Microchannel structure of ceramic membranes for oxygen separation

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Abstract

Microchanneled ceramic membranes have demonstrated superior performance in oxygen separation from air over conventional membranes. In this study, the contributions of the microchannel structure to the superior performance were investigated. Compared with supported membranes, the microchanneled membranes provide fast pathways within the channels for gas diffusion as compared to the tortuous interconnection of pore channels in the supported membranes. The walls of the numerous channels provide a large surface for facilitating oxygen dissociation, which was confirmed by varying the channel wall surface using mesh templates with different aperture sizes. In summary, the microchannel structure facilitates gas diffusion and provides a large membrane active surface, resulting in high performance in oxygen separation.

1. Introduction
Ceramic membranes for oxygen separation from air have attracted significant attention because the ceramic membranes operate via oxygen ion diffusion through dense solids and show theoretically 100% selectivity of oxygen separation. It is also considered as a cheaper technology than cryogenic separation, especially when the oxygen separation using ceramic membranes at high temperatures is coupled with high temperature reactions, such as natural gas conversion, coal gasification and oxyfuel combustion [1-5].

Two major issues have hindered the commercialisation of oxygen permeation membranes: low oxygen permeation rate and poor membrane material stability. Some materials with high intrinsic oxygen ion transfer rates show poor stability. For example, $\text{Ba}_{0.5}\text{Sr}_{0.5}\text{Co}_{0.8}\text{Fe}_{0.2}\text{O}_3-\delta$ and $\text{SrCo}_{0.8}\text{Fe}_{0.2}\text{O}_3-\delta$ membranes could produce an oxygen flux above 5 ml cm$^{-2}$ min$^{-1}$ at 900 °C, but they are unstable in the presence of CO$_2$ or reducing atmosphere due to chemical decomposition [6-8]. Many materials having strong chemical toughness exhibit low oxygen permeation rates; a typical example is the dual phase membrane of 40 wt % NiFe$_2$O$_4$ and 60 wt % Ce$_{0.9}$Gd$_{0.1}$O$_{2-\delta}$ [9].

One approach to overcoming these issues is to develop new membrane geometries and microstructures that can bring significant improvements in oxygen permeation because the membrane geometry and microstructure are not limited by materials. Hence, the materials with good chemical stability can be used to improve the oxygen permeation by modifying membrane geometry and microstructure. In our previous work, microchanneled membranes were fabricated using a mesh-templating phase inversion process [10]. With a superior structure to conventional supported membranes, the microchanneled membranes possess well-organized microchannels sitting on a thin dense layer with the other end open on the other side of the membranes. The membranes were formed by a one-step phase inversion process, and the integrated structure has high resistance to thermal shock. This
microchanneled membrane produced high oxygen fluxes. For example, La$_{0.6}$Sr$_{0.4}$Co$_{0.2}$Fe$_{0.8}$O$_{3-\delta}$ microchanneled membranes achieved oxygen fluxes of 3.2 ml cm$^{-2}$ min$^{-1}$ at 1000 °C and 2.45 ml cm$^{-2}$ min$^{-1}$ at 950 °C. In addition, applying Pt catalyst on the channel wall surface effectively improved the oxygen fluxes [11]. However, it is not clear precisely how the microchanneled structure contributes to the high oxygen fluxes.

In this study, we designed experiments to explore the advantages of the microchannel structure for oxygen separation from air. Conventional supported membranes were prepared to compare gas diffusion through their tortuous pore channels with that through the straight channels within the microchanneled membranes. The contribution of the channel walls for surface reaction (oxygen dissociation) is also investigated by varying the channel size using different mesh templates.

2. Experimental

2.1 Fabrication of microchanneled membranes

The fabrication process of microchanneled membranes is shown in described in Fig. 1a. It starts with slurry preparation using a planetary ball mill (MTI corporation, USA) by mixing 30 g of La$_{0.6}$Sr$_{0.4}$Co$_{0.2}$Fe$_{0.8}$O$_{3-\delta}$ (LSCF, Fuel Cell Materials, Ohio, USA), 1.87 g of polyethersulfone (PESF, Radel-A300), 0.21 of g polyvinylpyrrolidone (PVP, MW=40000) and 11 g of 1-methyl-2-pyrrolidinone (NMP, Sigma-Aldrich, Australia) at a speed of 300 RPM for 48 hours to form a homogeneous slurry. Then, the degassed slurry was cast into a mould and then a piece of stainless steel mesh (SEFAR, Australia) was immersed just below the surface of the slurry. After phase inversion for 30 mins induced by water on the top of the slurry, the mesh was lifted up to remove the skin layer. The formed membrane green body
was rinsed with water several times to remove the solvent residue and dried at 105 °C. The prepared green bodies of the microchanneled membranes were then sintered at 1300 °C in a box furnace (ModuTemp Pty. Ltd., Australia) for 5 hours to densify the separation layer. The membranes with different channel diameters were prepared using meshes with different aperture sizes.

Fig. 1. Schematic representations of fabrication processes for microchanneled membrane (a) and supported membrane (b).

2.2. Fabrication of dense and supported membranes

Dense membranes were fabricated using a hydraulic press, followed by sintering at 1300 °C. The membrane thickness was controlled by the amount of powder used to form the
membrane, and thickness of about 1.4 mm was produced. The supported membrane was made by press forming a porous support and subsequently dip-coating a separation layer. The fabrication process is shown in Fig. 1b. Firstly, Pore former potato starch (Sigma-Aldrich, Australia) and LSCF powder were mixed in the mass ratio of 1:4 in ethanol using a ball mill at a speed of 400 RPM for 24 hours [12]. After drying, the mixed power was hydraulically pressed at a pressure of 300 MPa to form the green body of the porous support. The green body was pre-sintered in a box furnace at 800 °C for 2 hours. The slurry for dip-coating was prepared by mixing 3 g of LSCF powder, 27 g of ethanol and 0.1 g of PVP using a ball mill at the speed of 400 RPM for 24 hours. The pre-sintered porous support was dipped into the slurry and then dried. The process was repeated for several times to achieve the desired dense layer thickness. Finally, the membrane was sintered at 1200 °C in the box furnace for 5 hours to densify the coating layer.

2.3. Membrane characterisation and testing

The membrane microstructure was observed by scanning electron microscopy (SEM, Zeiss Evo 40XVP). Membrane porosities were measured using the Archimedes method. In this study, the microchannel side of the microchanneled membranes and the porous side of the supported membranes were faced to the feed gas in all experiments because this test configuration can achieve higher oxygen permeation rates compared to the arrangement where the dense sides were faced to the feed gas [10, 13]. The gas flow rates were controlled by mass flow controllers (MFC, AALBORG). Oxygen concentration in the sweep gas was measured by online gas chromatography (GC, GC-2014, Shimadzu) with a packed column of molecular sieve 5Å. The oxygen permeation fluxes were calculated using the following equation:
\[ J_{O_2} = \frac{F_S(C_{O_2} - C_{N_2}(21/79))}{A} \]

where \( J_{O_2} \) is the oxygen permeation flux of the membranes, \( F_S \) is the flow rate of the sweep gas, \( C_{O_2} \) is the oxygen concentration in sweep gas, \( C_{N_2} \) is the concentration of nitrogen leaking into the sweep gas; \( A \) is the effective area of the membranes. All testing membranes were 14-15 mm in diameter with an active area of about 0.7 cm².

3. Results and discussion

3.1 Gas diffusion through microchannels

Supported membranes with a thin dense layer on porous supports have been developed to shorten oxygen ion diffusion distances in dense membranes, and this membrane configuration has been employed to achieve high oxygen fluxes [14, 15]. Considering that the supported membranes and the microchanneled membranes have a similar dense layer through which oxygen ions diffuse, it is feasible to compare the gas diffusion through the conventional porous supports of supported membranes and the channels of the microchanneled membranes by measuring the oxygen permeation rates. The SEM images of a microchanneled membrane and a supported membrane are shown in Fig. 2. The dense layer of the porous supported membrane is about 150 μm thick. The distance from channel bottom to membrane dense surface varies from 80 to 200 μm, and the average oxygen ion diffusion distance is therefore about 140 μm. The similar oxygen ion diffusion distance makes it valid to compare gas diffusion within two different porous structures. Porous structure and microchannel structure are shown in Fig. 2c and d. The pores in the supported membrane are around 5-10 μm, and the diameters of the microchannels are about 30 μm. The channel walls are dense. The
The porosity of the microchanneled membrane is 23%, which is lower than that of the supported membrane (36%).

**Fig. 2.** SEM images of the cross sections of the supported membrane (a and c) and the microchanneled membrane (b and d) at low (a and b) and high (c and d) magnifications. The scale bars are 100 μm in a and b, 10 μm in c and 20 μm in d. The microchanneled membrane was templated by a mesh with an aperture size of 70 μm.
Fig. 3. Temperature dependence of the oxygen fluxes of the microchanneled membrane and the supported membrane using air (a) and pure oxygen (b) as a feed gas, respectively. The flow rates of the feeding gases were 20 ml min⁻¹, and the sweep gas flow rate was maintained at 100 ml min⁻¹.

As shown in Fig. 3a, the oxygen flux of the microchanneled membrane is higher than that of the supported membrane. As the operating temperature was increased, the difference in the oxygen flux increased. For example, the oxygen fluxes of the microchanneled membrane and the supported membrane are 0.13 and 0.12 ml cm⁻² min⁻¹ at 750 °C, and 1.5 and 0.76 ml cm⁻² min⁻¹ at 1000 °C, respectively. The difference enlargement might be caused by the difference of gas diffusion resistance through the two membranes. That is to say that the build up of concentration of nitrogen within the pores/channels, and its back diffusion out of the membrane will hinder the influx of oxygen in fresh air. Cao et al. reported the presence of gas diffusion resistance/concentration polarization within the porous supports of membranes at high oxygen fluxes [16]. The gas diffusion resistance becomes obvious as the oxygen flux is increased with operating temperature, resulting in the enlarged difference in oxygen flux between the two types of membranes.
To investigate this interpretation, the two membranes were tested using pure oxygen as a feed
gas to eliminate gas diffusion resistance. As the kinetics of oxygen surface exchange increase
with the oxygen partial pressure regardless of membrane structure, it is valid to compare by
feeding oxygen. Without nitrogen dilution, there would be no oxygen concentration gradient
and therefore no gas diffusion resistance. As shown in Fig. 3b, with oxygen as a feed gas, the
supported membrane showed a higher oxygen flux than the microchanneled membrane; for
example, the fluxes were 4.2 and 3.1 ml cm\(^{-2}\) min\(^{-1}\) for the supported membrane and the
microchanneled membrane at 1000 °C, respectively. The higher oxygen flux of the supported
membrane is attributed to its higher surface area within the porous support compared to that
of the microchannel walls. Overall, compared with conventional supported membranes,
microchanneled membranes provide straight channels as gas diffusion pathways instead of
the tortuous pore channels within the porous support, greatly facilitating gas diffusion and
therefore leading to high oxygen fluxes when the feed gas is air [17].

To further investigate the gas diffusion resistance, the activation energy of the oxygen
permeation through the supported membranes and microchanneled membranes were
measured. Compared with conventional dense membranes, the supported membranes and
microchanneled membranes involve gas diffusion through pores/channels during the oxygen
permeation. It is known that the activation energy of the gas diffusion (less than 20 kJ mol\(^{-1}\))
is much smaller than those of ionic bulk diffusion (about 115 kJ mol\(^{-1}\)) and surface reaction
(about 191 kJ mol\(^{-1}\)) [18, 19]. Therefore, it is possible to identify the presence of gas
diffusion resistance on the basis of activation energy measurements.
Fig. 4 a. Oxygen fluxes of the microchanneled membrane templated by a mesh with an aperture size of 70 μm, a supported membrane and a dense membrane. The oxygen partial pressure difference on two sides of the membranes was maintained at 0.21/0.004 atm and the air flow rate was 100 ml min\(^{-1}\). b. Arrhenius plots and activation energies calculated according to Fig. 4a.

Fig. 4a shows the oxygen permeation through a microchanneled membrane, a supported membrane and a dense membrane with the membrane thickness of about 1.4 mm with an oxygen partial pressure gradient of 0.21/0.004 atm, and the corresponding Arrhenius curves are plotted in Fig. 4b. As there is no gas diffusion involved in the oxygen permeation through dense membranes, the activation energy of the oxygen permeation through a dense membrane was also tested as a reference. Within the high temperature range (900-1000 °C), the activation energy of the dense membrane is close to that of oxygen ionic diffusion (115 kJ mol\(^{-1}\)) in LSCF membranes, which indicates that the oxygen permeation of the dense membranes is rate-limited by oxygen ion diffusion in the solid phase. However, the activation energy was increased to around 150 kJ mol\(^{-1}\) in the low temperature range (750-900 °C); these values are located between those of the activation energy of oxygen ionic diffusion and
surface reactions (191 kJ mol\(^{-1}\)), suggesting that the oxygen permeation through the dense membranes was jointly rate-limited by oxygen ionic diffusion and surface reactions.

As the activation energy of gas diffusion is less than 20 kJ mol\(^{-1}\), the existence of gas diffusion resistance in the oxygen permeation will decrease the overall activation energy of the oxygen permeation. In the high temperature range, the gas diffusion resistance is more likely to occur because of the large oxygen fluxes. From Fig. 4b, the activation energies for the oxygen permeation of supported and microchanneled membranes are 63 and 80 kJ mol\(^{-1}\) respectively, which are much smaller than the activation energy of ionic bulk diffusion (112 kJ mol\(^{-1}\)). This suggests the existence of gas diffusion resistance in the oxygen permeation through both the membranes. Moreover, compared with the supported membrane, the microchanneled membrane showed a higher activation energy, indicating less gas diffusion resistance and hence improved gas diffusion through straight microchannels. The gas diffusion resistances can also be observed in the low temperature range on the basis of activation energies, although the gas diffusion resistance is not as serious as it is in the high temperature range due to the lower oxygen fluxes. However, the activation energies of oxygen permeation though the dense membrane and microchanneled membrane at low temperature are similar (151 vs 148 kJ mol\(^{-1}\)). Compared with the dense membrane, the microchanneled membrane has shorter ionic diffusion distance, and therefore the oxygen permeation is more controlled by surface reaction, which increases the overall activation energy. At the same time, the existence of gas diffusion resistance decreases the activation energy. The two effects cancel each other out, leading to no net change in activation energies. For the supported membrane, gas diffusion resistance contributed more to the permeation process, resulting in the lower activation energy of 108 kJ mol\(^{-1}\).

3.2 The effect of membrane surface within channels
Apart from facilitating gas diffusion through membranes, the microchannel structure potentially increases the membrane active surface area as the surface of numerous channel walls can conduct oxygen dissociation, as illustrated in Fig. 5. The channel wall contribution decreases with the distance from the channel bottom. To identify the contribution, membranes with different channel surfaces were prepared by using mesh templates with different aperture sizes.
Fig. 6. SEM images of surfaces (a, b and c) and cross sections (d, e and f) of the membranes templated by the meshes with the aperture sizes of 55 μm (a and d), 70 (b and d) and 150 μm (c and f). The scale bars are 100 μm.

The mesh-templating phase inversion process can control the diameter of the channels by using meshes with different aperture sizes, which results in different channel wall surface areas. Narrow channels provide a large wall surface area, per unit area of membrane. Three steel meshes with the aperture sizes of 55 μm, 70 μm and 150 μm were used. As shown in Fig. 6, uniform channels were templated by the mesh apertures, and the channel diameter increases with mesh aperture size. Although the mesh aperture is square, the cross section of the templated channels is circular due to the surface tension of the slurry. For the mesh aperture size of 150 μm, the initial cross section of the templated channels is square. However, it becomes circular as the channels grow away from the mesh. The cross sections of the membranes showed that all the membranes have a similar dense separation layer of about 150 μm in thicknesses, which makes it feasible to compare the channel wall contribution by testing oxygen permeations.

The channel diameters were measured from the SEM images. The channel diameter distributions are shown in the Fig. 7, and it is noted the microchannel sizes tend to be more uniform when they are templated using meshes with smaller aperture sizes. The mean pore diameters were calculated to be 25 μm, 40 μm and 87 μm for the microchanneled membranes templated by the meshes with the aperture sizes of 55 μm, 70 μm and 150 μm, respectively.

Assuming the channels have perfect cylindrical shapes and have the same length, the theoretical surface area of channel wall can be calculated using the following equation:

\[ S = \pi dlN \]
where $N$ is the channel numbers in 1 cm$^2$ of membrane area

$N$ is calculated by dividing membrane pore/void volume with single channel volume:

$$N = \frac{4PL}{\pi ld^2}$$

where $P$ is porosity; $L$ is membrane thickness; $l$ is channel length; $d$ is channel diameter.

Fig. 7. Channel diameter distribution of the membranes templated by the meshes with different aperture sizes.
Fig. 8. Theoretical membrane surface areas for the microchanneled membranes templated by the meshes with the aperture sizes of 55 µm, 70 µm and 150 µm.

Fig. 9. Oxygen fluxes of the membranes templated by the meshes with different aperture sizes 55, 70 and 150 µm. Both sweep gas rate and air flow rate were 270 ml min⁻¹.

Fig. 8 demonstrates that the membrane surface area varies with mesh aperture sizes. The membrane surface area and porosities both decrease with increasing channel size. The
The microchanneled membrane templated by the mesh with an aperture size of 55 µm has a surface area of 58 cm², while it is only 13 cm² for the microchanneled membrane templated by the mesh with an aperture size of 150 µm. The membrane templated by the mesh with an aperture size of 55 µm has the largest channel wall surface area compared with other membranes and should therefore produce the highest oxygen flux; this is confirmed by the experimental results shown in Fig. 9. At 1050 °C, the membrane made by the mesh with an aperture size of 55 µm produced an oxygen flux of 3.4 ml cm⁻² min⁻¹, which is higher than that of the other membranes. For example, the membrane made by a mesh with an aperture size of 150 µm only achieved an oxygen flux of 2.6 ml cm⁻² min⁻¹. These results confirm that the channel wall surface contributes to the oxygen permeation and higher oxygen permeation rates can be achieved by reducing the channel diameter.

4. Conclusions

The microchanneled membranes have produced higher oxygen fluxes during oxygen separation from air compared with conventional dense membranes and supported membranes. The superior performance is attributed to the microchannel structure: straight channels provide fast pathway for oxygen molecular diffusion to the thin separation layer compared with the tortuous pore channels within porous supports; numerous microchannels also offer large active membrane surface for oxygen dissociation, which is confirmed by comparing the performance of membranes with different channel diameters. The advantages of the microchannel structure can potentially be applied in other applications such as solid oxide fuel cells and batteries [20-22].

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