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Facile and green synthesis of disordered mesoporous silica as biogenic filler for mixed-matrix membranes

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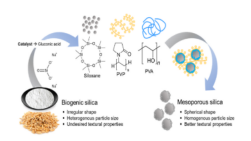
HIGHLIGHTS

GRAPHICAL ABSTRACT

- Mesoporous silica was synthesized using a facile and green sol-gel process.
- Gluconic acid was a worthy replacement for the existing acid catalysts.
- PVA, PVP, and starch had a role to enhance the surface area of particles.
- PVP could govern spherical shape of silica particles.
- PVA/PVP gave a similar effect with PVP but was accompanied by an excellent particle size distribution.

ARTICLEINFO

Keywords: Eco-friendly approach Sol-gel preparation Nanoparticles Disordered mesoporous silica Rice husk



ABSTRACT

Disordered mesoporous silica particles (MSNs) were successfully synthesized using facile sol-gel using rice husk as a silica precursor and water-soluble polymers as structure-directing agents (SDAs) under acidic conditions of gluconic acid solutions. The influence of polyvinyl alcohol/PVA, polyvinylpyrrolidone/PVP, and soluble starch on the textural properties and morph 2 pgies were studied. Besides, the impact of MSNs with considerable properties on the CO₂/N₂ separation performance of poly(ether-*block*-amide) or Pebax membranes was also investigated. SDAs could generally double the specific surface area of the original sample up to 400–600 m²/g, except PVP/starch and PVA/PVP. In addition, PVP governed spherical shape, while its combination with PVA established excellent per cle size distribution (100–200 nm). MSNs prepare 2 using PVA/PVP were even considered to upgrade Pebax membrane's CO₂/N₂ separation performance. CO₂ permeability and CO₂/N₂ selectivity were enhanced up to 58% and 21%, respectively, compared to pure Pebax membranes. This finding will be promising for encouraging sustainable filler development for membrane technology.

1. Introduction

Mesoporous silica nanoparticles (MSNs) are well-known valuable materials with several beneficial textural properties, excellent

biocompatibility, and high flexibility toward surface modification [1,2]. As a result, it has received massive reliability for comprehensive applications such as solid-state sensors [3], biomedical materials [4], catalysts [5], aquatic pollutants removal [6], and gaseous pollutants

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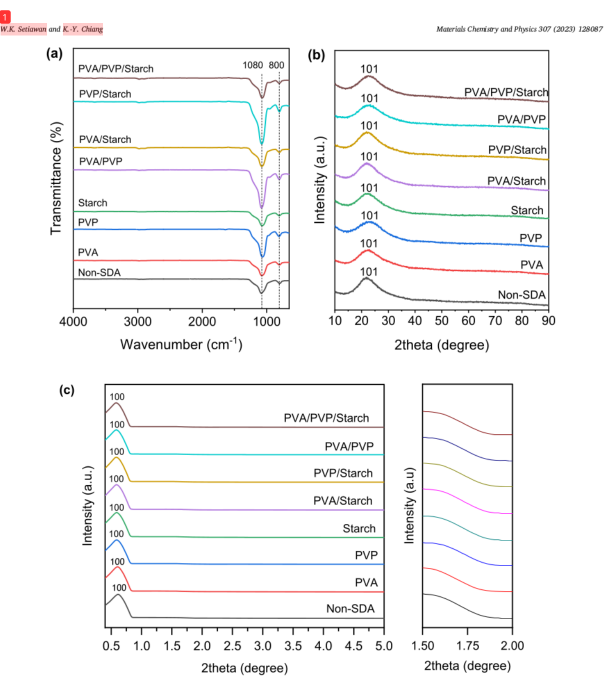


Fig. 1. FTIR spectra (a), wide-angle XRD pattern (b), and low-angle XRD pattern (c) of MSN with different structure-directing agents.

removal [7]. Moreover, the global mesoporous silica market is continuously growing and is predicted to value around USD 295.1 billion in 2027 [8]. Therefore, developing newly mesoporous silica with tunable properties is worth fulfilling the global market demand.

The Stober method recently prepared well-structured mesoporous silica [9–13]. However, it is challenging in large-scale production due to the utilization of high-cost silica precursors and the high-risk generation of harmful residues. Thus, seeking inexpensive and eco-friendly materials is necessary. As a result, waste-based materials from agricultural and industrial sectors, such as rice husk [14,15], bamboo leaf ash [16], banana peel ash [17], electronic waste [18], and ore tailings [19,20],

were currently utilized as precursors to synthesize mesoporous silica. Likewise, less harmful acid catalysts were also used in the synthesis process, mostly from carboxylic acid families such as citric acid [21] and acetic acid [22]. In addition, the water-soluble templates might also be preferable to avoid the excessive employment of toxic solvents. Hwang et al. [22] used polyethylene glycol (PEG) as a morphology-directing agent to prepare mesoporous silica from sodium silicate. It was found that PEG exposed an ability to govern particle shape and mesopores arrangement. Another water-soluble polymer, polyvinyl alcohol (PVA), was reported by Thahir et al. [23] to synthesize the high surface area of mesoporous silica. While promising, several previous studies indicated

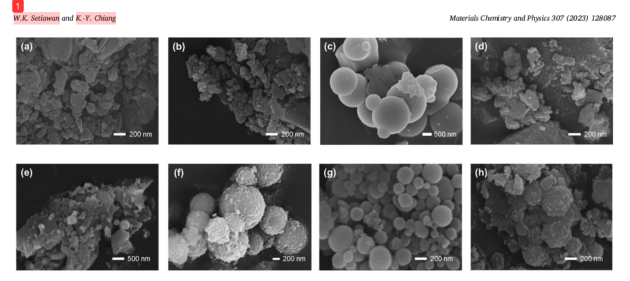


Fig. 2. SEM images of MSNs synthesized without SDA (a) and with PVA (b), PVP (c), starch (d), PVA/starch (e), PVP/starch (f), PVA/PVP (g), and PVA/PVP/ starch (h).

that the applications of water-soluble polymers (e.g., starch and polyvinylpyrrolidone) in mesoporous silica synthesis were limited as a co-template along with the surfactant [24–27]. Thus, a further investigation of these materials is necessary to establish.

Employing filler materials from inexpensive and natural resources became a highly rated alternative to improve the gas separation performance of polymeric membranes [28]. For example, Bhattacharya and Mandal [29] have synthesized mesoporous silica derived from rice straw using precipitation. The silica sol was then incorporated into a polyether-polyamide block co-polymer matrix to obtain mixed matrix membranes. Waheed et al. [30] also employed an identical technique to generate rice husk-derived silica as a filler in polysulfone membrane for CO₂ separation. However, further studies would be necessary concerning poor dispersion between polymer-filler, causing overall membrane performance decrement.

This study developed a one-pot green synthesis of mesoporous silica for the first time by a sol-gel process using rice husk as a silica precursor and three different water-soluble polymer structure-directing agents (SDAs) under acidic conditions of gluconic acid solutions. Remarkably, the ability of polyvinyl alcohol (PVA), polyvinylpyrrolidone (PVP), and soluble starch to govern silica's pore structure and morphology was comparably studied. In addition, the potential of the best MSN for improving CO_2/N_2 separation features of poly(ether-block-amide) membranes was also investigated.

2. Materials and method

2.1. Materials

The rice husk was obtained from a local rice mill in Taoyuan City, Taiwan. D-gluconic acid ($C_6H_{12}O_7$ 50% aqueous solution) and soluble starch were acquired from Alfa Aesar. Sodium hydroxide (NaOH) was purchased from Showa Chemical. Polyvinyl alcohol (Mw = 89,000–98,000, PVA) and polyvinylpyrrolidone (Mw = 40,000, PVP) were supplied by Sigma Aldrich.

2.2. Mesoporous silica synthesis

Rice husk silica (RHS) was obtained by the process described elsewhere [31]. RHS and 2 mol L^{-1} , NaOH at a ratio of 1:10 (w/v), were refluxed to get a sodium silicate (Na₂SiO₃) solution. Mesoporous silica nanoparticle (MSNs) synthesis procedures were modified from another study [22]. Initially, 20 ml Na₂SiO₃ solution was mixed with 20 ml polymer solution (5% w/v). The weight ratio was controlled at the same level for the SDA mixtures. Next, gluconic acid was gradually added d adjust the pH to 4. The sol was then aged at 60 °C overnight. Finally, it was filtered and calcined at 550 °C for 2 h to obtain MSNs.

2.3. Membrane fabrication and gas permeation analysis

Dried Pebax pellets were dissolved into ethanol/water (70/30) at 70 °C for 2 h to obtain 4 wt% polymer solution. At the same time, the filler was separately dispersed into ethanol/water and sonicated at 25 °C for 2 h. The dispersed filler was mixed with polymer solution for another 2 h reflux process. Consequently, it was cast on a 6 mm PFTE circular dish and evaporated under ambient temperature for 48 h. Finally, the Pebax membrane was dried at 50 °C for 24 h under vacuum conditions. The single gas permeation of membranes was measured by custom permeation equip.² ent at 25 °C and 1 bar operation. The single gas permeability, P (1 Barrer = 10^{-10} cm³ STP cm⁻²s⁻¹cmHg⁻¹) was determined using the following equation.

$$P = 10^{10} \frac{Q \times l}{\Delta P \times A}$$

Q is the permeate flow rate (cm³/s STP), *l* is the membrane thickness (cm), *A* is the effective area (3.069 cm²), and ΔP is the transmembrane partial pressure difference (cmHg). The ideal CO₂/N₂ selectivity was determined using the equation:

$$\alpha_{CO_{2/N_2}} = \frac{P_{CO_2}}{P_{N_2}}$$

where P_{CO2} and P_{N2} are the permeability of CO₂ and N₂, respectively.

2.4. Characterization methods

The wide-angle and low-angle X-ray diffraction (XRD) patterns of MSNs were characterized by XRD Bruker D8 Advance at $2\theta = 10-90^{\circ}$ and XRD D2 Phaser at $2\theta \equiv 0.4-10^{\circ}$, respectively. The MSNs surface properties were investigated using Fourier-transform Infrared (F1R) Frontier at 650-4000 cm⁺¹. The Brunauer EmmetteTeller (BET) and Barrett-Joyner-Halenda (BJH) analyses were executed by nitrogen adsorption measurement using an Autosorb iQ instrument. The morphology of MSNs was observed using JEOL JSM-7600F Field Emission Scanning Electron Microscope (FESEM) and JEOL JEM-2100

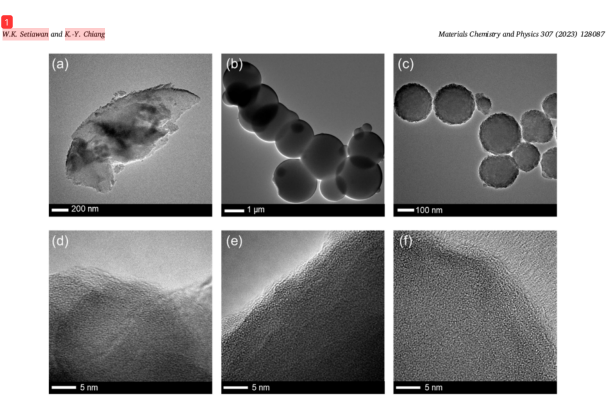


Fig. 3. TEM images of MSNs synthesized without SDA (a-d), with PVP (b-e), and with PVA/PVP (c-f).

Transmission Electron Microscope (TEM).

3. Results and discussion

FTIR analysis (Fig. 1a) indicated that all samples displayed identical spectra at around 1080 $\rm cm^{-1}$ and 800 $\rm cm^{-1}$, ascribed to the asymmetric

and symmetric stretching vibration of siloxane bonds \equiv Si–O–Si \equiv , respectively [32]. The WAXRD analysis (Fig. 1b) revealed a broad peak at $2\theta = 22^{\circ}$ (1 0 1) for all MSNs, corresponding to amorphous silica nature. Moreover, the LAXRD pattern (Fig. 1c) showcased an intense (1 0 0) reflection at $2\theta = 0.58^{\circ}$ and a broad weak shoulder in the 2θ range 1.5° – 2.0° for all MSNs samples, referring to wormholes pore

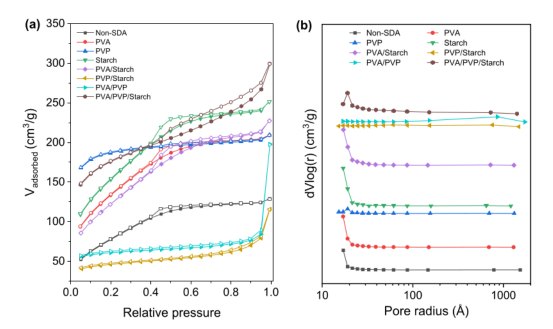


Fig. 4. The nitrogen adsorption/desorption isotherms (a) and pore size distribution (b) of MSN at different structure-directing agents.

Table 1

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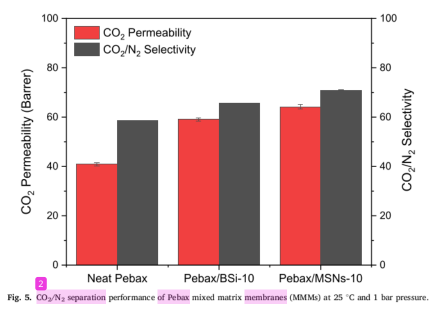
| Sample | Specific surface area (m ² /g) | Total pore volume (cm³/g) | Average pore radius (Å) | BJH deportion summary | | |
|------------|---|---------------------------|-------------------------|---------------------------------------|----------------------------------|-----------------|
| | | | | Pore surface area (m ² /g) | Pore volume (cm ³ /g) | Pore radius (Å) |
| Non-SDA | 293 | 0.20 | 13.7 | 57 | 0.063 | 17.1 |
| PVA | 486 | 0.32 | 13.3 | 96 | 0.100 | 17.0 |
| PVP | 574 | 0.32 | 11.3 | 23 | 0.036 | 19.1 |
| Starch | 553 | 0.39 | 14.1 | 131 | 0.145 | 17.1 |
| PVA/Starch | 451 | 0.35 | 15.6 | 139 | 0.165 | 17.1 |
| PVP/Starch | 146 | 0.18 | 24.5 | 28 | 0.114 | 17.1 |
| PVA/PVP | 298 | 0.43 | 28.9 | 30 | 0.287 | 15.3 |
| PVA/PVP/ | 570 | 0.46 | 16.3 | 127 | 0.221 | 19.1 |
| Starch | | | | | | |

arrangement. Park and Pinnavaia [33] argued that these patterns indicated a correlated pore network distribution lacking a regular pore structure. Regarding FTIR and XRD results, it was then implied that the employment of PVA, PVP, starch, and their mixtures could not alter the pore framework of mesoporous silica.

SEM and TEM imaging were performed to assess the morphology of the prepared MSNs, as seen in Figs. 2 and 3. In the absence of SDAs, irregular shape and heterogenous particle size were observed. The employment of SDAs was expected to avoid particle aggregation and govern into uniform morphologies. PVA and starch, however, could not effectively configure the morphology despite the smaller particle size (see Fig. 2b and d). Moreover, their mixture was not even better with the aggregation phenomenon (Fig. 2e). In contrast, PVP became a better SDA concerning its ability to arrange micro-sized spherical particles of MSN with smooth surfaces (see Figs. 2c and 3b). The combination of PVP with other SDAs promoted rough surfaces with smaller particle sizes. Furthermore, PVP/PVA interestingly offered dispersed-spherical particles with uniform smaller particle sizes (100-200 nm), as illustrated in Figs. 2g and 3c. TEM images (Fig. 3d-f) were also beneficial to demonstrate the pore geometry of MSN. The wormhole-like pores were more visibly found on the outer surface of the MSN sample. It was because the inner part of pores was predominantly arranged by microsized gaps, which hindered the appearance of pore geometry.

Fig. 4a displays the nitrogen adsorption-desorption isotherms of MSN samples. All models exhibited type II closely related to type IV adsorption isotherm, confirming the mixture of micro-mesoporous structure. In addition, different hysteresis loops were observed, suggesting the exist 1 ce of pores with non-identical pore shapes and sizes. Non-SDA MSNs showed a type-H4 hysteresis loop from $P/P_0 = 0.35$ to 0.75, indicating a limited amount of mesopores with slit-shape. Likewise, the PVA-MSN, PVP-MSN, Starch-MSN, PVA/Starch-MSN, and PVA/PVP/Starch-MSN showed an identical profile at $P/P_0 = 0.40$ to 0.99. Meanwhile, the PVP/Starch-MSN and PVA/PVP-MSN revealed a type-H3 hysteresis loop at $P/P_0 = 0.05$ to 0.99. Those indicate 1 wedge-shaped pores with broad size distribution. Fig. 4b depicts the pore size distributions (PSD) calculated by the Barrett-Joyner-Halenda (BJH) model. Most MSN samples presented an identical unimodal distribution with a mean pore radius of 13.00–16.25 Å (see Fig. 4b and Table 1). Only PVP/Starch-MSN and PVA/PVP-MSN demonstrated a wide distribution with a larger mean pore radius at 24–29 Å.

BJH desorption results, as listed in Table 1, showed that single employment of SDA could significantly alter cumulative pore volume and surface area. PVA and starch positively impacted these pore properties, whereas PVP gave smaller values than non-SDA-MSN. The effect of PVP was also observed in the mixed SDAs (PVP/starch and PVA/PVP). The other SDA mixtures (PVA/starch and PVA/PVP/starch) likewise promote identical influences following their single dominant components. Furthermore, BET analysis implied that the presence of SDAs in the synthesis process could generally build up enormous narrow pores, elevating their specific surface area ($400-600 \text{ m}^2/\text{g}$). On the contrary, PVP/starch and PVA/PVP presented relatively unchanged or even lower specific surface area ($150-300 \text{ m}^2/\text{g}$) due to larger pore size with low



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occupancy.

Considering morphology, particle size distribution, and textural properties, mesoporous spherical biogenic silice 2 anoparticles (MSNs) prepared by PVA/PVP could be chosen as a filler to improve the CO₂/N₂ 2 formance of Pebax membranes. MSNs exhibited a better impact on CO₂/N₂ separation features of Pebax membranes compared to original biogenic silica (BSi) from rice husk (as shown in Fig. 5). In particular, MSNs could homogenously spread out without severe tortuosity and have an excellent interfacial link-up within the polymer matrix. Hence, the appropriate pore aperture of MSNs (average pore size = 2.89 nm, as indicated in Table 1) could enable Knudsen diffusion transport for CO₂/N₂. As a result, Pebax membrane comprising 10 wt% MSNs exhibited 64.19 Barrer of CO₂ permeability (1.6 higher than pure Pebax) and CO₂/N₂ selectivity of 71 (1.2 higher than pure Pebax).

4. Conclusion

Disordered mesoporous silica was successfully synthesized by facile sol-gel process rice husk as a silica precursor and water-soluble polymers as structure-directing agents under acidic conditions of gluconic acid solutions. PVP could effectively control particle morphology into spherical shapes, whereas PVA and starch enhanced specific surface area. Moreover, PVA/PVP showed a remarkable particle size distribution without severe agglomeration 2 MSNs prepared using PVA/PVP exhibited more significant effect on CO_2/N_2 2 paration performance of Pebax membranes compared to original BSi. CO₂ permeability and CO₂/ N_2 selectivity were upgraded to 58% and 21% by incorporating MSNs into Pebax membranes. This finding may incubate the development of biogenic filler in membrane technology for CO₂ separation.

CRediT authorship contribution statement

Wahyu Kamal Setiawan: Conceptualization, Writing – original draft, Writing – review & editing. Kung-Yuh Chiang: Methodology, Reviewing; Revising, Supervision.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

The data that has been used is confidential.

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