Zeolites are crystalline microporous aluminosilicates whose unique pore and channel systems are the reason for their important role in catalysis, separation, and ion exchange. This work focuses on the morphology and structure of a natural zeolite, mordenite. Our samples were collected at Lengyendi-galya (Gyökereszto) in the Mátra Mountains (NE-Hungary). Zeolite samples were investigated by X-ray powder diffraction, scanning, and transmission electron microscopy. Mordenite has typically fibrous appearance and occurs in association with other zeolites such as heulandite and chabazite. Based on intense streaks and superlattice reflections in selected area electron diffraction patterns, we identified planar faults in the structure. A single fault produces a dachiardite-type structural slab within the mordenite lattice, reducing its channel size.

Keywords: Natural mordenite, structure, diffraction, faults

1. Introduction

Zeolites have complex aluminosilicate frameworks. These frameworks contain large channels and interconnected cages. We use zeolites in gas and petroleum industry, water softening, sewage treatment, agriculture, paper production, and radioactive waste treatment, as well as in construction materials. The structural channels and voids are occupied by loosely bound cations and water molecules that can be removed and replaced without disrupting the tetrahedral framework. This unique – so-called, zeolitic – property makes a zeolite structure the best candidate to perform cation exchange, adsorption, and molecular sieving (passing a gas or liquid through a zeolite). Zeolites can be used in dehydration and rehydration processes and can be resistant to high energy radiation [1].

The mordenite group includes the minerals mordenite, dachiardite, epistilbite, ferrierite, and bikitaite [2, 3]. Zeolites in the mordenite group are built by five-membered rings of tetrahedra – this is called the secondary building unit – and puckered sheets made of six-membered rings of tetrahedra parallel to (100), which ensure a perfect prismatic cleavage (Figure 1). Parallel to the c axis, there are channels formed by twelve-membered rings (12MRc: aperture, 6.5 × 7.0 Å) and compressed eight-membered rings (8MRc: aperture, 5.7 × 2.6 Å). There are another eight-membered rings, and channels also occur parallel to b (8MRb: aperture, 3.4 × 4.8 Å) (Figure 2). In dachiardite, ten-membered rings are parallel to b (Figure 1) and there are also compressed eight-membered rings
parallel to the $c$ axis [4]. Extraframework cations in mordenite occupy mainly three sites. Two of these sites are close to the four-membered Al-enriched rings (light-gray shaded tetrahedra in Figure 1) and are located in the connecting channels parallel to the $b$ axis; the A site is in the center of the compressed eight-membered ring channels, whereas the D site is near the center of the eight-membered ring, connecting to the wide channel. The E site is in the large channels (Figure 2).

The ideal mordenite composition is $(Na_2,K_2,Ca)_4[Al_8Si_4O_96] \cdot 28H_2O$; the structure is orthorhombic with space groups of $Cmc2_1$ ($a = 18.11$, $b = 20.46$, $c = 7.52$ Å, $Z = 1$) [5]. The symmetry lowering from $Cmcm$ to $Cmc2_1$ occurs when locations of extraframework cations are asymmetric [6].

Mordenite is one of the zeolites which have very good absorption and separation properties. For this reason, Barrer [7] examined this type of zeolite framework and synthesized mordenite with a small port (“port” meaning the channels in the structure). Therefore, two different structural varieties are now distinguished – large-port and small-port mordenite – by the accessibility of 12-membered ring channels (Figure 1). In the small-port mordenite, molecules fit with sizes less than 4.2 Å in diameter in the channels, and in the large-port mordenite, the channels are accessible for molecules larger than 4.2 Å [8]. Mordenite as a mineral typically has the properties of small-port mordenite. The widely accepted explanation for the different port sizes is that stacking faults disrupt the large port structure, resulting in channels of smaller diameter [9]. In the present study, we examine a mordenite mineral specimen, in order to understand the structural difference between the large- and small-port variants.

2. Experimental

The locality of the measured samples is Lengyendi-galya (Gyökeres-tető), Mátra Mountains (NE-Hungary). The mordenite
bundles occur in hydrothermally highly altered andesite. Pure mordenite sample was prepared from the andesite.

X-ray diffraction measurements were carried out using Bruker D8 Advance diffractometer (Cu-Kα source) with Göbbel-mirror, parallel beam geometry, equipped with Vantec1 position sensitive detector, with symmetrical 2.5° axial Soller slits. Measurements were recorded in the 2°–100° (2Θ) angular range, with 0.007° (2Θ)/140 s goniometer speed. XRD patterns were evaluated by Search/Match in DiffracPlus EVA using ICDD PDF2 (2005) for crystalline components. Quantitative evaluation was performed in TOPAS4 with fundamental parameters approach, using crystal structures from ICSD database.

Morphological and textural observations were made using a Wild M400 stereomicroscope and an AMRAY 1830 scanning electron microscope (SEM). The crystal structure of mordenite was investigated by selected area electron diffraction (SAED), using a Philips CM20 transmission electron microscope operated at 200 kV. The composition was measured using Nanoprobe X-ray microanalysis (EDS) with a Bruker system.

3. Results and Discussion

3.1. Appearance of mordenite and associated minerals. Using X-ray powder diffractometry, we measured the hydrothermally altered andesite, composed dominantly by mordenite and heulandite; other minerals include quartz, andesine, orthoclase, hematite, and smectite.

X-ray diffraction patterns recorded on 1 mg powder sample. Rietveld measurements were carried out in TOPAS software in order to verify the powder purity. In the next step, peak search was done to prepare indexing data. Based on the measured reflections, we identified the Laue-group 2/m2/m2/m with C-centering. We observed forbidden reflections in the Cmc21 space group, so a reduced symmetry is evident for the bulk material.

Stereomicroscopy and secondary electron (SE) images revealed that the first product of hydrothermal alteration is mordenite, followed by heulandite (Figures 3 and 4). Secondary electron images in Figure 4A and B show heulandite crystals speared by mordenite fibers. Apparently, mordenite formed as the first zeolite and then heulandite grew around it.

3.2. Transmission electron microscopy. Features in SAED patterns from the mordenite fibers suggest a reduced symmetry compared to the Cmc21 space group and the presence of planar faults. We observed forbidden h + l odd reflections in the Cmc21 space group. For example, in the [010] projection, 10l, 30l, and 50l reflections are present (red arrows show some h0l lines in Figure 5A). Another example is the Benchmark.
[00] projection in which \( l = 2n + 1 \) reflections are also visible (Figure 5B).

Intense streaks and superlattice reflections in SAED patterns of mordenite suggest the presence of planar faults. Figure 6 shows \([110]\) and \([110]\) projected SAED patterns of mordenite, in which the lines of \( hkl \) reflections with \( l = 2n + 1 \) are interconnected by intense streaks, some of them marked with a red arrows.

The composition of mordenite was measured with EDS as \((Na_{1.25}K_{0.42}Ca_{1.67}Mg_{0.21}[H_2O]_{28.9})[Al_{8.38}Si_{40}O_{96}]\). The correct \( k \) factors were confirmed successfully on quartz and corundum earlier.

4. Discussion

Song et al. [11] observed different defect types in mordenite such as dislocations, planar defects, and local intergrowths using high-resolution electron microscopy (HREM) [11]. Planar faults with a \( \sim 1/4 [010] \) displacement vector result in the presence of dachiardite moduls in the structure. Based on intense streaks and superlattice reflections in SAED patterns, we also identified planar faults. Figure 6 shows examples of the mordenite reciprocal lattice in which the rows of \( hkl \) reflections with \( l = 2n + 1 \) are connected by intense streaks. This diffuse scattering is interpreted as an evidence of faults, where a fault produces a dachiardite-type structural slab within the mordenite lattice.

As shown by Figure 1, the puckered sheets are the same in mordenite and dachiardite, but the connection of these puckered sheets by four-membered rings (light shading) differs. The presence of the dachiardite domains affect the mordenite channel capacity by disrupting the system. Figure 7 shows the mordenite framework along [001]. With a displacement by a \( \sim 1/4 [010] \) vector (marked with a black arrow), dachiardite cages appear in the mordenite framework. This lattice displacement causes a surplus and a deficit of bonds which will connect by flipping (Figure 8B). The modified bonds are then relaxed in order to transform into a more stable structure (Figure 8C).

5. Conclusions

Space group determination from SAED patterns obtained from mordenite fibers suggested a symmetry reduction from \( Cmc2_1 \) to \( 2/m2/m2/m \) Laue group by the forbidden \( h + l \) odd reflections and the evident C-centering. Possible space groups include \( C222, Cnm2, C2nm, C2m2, \) and \( Cnmn \). This symmetry is also confirmed by powder diffraction data of mordenite.

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