### 1. Introduction

Zeolites are crystalline aluminosilicate materials and possess intracrystalline pores and nanospaces of similar sizes to the molecules of the lighter hydrocarbons. Moreover, zeolites have strong acid sites on the nanopore surfaces within and on the external surfaces of the crystalline structure. These properties enable zeolites to be used as in reaction and separation processes. The use of zeolites as membranes combines the advantages of inorganic membranes (high temperature stability and solvent resistance) with the zeolitic properties. Such zeolite membranes have potential uses as separation and reaction membranes, and as simultaneous reaction/separation membranes. A number of approaches have proved effective for modifying the properties of zeolite membranes, including the use of nano-crystalline zeolites for membrane preparation, regioselective deactivation of acid sites, and modification of zeolitic pore size. This review describes the synthesis of nano-crystalline zeolites by an emulsion method and the regioselective deactivation of acid sites by the catalytic cracking of silane (CCS) method. The effects of the crystal size and CCS treatment on the performance of the zeolite membrane are reviewed.

### Keywords
- Nano-sized zeolite, Emulsion method, Catalytic cracking of silane, Zeolite membrane, Membrane reactor, Membrane separation

### 2. Synthesis of Nano-sized Zeolite Crystals


Zeolites are normally prepared in an aqueous alkaline solution containing Si and Al sources and alkaline metal ions (sodium or potassium). An organic structure directing agent (OSDA), such as an ammonium alkyl cation, is also necessary to form the zeolite framework in the synthesis of some types of zeolite. To prepare the zeolite, solutions containing these inorganic...
materials and the OSDA are simply poured into a Teflon-sealed stainless bottle and heated to the required temperature.

Nano-crystalline zeolites can be prepared by quenching the hydrothermal treatment on nucleation. However, unreacted Si and Al species still remain in the solution, so an amorphous phase tends to form on the surface of the zeolite crystals. Another method involves increasing the nucleation rate by using higher concentrations of the Si and Al sources. By increasing the nucleation rate, the crystal size of the zeolite decreases. However, zeolites obtained by this method have a broad size distribution due to simultaneous nucleation and crystal growth during the hydrothermal treatment. Accordingly, it is important to separate the nucleation and growth stages to obtain nano-crystalline zeolite. Recently, nano-crystalline zeolites have been synthesized with the addition of surfactants, including in a water/surfactant/organic mixture (emulsion method). Research has focused on how to control the size and morphology of the crystals, because of the importance of these parameters in applications such as separation and catalytic reactions.

In the emulsion method, two solutions are prepared; an aqueous solution containing the Si and Al sources diluted with an OSDA aqueous solution, and a solution of the surfactant in an organic solvent. Zeolite crystals with different properties can be prepared by varying the concentrations of the Si and Al sources in the aqueous solution, and the molar ratio of Si to OSDA, and by changing the type of OSDA. The aqueous solution is added to the surfactant organic solution, and the mixture is magnetically stirred at 323 K to obtain a homogeneous solution. The resulting mixture of water/surfactant/organic solvent is then placed in a Teflon-sealed stainless steel bottle, heated to 373 to 423 K, and held at the required temperature for 12 to 120 h with stirring to yield the nano-crystalline zeolites. In the emulsion method, two additional parameters are important; the concentration of surfactant in the organic solvent and the molar ratio of water to surfactant, because these parameters affect the morphology and crystal sizes.

2.2  Effect of Ionicity of Surfactant on Preparation of Nano-crystalline Zeolites

The effect of the surfactant on the morphologies for MFI zeolite (silicalite-1) was examined using various surfactants at a hydrothermal temperature of 140°C and a surfactant concentration in the organic solvent of 0.50 mol/L. In general, there are four types of surfactants; anionic, cationic, non-ionic, and bipolar surfactants. The present study used nonionic surfactants, polyoxyethylene(15)oleylether (O-15), polyoxyethylene(15)nonylphenylether (N-15), and poly(oxyethylene)(15)cetylether (C-15), and ionic surfactants, sodium bis(2-ethylhexyl) sulfosuccinate (AOT) and cetyltrimethyl ammonium bromide (CTAB). Cyclohexane or 1-hexanol was used as the organic solvent.

Figures 1 and 2 show the X-ray diffraction (XRD)
patterns and FE-SEM images of the obtained samples, respectively. For the sample prepared in the AOT/cyclohexane solution, the XRD pattern included peaks corresponding to sodium sulfate rather than silicalite-1. Moreover, the sample showed an irregular morphology. In this case, the tetra-n-propyl ammonium hydroxide (TPA-OH) molecules did not act as an OSDA in the synthetic solution, possibly because the AOT surfactant and the TPA-OH used as the OSDA have opposite ionic charges.

For the sample prepared in CTAB/1-hexanol, scanning electron microscope (SEM) showed silicalite-1 crystals of about 1.0 μm embedded in the amorphous SiO2. XRD also showed the coexistence of silicalite-1 crystals and amorphous SiO2. Since the pH of the synthetic solution is alkaline in this case, the surface of the SiO2 had a negative charge. Accordingly, CTA+ and TPA+ were independently adsorbed on the surface of SiO2 because of their cationic charges. Consequently, silicalite-1 crystals and amorphous SiO2 were formed separately due to the independent adsorption of TPA+ and CTA+, respectively.

For the samples prepared in cyclohexane with the nonionic surfactants, C-15, NP-15, and O-15, XRD showed peaks corresponding to MFI zeolite (the reference silicalite-1), and SEM showed mono-dispersed silicalite-1 nanocrystals. In contrast, the silicalite-1 crystals prepared in water (without surfactant) had a heterogeneous structure and were present as smaller crystals (diameter of approximately 30 nm) formed on larger crystals (approximately 120 nm), indicating that nucleation, crystallization, and crystal growth occurred simultaneously. However, in the emulsion method, aggregation of the silicalite-1 nuclei is inhibited by the surfactants adsorbed on the surface during hydrothermal treatment, and mono-dispersed nanocrystals can be obtained. These results indicate that the ionicity of the hydrophilic groups in the surfactant molecules is important in the formation and crystallization processes of the nano-crystalline zeolites.

2.3. Crystal Size Control

In the preparation of zeolite membranes, zeolite seed crystals are typically immobilized on porous supports before secondary growth. The crystallinity and the size of the zeolite seed crystals affect the secondary growth of the zeolite during membrane formation, so preparation of mono-dispersed silicalite-1 nanocrystals is very important. In the emulsion method, adsorption of non-ionic surfactants on the surfaces of the zeolite precursors suppresses re-dissolution of the precursor in solution, thus inducing and enhancing nucleation of the zeolite. Control of the surfactant concentration is quite effective to suppress re-solution of the zeolite precursor. Accordingly, the effects of varying the O-15 surfactant concentration on the morphology of MFI (silicalite-1) and MOR zeolites were examined. Figure 3 shows the FE-SEM images of the obtained samples. The Si/Al ratio of the MOR zeolites was 12.5. The surfactant concentrations were varied in the range from 0.5 to 0.75 mol/L, and preparation of zeolites from water solutions without surfactant/organic solvent (the conventional method) was also carried out for comparison. Interestingly, the crystal size and morphology depended on the surfactant concentration, despite use of the same concentrations of the Si, Al and OSDA throughout. Nano-crystalline zeolites with mean size of below 100 nm were obtained with a surfactant concentration of 0.5 mol/L. At higher concentrations, growth of MFI and MOR zeolites was clearly observed, with the crystal size reaching ~1.0 μm at 0.75 mol/L. The BET surface areas of the MFI and MOR zeolites were about 370 and 430 m²/g, respectively, regardless of the crystal sizes[19],20),32). Moreover, temperature-programmed desorption (TPD) of the NH₃ profiles of the obtained MOR zeolites confirmed that these MOR zeolites with Si/Al ratio of 12.5 had almost the same acidity[32].

3. Deactivation of Acid Site and Modification of Pore Size

3.1. Catalytic Cracking of Silane Method

Because zeolites contain micropores of a specific diameter almost equal to the molecular sizes of the lighter hydrocarbons, they exhibit a remarkable molecular sieving effect for these hydrocarbons in reaction and adsorption processes, in which the spatial limitations within the zeolitic pores are expected to provide effective reaction/separation fields. Unfortunately, zeolites exhibit poor spatial limitation effects in some cases. For example, reactions over acid sites located on the external surface of the zeolite crystals can result in coke deposition, leading to a short catalyst lifetime. To overcome these problems, selective deactivation of acid sites located near the external surface of the zeolite crystal is required. Another problem occurs if all mol-
ecules have sizes smaller than the pore diameter. In this case, methods for modifying the pore wall and/or decreasing the pore size are needed for effective separation. Therefore, methods that allow regioselective deactivation and/or pore-size modification are required to improve the molecular sieving effect of zeolites.

Niwa et al. 33,34 have developed a method utilizing the chemical vapor deposition (CVD) technique with tetraethoxysilane and/or tetramethoxysilane precursors to form a thin silica (SiO2) layer on the external surface of zeolite crystals. We have also proposed a method for SiO2 formation based on the catalytic cracking of silane (the CCS method) using organic silane compounds, in which SiO2 formation selectively occurs on the acid sites of the zeolite 22,31. The CCS method can use several types of silane compounds to deactivate the acid sites of zeolite, including silanes with methyl, ethoxy, and phenyl substituents. Because the molecular size of the silane compounds depends on the organic groups bonded to the Si atom, it is expected that selective deactivation of the acid sites located on the external surface of the zeolite crystals can be achieved by utilizing the molecular sieving effect of the zeolites to separate the silane compounds depending on their size 31. Moreover, if silane compounds smaller than the zeolite pores are used, SiO2 formation may occur on the acid sites located on the pore surfaces within the crystals, leading to modification of the pore size 22.

3.2 Adsorption of Silane Compounds on MFI Zeolite

Figure 4 shows typical results for the weight changes observed during exposure of powdery MFI zeolite (proton-type ZSM-5, Si/Al = 15) to diethoxymethyl silane (DEM-Silane) vapor in a thermo-balance. A rapid weight increase was observed up to 400 K caused by chemisorption and physisorption of the silane compounds on the zeolite surface. The weight of the sample slightly decreased from 400 to 475 K, resulting from desorption of the physically adsorbed silane compounds on zeolite surface. No further weight changes were observed up to 700 K, indicating that the silane compounds chemisorbed on the acid sites reached an equilibrium state, which was confirmed by comparison of the number of adsorbed silane molecules with the number of acid sites of the zeolite 22. The weight of the zeolite rapidly increased above 700 K, due to thermal CVD and coke deposition by decomposition of the silane compounds.

In the CCS method, modification of the zeolite acidity and pore size can be achieved by SiO2 unit formation, which occurs via a silane compound chemisorbed on the acid site of the zeolite. Accordingly, the procedure for the CCS method is outlined as follows 22,31,32. After air calcination, the zeolite is exposed to the silane compound vapor at 373 to 393 K in a N2 stream, and the feed of the silane compound is then stopped to remove the physically adsorbed silane compounds on the zeolite surface. Therefore, the remaining silane compounds are chemisorbed on the acid sites of the zeolite. Next, the sample is heated to 823 K in a N2 stream to decompose the silane compounds adsorbed on the acid sites, resulting in the formation of silicon-coke composites on the acid sites. Finally, the gas feed is switched from N2 to air, and the temperature of the reactor is held at 823 K for 60 min, allowing the silicon-coke compounds formed on the acid sites to be oxidized to SiO2 units. Therefore, the formation of SiO2 units occurs on the acid sites which initially chemisorbed the silane compounds. By repeating this sequence, the formation of SiO2 units occurs again on the acid sites, leading to decreases in the zeolite acidity and pore size.

3.3 Regioselective Deactivation of Acid Sites by the CCS Method

The CCS method can use various silane compounds, such as diethoxymethylsilane (DEM-silane), phenylsilane (P-silane), diphenylsilane (DP-silane), diphenylethoxymethylsilane (DPM-silane), and triphenylsilane (TP-silane). The order of the molecular sizes of the silane compounds is as follows (with the pore size of the MFI zeolite included for comparison): TP-silane > DPM-silane = DP-silane = pore diameter of MFI (10-membered rings) > P-silane > DEM-silane. With the phenylsilane compounds, the number of phenyl groups bonded to the Si atoms affects the molecular size of the silane compounds. Figure 5 shows a schematic of the selective deactivation of acid sites by the CCS method, according to the molecular diameter of the precursor silane compounds. For organic silane compounds with almost the same molecular size as the pore size of the zeolite (e.g. DP-, DPM- and TP-silanes), selective deactivation occurs at acid sites located near the external surface of the MFI zeolite (ZSM-5) crystal. In contrast, for silane compounds with molecular sizes smaller than the...
pore size of the zeolite (e.g. P-silane and DEM-silane), acid sites are accessible on the pore surfaces within the crystal as well as near the external surface, enabling modification of the pore size. As a result, different distributions of acid sites on the MFI zeolite (ZSM-5) can be achieved by regioselective deactivation using silane compounds with different molecular sizes.

4. Applications of Zeolite Membrane for Gas-phase Reaction/Separation

4.1. Experimental Apparatus for Gas-phase Reaction/Separation Using Zeolite Membrane

Zeolite membranes have been applied to simultaneous reaction and separation systems. These zeolite membrane reactors can be divided into two configurations: catalyst placed on top of a zeolite layer, and homogeneous membranes containing zeolite crystals. Zeolite membrane reactors based on a catalytic zeolite layer are less common. In such a reactor, molecules are transported to the external surface of the zeolite layer and penetrate into the layer, then react over the active sites of the zeolite. The reactions cease after the molecules have passed through the layer.

In the present study, an MFI zeolite (ZSM-5) membrane was prepared by forming a MFI zeolite layer on the surface of a cylindrical porous alumina ceramic filter with an inner diameter of 6 mm, an outer diameter of 11 mm, and a length of 50 mm. The filter pores had a diameter of 0.1 μm. Three reactant solutions were mixed for the hydrothermal preparation of the MFI zeolite membrane: sodium chloride solution, sodium silicate solution and aluminum sulfate solution containing tetra-n-propyl ammonium bromide (TPABr). The alumina ceramic filter with seed MFI zeolite crystals (ZSM-5 with crystal size of about 1000 nm) was immersed into the mixture followed by hydrothermal treatment to form the MFI zeolite membrane on the surface of the filter. After ion-exchange treatment from Na⁺ to NH₄⁺ and air calcination to remove the TPABr molecules, the membrane was used in further experiments.

Figure 6 shows the overview and cross-sectional area of the prepared membrane (ZSM-5 zeolite layer with Si/Al ratio of 15). XRD showed that the membrane layer had the same structure as MFI zeolite, indicating that the zeolite membrane consisted of MFI zeolite (ZSM-5) crystals. Moreover, the zeolite layer formed on the outer surface of the alumina ceramic filter was dense and uniform. Applications of zeolite membrane to gas-phase separation and reaction require a dense membrane without physical holes for optimum separation properties of gaseous molecules, based on the molecular sieving effect of the micropore of the zeolite. Accordingly, the dense membrane shown in Fig. 6 was...
suitable for further applications.

**Figure 7** shows a schematic view of the apparatus used for gas-phase reaction/separation in a flow system. One end of the cylindrical zeolite membrane was sealed, and a gas mixture of H₂ and N₂ (for gas separation) or methanol and N₂ (for reaction) were fed to the outer side (feed side) of the membrane. Helium (He) or N₂ gas was fed to the sealed end of the membrane from a pipe inserted in the inner side (permeate side of the membrane) of the sample holder to allow backflow to sweep out the permeating molecules. The gas compositions in the feed and permeate sides were measured using a quadrupole mass analyzer and gas chromatography (GC-8A, Shimadzu Corp., Japan).

4.2 MTO Reaction Using MFI Zeolite Membrane Treated by the CCS Method

This study showed that SiO₂ units are selectively formed on the acid sites of the zeolite by the CCS method. Moreover, regioselective deactivation of acid sites can be achieved by the CCS method using silane compounds with different molecular sizes. Next, the methanol to olefins reaction (MTO reaction) was investigated using the MFI zeolite (ZSM-5) catalytic membrane to assess the effect of CCS treatment on the product yields of MTO reaction.

The MTO reaction using the catalytic zeolite membrane was expected to form light olefins such as ethylene and propylene followed by formation of paraffins and aromatics at the feed side of the membrane, due to the nonshape-selective reactions over the acid sites of the membrane facing the feed side. In order to increase the olefin yield, such nonshape-selective reactions must be limited without any reduction in olefin formation. For this purpose, regioselective deactivation of the acid sites by the CCS method may be achieved using silane compounds with larger molecular sizes than the pore size of the zeolite, providing selective deactivation at acid sites located on and/or near the external surface.

In this study, selective deactivation of the acid sites on the external surface of the zeolite membrane was achieved without deactivation of acid sites in the zeolite pores in an attempt to increase the olefin yields. **Figure 8** shows the typical production selectivity for the MTO reaction using the ZSM-5 zeolite membranes before and after CCS treatment using TP-silane. Membranes before CCS treatment formed olefins in the products recovered from the permeate side, but the selectivity for paraffins at the feed side was much higher than the selectivity for olefins at the permeate side. Paraffin production at the feed side resulted from the conversion of methanol to paraffins over acid sites on the external surface of the membrane facing the feed side. Moreover, the paraffins formed at the feed side had longer chain lengths than the olefins and paraffins formed at the permeate side. These longer chain products result from the nonshape-selective reactions of the hydrocarbons over the acid sites on the surface of the membrane facing the feed side. In contrast, the zeolite
membrane after CCS treatment using TP-silane showed markedly reduced selectivity for paraffins at both the feed and permeate sides. Moreover, the olefin selectivity at the permeate side was higher than that at the feed side. The acid sites on the surface facing the feed side of the zeolite membrane were selectively deactivated by the CCS method using TP-silane, so suppressing the formation of undesirable products through reactions with ethanol, such as paraffins and aromatics.

4.3. H₂ Separation Using MFI Zeolite Membrane Treated by the CCS Method

The MFI zeolite (ZSM-5) membrane treated by the CCS method was applied to the separation of H₂ from mixtures of gases containing H₂ and N₂ or O₂. The experimental apparatus shown in Fig. 6 was also applied for the gas-phase separations. All molecules permeating from the feed side to the permeate side of the membrane must pass through the pores of the MFI zeolite. The CCS method using DEM-silane and P-silane, which are smaller than the pore size of the zeolite, is expected to form SiO₂ units on acid sites inside the pores of the crystal as well as on the external surface. Accordingly, the pore sizes of the MFI zeolite (ZSM-5) are decreased to a size only slightly larger than the molecular sizes of N₂ and O₂, but significantly larger than that of H₂. Therefore, the diffusion rate ratio of H₂ to N₂ or H₂ to O₂ through the membrane is expected to be greater for the MFI zeolite membrane after CCS treatment than for the membrane before CCS treatment.

Figure 9 shows the overall mass transfer coefficient of the membrane (permeance) for H₂ at different molar fractions for a mixture of H₂ and N₂ at the feed side. To investigate the gas-separation behavior, MFI zeolite (ZSM-5) membranes before and after CCS treatment using DEM-silane were used. Over a wide range of H₂ molar fractions, the permeance of N₂ decreased markedly (to about 1/500) for the membrane after CCS treatment, whereas that of H₂ was much less decreased (to about 1/10). Based on the measured permeances and molar fractions of H₂ and N₂ at the feed and permeate sides, the separation factor for H₂ was calculated as 90 to 140 for the membrane after CCS treatment, whereas that of H₂ was much less decreased (to about 1/10). Based on the measured permeances and molar fractions of H₂ and N₂ at the feed and permeate sides, the separation factor for H₂ was calculated as 90 to 140 for the membrane after CCS treatment, whereas that of H₂ was much less decreased (to about 1/10). Similar results were obtained for mixtures of H₂ and O₂ (H₂ separation factor of 1.5-4.5).

5. Applications of Zeolite Membrane for Liquid-phase Separation

5.1. Pervaporation Membrane

Separation and purification processes of liquid mixtures are among the most important unit-operations in the petrochemical industry, and are primarily carried out using distillation columns. However, the distillation process requires huge amounts of energy due to the large number of plates in the distillation columns and the high reflux ratio, which is above 20. Accordingly, alternative methods for high purity separation are necessary, with pervaporation emerging as a promising solution.

In the pervaporation method, the feed liquid mixtures are contacted with one side of a membrane (feed side), and the target components are removed from the mixture as vapor from the other side of the membrane (permeate side) after permeation through the membrane. The relative affinities (e.g. hydrophilic or hydrophobic interaction) between the membrane and the various components in the liquid solution allow for selective permeation that is superior to the separation obtained by distillation, which relies on the vapor-liquid equilib-
rium. Therefore, several types of zeolite membranes have been prepared and studied as potential pervaporation membranes for dehydration of organic solvents such as ethanol, isopropanol, and acetone, due to the high stabilities at high temperatures and strong resistance to organic solvents. The function of zeolite membranes to perform selective dehydration by pervaporation depends on the hydrophilic properties, which in turn depend on the composition of elements within the constituent components, such as the Si/Al ratio. However, dealumination may occur in acid solutions with zeolites containing Al₂O₃, resulting in a reduction in the separation factor due to the generation of defects after long-term use. Therefore, Al₂O₃-free zeolite membranes should be used for such applications.

5.2. Preparation of Hydrophilic and Al₂O₃-free Zeolite Membrane

Silicalite-1, which has an MFI structure, is a suitable Al₂O₃-free zeolites with the potential to avoid generation of defects caused by dealumination. Prepared MFI zeolites are usually calcined in air at 803 K, to remove any OSDA molecules from within the zeolite. However, during this procedure, silanol groups undergo dehydration to form siloxane bonds, resulting in a hydrophobic surface. Accordingly, the preparation of hydrophilic silicalite-1 membranes is not well characterized. To prepare silicalite-1 membrane with high hydrolytic properties retaining the silanol groups, we have proposed a liquid phase oxidation technique by modifying the method reported by Fukuoka.

Briefly, silicalite-1 membranes prepared without air calcination were immersed into a nitric acid solution (300 cm³, 1 mol/L) and the solution heated to a reflux condition. Then, hydrogen peroxide solution (30 cm³, H₂O₂, 30 wt%) was slowly added to the solution, and maintained for 24 h under reflux. The treated membrane was washed by distilled water, dried under an air stream, and then used in the experiments.

The silicalite-1 membranes obtained after liquid-phase oxidation using H₂O₂ retain considerable numbers of hydrophilic silanol groups on the silicalite-1 surface, to which water molecules selectively adsorb. Moreover, the water–silanol bonding networks that form on the non-zeolitic pores of the silicalite-1 crystals are the dominant channels for permeation of water molecules, allowing water molecules to selectively diffuse through the network, resulting in a high separation factor. As the size of the crystals decreases, the number of hydrophilic channels for water permeation is expected to increase, leading to improvement of the water flux in addition to the separation factor. Accordingly, membranes were prepared using nanocrystalline silicalite-1 with different crystal sizes, and the effect of the crystal sizes on the separation of water from water-acetone solution was examined.

5.3. Hydrophilic Silicalite-1 Membrane

Silicalite-1 nanocrystals were ultrasonically dispersed in alkaline water solution at a pH of approximately 10. The dispersed nanocrystals were layered on the outer surface of a cylindrical alumina ceramic filter (as described in section 4.1) by a filtration method with evacuation on the inner side. The thickness of the silicalite-1 nanocrystal layer deposited on the ceramic filter could be controlled by changing the concentration of the silicalite-1 nanocrystal in the water solution. Finally, a silicalite-1 layer (protective layer) was hydrothermally formed on the nanocrystal layer, as described previously.

Figures 11(a) and 11(b) show SEM micrographs of the cross-sectional area and top view, respectively, of the silicalite-1 nanocrystal-layered membrane. The nano-crystalline silicalite-1 used for the preparation of the membrane had a crystal diameter of 60 nm. The nanocrystalline layer with a thickness of approximately 4.0 μm could be clearly observed on the alumina filter, together with a protective layer of columnar crystalline silicalite-1 with a thickness of approximately 1.2 μm on top of the nanocrystal layer. As shown in Fig. 11(b), many large pores can be observed among the silicalite-1 crystals of the protective layer. The crystalline size in the protective layer was approximately 1000 nm, and was much bigger than that of the nano-crystalline silicalite-1. The morphology of the top view suggests that the protective layer had almost no separation function. In contrast, secondary growth of the nanocrystals can be observed around the interface between the protective and nanocrystalline layers. Such growth of nanocrystals occurred during the formation of the protective layer.
al.\(^{15,16}\) have reported that the amount of seed crystals on the support affected the secondary growth of seed crystals and the membrane characteristics for pervaporation and gas separation. Therefore, the effect of the thickness of the nanocrystalline layer on the separation properties was investigated.

The thicknesses were a 60 nm for the nanocrystal monolayer and 4.0 \(\mu\)m for the nanocrystal layer, and after depositing the nanocrystals, the silicalite-1 protective layer was formed at a hydrothermal temperature of 140 \(^\circ\)C. Figure 12(a) shows the effects of the thickness of the nanocrystal layer on the separation of water through the layered membrane. In the membrane prepared without the nanocrystal layer, the silicalite-1 protective layer was directly formed on the alumina filter by hydrothermal synthesis. This membrane without the nanocrystal layer showed no separation function, indicating that the silicalite-1 nanocrystal layer was essential for the separation function. In the membrane with a nanocrystal monolayer, excessive secondary growth of the nanocrystals had occurred, leading to formation of a dense silicalite-1 layer, so that both water and acetone molecules could not permeate the membrane. In contrast, in the membrane with a nanocrystal layer of 4.0 \(\mu\)m thickness, appropriate pores for selective permeation of water were formed among the nanocrystals by secondary growth, so that the membrane exhibited high water flux with high separation factor.

The dependency of the separation performance on the amount of silicalite-1 nanocrystals was ascribed to the secondary growth during hydrothermal synthesis to form the protective layer\(^{27}\), which was in good agreement with the previous reports\(^{14\sim16}\), and since the protective layer was formed by secondary growth of the silicalite-1 nanocrystals in the layer, the active separation region was considered to be the interface between the nanocrystal and protective layers. Zeolite mem-

![Fig. 11 SEM Images of (a) Cross-section and (b) Top View of the Silicalite-1 Nanocrystal-layered Membrane](image)

The acetone concentration was 90 wt%.

![Fig. 12 Effects of (a) Layer Thickness of and (b) Crystal Size of Nano-crystalline Silicalite-1 on the Water Flux through the Layered Membrane](image)
branes formed on porous ceramic filters in this way usually suffer from the serious problem of thermal mismatch caused by the different expansion coefficients of the zeolite layer and the alumina filter. In contrast, the thick nanocrystal layer on the alumina filter compensates for this mismatch because the active separation region is located at the interface between the nanocrystal and protective layers.

Figure 12(b) shows the effects of the crystal size of the nano-crystalline silicalite-1 on the separation of water from a water-acetone solution passed through the layered membranes, where \( D \) and \( \alpha \) represent the crystal size and the separation factor, respectively. The flux of water through the layered membranes increased with higher permeation temperature. Decreased crystal sizes of the silicalite-1 resulted in dramatically increased water flux as well as separation factor, which was considered to result from reduction in the mesopore spaces between the initial crystals and the spaces remaining after secondary growth. Accordingly, the separation factor increased with smaller crystal size of the nano-crystalline silicalite-1. Moreover, the decrease in crystal size also caused higher numbers of hydrophilic channels between the crystals, allowing the water flux to increase through the channels of water-silanol networks for the selective permeation of water molecules.

6. Conclusion

Reaction and separation processes are very important petrochemical processes, for which zeolites are used widely as catalysts and adsorbents. The surface properties of zeolite crystals can be easily modified by post-synthesis treatments such as ion-exchange and using organic silane compounds, so zeolites are expected to be useful in a wide range of new applications. One promising application is the use of zeolites in membranes, which combine the properties of zeolites with those of inorganic membranes. The present review introduces the use of nano-crystalline zeolites in membrane preparation, and modification of the acid sites and pore sizes of the zeolite membranes. The size-controlled synthesis of nano-zeolites was achieved by an emulsion method, and reduction in crystal size led to improved separation properties for a pervaporation membrane. The CCS method could be used to selectively deactivate acid sites located on the feed side of the zeolite membrane. The membrane treated by the CCS method exhibited high olefin selectivity in an MTO reaction. The use of nano-crystalline zeolites and regioselective deactivation of the acid sites in the zeolite membrane are effective for controlling the zeolite properties in a membrane configuration.

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References


要 旨

ナノサイズゼオライトの合成法とゼオライト膜への展開

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ナノサイズゼオライトの合成法とゼオライト膜への展開について、ナノサイズゼオライト膜の特性（耐熱、耐薬品性）を考慮して、ゼオライト膜の製造を検討した。ナノサイズゼオライトの粒子径が細孔径の約1/10以下の細孔径を持つナノサイズゼオライトを用いることにより、ゼオライト膜の特性が大きく改善する。これはゼオライト膜の製造において重要な手段である。ナノサイズゼオライト膜の製造において、ナノサイズゼオライトの粒子径が細孔径の約1/10以下の細孔径を持つナノサイズゼオライトを用いることにより、ゼオライト膜の特性が大きく改善する。これはゼオライト膜の製造において重要な手段である。