments. The electrochemical experiments were performed using a Versastat 3 electrochemical workstation. Two 1 cm$^2$ AISI-304 stainless steel meshes, mesh number 300 and 0.035 mm wire diameter, coated by drop-casting with a suspension of the material (20 wt % of active material and 3.6 wt % polyvinylpyrrolidone in ethanol) were used as electrodes (125 $\mu$g of active material on each electrode, 250 $\mu$g of total active material mass). Electrochemical studies were carried out in 300 $\mu$L of 1 M lithium perchlorate solution in propylene carbonate and using an EL-CELL PAT-CELL commercial electrochemical cell in a two-electrode configuration with gold current collectors; both electrodes were separated by a 21 mm Whatman glass microfiber filter GF/A. The cyclic voltammograms (CVs) were recorded at a scan rate of 50 mV/s. The galvanostatic charge/discharge (GCD) curves for EDLCs were obtained at 2 A/g.

3. RESULTS

3.1. Synthesis and Characterization of Ge-Based Hybrid Composites from Ge-Rich Zeolites. Ge atoms in tetrahedral coordination within the zeolite frameworks are very labile, particularly for Ge-rich zeolites with Si/Ge ratios below 5.$^{22}$ While typically considered unfavorable for catalytic applications, this feature is optimal for creating well-dispersed extra-framework Ge particles for electronic applications since their size and dispersion can be readily tuned depending on the initial physicochemical properties of the as-prepared Ge zeolites and the postsynthetic treatments carried out starting from the as-prepared Ge zeolites.

We selected ITQ-22 (IWW)$^{26}$ and ITQ-33 (ITT)$^{27,28}$ for this study because both zeolites can be prepared under identical Ge-rich conditions (as low as Si/Ge~2).$^{28}$ In addition, both frameworks feature multipore systems involving interconnected large and medium pores for IWW (12 x 10 x 8 rings, with pore openings of ~6.5 x 5.0 x 3.5 Å) and extra-large and medium pores for ITT (18 x 10 x 10 rings, with pore openings of ~12.3 x 5.5 x 5.5 Å), thereby resulting in similar pore connectivities but very different framework densities (16.1 and 12.4 T/1000 Å$^3$, respectively) (see Figure 1).

The as-prepared zeolites with Si/Ge = 2 (denoted IWW-2_a.p. and ITT-2_a.p.) showed the characteristic PXRD patterns of the IWW and ITT topologies, respectively (see Figures 2 and 3). The chemical and elemental analyses of both materials revealed similar Si/Ge molar ratio (~2.5, see Table 1) and organic content (~6.6~7.8% C and ~1.4~1.6% N), which corresponds to intact hexamethonium molecules trapped within the zeolite crystals during the hydrothermal crystallization process. Scanning electron microscopy (SEM) images showed crystal aggregates formed by stacked thin layers of
of ∼3–4 μm for IWW-2_a.p. (see Figure 4) and arranged needles of ∼2–3 μm for ITT-2_a.p (see Figure 5).

Optimal Ge-based hybrid devices require uniform metallic Ge nanoparticles dispersed over a conductive polycyclic aromatic C source, such as graphitic or graphenic species, to favor the electron transport and electrical contact between the Ge nanoparticles. Considering the presence of the occluded organic molecules within the pores of the as-prepared IWW and ITT zeolites, we hypothesized that a simple thermal reduction at high temperatures (i.e., 580–700 °C) would fulfill both requirements by simultaneously forcing the extraction and aggregation of framework Ge species into well-dispersed metallic nanoparticles and the generation of aromatic C-like entrapped species from ODSA degradation. To evaluate the most efficient postsynthetic method to form well-defined Ge particles from the above as-prepared materials, we have first focused on the IWW-2 material. The crystalline structure of the IWW-2 zeolite remained almost unaltered when treating it at 580 °C with N₂ (see IWW-2_N₂-580 °C in Figure 2), while a partial amorphization was observed when treated with N₂ at higher temperatures (i.e., 700 °C, see IWW-2_N₂-700 °C in Figure 2). We note that the final organic content in the treated material was only ∼1 C wt % (see IWW-2_N₂-580 °C and IWW-2_N₂-700 °C in Table 2), which is much lower than that in the as-prepared IWW-2 material (∼7.8 wt % C, see IWW-2_a.p. in Table 2), even though the thermal treatment was performed under nonoxidizing conditions. This observation has also been reported in the literature for other medium- and large-pore silicates and is

<table>
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<tr>
<th>Zeolite</th>
<th>wt % N</th>
<th>wt % C</th>
<th>Resistivity (Ω·m)</th>
<th>Conductivity (S/m)</th>
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<tr>
<td>IWW-2_a.p.</td>
<td>1.58</td>
<td>7.76</td>
<td>1.49 × 10⁵</td>
<td>6.72 × 10⁻⁶</td>
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<tr>
<td>IWW-2_N₂-580 °C</td>
<td>0.00</td>
<td>0.93</td>
<td>5.15 × 10⁴</td>
<td>1.94 × 10⁻⁵</td>
</tr>
<tr>
<td>IWW-2_C₂⁺-580 °C</td>
<td>0.00</td>
<td>1.72</td>
<td>1.11 × 10⁶</td>
<td>9.04 × 10⁻⁷</td>
</tr>
<tr>
<td>IWW-2_N₂-700 °C</td>
<td>0.00</td>
<td>1.35</td>
<td>1.08 × 10⁵</td>
<td>9.25 × 10⁻⁸</td>
</tr>
<tr>
<td>IWW-2_C₂⁺-700 °C</td>
<td>0.00</td>
<td>7.65</td>
<td>0.45</td>
<td>2.24</td>
</tr>
<tr>
<td>ITT-2_a.p.</td>
<td>1.37</td>
<td>6.64</td>
<td>∞</td>
<td>0.00</td>
</tr>
<tr>
<td>ITT-2_C₂⁺-700 °C</td>
<td>0.00</td>
<td>22.94</td>
<td>0.81</td>
<td>1.23</td>
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<tr>
<td>IWW-4_a.p.</td>
<td>1.42</td>
<td>7.04</td>
<td>3.50 × 10⁵</td>
<td>2.85 × 10⁻⁶</td>
</tr>
<tr>
<td>IWW-4_C₂⁺-700 °C</td>
<td>0.00</td>
<td>12.95</td>
<td>0.53</td>
<td>1.90</td>
</tr>
<tr>
<td>IWW-4_C₂⁺-700 °C (6 months)</td>
<td>0.00</td>
<td>12.95</td>
<td>0.56</td>
<td>1.77</td>
</tr>
<tr>
<td>MFI-∞_C₂⁺-700 °C</td>
<td>0.00</td>
<td>6.65</td>
<td>3.17</td>
<td>0.32</td>
</tr>
<tr>
<td>Graphene</td>
<td>0.34</td>
<td>2.98</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ge/graphene</td>
<td>1.06</td>
<td>0.94</td>
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<td>Ge-SiO₂/graphene</td>
<td>348.4</td>
<td>2.87 × 10⁻⁵</td>
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</tbody>
</table>
attributed to the combination of pore dimensions and the structure of the OSDA used as template.29

To increase the carbon content in the final materials, we decided to perform the thermal treatments in presence of a light olefin, such as 1-butene, which is known to undergo oligomerization to form carbon deposits within porous materials at these conditions (see Experimental Section for details).30,31 As seen in Figure 2, the crystalline structure of the IWW-2 sample was mostly retained after being exposed to 1-butene at 580 °C (see IWW-2_C4-580 °C), but it fully collapsed after being treated with 1-butene at 700 °C (see IWW-2_C4-700 °C). In both cases, we observed a very intense peak centered at 2θ of 27.5° in their PXRD patterns that corresponds to the formation of metallic germanium particles.19 Nanoparticle formation was confirmed with FE-SEM, which revealed Ge particles ranging between 50 and 80 nm on the external surface of the composites (see IWW-2_C4-580 °C and IWW-2_C4-700 °C in Figure 4), with a significantly higher concentration for the sample treated at 700 °C.

Interestingly, a deeper inspection of the IWW-2_C4-700 °C sample by HR-TEM microscopy revealed a dual Ge distribution, consisting of very small Ge nanoparticles of ~5 nm encapsulated within the silicate particles, and larger Ge nanoparticles of 50–80 nm deposited on the external surface of the silicates (see Figure 6a). The larger Ge nanoparticles presented a Ge@C core–shell structure with a uniform carbon-coating layer of 5–10 nm thickness (see Figure 6b,c). This carbon overlayer is critical to prevent their undesired agglomeration and oxidation to GeO2 as well as to minimize the mechanical stress induced by the volume change of Ge in particular applications (i.e., Li-ion batteries).13,32

Elemental analysis on the IWW-2_C4-700 °C sample revealed a carbon content of ~7.7 wt %, which is substantially higher than the organic content obtained for the IWW-2 samples treated under inert conditions (~1 wt % C, see Table 2). Raman spectra of the carbonaceous species formed in the IWW-2_C4-700 °C sample showed two well-defined signals centered at ~1580 and 1350 cm\(^{-1}\), which are characteristic features of the sp\(^2\) carbon atoms of graphitic layers (G band) and disordered carbon or defective graphitic structures (D band), respectively (see IWW-2_C4-700 °C in Figure 7).33 The higher intensity observed for the G band in the Raman spectrum suggests a larger graphitic order degree within the IWW-2_C4-700 °C sample.

From the previous treatments on the IWW-2 material, one could conclude that the thermal treatment at 700 °C in presence of 1-butene allows obtaining a very interesting dual Ge dispersion with relatively high graphitic C content along the entire sample. Thus, we performed the same postsynthetic treatment at 700 °C in the presence of butene to evaluate the properties of the ITT-2 zeolite. Akin to IWW-2, the crystalline structure of ITT-2 material also collapsed and its PXRD pattern featured an intense peak centered at 2θ of 27.5°, indicating the formation of metallic Ge particles (see the PXRD pattern for ITT-2_C4-700 °C in Figure 3). FE-SEM and HR-TEM images show the formation of larger Ge particles than those observed before for IWW-2, with average sizes ranging between 80 and 120 nm (see ITT-2_C4-700 °C in Figures 5 and 8). It could be envisioned that the combination of larger pores and lower framework density in the ITT zeolite (∼12 Å and 12.4 T/1000 Å\(^3\)) would offer fewer diffusion limitations for the extra-framework Ge atoms to move and agglomerate once extracted from the zeolite framework compared to IWW zeolite (∼6.5 Å and 16.1 T/1000 Å\(^3\)). However, the different distribution of the secondary building units (SBUs) that preferentially host Ge atoms (i.e., D4Rs and 3 rings) along the ITT and IWW frameworks could also present an important influence on the achieved Ge nanoparticles. It is important to note that the extra-large-pore ITT zeolite presents almost equally distributed D4Rs along the entire framework, whereas the IWW zeolite contains exclusively these D4Rs in the interlayer region of high-silica layers. Furthermore, the more open pore topology of the ITT zeolite favors olefin oligomerization reactions, resulting in a remarkably larger C contents (∼23 wt % C, see ITT-2_C4-700 °C).
700 °C in Table 2) compared to those of the treated IWW-2 sample.

Given that cost is a very important issue when designing Ge-based hybrid materials for electronic devices, we have investigated the use of IWW precursors presenting half of the Ge content (Si/Ge~4, IWW-4, see synthesis details in the Experimental Section). Although a decrease in Ge content could negatively impact the conductivity properties, we reasoned that a better dispersion along the IWW framework could result in an improved uniformity after the heat treatment, ultimately enhancing electronic conduction. The as-prepared IWW-4 material showed the characteristic PXRD pattern of the IWW framework (see IWW-4_a.p. in Figure 3), and, similar to IWW-2, the crystals were also mainly formed by stacked thin layers of ~4 μm (see FE-SEM images for IWW-4_a.p. in Figure 5). After treating the IWW-4_a.p. material at 700 °C in the presence of 1-butene, the initial crystalline framework collapsed and revealed the peak at 2θ of 27.5° associated with metallic Ge species (see IWW-4_C4°-700 °C in Figure 3). HR-TEM images showed the formation of homogenous Ge@C particles with average sizes between 30 and 60 nm (see FE-SEM and HR-TEM images in Figures 5 and 8), together with the presence of very small trapped Ge nanoparticles of ~5 nm within silicate particles (see Figure 8).

3.2. Conductivity Performance. We surmised that the synergy between large and small metallic Ge nanoparticles dispersed in a conductive graphitic matrix would induce remarkable conducting properties to the final material, even if the original crystalline structure of the zeolite was not maintained. As seen in Table 2, the measured electrical conductivities at a given voltage (1 V) clearly show an influence of the postsynthetic treatments. Unsurprisingly, the as-prepared Ge-containing zeolites behaved as insulators rather than conductors (0.1−0.3 MΩ·m), with negligible measured conductivity values (see IWW-2_a.p., ITT-2_a.p. and IWW-4_a.p. in Table 2), given that isolated Ge species in the zeolite

Figure 8. STEM and HR-TEM images of the Ge-containing ITT-2 (Si/Ge~2) and IWW-4 (Si/Ge~4) zeolites after being postsynthetically treated with butene at 700 °C.

Figure 9. I/V plot comparing the electrical conductivity of the Ge-rich IWW-2 zeolite (Si/Ge~2) after being treated with N2 or butene at 700 °C.
frameworks together with the occlusion of nonaromatic organic molecules within the zeolite pores should not conduct charge. The resistivity values for the Ge-containing zeolites after being treated at 580 °C (either in N2 and 1-butene) are remarkably lower compared to those of the as-prepared materials (<5 MΩ·m, see IWW-2_N2-580 °C and IWW-2_C4−780 °C in Table 2), but their conductivity values still remained pretty low (<1 × 10−6 S/m, see IWW-2_N2-580 °C and IWW-2_C4−780 °C in Table 2). The slight improvement in their electrical conductivity could be ascribed to the presence of some aromatic-based C species (~1–2 C wt %) after the thermal treatments at 580 °C.

In contrast, the measured conductivities on the samples treated at 700 °C, in particular, those treated in presence of 1-butene, are substantially higher, achieving values of 1−3 S/m (see IWW-2_C4−700 °C, ITT-2_C4−700 °C and IWW-4_C4−700 °C in Table 2), which are comparable to well-established electrical conductors, such as commercially available graphene (~3 S/m, see Table 2). In fact, significant differences can be observed when comparing the electrical conductivity profiles for the same Ge-containing sample, IWW-2, after being treated at 700 °C, but under different atmospheres (N2 or 1-butene). As seen in Figure 9, the I/V plot for the IWW-2 material treated at 700 °C with butene performs as an outstanding conductor material, whereas the IWW-2 material treated at 700 °C with N2 mostly performs as an insulator material. However, despite its low conductivity, a deeper inspection of the I/V curve for the IWW-2_N2-700 °C revealed a semiconductor-type behavior, where the presence of two well-defined regimes indicates the performance of a resistor at low voltages and, at least partially, the performance of a conductor material at high voltages. Similar semiconductor behaviors have been described for other metal-containing zeolites, such as Co-FAU.34 We hypothesize that a better metallic Ge nanoparticle dispersion along the sample combined with the higher graphitic C content is responsible for the higher conductivity measured for the IWW-2_C4−700 °C compared to the one for IWW-2_N2-700 °C.

To unravel the influence of these two parameters, we compared the electronic behavior of the other two Ge-containing materials, ITT-2 and IWW-4, after being treated with 1-butene at 700 °C. As seen in Figure 10, the I/V profile for ITT-2_C4−700 °C also performed as an excellent conductor material, but with lower current intensities at a given voltage than IWW-2_C4−700 °C, despite the remarkably higher aromatic C content within the treated ITT-2 material (22.9 and 7.6 wt % C for ITT-2_C4−700 °C and IWW-2_C4−700 °C, respectively, see Table 2). This result clearly indicates that the better Ge dispersion along the Ge-based hybrid material has a stronger influence on the electronic properties of the Ge-composite than graphitic C content. This conclusion is further supported by the very similar I/V characteristics obtained with the sample prepared with much less Ge content, IWW-4, compared to the Ge-rich IWW-2 sample after being treated with 1-butene at 700 °C (see IWW-2_C4−700 °C and IWW-4_C4−700 °C in Figure 10). The enhanced Ge dispersion in IWW-4_C4−700 °C that was observed by microscopy (see Figures 5, bottom and 8, bottom) is consistent with the high conductivity values even when containing only half of the Ge. Notably, this sample maintained virtually identical conductivity profiles even after 6 months of being treated and stored under ambient conditions, thus revealing an outstanding stability. Based on these experimental observations, the ability to maximize and control the dispersion of the Ge particles within the nanometer scale along the Ge composites is currently a main research topic in our laboratories to improve their conductivity performances.

For comparison purposes, a medium-pore Ge-free Silicalite-1 material (MFI structure) was synthesized to evaluate its conductivity behavior after being treated with 1-butene at 700 °C (see synthesis details in the Experimental Section and the PXRD pattern in Figure 11). This sample had a C content of ~7 wt %, predominantly in the form of graphitic species with high-order degree, as revealed by Raman spectroscopy (see Figure 7). However, this material showed a considerably lower conductivity compared to that of the treated Ge-containing zeolites (see Figure 10), further demonstrating the necessity of having well-dispersed metallic Ge nanoparticles to obtain efficient conductor materials.

In addition, two Ge-based graphene materials have been prepared also for comparison purposes. On the one hand, we have prepared a Ge-SiO2/graphene, with a similar chemical composition to that of the IWW-4_C4−700 °C sample (21.4 wt % GeO2, 65.7 wt % SiO2, and 12.9 wt % graphene, see the Experimental Section for details). This Ge-SiO2/graphene sample gives a conductivity value of ~2.87 × 10−3 S/m, which is substantially lower than that measured for IWW-4_C4−700 °C (~1.9 S/m, see Table 2). On the other hand, we have also prepared a Ge/graphene sample containing an overall Ge content similar to the one present in IWW-4_C4−700 °C, but without including SiO2 in the formulation (21.4 wt % GeO2 and 78.6 wt % graphene, see the Experimental Section for details). This Ge/graphene sample gives a conductivity value of ~0.94 S/m, which is lower than the values measured for the pure graphene (~2.98 S/m) and for the IWW-4_C4−700 °C sample (~1.90 S/m) (see Table 2). These results put in relevance the importance of the initial Ge-rich zeolite precursor to properly allow the formation of Ge composites with unique physicochemical properties and, consequently, improved conductivity performances.

![Figure 10. I/V plot comparing the electrical conductivity of the different Ge-based hybrid composites and pure silica MFI after being treated with butene at 700 °C.](image-url)
As a preliminary study to evaluate the reversible capacity and cycling capability in a battery cell of these Ge composites, the IWW-4_C4=-700 °C sample has been studied as an electrode in electrochemical double layer capacitors (EDLCs). In this regard, the electrochemical performance of the material has been studied by cyclic voltammetry at room temperature, using 1 M lithium perchlorate in propylene carbonate as an electrolyte. As seen in Figure 12, the shape of the curves is similar after 30 cycles and only a slight change in the current is observed, indicating good stability of the material for application in electrochemical energy storage devices.

4. CONCLUSIONS

We subjected two different Ge-rich zeolites, ITQ-22 (IWW) and ITQ-33 (ITT) to postsynthetic treatments at different temperatures (580 and 700 °C) and controlled atmospheres (N₂ and 1-butene) and studied their conductivity behavior. The resulting Ge-based hybrid composites were analyzed by different techniques to reveal the nature, size, dispersion, and stability of the metallic Ge nanoparticles in the final structure. The dispersion of metallic Ge nanoparticles was influenced by the postsynthetic thermal treatment, the pore topology of the parent zeolite, and the initial Si/Ge molar ratio. We showed that the IWW zeolite with an initial Si/Ge molar ratio of ∼4 featured electrical conductivity performance commensurate to that of graphene after being treated with 1-butene at 700 °C. The postsynthetic treatment of Ge-based zeolite precursors opens an interesting research pathway to design efficient Ge-based hybrid composites, which could have important implications in the preparation of electrodes for lithium-ion batteries.

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Notes
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