Organo Silica Membranes for Wetland Saline Water Desalination: Effect of membranes calcination temperatures

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Abstract. Wetland saline water has great potential to overcome water scarcity due to high salinity of intruded seawater. This work determines performance of silica membranes using organo catalyst applied for wetland saline water desalination via pervaporation and investigates the effect of calcination temperatures. These membranes were prepared from precursor tetraethyl orthosilicate (TEOS) for 3 h through sol-gel process refluxed at 0°C (membrane A) and 50°C (membrane B). The sols were dipcoated onto alumina (Al2O3) support for 4 layers followed by calcination for 1 h. Performance of membranes were evaluated by feeding wetland saline water through desalination at room temperature. Results show the water flux for membrane A were 0.27 & 0.15 kg.m^{-2}.h^{-1} and salt rejection were 97.5 & 99 % as a function of calcination temperature (200 & 250°C), respectively. Furthermore, water flux of membranes B were 0.90 & 0.93 kg.m^{-2}.h^{-1} and excellent salt rejection (>99.9 %) for both calcination temperatures. The highest water flux and salt rejection were found for membranes B. For both using citric acid under refluxed and calcination process, it gives more vibration of Si-C formation and membrane pores. This membrane is the very first and mesoporous organo silica membranes which is successfully fabricated from organo catalyst.

1 Introduction

Increasing of water scarcity issues when the earth consist of 97.5 % water but only 2.5 % of proper consumption gives the big impact for fulfillment of life’s needs. Indonesia is in high stress level (40-80%) representing the ration of withdrawal to supply of water [1]. The greatest issue of world’s water scarcity has been encountered in Indonesia which is as the world’s largest archipelago. South Kalimantan as the largest island in Indonesia characterized by wetland whereas the height of soil at 0.16 under sea level is important in hydrology and big potential as water reservoir. The rivers connect to the sea. During rainy season the seawater intrudes into rivers and salty water formed. However, wetland saline water as a big source of clean water requires for contaminant removal process.

Desalination using membrane is one of technology to increase quality of wetland saline water. Desalination technology enlarged widely through varying method by using membranes, such as reverse osmosis, membrane distillation and pervaporation [2]. Pervaporation comes as other method accompanying with high performance, lower cost and high level operating [3]. Pervaporation operates on changing phase to separate between water molecule and hydrated ion under vacuum condition employing selective semi-permeable membrane. The mechanism shows in Figure 1 (b).

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Commonly, inorganic membrane from silica based is affordable to be used for pervaporation process due to its simple fabrication technique, more stable for high temperature, robustness, offering high molecular sieving and relatively low cost [4, 5]. Besides, having 5 Å silica amorf as structural molecular sieving properties is suitable particularly when applied in desalination. Thus, membrane has high potential to separate small molecule water and hydrated salt ion through pore small pathway [6, 7]. TEOS (tetra ethyl ortho silicate) as hydrophobic inorganic material from metal alkoxide (precursor) is skillfully utilized for membrane synthetic[8].

Desalination using silica membrane has been investigated to remove the salt content in the raw water. However, most of separation technology using membrane has big problem in hydrostability. Consequently, membrane becomes dense and swell earlier thus decreasing on water flux and salt rejection[9]. Membrane damage because of silanol (Si-OH) content which is high reacted with water. Solving for decreasing performance of membrane due to hydrophilic, some strategies was run for membrane modification. The strategies consist of surface functionalization using metal doping [10], carbonized templetting [11]and induced organo material into silica matrices [12].
Previous work has been reported that organosilica membrane was prepared by templating carbon material such as P123 and pectin which is approve to enhance silica membrane performance [13, 14]. However, that method was spent much time to templating carbon agent into silica thin film. In this work, fabrication of organosilica membrane takes short time relatively than templating method by using organo catalyst. Organo catalyst from citric acid acts as carbon sources which able to increasing hydrostability of silica matrices [15].

In addition, rapid thermal processing (RTP) technique is more attracted to applies for membrane calcination also in this work. RTP technique is offering shorten time to production of membrane and lessen the cost [16]. This work aims to evaluate the performance of organosilica membrane using precursor TEOS and citric acid catalyst for wetland saline water desalination via pervaporation and investigate effect of calcination temperature.

Figure 1. Mass transfer mechanisms of (a) separation technology using membrane; (b) desalination via pervaporation

2 Experimental

2.1 Materials and Synthesis of Membrane

In this section, some of material need initial treatment before mixed. Firstly, pretreatment was prepared to precursor (TEOS, 99.0%, Sigma-Aldrich), 0.0007 N citric acid, ethanol solution (EtOH 70%), water and support membrane (-Al₂O₃). Organosilica membrane was synthesized through sol-gel process including hydrolysis and condensation reaction for 3 h by employing citric acid catalyst. Silica sol was refluxed at 0℃ and 50℃. Ratio molar TEOS:EtOH:C₆H₅O₇:H₂O is 1: 38:0.0007:5.

Then, for dip-coating step silica sol was poured into centrifuge cube putting on Erlenmeyer. Support membrane was stapled on dip-coater (run in 5 cm/s) for 2 minutes in soaked. Coated membrane on support membrane was calcined at 200℃ and 250℃ for 1 hours each. Coated membrane was cooled at room temperature and continue to repeating dip-coating for 4 times to form 4 layers of membranes.

2.2 Material and Characterization of membrane

This process employed citric acid as organic catalyst. Firstly, EtOH was added into reagent bottle followed by cooling process at 0℃ and stirring at 250 rpm for 5 minutes. Then TEOS was dropped wise and resting. After resting for 5 minutes TEOS and EtOH mixture was dropped wise by C₆H₅O₇ to produce sol through one-step sol gel process.

Silica sol was dried in oven at temperature controlled 60℃ for 24 hours to create dried gel. Dried gel (xerogels) was subsequently grounded into fully homogeny powder and calcined in temperature controlled at varying temperature 200℃ and 250℃ for 4 h. Xerogels were characterized using FTIR at wavelength range 1400-600 1/cm.

2.3 Characterization Using FTIR

FTIR as one method of indirect analysis can assist to provide information of functional group within the silica network[17, 18]. According to presence hypothesis of silanol, siloxane and Si-C bonds in matrix, Infrared spectra was carried out along with 1400-600cm⁻¹ wavelength. Socrates (2011) listed some peak relates to organic compound and silicone alkoxide. Wavenumber for Si-C, Si-O-Si and Si-OH are 1260 cm⁻¹, 1000-1100 cm⁻¹, and 940 cm⁻¹, respectively [19].

3 Results and Discussion

3.1 Organosilica Membrane and Hydrostability

Organosilica membrane was prepared through sol gel process including hydrolysis and condensation reaction. Sol gel is a process to synthesize inorganic compound through chemical reaction in solution under low temperature converting sol to gel. Inorganic acid was rarely used for sol gel reaction, such as HCl, HNO₃ and H₂SO₄. On the other hand, organic acid was also used as catalyst for gel forming and its physical properties [20]. TEOS is in solution phase and as major material to synthesize inorganic membrane. Precursor TEOS will
be reacted during hydrolysis and condensation process to form silica network. During hydrolysis reaction, alkoxide group (OR, whereas R is alkyl group, C$_{x}$H$_{2x+1}$) was replaced by hydroxyl group (OH). Then, silanol group (Si-OH) reacted and produced siloxane (Si-O-Si), alcohol (R-OH) and water. Equation of hydrolysis, alcohol condensation and water condensation were described in equation (1), (2), and (3), respectively.

\[ \text{Hydrolysis: } \equiv \text{Si} - \text{OR} + \text{H}_2\text{O} \rightarrow \equiv \text{Si} - \text{OH} \]

\[ \text{Alcohol Condensation: } \equiv \text{Si} - \text{OR} + \text{Si} - \text{OH} \rightarrow \equiv \text{Si} - \text{O} - \text{Si} - \equiv + \text{R} - \text{OH} \]

\[ \text{Water Condensation: } \equiv \text{Si} - \text{OH} + \text{HO} \rightarrow \equiv \text{Si} - \text{O} - \text{Si} - \equiv + \text{H}_2\text{O} \]

Due to affinity of amorphous silica, structural degradation happen when exposing into water. Briefly, silica surface has big tendency on physisorption of water molecule on silanol groups (Si-OH) during hydrolysis step as shown in equation (1). Hence, acid and base catalyst help for up taking of water. Big problem of using hydrophobic membrane is hydrostability. There are many researchers have been delivered for surface silica modification. In particular, there are three method, such as carbon template silica, hybrid organic-inorganic silica and metal oxide silica[21]. During on latest decade, technique to increase hydrostability was proposed on existence of organic compound that tailored into silica network [22]. Existence of carbon side effected in hydrophobic properties. Some of modification to improve existence carbon compound described into some methods. Modification in templating carbon using Pluronic triblock copolymers [11] and surfactant. Besides, Elma et al (2015) synthesize silica membrane from precursor TEOS and Pluronic P123 as templating agent through sol gel process.

Micro structure from silica membrane was determined by reactivity, precursor size and exacting composition among precursor concentration, alcohol and catalyst. For silicon alkoxide, acid condition was particularly produced structural fractal of sols and encouraged to mesopored silica [3].

In this work, membrane was prepared through sol gel process employing citric acid catalyst for two types of reflux condition, 0°C and 50°C followed by calcination at 200°C and 250°C. Organic catalyst has two responsibilities that are for supporting high kinetic reaction in sol gel process and as carbon source to assist silica-carbon bond into silica network.

3.2 Characterization of Xerogels

Hydro-stability and pore size are the fundamental qualifications in determining the characterizations of a membrane applied to water-contaminant separation techniques. In order to increase hydro-stability and robustness of membrane against both physical and chemical resistances, carbon silica configuration is an important group tailored into silica network. In Figure 2 wavelength at 1068 cm$^{-1}$, 926 cm$^{-1}$, and 788 cm$^{-1}$ are wavenumber of siloxane, silanol and silica-carbon, respectively.

As shown in Figure 2, there are two differences trend line between two types of membrane, type A for 0°C refluxed and type B for 50°C refluxed. Sol gel process systematically includes hydrolysis, condensation, aging and drying. Reflux temperature effects polymerization process particularly in aging step [23]. Particles grow fast under reflux condition thus physically effects in ratio of bond to form silica network.

After sol gel process finished, then continued to calcination step as endothermic decomposition process to discharge gas contaminant, for example carbonate or hydroxide ion. Calcination process consists of (a) discharging of free water (H$^2$O) and bound water (OH) at 100-300°C; (b) discharging of gas at 600°C followed by significant mass decreasing and at 800°C assist to network damaging [24]. For both variation low temperature (200 and 250°C) and types of catalyst do not show significant differences of trend line. Interestingly, for silica carbon is able to stick out into silica network. Existence of silica carbon into silica network support to high robust of membranes.

3.3 Permeation Performance

Duke et al (2009) reported that high salt rejection was resulted during desalination via pervaporation using only oxide silica membranes. Elma (2013) prepared silica membrane through two step sol gel reaction using TEOS by employing nitric acid-ammonia (as catalyst). In order to investigate membranes quality, it was applied to desalinate 0.3–15wt% sodium chloride feed solution, respectively [25].
The pervaporation process gives many variables for execute desalination performance. It takes two aspect: (1) flux of water and (2) salt rejection. Also, permeate water is captured in a condenser and flux (kg. m^{-2} h^{-1}), F, of water during given time period is calculated as $F = \frac{M}{S.t}$, where $M$ is mass of permeate (kg), $S$ is the surface area of membrane (m²) and $t$ is the performance test time (h). Also, percentage of salt rejection was described in mathematical formulation as $R = \frac{(C_f - C_p)}{C_f} \times 100\%$, where $C_f$ and $C_p$ are the concentrations of salt in the feed and permeate solutions, respectively, using conduct meter measurement.

**Table 1. Desalination performance comparison for silica membranes**

| Types of silica membrane | Desalination Method | Feed Temp. (°C) | Feed concentration (%) | Water Flux (kg.m^{-2}.h^{-1}) | Salt Rejection (%) | References |
|--------------------------|---------------------|-----------------|------------------------|-------------------------------|-------------------|------------|
| Organo silica            | Pervaporation       | Room temp.      | Wetland saline water   | 0.93                          | >99               | This work  |
| Organo silica            | Reverse osmosis     | 20              | 0.3-3.5                | 4.7-2.5                       | 93.7-83           | [26]       |
| Organo silica            | Non-osmotic         | 30              | 0.2                    | 3.0                           | 99                | [27]       |
| Pure silica (TEOS pH 6)  | Pervaporation       | 22              | 3.5-15                 | 6.8-1.6                       | 98-89             | [25]       |
| Organo silica            | Pervaporation       | 60              | Brackish (1%-w) Brine (15%-w) | 26.5 | 99.5 | [28] |

![Figure 3 (a) schematic illustration for salt and water transport during pervaporation, (b) representation of silica matrix](https://doi.org/10.1051/e3sconf/202014807006)

Performance in separation using membrane aims to measure membrane capability for separating salt ion and water molecule. High water flux and salt rejection effect in high membrane performance. On the other hand, performance was effected in membrane structure (mesopore).

In Figure 4 shows the water flux for membrane A (refluxed at 0°C) were 0.27 & 0.15 kg.m^{-2}.h^{-1} and salt rejection were 97.5 & 99 % as a function of calcination temperature (200 & 250°C), respectively. Additionally, water flux of membranes B were 0.90 & 0.93 kg.m^{-2}.h^{-1} and excellent salt rejection (>99.9 %) for both calcination temperatures. The highest water flux and salt rejection were found for membranes B.

Calcination affects membrane performance basically due to existence of silica-carbon in silica network. Table 1 shows comparison among similar research about desalination. Organosilica employing organic compound for sol gel process has a big contribution in desalination.

### 4 Conclusion
By employing citric acid catalyst through sol gel reaction refluxed at 50°C, organo silica membranes were successfully synthesized. Compared to many other method for existing carbon compound, employing acid catalyst could be able to eliminate one step process in templating carbon for cost decreasing relevance.

Organo silica was applied into desalination via pervaporation feeding by wetland saline water at room temperature. The water flux and salt rejection values show an excellent value for membrane refluxed at 50°C. Interestingly, the IR spectra shows carbon chains in the silica matrices. It comes from organo catalyst employed during sol gel process.

The authors gratefully thank the facilities of Process Technology Laboratory, Engineering Faculty, University of Lambung Mangkurat. Riani Ayu Lestari thanks to KURITA-AIT Research Grant 2018. Muthia Elma thanks to Alumni Grants Scheme Batch 2 (2018-2019); Applied Research Universities Grant (2019-2020); Grant of Thesis Magister 2019 and Grant of Post-Doctoral (2017-2018) Directorate of Research and Community Service, the Ministry of Research, Technology and Higher Education Republic of Indonesia.

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