Tailoring Particle Size and Agglomeration State of Mesoporous MCM-48 via Optimisation of Sol-gel Silica Process

Nurul Afiqah Mokri, Oh Pei Ching,* Hilmi Mukhtar and Chew Thiam Leng

CO₂ Research Centre (CO2RES), Institute of Contaminant Management, Department of Chemical Engineering, Universiti Teknologi PETRONAS, 32610 Seri Iskandar, Perak, Malaysia

*Corresponding author: peiching.oh@utp.edu.my

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ABSTRACT: Mobil Composition of Matter no. 48 (MCM-48) is a promising inorganic particle due to its unique crystal structure and 3-dimensional porous network. Incorporating a discrete amount of nanosized MCM-48 for material development would give significant impact to the material’s performance and mechanical robustness. Thus far, the synthesis of MCM-48 in the nanoscale regime has been achieved via chemical modification techniques. However, they are often not reproducible due to limited understanding of surfactant behaviour. On the other hand, physical modification technique is more stable, thereby can reduce the possibility of producing other undesired mesophase. Nevertheless, this technique has not been adapted for nanosized MCM-48 synthesis. In this work, the physical modification technique was adapted to tailor MCM-48 particle size by varying stirring speed (i.e., 1000 rpm, 3000 rpm and 5000 rpm) and using different types of washing medium. Particle size distribution (PSD) studies showed mean particle size of 250 ± 50 nm. XRD analysis displayed several crystalline peaks indexed to cubic crystal mesophase. Transmission electron microscopy (TEM) and Brunauer-Emmett Teller (BET) analyses also showed a unique 3D cubic interconnecting porous network with specific surface area of 1030 m² g⁻¹. It was revealed that the degree of agglomeration increased as relative polarity of alcohol increased, CH₃OH > CH₃CH₂OH. Therefore, particle size and agglomeration state of MCM-48 can be tailored by varying the physical modification techniques while maintaining its intrinsic pore morphology.

Keywords: Sol-gel silica, agglomeration, MCM-48, nanosized, stirring speed
1. INTRODUCTION

In the past two decades, ordered mesoporous silica (OMS) has inspired prominent research interest due to its unique attributes, facile synthesis and ease of functionalisation. Although OMS is considered a conventional filler, it is found to be widely explored in numerous fields due to its tunable pore and surface properties.\textsuperscript{1-3} OMS can be integrated into an organic polymer matrix to fabricate polymer nanocomposites or mixed matrix membranes (MMMs). OMS can be categorised into a class of mesoporous silica member including: Mobil Composition of Matter (MCM-\(n\)), \(n = 41, 48\) and \(50\); Santa Barbara (SBA-\(n\)), \(n = 15, 16\); and Korea Advanced Institute of Technology (KIT-\(n\)), \(n = 6, 46\) OMS is driven by two primary factors: (1) self-assembly of surfactant micelles in lyotropic liquid crystals (LLCs); and (2) cationic exchange between negatively charged silicate species and positively charged head group of surfactants.\textsuperscript{7-11} M41S family can be differentiated according to the space group such as 2D-hexagonal (\textit{p6mm}) and gyroidal cubic \textit{Ia3d} for MCM-41 and MCM-48, respectively.\textsuperscript{6,12,13} Highly crystalline cubic mesoporous materials possess a perfectly ordered and symmetric pore geometry of cubic gyroidal structure (\textit{G}-surface).\textsuperscript{14} The pore structure of hexagonal and gyroid cubic ordered porous arrays are represented in Figure 1.

![Diagram of mesoporous silica pore wall](image)

2D-hexagonal pore structure (\textit{p6mm})

3D-gyroid cubic pore structure (\textit{Ia3d})

Figure 1: Structures of ordered mesoporous materials, M41S: (a) MCM-41 (2D hexagonal, space group \textit{p6mm}), and (b) MCM-48 (cubic, space group \textit{Ia3d}).

To date, both hydrothermal and room temperature synthetic condition can be applied for the synthesis of mesoporous silica. The primary difference between the hydrothermal and room temperature syntheses are reaction temperature, reactant composition, stirring condition, aging time, and the use of co-surfactant as structure
directing agent. The formation of mesoporous silica is dependent on the nature of surfactants such as alkyl chain length and polarity. The surfactant(s) is crucial in determining surfactant packing parameter $g = V/a_o l$, where $V$ is the volume of the hydrophobic chain, $a_o$ is the effective area of the polar head group and $l$ is the alkyl chain length of surfactant. For instance, the packing parameter of gyroidal cubic mesophase surfactant ($G$-surface) should be inherently controlled at surface curvature, $g = 1/2 < g < 2/3$. Generally, silica growth is governed by the rate of silica hydrolysis and condensation. These parameters should be controlled to prevent the formation of secondary particles with larger particle size. Besides that, during silica growth, modification such as the addition of co-surfactant or pH adjustment can be performed to tailor the growing silicate species. However, other complications might arise such as the presence of other undesired mesophase or structure collapse. Therefore, tailoring of MCM-48 via physical modification provides a better option to reduce the particle size. For instance, hexagonal mesoporous silica with reduced particle size has been successfully achieved by tailoring the stirring speed up to 500 rpm. In comparison with MCM-48, stirring speed up to 850 rpm has been applied to single surfactant system at room temperature. It was reported that particle size of MCM-48 synthesised at 450–850 rpm produced particle size distribution between 200 nm and 1000 nm. It is hypothesised that longer duration of mechanical stirring can induce the formation of larger secondary silica particle due to interparticle collision. To date, Kim and his co-workers successfully synthesised nanosized MCM-48 spheres using double surfactant system, i.e., cationic surfactant and non-ionic block-copolymer. The usage of optimum stirring speed and stirring time could minimise the agglomeration of particles. By adapting to this method, synthesis of nanosized MCM-48 can be achieved using minimal amount of silica precursor at a controlled rate of silica addition, under single cationic surfactant system. In this work, stirring speed of disperser was manipulated to tailor the particle size of MCM-48 via single surfactant system. The study was extended to evaluate the effect of washing medium towards the size of MCM-48 particle agglomerates.

2. EXPERIMENTAL

2.1 Materials Selection

Cetyltrimethylammoniumbromide (CTAB), 35% aq. NH$_4$OH, ethanol, tetraethylorthosilicate (TEOS) were purchased from Merck, KGaA, Germany. Polyethylene glycol tert-octyl phenyl ether (Triton™ X-100) was obtained from Sigma-Aldrich. All chemicals were used without further purification.
2.2 Synthesis of MCM-48

2.2.1 Effect of surfactant type on formation of MCM-48

Synthetic solutions with single and double surfactant system were prepared to study favourable condition to synthesise cubic mesostructured siliceous MCM-48. The reactant composition is shown in Table 1.

Table 1: Molar composition at room temperature for single and double surfactant system.

| Sample | Synthesis condition | Surfactant composition | Molar composition mixture |
|--------|---------------------|------------------------|---------------------------|
| S1     | Single surfactant   | 0.41 CTAB              | 0.41 CTAB/11 aq. NH₃/1.0 TEOS/53 EtOH/344 H₂O |
| S2     | Double surfactant   | 0.139 CTAB: 0.028 Tx-100 | 1.0 TEOS/0.139 CTAB/0.028 Triton™ X-100/ 0.468 NaOH/ 54.7 H₂O |

2.2.2 Effect of synthesis condition (room temperature vs. hydrothermal) on MCM-48 formation

The experimental study on the effect of synthesis condition towards the formation of MCM-48 silica was executed at room temperature and hydrothermal condition. The system was designed to determine the optimum reaction temperature for gyroidal cubic MCM-48 formation. Referring to Table 2, single surfactant system was prepared by using cetyltrimethylammonium bromide (CTAB) as templating medium, ethanol (co-solvent), ammonium hydroxide (morphology catalyst), sodium hydroxide (catalyst) and deionised water (solvent). Particle shape and size distribution for the as-synthesised MCM-48 were studied via scanning electron microscopy (SEM) and particle size analyser (PSA).

Table 2: Molar composition of mixture under single surfactant system at different synthesis conditions.

| System | Synthesis condition | Molar composition of mixture (M) |
|--------|---------------------|----------------------------------|
| S3     | Room temperature    | 1 M TEOS/12.5 M aq. NH₃/54 M EtOH/0.4 M CTAB/174 M H₂O |
| S4     | Hydrothermal        | 1 M TEOS/0.25M Na₂O/0.65 M CTAB/0.62M H₂O |
2.2.3 Effect of synthetic solution composition on MCM-48 formation

Under single surfactant system, four different samples were prepared by varying the reactant ratios of surfactant/silica, water/silica, water/ammonia (aq.) and ethanol/surfactant as displayed in Table 3.

| Sample no. | CTAB/TEOS | H$_2$O/TEOS | H$_2$O/NH$_3$ aq. | EtOH/CTAB |
|------------|-----------|-------------|-------------------|-----------|
| S5         | 0.4       | 174         | 13.92             | 135       |
| S6         | 0.38      | 935         | 14.60             | 101       |
| S7         | 0.41      | 344         | 31.27             | 129.3     |
| S8         | 0.64      | 1668        | 33.36             | 135       |

2.2.4 Effect of stirring speed and type of washing solvent

Based on predetermined reactant composition, S8 experimental condition was adopted to tailor MCM-48 by varying the stirring speed (Table 4). The experimental setup and process flow were displayed in Figure 2. First, the synthetic solution was prepared by dissolving CTAB in deionised water at 250 rpm before adding ethanol and aqueous ammonium hydroxide. After 15 min, the solution was transferred to disperser and TEOS was added dropwisely at 0.15 cm$^3$ s$^{-1}$. Then, the homogeneous mixture was aged for 24 h. The solid precipitates were recovered and the CTAB templates were removed at 550°C for 6 h. Post treatment of MCM-48 was executed using different washing medium to reduce the agglomeration of particles. It is hypothesised that strong polarity washing medium such as water, normally causes the particles to stick and clump together after complete drying.

| Sample trial | Stirring speed (rev min$^{-1}$) |
|--------------|---------------------------------|
| S9           | 1000                            |
| S10          | 3000                            |
| S11          | 5000                            |
2.3 Effect of Rotational Motion and Vortex Pattern for Formation of MCM-48 Nanoparticles

T-25 Ultra Turrax Disperser with 1000–5000 rpm is employed. This disperser is expected to homogenise and ensure better dispersion of silicates during sol-gel silica process. By controlling the stirring speed and stirring time, stable nanoparticles will be produced. Additionally, the unique sawtooth design allows large particles (agglomerates) to dissociate at high circumferential speed as shown in Figure 3. However, the resulting shear force and inward rotating mechanism might disrupt the micelle-templating solution. Therefore, stirring time must be kept at a minimum to allow formation of micelle-silicate complex and silica deposition by ion-exchange mechanism (i.e., Br\(^-\) is substituted by Si-O\(^-\)).
2.4 Characterisation of MCM-48

X-ray diffraction data of the MCM-48 spheres were obtained using X’Pert³ Powder, PANalytical, with Cu Kα radiation wavelength of $\lambda = 0.154$ nm in the 2-theta range of $2^\circ$–$8^\circ$ with a scan speed of $1^\circ$ min$^{-1}$. The morphology of MCM-48 was examined from SEM and field emission SEM (FESEM) images using Bruker Nano EDS-Hitachi TM3000 Tabletop Microscope (GmbH, Germany) and VPFSEM, Zeiss Supra55 VP. For the imaging analysis, the sample was deposited on a sample holder with an adhesive carbon foil and sputtered with platinum. Mean particle size and particle size distribution analyses were conducted using PSA, Mastersizer 2000. BET surface analysis was performed using Micromeritics TriStar II. The samples were degassed at 250°C, 1.382 atm for 6 h prior to measurements. Nitrogen adsorption and desorption isotherms were measured at liquid nitrogen temperature of $-195.8^\circ$C. TEM (Tecnai TF20 X-Twin) FEI was employed to analyse the pore geometry of MCM-48 at different incidence planes. The pore geometry and crystal structure of tailored MCM-48 were investigated by tilting the crystals along [111] and [110] zone axes.
3. RESULTS AND DISCUSSIONS

3.1 Effect of Reaction Parameters: Surfactant Type and Synthesis Condition

3.1.1 The effect of surfactant type towards formation of gyroidal cubic MCM-48

The XRD of MCM-48 synthesised via single and double surfactant systems were displayed in Figure 4. The result for single surfactant system (S1) is in agreement with Bragg’s peaks definition for bicontinuous pore network ascribed to gyroidal cubic mesophase. In contrast to S1, S2 which represents double surfactant system showed lack of observable peak at (220), as shown in Figure 4(b). It is hypothesised that the major peak (220) did not appear due to co-existence of other mesophases like hexagonal or lamellar or both. S1 showed well-resolved major Bragg’s peaks at $2\theta = 2.65^\circ$, $3.07^\circ$, and minor peaks at $4.33^\circ$, $4.52^\circ$, $4.94^\circ$ and $5.07^\circ$, as shown in Figure 4(a). This clearly indicated that S1 has a highly ordered cubic mesoporous structure, defined at (211), (220), (400), (421), (332) and (431) and consistent with $Ia3d$ cubic space group.\textsuperscript{22–26} It showed that single surfactant system was favorable because higher degree of crystallinity for cubic mesoporous silica was obtained.

![Figure 4: XRD diffraction pattern for MCM-48 using (a) single surfactant system S1, and (b) double surfactant system S2.](image)
3.1.2 Physical morphology of MCM-48 via single and double surfactant system

Electron microscopy image of MCM-48 shows the distribution of spherical MCM-48 particles formed at room temperature and hydrothermal condition (Figure 5). It was found that double surfactant system shows enhanced monodispersity and particle size compared to single surfactant. It can be deduced that the growth of particles in single surfactant system is due to deposition of primary particles to the stable particle and formation of agglomerates. It may also be due to interparticle contact by collision and aging duration. Longer stirring time and interparticle contact lead to permanent aggregation due to hydrophilic surface of silica.

Figure 5: Electron microscopy images of MCM-48 at 10K magnification using (a) single surfactant, S1, and (b) double surfactant, S2.

3.1.3 The effect of surfactant type on the structure of cubic MCM-48

TEM images confirmed the structure of cubic mesopores for S1. Referring to Figure 6, S1 displayed the intrinsic cubic $Ia3d$ mesopores from incidence plane [111]. As compared to S1, S2 barely displayed any pore formation and no structural order of cubic $Ia3d$ was observed as well. In spite of that, a lettuce-like or layered form of pore structure possessed by the lamellar material was observed. This might be due to several factors such as the use of co-surfactant, pH changes, reaction temperature and aging time.27–29 Even though it results in monodispersed silica sphere and uniform particle size, double surfactant system is not favourable for MCM-48 formation. This is mainly due to extremely sensitive templating solution, wherein a slight change in surfactant volume or chain length will cause the formation of undesired mesophase. Hence, single surfactant system is easier to control in order to further optimise the synthetic condition.
3.1.4 Effect of synthesis condition (hydrothermal vs. room temperature) on MCM-48 via N$_2$ isotherm studies

Referring to Figure 7, S3 and S4 showed a typical Type IV nitrogen adsorption-desorption isotherm. However, capillary condensation accompanied with adsorption hysteresis was observed at p/p$^o$ 0.6 – 0.9 for S3. This occurs when the pore diameter of 43.84 Å exceeds a certain critical width as revealed by BJH desorption at 196.15°C. In principle, hysteresis occurs when the pores are wider than 40 Å. In contrast, S4 revealed a reversible isotherm of Type IV.$^{4,30,31}$ S4 has smaller pore size (28.36 Å) than S3. The pore size of S4 is too large for mesoporous silica, which occurs as a result of change in surfactant volume. The double surfactant system which employed cationic and non-ionic surfactant, experienced a drastic change in hydrophobic surfactant chain length ($l$), leading to pore-expansion of cubic silica mesophase. It is hypothesised that this phenomenon might be induced by addition of hydrochloric acid during micelles-silicates complex which alters the system equilibrium. Overall pore properties such as BET surface area, pore volume, and pore size are presented in Table 5. S4 has larger surface area than S3, along with a slight increase in pore volume and smaller pore size or diameter. Hence, this concludes that mesoporous system of gyroidal cubic $Ia3d$ silica is favoured at room temperature.
Figure 7: Nitrogen adsorption-desorption isotherms for MCM-48 synthesised at (a) hydrothermal S3, and (b) room temperature S4.

Table 5: BET surface area, pore volume and pore size of MCM-48.

| Sample trial                | BET surface area (m² g⁻¹) | BJH desorption pore volume (cm³ g⁻¹) | BJH desorption pore size (Å) |
|-----------------------------|----------------------------|-------------------------------------|-----------------------------|
| Hydrothermal, S3            | 550.79                     | 0.766                               | 43.84                       |
| Room temperature, S4        | 1108.24                    | 0.797                               | 28.36                       |
3.1.5 Effect of synthesis condition (hydrothermal vs. room temperature) on particle size distribution of MCM-48

The particle size distribution (PSD) for MCM-48 synthesised at hydrothermal, S3 and room temperature, S4 revealed bimodal distribution. The particles size range for S3 and S4 are 0.7–1.0 µm and 0.9–2.0 µm, respectively. Although S3 exhibits smaller particle size, BET isotherm in Figure 8(a) showed undesirable pattern indicating weak adsorbent-adsorbate interaction.

Figure 8: Particle size distribution by volume (%) of MCM-48 synthesised via (a) hydrothermal S3, and (b) room temperature S4.
3.1.6 Effect of synthesis condition (hydrothermal vs. room temperature) on particle morphology of MCM-48

SEM images of MCM-48 prepared via room temperature (RT) synthesis showed the formation of MCM-48 with uniform spherical morphology whilst hydrothermally (HT) synthesised MCM-48 showed irregular particles (Figure 9). It was hypothesised that spherical morphology is due to the presence of ammonium hydroxide which not only acts as a basic catalyst but also as a morphological catalyst. To date, spherical shaped MCM-48 is easier to disperse in a wet medium like polymer dope solution, thus making it feasible as an inorganic filler for fabrication of filled polymer or composite membrane film. Furthermore, by reducing the possibility of surface contact within the particles, surface cohesiveness could be diminished due to polar and van der Waals attraction of hydrophilic silica.

![SEM images of MCM-48 (6K magnification) at (a) hydrothermal S3, and (b) room temperature S4.](image)

3.1.7 Effect of synthetic solution composition on MCM-48 particle morphology

The physical morphology of MCM-48 was studied by using SEM and FESEM to ascertain the particle shape, size and aggregation states of each sample. Based on the water to silica composition ratio and cetyltrimethylammonium bromide to tetraethylorthosilicate ratio, different morphological patterns of MCM-48 were observed. In Figure 10(a–d), MCM-48 particle morphologies were seen as: (a) smaller and monodispersed spherical particles; (b) a combination of large and
small particles with fused-like shapes and aggregated particles; (c) monodispersed particles with the presence of smaller unresolved and irregular shapes; and (d) spherical particles with ellipsoidal shaped-like structures in uniform size ranges. Based on microscopic images, it can be concluded that S8 yields the most desired MCM-48 morphology, indicating that reactant molar ratio played a crucial role in silica growth.

Figure 10: Electron microscopy images of (a) S5, (b) S6, (c) S7, and (d) S8 for MCM-48 at different reactant molar composition ratio synthesised in single surfactant system at room temperature.

3.2 Tailoring MCM-48 via Optimised Sol-gel Silica Process

3.2.1 Effect of stirring speed on physical morphologies of MCM-48

Figure 11 displays the morphology of tailored MCM-48. The as-synthesised and calcined MCM-48 tailored at 5000 rpm showed uneven silica growth on stable particles. In contrast, MCM-48 synthesised at 1000 rpm and 3000 rpm yield similar
particle morphology and aggregation states. PSD analysis presented in Figure 12 shows that 3000 rpm is the optimum stirring speed to produce well-defined morphology of silica particles. Beyond that, it will cause undesired irregular silica particles formation since extreme agitation could disrupt the arrangement of micelle aggregates and facilitate agglomeration of siliceous MCM-48.

![Figure 11: FESEM images of calcined MCM-48 viewed at 30K magnification for (a) S9: 1000 rpm, (b) S10: 3000 rpm, and (c) S11: 5000 rpm washed with ethanol as washing medium.](image)

MCM-48 silica tailored at 1000 rpm (S9) and 3000 rpm (S10) showed a narrow particle size distribution as compared to 5000 rpm (S11). At 5000 rpm, MCM-48 shows a broad range of particle size between 150 nm and 1 µm. Whilst for S9 and S10, the smallest particle sizes were determined at approximately 200 nm and 150 nm, respectively. It was concluded that rigorous stirring speed possibly result in significant impact to silica growth and agglomeration of particles whereas the reduction in particle size was observed as stirring speed increased from 1000 rpm to 3000 rpm.
Figure 12: PSD of MCM-48 stirred at (a) 1000 rpm, S9, (b) 3000 rpm, S10, and (c) 5000 rpm, S11 washed with ethanol.
3.2.2 Effect of stirring speed to gyroidal cubic mesoporous structure

Figure 13 shows the XRD diffraction pattern of the as-synthesised MCM-48 stirred at 3000 rpm. Two major significant peaks at 2θ = 3.05 and 3.42 were revealed at (211) and (220). The distortion and unresolved peak of (220) might be due to the presence of distorted portion of pore arrangement in MCM-48 because of rigorous stirring procedure applied.

![XRD diffraction pattern](image)

Figure 13: XRD diffraction of as-synthesised MCM-48 stirred at 3000 rpm and washed with ethanol, S10.

3.2.3 Pore mesostructure analyses of MCM-48 tailored at optimum stirring speed

The HRTEM images for selected particles along the [111] and [110] plane axes are presented in Figure 14(a) which matched well with the reported MCM-48 images.\textsuperscript{32,33} The [111] plane exhibited hexagonal-like porous structure which ascribed to the 211-type cubic structure. The plane [110] plane showed tetragonal crystal surface due to intersection of bicontinuous porous arrays.\textsuperscript{7} Both hexagonal and tetragonal structure represent a body-centred cubic (BCC) structure. As-synthesised MCM-48 at 3000 rpm showed a rhombohedral crystal surface morphology as seen in Figure 14(c). The SAED pattern in Figure 14(d) indicating a BCC mesostructure was formed. It is concluded that stirring speed variation within 1000 to 3000 rpm is able to produce orderly arranged gyroidal cubic mesoporous silica arrays (Ia3d).
Figure 14: Morphology of MCM-48 tailored at 3000 rpm (ethanol washed), for S10 viewed via HRTEM show (a) pore geometry from axis [110] and [111], (b) rhombohedral crystal surface, and (c) SAED image from plane [111].

3.2.4 Surface area and porosity (SAP) analysis of tailored MCM-48

The presence of silanol group density, Si-OH on mesoporous silica wall and N\textsubscript{2} adsorption-desorption isotherm of calcined MCM-48 were determined from SAP analysis via BET method. The specific surface area, pore diameter and pore volume were calculated using Barrett, Joyner and Halenda (BJH) desorption method. Referring to Figure 15, BET surface analysis revealed that N\textsubscript{2} isotherm pattern follows Type IV isotherm, indicating strong sorbate-adsorbate interaction in mesoporous solid, specifically for MCM-\textit{n}, \textit{n}=41,48.\textsuperscript{33,34} It showed that hysteresis peaks occurred at relative pressure of 0.11–0.42 (p/p\textsubscript{0}) due to multilayer adsorption followed by capillary condensation in mesopores. BJH pore analysis revealed an average pore diameter and pore volume of 2.79 nm and 0.545 cm\textsuperscript{3} g\textsuperscript{-1}, respectively.
3.2.5 Effect of washing medium polarity on MCM-48 agglomeration degree and particle size distribution

The effect of washing solution (i.e., water, ethanol and methanol) on tailored MCM-48 were observed via FESEM as seen in Figure 16. MCM-48 particles in S12 and S13 showed a similar pattern of dispersity and surface morphology as compared to S14. It is hypothesised that silicate species polymerise better in the presence of methanol which then leads to the formation of larger silica particles. Furthermore, physical defect was also observed on the surfaces of silica particles. This may occur during washing whereby unreacted silicates present in the mother solution further polymerised and deposited on the surface of nearby particles. Therefore, methanol might not be effective in reducing agglomeration of nanosized MCM-48. Since the polarity of methanol is slightly 0.01 D higher than ethanol (0.69 D), therefore it is suggested that solvent with ≤ 0.5 D might be a better alternative for washing siliceous MCM-48.
Upon further study on PSD, MCM-48 washed using deionised water and ethanol yielded the highest number of particle size within the range of 200–250 nm. Two different patterns of PSD were observed as clearly seen in Figure 17. For ethanol-washed MCM-48, it showed uniform distribution whilst water-washed MCM-48 showed bimodal distribution. It is believed that nanosized range of MCM-48 can be obtained by using water and ethanol as washing medium.
Figure 17: Particle size distribution of tailored MCM-48 washed using (a) deionised water, and (b) ethanol.
4. CONCLUSION

Using a single surfactant system, nanosized MCM-48 in the range of 100–450 nm was successfully formed. We revealed that stirring speed from 1000–3000 rpm can provide better silica dispersion and nucleation of particles during sol-gel silica process. The stirring speed applied during silica addition has facilitated the hydrolysis and condensation reaction within a very short stirring time. Herein, small mean particle size was discovered at 1000 rpm and 3000 rpm, ca. 200 nm and ca. 150 nm respectively. Study using different types of washing medium has shown that ethanol and deionized water are preferable than methanol for stabilising interparticle interaction, thus minimising agglomeration of hydrophilic MCM-48 silica. Furthermore, intrinsic pore properties of cubic $\text{I}_a\text{d}$ MCM-48 were retained even at high stirring rate as revealed by XRD, TEM and $\text{N}_2$ adsorption-desorption analysis. For future recommendation, kinetic of sol-gel silica process such as hydrolysis and condensation could be further controlled by tailoring the rate of silica addition and type of basic catalyst used.

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