Na-A (LTA) zeolite synthesis directly from alumatrane and silatrane by sol-gel microwave techniques

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Abstract

Na-A (1 \textmu m crystal size) zeolite was successfully synthesized via a sol-gel process and microwave heating technique using alumatrane and silatrane as precursors. After fixing the SiO\textsubscript{2}:Al\textsubscript{2}O\textsubscript{3} ratio at 1:1 and microwave heating temperature at 110 °C, increasing the Na\textsubscript{2}O concentration by adding more NaOH exponentially reduces the microwave heating time from 160 min at a Na\textsubscript{2}O:SiO\textsubscript{2} ratio of 3:1 to 5 min at a Na\textsubscript{2}O:SiO\textsubscript{2} ratio of 9:1. The increase of Na\textsubscript{2}O concentration strongly affects the particle size and particle size distribution, but does not affect the product composition. Small crystallite sizes are obtained from a high Na\textsubscript{2}O:SiO\textsubscript{2} ratio (10:1) while a low Na\textsubscript{2}O:SiO\textsubscript{2} ratio (3:1) gives large crystallite sizes (~4.5 \textmu m). The analyzed Si:Al:Na ratio of synthesized Na-A zeolite is 1:1:1.25. The moisture absorption ability of the synthesized Na-A zeolite is higher than that of the commercial one by approximately 20%. The increase of water ratio also affects the crystal size. As the water ratio increases, larger crystallites with a higher degree of irregularity are formed.

Keywords: Microwave techniques; Organic precursors; Sol-gel processes; Zeolites

1. Introduction

Zeolites are versatile materials, which are used in many applications, such as household products, aquaculture, agriculture, water treatment, etc. due to their absorption, ion exchange and size selectivity properties.\textsuperscript{1} Na-A zeolite has also been employed in gas separation membranes, to enhance the selectivity of the support.\textsuperscript{2} Commercial Na-A zeolite is synthesized by both conventional and microwave heating techniques and mostly silicates and aluminates are used as starting materials.

A Na-A zeolite membrane was produced on zAl\textsubscript{2}O\textsubscript{3} support by dipping the support into the gel and then crystallizing the gel by hydrothermal heating at 90 °C.\textsuperscript{3} The results indicated that microwave heating provided more uniform and smaller particles with a shorter heating time (4–12 times shorter than conventional heating) and gave four times higher gas permeability with equal selectivity in permeation. A second technique of making Na-A zeolite membrane is to seed the support with Na-A zeolite crystals followed by an one-time-only hydrothermal synthesis at 100 °C. The seeding method yielded a dense inter-grown zeolite crystal layer of about 30 \textmu m in thickness on the outer surface.\textsuperscript{4} The resulting membranes were highly permeable to water vapor but impermeable to other gases unless dried completely. They exhibited excellent water perm-selective performance in water/organic liquid mixtures. The microwave heating times can be shortened by increasing the heating temperature to 100 °C. Additionally by increasing the Na\textsubscript{2}O concentration, the heating time can be reduced to 5 min.\textsuperscript{5} However, the resulting particles had knotted edges due to the fact that pre-nuclei were formed during aging. It was also found that the aging time strongly influences the crystallization time, particle size and yield. Ultrasound and XRD revealed that an amorphous gel is first observed before nucleation and crystallite growth.\textsuperscript{6} A similar observation was made by using dynamic light scattering (DLS) to follow zeolite synthesis.\textsuperscript{7} Even though Na-A zeolite synthesis has been studied in great detail, it is still a challenge to find a new synthetic
route, which results in better crystallinity. The use of small amines, such as ethylamine, isopropylamine or diethylamine (short or long primary or secondary amine) as a pore-filler during the crystallization stage, has been successful in siliceous zeolite synthesis and in controlling the framework structures. However, the obtained product contained charge-balancing protonated amine (despite a gel pH often above 13) and had a Na/Al ratio substantially below unity (Na/Al < 1). Our previous work used atrans (metal alkoxides), see Fig. 1, as precursors for Na-A and GIS zeolite synthesis via the sol-gel process and resulted in thermodynamically stable materials in an aqueous-base system. Atrane is an amine-trialkoxo complex (with a diversity of elements), in which tertiary amine acts as anionic (tetradenate) tripod ligand and a neutral nitrogen-donor while trialkoxo groups function as polyalcohols making it capable in acting as reagent in sol-gel process. These properties increase the possibility of expanding its coordination sphere of inorganic-organic micelle formation during the sol-gel process and hydrothermal treatment. Nevertheless, they moderate the alkoxide reactivity towards the nucleophilic attack of water, resulting in slow precipitation of crystal and nuclei. This paper reports further results of our studies on the synthesis of Na-A or LTA zeolite by using the same method with the expectation of a higher uniform crystallinity and a narrower particle size distribution.

2. Experimental

2.1. Materials

Fumed silica (SiO₂, surface area 474 m²/g, average particle size of .007 μm) and aluminum hydroxide hydrate [Al(OH)₃], surface area 51 m²/g were purchased from Sigma Chemical Co. and used as received. Triethanolamine [TEA, N(CH₂CH₂OH)₃], and triisopropylamine (TIS, N(CH₃CH(CH₃)₂)OH)₃) were supplied by Carlo Erba Reagenti and Fluka Chemical AG., respectively, and used as received. Ethylene glycol (EG, HOCH₂CH₂OH) was obtained from J.T. Baker Inc. and distilled using fractional distillation prior to use. Sodium hydroxide (NaOH) and sodium chloride (NaCl) were purchased from EKA Chemicals and AJAX Chemicals, respectively. Both were used as received. Acetonitrile (CH₃CN) was obtained from Lab-Scan Co., Ltd. and distilled using standard purification methods prior to use.

2.2. Instrumental

FTIR spectroscopic analysis was conducted using a Bruker Instrument (EQUINOX55) with a resolution of 4 cm⁻¹ to measure functional groups of materials. The solid samples were prepared by mixing 1% of the sample with dried KBr, while the liquid samples were analyzed using a Zn–Se window cell. To measure the molar mass of precursors, mass spectra were obtained by means of a VG Autospec model 7070E from Fison Instruments with a VG data system, using the positive fast atomic bombardment (FAB⁺-MS) mode and glycerol as a matrix. CsI was used as a reference, while a cesium gun was used as an initiator. The mass range used was from m/e = 20 to 3000. Thermal properties and stability were analyzed by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC). TGA was performed using a Perkin Elmer TGA analyzer while DSC was conducted with a Netzsch instrument: DSC200 Cell at a heating rate of 10 °C/min under nitrogen atmosphere. Aluminum pans containing 5–10 mg of sample were used in DSC analysis, while a platinum pan containing 10–20 mg of sample was used in TGA. For the liquid and gel samples, a high-pressure gold cell containing 10–20 mg of sample was used. The crystallinity of products were characterized by Rigaku X-ray diffractometer at a scanning speed of 5 °/s using CuKα as a source and CuKβ as a filter. XRD spectra were recorded after 0/2θ scans in the range 5–50°. SEM micrographs were obtained with a JEOL 5200–2AE scanning electron microscope. Electron probe microanalysis (EPMA) was used for both qualitative and quantitative elemental analysis using the X-ray mode detector (SEM/EDS) to obtain product compositions. Particle sizes and particle size distributions were determined using a Malvern Instruments Mastersizer X Ver.2.15 analyzer. Water was used as a mobile phase. Hydrothermal treatment by microwave heating technique was conducted using MSP1000, CME Corporation (Spec. 1000 W and 2450 MHz). Samples were heated in a Teflon-lined digestion vessel using inorganic digestion mode with time-to-temperature program.

2.3. Precursors synthesis

Silatrane synthesis was performed by mixing silicon dioxide, 0.10 mol, and triethanolamine, 0.125 mol, in a simple distillation set using 100 ml of ethylene glycol, as solvent. The reaction was carried out at the boiling point of ethylene glycol under flowing nitrogen to remove the by-product water and ethylene glycol from the system. The reaction was carried out for 10 h and...
the rest of ethylene glycol was removed under vacuum (10⁻² torr) at 110 °C for 8 h. The brownish white solid was washed three times with dried acetonitrile (or until the filtrate had no color) to remove undesired organic residues. Approximately 95% yield white powder product was obtained and characterized using FTIR, TGA, DSC and FAB⁺-MS.

A similar process was used to synthesize alumatrane from aluminum hydroxide (0.1 mol) and trisopropanolamine (0.125 mol). The crude product was washed with dried acetonitrile for three times to give approximately 90% yield of the final product, which was characterized using FTIR, TGA, DSC and FAB⁺-MS.

2.4. Sol-gel process and microwave technique

SiTEA and AlTIS were mixed with NaOH or NaCl solution at room temperature at a ratio of SiO₂:Al₂O₃:xNa₂O:yH₂O (where 0 ≤ x ≤ 10 and 63 ≤ y ≤ 1000). The solution mixture was aged for at least 12 h to fully obtain gel formation and then placed into a Teflon vessel for further hydrothermal treatment using the microwave technique. The solution mixtures containing different ratios of SiO₂:Al₂O₃:xNa₂O:yH₂O were treated for various times and the resulting white powder products were washed three times using distilled water. The products were finally dried at 75 °C for 15 h prior to characterization using DSC/TGA, FTIR, XRD, N₂ porosimetry (BET is the analysis method) and SEM.

2.5. Moisture absorption

Crucibles were heated at 350 °C for 10 h and cooled to 120 °C in a hot-air oven. They were then moved and kept in a dessicator until use. The samples and the dried crucibles were weighed and then heated at 350 °C for 10 h. They were cooled to 120 °C in oven and then kept in dessicator until they cooled to room temperature. The dried samples were weighed and dried weight of sample (W₁) was calculated. The dried samples were placed in a closed humidification chamber for 15 h and then weighed to determine the mass of water adsorbed (W₂). The moisture absorption ability was calculated per unit mass of the sample (W₂/W₁).

3. Results and discussion

3.1. Precursor synthesis

Via the oxide-one-pot-synthesis (OOPS) process, silatrane and alumatrane were successfully synthesized directly from silica and alumina. Due to the condensation reaction generating water as a byproduct, the reactions were carried out under a nitrogen atmosphere and water was removed from the system to drive the reactions forward. The products can slowly absorb moisture and then undergo the hydrolysis process and thus need to be kept under vacuum.

The FTIR spectrum of the synthesized alumatrane shows the following significant peaks: 2860–2986 cm⁻¹ (m, nC–H), 1244–1275 cm⁻¹ (m, nC–N), 1130 cm⁻¹ (m, nC–O), 1102 cm⁻¹ (s, nAl–O–C), 1037 cm⁻¹ (m, nC–O), 890 cm⁻¹ (s, dAl–O–C) and 649–400 cm⁻¹ (s, dAl–O). The TGA results give two mass loss transitions at 139 and 393 °C with 24% ceramic yield corresponding to the structure of Al(OCHCH₃CH₂)₃N, which has 23.7% theoretical ceramic yield. Coincidently, DSC also
shows 145 °C (endothermic) and 380 °C (exothermic) referring to decomposition of organic ligand and oxidation of carbon residue, respectively, since a second heating cycle (after the first heat-cool cycle) revealed no endothermic transition as the material had undergone degradation. The other method to confirm the structure of the desired product is FAB+-MS giving the base peak at m/e 216 belonging to the structure of Al(OCHCH3CH2)3 NH+ and approximately 5% of the molecular peak at m/e 1292, which belongs to (Al(OCHCH3CH2)3N)6H+.

Concerning the synthesized silatrane product, its FTIR spectrum showed the following peaks: 3000–3700 cm⁻¹ (w, νO–H), 2860–2986 cm⁻¹ (s, νC–H), 1244–1275 cm⁻¹ (m, νC–N), 1170–1117 cm⁻¹ (bs, νSi–O), 1093 cm⁻¹ (s, νSi–O–C), 1073 cm⁻¹ (s, νC–O), 1049 cm⁻¹ (s, νSi–O), 1021 cm⁻¹ (s, νC–O), 915–940 cm⁻¹ (m, δSi–O–C), 785 and 729 cm⁻¹ (s, δSi–O–C) and 579 cm⁻¹ (w, Si–N). In this case, TGA thermogram provides only one mass loss transition at 390 °C with 18.5% ceramic yield corresponding to Si((OCH2CH2)3N)2H2, whose theoretical ceramic yield is 18.6%. DSC thermogram shows two peaks at 349 °C (endothermic) and 373 °C (exothermic). Again, these two peaks are attributed to the decomposition of the organic ligand and oxidation of the carbon residue, respectively. FAB+-MS gives the molecular as well as base peak at m/e 323 of Si((OCH2CH2)3N)2H3⁺.

3.2. Sol-gel process

The hydrolytic reaction of the silatrane and alumatrane mixture in NaOH or NaCl/H2O solution was followed by FTIR, as shown in Fig. 2. The NaOH system showed significant change at the Si–O–C and Al–O–C regions, 1000–1170 cm⁻¹. The peak at 1049 cm⁻¹ referring to Si–O–Si was higher and broader, while the peak at 1102 cm⁻¹ corresponding to Al–O–Al was slightly shifted to higher frequency and became broader. The C–O–M (M = Si or Al) peak at 1021 cm⁻¹ was also reduced due to the cleavage of the organic ligand from the system. This was confirmed by the reduction of the C–N peak at 1275 cm⁻¹ referring to Si–O–Si was higher and broader, while the peak at 1102 cm⁻¹ corresponding to Al–O–Al was slightly shifted to higher frequency and became broader. The approximate rate of hydrolysis during the sol-gel process was determined from Fig. 3. As can be seen in this figure, the hydrolysis rate was two times faster with hydroxyl anion (OH⁻). The hydroxyl ion attacked Si- or Al-atoms faster and easier to form hydroxide in the hydrolysis step due to the lower electronegativity of M–Cl bond and the weaker nucleophilicity of Cl⁻.22 The hydroxyl group of metal hydroxide part attacked Si- or Al-atom of other groups to form a metal–oxygen–metal bridge and the organic part was released in the condensation step. With both anions, OH⁻ and Cl⁻, the hydrolysis rate was rapid during the first hour and then became much slower.

3.3. Gel transformation to aluminosilicate

Both alumatrane and silatrane underwent hydrolysis and condensation reactions in a system of NaCl/water or NaOH to form amorphous metal oxide gel. The amorphous gel was aged for 12–15 h to obtain complete gelation. The gel was transformed to a crystalline aluminosilicate by hydrothermal treatment using the microwave-heating technique. The nucleation occurred first at this step and was followed by the formation of the crystalline product. We found that the gel started to form at the SiO2:Na2O ratio of 1:0.0069 and the ratios of SiO2:Na2O to start with the synthesis of LTA were in.
the range of 1:3–1:10. The amount of water started from \( y = 410 \) to \( y = 510 \) due to the homogeneous dispersion of the powder product in solution mixture and the suitable SiO\(_2\):H\(_2\)O ratio for synthesizing LTA was 1:410. By fixing the ratio of starting materials SiO\(_2\):Al\(_2\)O\(_3\):3Na\(_2\)O:410H\(_2\)O at 1:1:3:410 and the microwave heating temperature at 110 °C for 3 h, XRD results indicated that only the NaOH:H\(_2\)O system can provide crystalline aluminosilicate, as illustrated in Fig. 4. The XRD results were matched with the Linde (A-LTA) structure having PDF# 39–0222 and Na\(_{96}\)Al\(_{96}\)Si\(_{96}\)O\(_{384}\).216H\(_2\)O ~ Si:Al:Na = 1:1:1. SEM confirmed (Fig. 5) that the building unit was in cubic form, developed from the connection of sodalite cage (\( \beta \)-cage or T24 unit) along the cube axes through double T4-rings. The crystallographic data are listed in Table 1. Moreover, the EDS-SEM results showed that the synthesized product contained only Si-, Al- and Na-atoms, and the ratio of Si:Al:Na ratio was 1:1:1.25 due to the interaction of OH\(^-\) anions with the zeolite framework in order to form the siloxy groups (mSiO\(^-\)), which must be accommodated with extra Na\(^+\) cations.\(^{23}\) The TGA and DSC results of synthesized zeolite also indicated no

![Fig. 4](image1.png) XRD spectra of aluminosilicate synthesized from SiO\(_2\):Al\(_2\)O\(_3\):3Na\(_2\)O:410H\(_2\)O and 110 °C for 180 min in NaCl/H\(_2\)O and NaOH/H\(_2\)O systems.

![Fig. 5](image2.png) Unit cell structure and crystal morphology of Na-A zeolite synthesized from SiO\(_2\):Al\(_2\)O\(_3\):3Na\(_2\)O:410H\(_2\)O and 110 °C for 180 min in a NaOH/H\(_2\)O system.

**Table 1**

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mass loss of residue carbon and no phase-transformation in the range of 150–600 °C. The thermal stability limit of the synthesized product was 600 °C. XRD results also indicated the same phenomena as those obtained from TGA and DSC. Even at 700 °C LTA was found as a major product. The baseline was changed to curve and the signal to noise ratio was higher indicating the morphology change of LTA (Fig. 6).

By following the reaction using XRD, the amorphous gel was firstly transformed to crystalline aluminosilicate at 60 min and the transformation was complete at 160 min, as illustrated in Figs. 7 and 8. The SEM micrographs also show the same results as the XRD spectra, and the Na-A zeolite was started to form at microwave heating time of 60 min. The observed Na-A zeolite was in a sharp edge cubic shape which indicated that pre-nuclei might not be formed due to the organic-inorganic micelle formation during the sol-gel process.\textsuperscript{18,24–28} The complex formation not only prevented the precipitation to occur, but also retarded the nucleophilic attack of water, which enhanced the metal–oxide–metal formation (condensation reaction), allowing a three-dimensional network to form. Additionally, the formation of complexes also helped dissolving amorphous aluminosilicate.

![Fig. 6. XRD spectra of calcinated Na-A zeolite at various temperature (room temperature to 1000 °C).](image)

![Fig. 7. XRD spectra of Na-A zeolite synthesized from SiO$_2$:Al$_2$O$_3$:3Na$_2$O:410H$_2$O and 110 °C for $x$ min ($x = 30–180$ min).](image)
gel and slowly precipitating crystalline aluminosilicate resulting in the low defect crystallization. However, we also observed some irregular cubic crystallites of Na-A zeolite, most likely due to the growing of new nuclei after the initial nucleation and precipitation. The Si/Al ratio of the synthesized products versus microwave heating time was reduced from 1.53 to ~1.00, as shown in Fig. 9. Even when we used a Si/Al ratio less than 1, some aluminum became sodium aluminate dissolved in NaOH solution. The Si/Al ratio dropped significantly at 140 min at which almost all of the crystalline products was formed. This is confirmed by the SEM results (Fig. 8). After crystalline formation, the Si/Al ratio approached unity while the Na/Al ratio was greater than one (~1.25).

3.4. Effect of NaOH concentration

In a previous paper, we had reported that Na₂O concentration and temperature affected the microwave heating time. In this work, the source of Na₂O is NaOH, and thus the use of more Na₂O implies
increased OH\(^-\) concentration (OH\(^-\)/SiO\(_2\) ratio = 2(Na\(_2\)O/SiO\(_2\) ratio)). In this investigation, we fixed the microwave heating temperature at 110 °C and the ratio of SiO\(_2\):Al\(_2\)O\(_3\):410H\(_2\)O. The Na\(_2\)O/SiO\(_2\) concentration ratio was varied from 3 to 10. It was found that as the Na\(_2\)O concentration increased, the microwave heating time was reduced dramatically in the range of Na\(_2\)O:SiO\(_2\) ratio from 3:1 to 5:1, as illustrated in Fig. 10. We speculate that this is due to the fact that the increase in Na\(_2\)O concentration increased the hydroxyl concentration, which in turn enhanced the dissolving rate of amorphous gel and the nucleation rate leading to the growth of more crystals. At Na\(_2\)O:SiO\(_2\) ratios higher than 5:1, the reaction time decreased slightly and at the Na\(_2\)O:SiO\(_2\) ratios of 9:1 and 10:1, the microwave heating time was reduced to 5 min. However, high concentration of Na\(_2\)O did affect the particle size and particle size distribution, as shown in Fig. 11. For Na\(_2\)O:SiO\(_2\) ratios of 3:1 to 5:1, the particle size distribution was bimodal with peaks at 2.16 and 7.84 μm, respectively. As the Na\(_2\)O:SiO\(_2\) ratio increased to 6:1 and 7:1, the average particle size of each modal was shifted to lower, 1.62 and 7.08 μm, respectively, whereas at Na\(_2\)O:SiO\(_2\) ratios higher than 7:1, the particle size distribution was changed to mono-modal with smaller particle sizes (less than 4 μm). For Na\(_2\)O:SiO\(_2\) ratios equal to 9:1 and 10:1, the particle size distributions were the same. However, from SEM results (Fig. 12), the crystal size was different. It was found that the crystals agglomerated due to the presence of much more nuclei in the system. As compared to the Na\(_2\)O:SiO\(_2\) ratio of 3:1, the average crystal size was ~4.5 μm which was half of the result obtained from the particle size analyzer. Obviously, the particles tended to agglomerate more than be freeform. Moreover, increasing the Na\(_2\)O concentration did not affect the product composition. All synthesized Na-A zeolites have the Si:Al:Na ratio of ~1:1:1.25.

Since the Na\(_2\)O concentration and hydroxyl group concentration were proportional, when increasing the Na\(_2\)O concentration, the HO\(^-\) concentration increased as well, introducing thus higher pH for the system.

Fig. 11. Agglomerated particle size distribution of Na-A zeolite synthesized at various Na\(_2\)O concentration at SiO\(_2\):Al\(_2\)O\(_3\):xNa\(_2\)O:410H\(_2\)O (x = 3–10) and 110 °C/180 min.

Fig. 12. SEM micrographs of Na-A zeolite synthesized using SiO\(_2\):Al\(_2\)O\(_3\):xNa\(_2\)O:410H\(_2\)O and 110 °C, with x = (a) 3, (b) 7, (c) 8 and (d) 10.
When the pH was fixed by controlling the OH\(^-\) loading, and adding NaCl to increase the Na\(_2\)O concentration was the same with the syntheses carried out under the same OH\(^-\) concentration, implying that the major influence on crystalline formation was OH\(^-\) concentration.

### 3.5. Effect of water quantity

Upon increasing the amount of water from 410 to 510 at the SiO\(_2\):Al\(_2\)O\(_3\):10Na\(_2\)O ratio and using a microwave heating temperature of 110 °C for 5 min, the agglomerate particle size distribution of synthesized Na-A zeolite was almost the same. However, the results from SEM were slightly different, as shown in Fig. 13. Higher water quantity leads to slightly bigger particle sizes (~1.5 μm for 510 H\(_2\)O and ~1 μm for 410 H\(_2\)O). Moreover, more irregular cubic particles were observed. The reason is that when increasing the amount of water the pressure of the hydrothermal system is increased, and as a result, the crystallization process is accelerated. Higher crystallization rates lead to less time to reach the most thermodynamically favored lattice positions resulting in irregular particle sizes and shapes. By fixing the microwave heating temperature at 110 °C and H\(_2\)O quantity at 410, we were able to construct the three-phase diagram, illustrated in Fig. 14. This phase diagram was constructed from the loading condition that can provide LTA product. Only Si:Al loading ratio of 1:2 provided the LTA type zeolite. At Si:Al ratio of 1:1, the mixture of LTA and GIS type zeolite were obtained while at 1:3, a mixture of LTA and amorphous alumina was produced. This is attributed to the enhanced formation of octahedral aluminium atoms at high Al-concentration.

### 3.6. Moisture absorption testing

Comparing moisture absorption ability of two synthesized zeolites at SiO\(_2\):Al\(_2\)O\(_3\):410H\(_2\)O ratio and

![Graph](image1.png)

![Graph](image2.png)

**Fig. 13.** Effect of water quantity on the Na-A zeolite synthesized from SiO\(_2\):Al\(_2\)O\(_3\):10Na\(_2\)O:xH\(_2\)O (x = 410 (a) and 510 (b)) and 110 °C/5 min.
Moisture absorption of synthesized zeolites at SiO\(_2\):Al\(_2\)O\(_3\):Na\(_2\)O = \(x\)Na\(_2\)O\(\times\)Al\(_2\)O\(_3\)\(\times\)Si\(_4\)O\(_{10}\)H\(_2\)O (\(x = 3\) and 10) and 110 °C as compared with commercial Na-A zeolite (purchased form Aldrich)

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4. Conclusions

Na-A zeolite can be synthesized by using alumatrane and silatrane as precursors via the sol-gel process and microwave technique. Due to organic-inorganic micelle formation, the sharp edged crystals were formed indicating a better crystallization. Increased Na\(_2\)O concentration (by increasing NaOH) leads to decreased microwave heating times and smaller particle sizes. The shortest microwave heating time was 5 min when the Na\(_2\)O:Si\(_2\)O\(_2\) ratios of 9:1–10:1 were used. Moreover, the agglomerated crystals were increased as Na\(_2\)O concentration increased while the composition of all synthesized Na-A zeolites was the same (Si:Al:Na ratio of \(\sim 1:1:1.25\)). Increased water quantity influenced the particle size and number of irregular particles due to the pressure built up in the hydrothermal system, which enhanced the nuclei growth and crystallization rates. The moisture absorption ability of synthesized Na-A zeolite was found to be higher than that of commercial Na-A zeolite.

Acknowledgements

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References


Fig. 14. Phase diagram of Na-A zeolite synthesized at microwave heating temperature of 110 °C.