Hierarchically organized porous materials can provide multi-dimensional spatial networks on different length scales with improved characteristics relevant to molecular diffusion.[13] Zeolites that are microporous crystalline materials having pores and channels at molecular dimensions are of great importance for industrial applications. However, the presence of only micropores in zeolite frameworks often limits the molecular diffusion and therefore, restricts the transport of bulky molecules. This problem can be resolved by shortening the effective diffusion path lengths, which has been achieved by miniaturizing zeolite crystals,[3] delaminating or exfoliating the zeolite particles,[4] and introducing mesopores into zeolite particles.[5–11] Among these solutions, the fabrication of hierarchical zeolites with meso- and microporosity is of interest because it combines intrinsic micropores with bypass-interconnected mesopores, and therefore, enhances both the micropore accessibility and molecular traffic within zeolite particles.[1,7b,12]

Hierarchical zeolites have been produced using several techniques, including top-down desilication by alkali post-synthetic treatment[6] and bottom-up directed assembly by hard[7] or soft[8–11] templates. The hard-template method requires multistep procedures and is therefore unfavorable for large-scale production. Alternatively, the direct introduction of organic mesopore-generating agents (mesoporogens) during zeolite synthesis can create uniform mesopores. The use of such mesoporogens is currently one of the most promising methods for the single-step construction of hierarchical zeolites. Progress has been made using well-designed mesoporogens composed of long hydrophobic chains and hydrophilic zeolitic structure-directing groups to generate zeolites with tunable mesoporosity[9a,10] and to direct the hierarchical assembly of zeolite nanosheets, yielding mesoporous zeolites with house-of-cards-like structures.[5a,9c,11] These hierarchical nanosheets showed excellent catalytic performance in several important reactions because the presence of thin layers with specific crystalline faces facilitates catalysis at the exteriors or pore mouths.[13] Such mesoporogens are probably necessary for the direct, single-step synthesis of hierarchical zeolites.

Herein we report an alternative, mesoporogen-free approach for the construction of hierarchically organized MFI zeolites by sequential intergrowth using a simple organic structure-directing agent (OSDA). The selection of an appropriate OSDA and optimized synthesis conditions that can form plate-like zeolites with enhanced 90° rotational intergrowths seems to be a key to achieving a hierarchical structure with three classes of porosity in one structure: the intrinsic microporosity of the zeolite framework together with mesoporosity existing within the zeolite plates and macro-porosity stemming from the complex intergrown structure.

Epitaxial and rotational intergrowths are commonly observed in many zeolites.[14] We have hypothesized that by engineering the zeolite intergrowths, hierarchically organized zeolites can be constructed without the need for mesoporogens. In particular, we have focused on the MFI zeolite because it is an excellent catalyst in many industrial processes[2b,15] and a promising material for membrane separations.[16] MFI zeolite that contains sinusoidal channels along the a axis interconnected with straight channels along the b axis is often formed with 90° rotational intergrowths, in which substantial (h00) faces are epitaxially overgrown on the (0k0)
faces. The tetrabutylphosphonium (TPA) cation is the most common and probably the most energetically favored OSDA for MFI zeolite formation. TPA cations are situated at the intersection of the straight and sinusoidal channels. Therefore, we have considered that organic molecules with slight differences from TPA molecules could still direct the crystallization of MFI zeolites but could also enhance the formation of rotational intergrowths. Very recently, in a similar strategy, self-pillared zeolite nanosheets containing micro- and mesoporosity were synthesized by repetitive branching using tetrabutylphosphonium cations as an OSDA. Although the hierarchical zeolites could be obtained without the use of mesoporogens, the use of a phosphonium-based OSDA would affect the physicochemical properties of the resultant zeolites because the decomposition of the OSDA by conventional thermal calcination cannot totally remove phosphorus from the zeolite structure, and therefore, a certain amount of phosphorus debris may remain in the zeolites.

Among the many OSDAs for MFI zeolite, we have focused herein on dimers of the TPA cation, (C₅H₁₁)₃N⁺-(CH₅)₅N⁻(CH₃)₃, because such diquaternary ammonium cations with n = 5–7 are situated in the framework with the N-N chains being better fitted along the straight channels than along the sinusoidal channels. This limits the degree of molecular motion within the framework, particularly in the case of OSDAs with shorter alkyl-chain spacers. The TPA dimer with n = 5, hereafter denoted as diquat-C₅, possesses an intramolecular N-N distance of 0.7 nm, which is much shorter than the distances between two intersections along the straight (1.0 nm) and sinusoidal (1.2 nm) channels. Although previous molecular mechanics simulations suggested that diquat-C₅ is not readily accommodated in the zeolite channels, it can still direct the formation of MFI zeolites. Accordingly, the diquat-C₅ cations have to adapt their intramolecular N-N distance to be occluded intact within the MFI framework, perhaps along the straight channel, generating structural constraints or stresses, and thereby yielding framework distortion, as suggested by synchrotron X-ray analyses. This event limits crystal growth along the b axis, resulting in an MFI zeolite with a plate-like morphology but with a few intergrowths. To generate hierarchical zeolites composed of such plate-like crystals, zeolites should be synthesized under conditions with more pronounced surface nucleation. In a previous report, zeolites were synthesized from very dilute solutions in the presence of seed materials. In addition, both SAED patterns reveal that the crystals probably constructed by the aggregation of nanoparticles (Figure 1c,d). The dot patterns of selected-area electron diffraction (SAED) of both the main plate (Figure 1e) and the crystal flake (Figure 1f) indicate single-crystalline materials. In addition, both SAED patterns reveal that the crystals are viewed along the [010] zone axis, suggesting that the b axis (i.e., the straight channels) runs perpendicular to the surface of each crystal plate.

To obtain further insights into the organization process, the textural evolution of the obtained silicalite-1 crystals was investigated, as shown in Figure 2. The first visible Bragg diffraction appears after 135 hours of crystallization (see Figure 2a). Before the onset of crystallization, the FE-SEM image shows that the products are nanoparticle aggregates (Figure 2b). No crystal-like, faceted particles were observed at this stage, consistent with the corresponding XRD pattern. After 135 hours (Figure 2c), the plate-like particles were first seen together with the nanoparticle aggregates. These plate-like particles were probably silicalite-1 crystals, as suggested...
by the XRD pattern. The unusual intergrown structure developed gradually during the course of crystallization (Figure 2d,e).

The calcined products synthesized at 130 °C for 150 hours were further characterized by TEM. The SAED patterns of representative regions of the nanoparticle aggregates are shown in the insets of Figure 3a and Figure S1a, confirming that the resulting nanoparticles are amorphous. High-resolution TEM (HR-TEM) images (Figure 3b; Figure S1b) further supported the amorphous nature of these nanoparticles. As expected, in contrast, plate-like particles exhibited SAED patterns common to MFI zeolite (Figure 3c; Figure S1c; insets). Lattice fringes indicative of crystalline orders were clearly observed in the HR-TEM images (Figure 3d; Figure S1d), corroborating that the plate-like particles are silicalite-1 zeolites.

A scheme of the formation of hierarchically organized MFI zeolites based on the above results is shown in Figure 4. In the early stage of crystallization, the amorphous nanoparticles, probably hydrated diquat-C5-silica composites, are formed. These composite nanoparticles aggregate to form very thin MFI zeolite plates. Although the formation process is still unclear, viable zeolite nuclei may be formed by the oriented-aggregation mechanism, the solid-state-like disorder-to-order transformation, reorganization at the solid-liquid interface, or a combination thereof. Under the present synthesis conditions, the 90° rotational intergrowths are enhanced and developed during the crystal growth process. The intergrown crystal flakes are also made of nanoparticle aggregates. The composite nanoparticles and the soluble species together supply the precursors for crystal growth, and subsequently, zeolite crystals evolve into the hierarchical structure.

We considered that increasing the synthesis temperature could enhance the crystal intergrowth, thereby enhancing the formation of the hierarchical structures. Therefore, silicalite-1 zeolites were also synthesized at 150 °C and 175 °C. Clearly, the hierarchical structure is more pronounced after synthesis at 150 °C, as shown in Figure 5a-c. The powder

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**Figure 2.** a) Powder XRD patterns of the products obtained at 130 °C after 120–264 h. FE-SEM images of the products after reaction times of b) 120 h, c) 135 h, d) 150 h, and e) 184 h.

**Figure 3.** a,c) TEM and b,d) HR-TEM images of the calcined products synthesized at 130 °C for 150 h. HR-TEM images in (b,d) correspond to those in (a,c), respectively. Insets in (a,c) show the corresponding SAED patterns taken from the circled areas.

**Figure 4.** Schematic representation of the formation of hierarchically organized MFI zeolite. a) Amorphous nanoparticles, probably hydrated diquat-C5-silica composites, form from a solution. These nanoparticles aggregate to form MFI zeolite plates. b) The 90° rotational intergrowths develop during crystal growth, in which each intergrown crystal flask also forms from nanoparticle aggregates. c) The nanoparticles and the soluble species in solution together supply the precursors for crystal growth, and finally, zeolite crystals evolve into the hierarchical structure.
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XRD patterns (Figure 5d) confirmed that the products are MFI zeolite. Interestingly, plate-like silicalite-1 zeolites with less-intergrown structures were obtained when the synthesis temperature was further raised to 175°C (Figure S2). We speculate that Ostwald’s law of successive reactions is, at least in part, responsible for the formation of the less-intergrown silicalite-1 at 175°C. This law essentially describes the consumption of the less-stable phase, the subsequent formation of a thermodynamically more stable phase, and so on, until the most stable phase is formed. At 175°C, such successive ripening proceeds rapidly, and therefore, the resulting zeolites did not form the hierarchical structure. Observation of such less-intergrown silicalite-1 synthesized at 130°C and 150°C after prolonged reaction times (Figure S3) additionally supported the theory of Ostwald ripening. These results also suggest that the obtained silicalite-1 was the result of a kinetically controlled synthetic pathway.

As is well known, quaternary ammonium cations can undergo a Hofmann elimination, particularly at high temperature. The solid-state $^1$H-$^1$C cross-polarization magic-angle spinning (CP/MAS) NMR spectrum of the silicalite-1 synthesized at 150°C reveals that the diquat-C5 cations are occluded intact in the zeolite product (Figure S4). In contrast, the product synthesized at 175°C showed additional signals, probably arising from Hofmann elimination products. For example, the signals at $\delta \approx 30, 118, \text{ and } 137$ ppm can be assigned to the center carbon of 1,4-pentadiene, the $\beta$-olefinic carbon (CH$_2$=CH=CH$_2$), and the $\alpha$-olefinic carbon (CH$_2$=CH=CH$_3$), respectively. This partial decomposition of the diquat-C5 SDA seems to affect the morphology of the resulting zeolite products in addition to Ostwald’s ripening.

Nitrogen sorption isotherms of the silicalite-1 sample show a large uptake at low relative pressures, a hysteresis loop at moderate relative pressures, and a slight uptake at higher relative pressures, indicating the presence of micro-, meso-, and macroporosity, respectively (Figure 5e). The apparent specific Brunauer–Emmett–Teller (BET) surface area and the total pore volume (at $P/P_0 = 0.99$) were calculated to be $490 \text{ m}^2 \text{ g}^{-1}$ and $0.35 \text{ cm}^3 \text{ g}^{-1}$, respectively. The external surface area and micropore volume were estimated by the $t$-plot method to be $215 \text{ m}^2 \text{ g}^{-1}$ and $0.11 \text{ cm}^3 \text{ g}^{-1}$, respectively. The pore-size distribution and cumulative pore volume calculated by non-local density functional theory (NL-DFT) are depicted in Figure 5f. The pore-size distribution was relatively narrow with a diameter centered at 3.3 nm. The SEM, TEM, and nitrogen physisorption results indicate that the silicalite-1 zeolite possessed the intrinsic microporosity of a zeolite framework together with mesoporosity within the zeolite plates and macroporosity stemming from the complex intergrown structure.

We also explored the synthesis of aluminosilicate MFI (ZSM-5) by adding an aluminum source to the starting synthesis solution at different Si/Al ratios. The powder XRD patterns confirmed the formation of MFI zeolite (Figure S5a). The obtained ZSM-5 zeolites showed a similar morphology, indicating that the presence of Al did not inhibit the formation of this unique structure (Figure S5c,d). Solid-state $^{27}$Al MAS NMR spectra of the calcined samples (Figure S5b) suggested that Al atoms are located at the tetrahedral positions of the framework, although non-framework, octahedral Al species also appear. The nitrogen sorption isotherms in Figure S6a show that the obtained ZSM-5 also possesses micro-, meso-, and macroporosity. NL-DFT analyses confirmed the presence of mesoporosity (Figure S6b), although the mesopore size distributions were less uniform than that of the silicalite-1 counterpart (see above). These hierarchical ZSM-5 zeolites are potentially promising for the catalytic conversion of bulky molecules and biomass processing. Such topics are vital and merit a detailed investigation in future studies.

In summary, we have demonstrated a new approach for the construction of hierarchically organized zeolites by sequential intergrowth. For the first time, a complex and unusual morphology was observed for MFI zeolites. These zeolites possessing micro-, meso-, and macroporosity have been produced by a simple, single-step, hydrothermal synthesis using a small OSDA without meso- or macroporogens. The use of an OSDA that is imperfectly fitted to the zeolite framework, yet can direct its formation, is thought to be responsible for the formation of very thin zeolite plates with enhanced 90° rotational intergrowth, thereby for the construction of hierarchically organized structure. This concept would be applicable to other zeolite frameworks and related...
porous crystalline materials that can form such rotational intergrowths.

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