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Development of selective Pd–Ag membranes on porous metal filters



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HIGHLIGHTS

- A method to fill the pores of Hastelloy filters via aspiration of α -Al₂O₃ water suspension has been developed.
- The effect of α -Al₂O₃ particle size on the surface quality has been studied with a statistical method.
- The asymmetrical filling method can reduce the average pore size of the filters from 1.1 μ m up to 200 nm.
- A boehmite-based layer is used to change the surface quality and as interdiffusion barrier.
- A highly H₂-selective Pd–Ag membrane (~40 000) has been obtained on the asymmetrically filled support.

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GRAPHICAL ABSTRACT



ABSTRACT

Metallic supports with sufficient surface quality to achieve highly selective thin Pd–Ag membranes require specific pre-treatments, are not readily available on the market and are generally very expensive. To reduce costs, rough and large media grade Hastelloy X filters have been acquired and pre-treated via polishing and chemical etching. The loss in gas permeance given by the polishing treatment proved fully recovered after chemical

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Keywords: Surface modification Metallic membranes Hydrogen separation Pd membranes Metallic supports etching. A method to fill the large pores of the filters via aspiration of α -Al₂O₃ water-powder suspension has been applied and characterized via imaging of the filled pores, inferential statistics, and capillary flow porometry measurements. The most suitable filler particle size for pore size distribution reduction has been identified as 18 µm, while a 5 µm filler proved optimal for further pore morphology improvement. The wide pore size distribution of the filters has thus been reduced up to 200 nm by filling with α -Al₂O₃ particles of decreasing size, similarly to the ceramic supports used for thin Pd–Ag membranes deposition. A boehmite based interdiffusion barrier has been deposited, achieving further surface roughness reduction. A highly H₂ selective membrane has been obtained via simultaneous Pd–Ag plating on the pre-treated filter.

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Introduction

Pd based membranes are well-known for their unique solutiondiffusion H_2 transport mechanism, which makes them the heart of membrane reactors (MRs) for in-situ selective hydrogen removal [1–3]. MRs have proven themselves as a promising alternative to conventional methane steam reformers by integrating both reaction and separation in a single unit and thus avoiding the purification steps required by a conventional process [4–6]. The introduction of membranes in the reaction environment promotes in fact the continuous removal of H_2 , resulting in equilibrium shift towards the H_2 production reaction owing to the Le Chatelier's principle. In this way, the process operating temperature is reduced, increasing overall energy efficiency and reducing operational costs [7–9].

As core of this intensification technology, Pd-based membranes have been subjected to optimization studies, mostly both in terms of their H₂ perm-selectivity performance and their suitability for integration in the reaction environment [10–12]. In particular, thin Pd films (<10 μ m) have been successfully deposited on tubular ceramic supports [13,14]. The deposition of Pd on an appropriate support allows in fact to reach outstanding H₂ perm-selectivity while ensuring mechanical stability of the thin films [12,15,16]. Asymmetric ceramic tubular supports prove in fact suitable for the electroless deposition of a thin and defect-free Pd-based layer thanks to their low surface roughness, narrow pore size distribution (PSD), and low resistance to gas permeation [17,18]. However, while they ensure adequate surface quality for thin layer deposition, ceramic supports prove to be fragile when introduced in the reactor environment. Particularly difficult is their connection and sealing to the steel parts of most reactors structures, making them prone to breakage and/or leaks and thus rendering the scale-up of membrane modules a difficult task [19]. For these reasons, a variety of steel and steel alloys supports have gained rising interest as possible substitutes. They in fact do not require sealing (which can be substituted by a simple weld) and are way less prone to breakage or crack formation thanks to their high mechanical stability. Metallic supported Pd films, however, are proved to be inclined to atomic migration within the support structure, a phenomenon known as intermetallic diffusion or Pd-support interaction [20,21]. In addition, steel-based supports display in fact large

surface roughness, large superficial potholes, and wide pore size distribution, making the preparation of Pd-based membranes with high H_2 permeation and selectivity a complex task [22,23]. In order to then acquire metallic supports with the desired characteristics, tailor made supplier treatments and extremely low media grades must be requested, clearly increasing their final costs [24]. In our work, we focus on operating suitable and cost-effective supports pre-treatments while starting from a large media grade, rough and unrefined Hastelloy filter. The focus is on acquiring a low-cost product and operating suitable pre-treatments to achieve a sufficient surface quality for deposition of a selective Pd–Ag layer.

To solve the intermetallic diffusion issue, an additional barrier is required between the Pd layer and the metallic support [25]. Several interdiffusion barriers have in fact been extensively investigated in literature, mostly ceramic based (Al₂O₃ [26], ZrO [27,28], CeO [29], SiO₂ [30]) or oxidation based (controlled metal oxides growth [31]). In our previous work [24], we proposed a boehmite-based layer with a dual function as interdiffusion barrier/smoothening layer. In fact, this layer, besides preventing intermetallic diffusion, if combined with a suitable polishing treatment is also able to reduce the surface roughness of the selected metallic supports. However, the deposition of this barrier alone proves insufficient to close the large superficial pore mouths of the filter, resulting in membranes with low perm-selectivity [24,26]. For this reason, this work focuses on the recovery of superficial porosity of polished filters via chemical etching [32], and the subsequential pore flow distribution narrowing via the introduction of a pore filler. In literature, several materials have been in fact introduced as fillers in an effort to improve the surface morphology of metallic supports [33-37]. Similarly, in the recent work of Kim et al. a selective membrane obtained via pore filling method and sol-gel barrier deposition has been tested for ammonia decomposition applications [38].

In our work, we propose asymmetrically layered α -Al₂O₃ inside the filter's superficial pore mouths. The sequential filling is carried out with α -Al2O3 of decreasing particle size via vacuum assisted aspiration. The surface roughness of the support is then lowered by the deposition of the boehmite-based barrier, covering the pores and preventing interdiffusion issues. The filling effect was studied by feeding imaging parameters to a Two-way ANOVA design, introducing an applicable method for statistical evaluation of membrane

preparation parameters. The outcomes in terms of support gas permeance, pore flow distribution, surface roughness and morphology are thoroughly investigated for each pretreatment step and, finally, the electroless deposition of a Pd–Ag layer is performed. The resulting membranes are then characterized in terms of ideal H_2/N_2 perm-selectivity.

Experimental

Preparation of porous Hastelloy X supports

Commercial unrefined porous Hastelloy X filters with an outer diameter of 1.2 cm, average surface roughness (Ra) of 6.1 μ m, and 0.5 µm nominal media grade (MG) were acquired by Hebei Golden Flame Wire Mesh Co, China. The supports are cut in samples of 10 cm length and welded to dense stainless steel (AISI316L) tubes, in order to achieve a one close end configuration. To preliminarily reduce the surface roughness of the filters, the sample supports were polished in an industrial surface finishing machine via wet-polishing mechanism (ERBA EVT-170). The chosen polishing time amounts to 6 h, as it allows for a suitable trade-off between surface roughness reduction and gas permeation preservation, which was determined in our previous work [24]. The polished supports are then etched by perpendicular immersion in aqua regia for 30 s and thoroughly rinsed with deionized water to remove all mordant residuals. The supports are then oxidized for 1 h at 750 °C in a furnace in static air atmosphere. Before further treatments, the supports are further rinsed both in ethanol and in deionized water in an ultrasonic bath, to remove all impurities resulting from the preparation pre-treatments and handling.

Filler introduction

The tubular supports are submerged in a powder-water suspension, which is pulled through the superficial pores via vacuum assisted dip-coating. The immersion time is set to 60 s per cycle. Between each cycle the support is gently rinsed with distilled water, and no calcination is required. The selected filler powder is α -Al₂O₃, which is evaluated in three different particle sizes (AA-1 Sumitomo 1 μ m, AA-5 Sumitomo 5 μ m, AA-20 Sumitomo 18 μ m). The powders are 10 wt% suspended in water with a magnetic stirrer. The suspension is improved by addition of HNO₃ (67%vol) dropwise. First, the supports undergo several immersion cycles in order to assess both the aspiration effect and the filler size effect on support's PSD and surface morphology. In a second study, supports are filled asymmetrically with α - Al₂O₃ of decreasing particle size.

Interdiffusion layer deposition

A smoothening interdiffusion barrier is deposited to finalize the improvement of support's surface roughness and ease Pd—Ag deposition. Solutions with boehmite loading 0,9 wt% are prepared in distilled water, incorporating a water-based solution of organic additives, namely 3.5 wt% polyvinyl alcohol (PVA) (MW 130000) and 1 wt% polyethylene glycol (PEG) (MW 400). The deposited layers are dried under rotation in a climate chamber at 40 °C and 60% relative humidity for 1 h. The layers are then sintered for 1 h at 550 °C in a static air furnace. These parameters yield to a mesoporous γ -Al₂O₃ layer about 540 nm thick [24].

Pd—Ag deposition

A layer of Pd–Ag alloy is deposited onto treated supports via electroless plating technique, reported in a previous work by Tanaka et al. [13,14]. To improve membrane selectivity, a consecutive plating procedure is carried out to achieve thicker Pd–Ag layers. After each plating step, the membrane is annealed at 550 °C in 10 vol% H₂ - 90 vol% Ar atmosphere for 4 h. At temperatures below 300 °C, only Ar was used to avoid fragilization of the Pd–Ag layer.

Characterizations

Capillary flow porometry

To evaluate pore size distribution (PSD) variations for each support pre-treatment, the capillary flow porometry technique (CFP, or gas-liquid displacement method) is employed. This technique relies on imposing a trans-sample pressure at which a suitable liquid is displaced from the pores of the examined sample. The displacement is detected by registering the permeating flow of a non-reactive gas through the media. In CFP tests, the sample is filled with a wetting liquid, assuming the filling of its entire accessible porosity. Pressure is applied to one side, while the other is kept at atmospheric pressure. This trans-sample pressure difference forces the wetting liquid out of the pores resulting in a permeating flow. Increasing the trans-sample pressure will promote pore clearance, increasing the permeating flow until the sample is fully cleared from the wetting liquid. Young-Laplace equation is then used to correlate the capillary pressure in the media with its pore diameter. If the capillary is assumed of cylindrical shape, Washburn equation can be applied, which is a typical assumption for indirect method CFP measurements [39,40]:

$$d = -\frac{4\gamma\cos\vartheta}{\Delta P}$$

where ΔP is the applied trans-sample pressure, d is the narrowest diameter of the capillary, γ is the surface tension of the chosen wetting liquid, and ϑ is the contact angle between the wetting liquid and the wet surface. The 10 cm porous tubes are measured via CFP technique in a geometry-specific setup (Fig. 1), which consists of.

- A. a tubular permeation box, which is able to withstand pressures up to 60 bara;
- B. An automatic mass flow controller for N₂ (Bronkhorst EL-FLOW Select F-221 M), which is chosen as inert displacement gas;





Fig. 1 - Graphical representation of the CFP setup for tubular membrane samples.

- C. An automatic backpressure regulator at the retentate side (Bronkorst EL-PRESS- P–502C);
- D. An automatic three way valve, which can switch between a low flow automatic flowmeter (Bronkhorst EL-FLOW Prestige FG-111B, range 0.004 ml/min - 0.2 l/ min) and a high flow automatic flowmeter (Bronkhorst EL-FLOW Prestige FG-111B range 0.2 l/min-10 l/min);
- E. An external computer with LabVIEW software.

The setup is fully automated as it follows.

- I. The controlling software allows the user to set a desired pressure ramp and/or step to be imposed within the permeation box;
- II. The correct feed flow is sent to the permeation box in order to increase the pressure according to the ramp set by the user (1);
- III. The permeating flow at each timestamp is registered at the permeate side by the flowmeter with the correct flow range, which can be automatically switched via the three way valve (2).

Laser confocal microscopy and profilometry: statistical approach

The surface morphology of the treated filters was studied using a dual optical-3D laser confocal microscope (VKX-3000, Keyence, Osaka, Japan). Said imaging method has been employed to verify and observe the presence of the selected filling particles into the superficial pore mouths of the supports. The extent of superficial pore filling is evaluated by imaging a superficial pore mouth and registering its highest point on the metal surface and its lowest point on the embedded filler. For each filler size and cycles combination, 10 random pore mouths have been evaluated. The result is a filling extent parameter Δ . Consequentially, a lower Δ indicates a fuller pore mouth.

 Δ has been employed for statistical considerations as dependent variable in a two-way ANalysis Of VAriance (ANOVA), in order to guarantee a more quantitative approach to the filling procedure optimization [41]. The average surface roughness (Ra) and average profile height (Rz) of the treated supports are evaluated with a portable contact profilometer (MarSurf PS 10) on 10 random positions on the tubular support. These parameters are normalized on their values for an untreated support and employed as dependent variables for a two-way ANOVA.

Being an inferential statistics method, ANOVA allows to infer on the whole population of support pores, while observing only a representative sample amount. The factorial design allows for full observation of the outcome variable (Δ , Ra, or Rz) while variating the two selected factors "Filling cycles" and "Filler size" combinations, making it especially suitable for a comprehensive observation of the filling phenomena.

The experimental designs are summed up in Table 1 and the full statistical documentation is attached in the supplementary material to this manuscript.

Gas permeance

The prepared Pd–Ag membranes are tested for N₂ permeance at 20 °C and 1 bar in a permeation box, described in our previous work [24]. If the N₂ permeance is lower than $1 \cdot 10^{-9}$ mol/ m²/s/Pa the membrane is selected for further high temperature short-term investigation. The selected membranes are activated at 400 °C with an air flow of 1 l/min for 2 min. They are then tested at 400, 450 and 500 °C for single-gas H₂ and N₂ permeance with an imposed pressure difference of 1, 2, 3 bar.

Results and discussion

Polishing and etching

The evolution of the filters surface with the chosen predeposition treatments is analyzed with the laser confocal microscopy images in Fig. 2. The untreated 0.5 µm media grade filters are characterized by the presence of high profile peaks (red) and large surface roughness (Fig. 2a), in agreement with the behavior observed in our previous studies [24]. At this stage, the pore mouths of the support are interconnected, showing a superficial diameter larger than 50 µm. After 6 h of polishing (Fig. 2b), the surface of the support is uniformly smoothened, erasing the presence of the large peaks by plastic deformation. However, the smaller superficial openings are mostly erased. A few large, isolated pore mouths remain, with superficial diameters close to 20 µm. After etching the support for 30s in aqua regia (Fig. 2c) the smoothened surface is broken down into smaller channels interconnecting the metallographic structure underneath: the surface roughness reduction achieved by polishing is preserved, while the support's valleys are uncovered. This promotes an increase in the number of superficial openings, leading to improved gas permeance after polishing, while preserving the treatment's smoothening effect. The evolution of N₂ permeance, mean flow pore and contact roughness parameters (Table 2) further confirms the behavior observed via microscopy. Firstly, the polishing treatment promotes both a gas permeance reduction of 76%, and a roughness decrease of 87%. Following the chemical etching, due to the presence of a larger amount of profile valleys, Ra is re-increased solely with an additional

Table 1 $-$ Two-way ANOVA design for each dependent variable Δ , Ra, Rz.					
Laser-confocal microscopy					
Definition	Variable type Name		Variable levels		
Outcome	Dependent	Δ [μm]	-		
Factor 1	Independent (A)	Filling cycles [–]	10, 15, 20		
Factor 2	Independent (B)	Filler size [µm]	1.5, 5, 18		
Average surface roughness (Ra)					
Definition	Variable type	Name	Variable levels		
Outcome	Dependent	Ra [μm]	-		
Factor 1	Independent (A)	Filling cycles [–]	10, 15, 20		
Factor 2	Independent (B)	Filler size [µm]	1.5, 5, 18		
Average profile height (Rz)					
Definition	Variable type	Name	Variable levels		
Outcome	Dependent	Rz [μm]	-		
Factor 1	Independent (A)	Filling cycles [–]	10, 15, 20		
Factor 2	Independent (B)	Filler size [µm]	1.5, 5, 18		



Fig. 2 – Laser confocal microscopy imaging and height distribution of a 0.5 μ m media grade Hastelloy filter (a)untreated (b) 6 h polished, (c) Etched 30s in aqua regia.

Table 2 – Ra, N₂ permeance, and CFP measured mean flow pore of a sample Hastelloy support for each pretreating step, compared with the values for an untreated Hastelloy filter and an α -Al₂O₃ support by Rauschert.

Pre-treatment	Ra	N ₂ permeance	Mean flow pore (CFP)
[-]	[µm]	$[mol/s/m^2/Pa \cdot 10^{-5}]$	[µm]
Ceramic, α -Al ₂ O ₃	0,5	8,0	0,1
Untreated Hastelloy	6,1	5,0	1,8
Wet polishing, 6 h	0,8	1,2	1,9
Chemical etching, 30s	1,1	8,9	1,1

20%, while the gas permeance surpasses the untreated support's original value. This behavior is well in agreement with the work of Xu et al. which carried out similar improving pretreatments on a disk-shaped stainless steel support, observing the same surface variations via Scanning Electron Microscopy [32].

Filler introduction

Taking into account the mean flow pore of a ceramic support (Table 2), which is proved to produce supported thin and ultrathin Pd based membranes without defects [42,43], the size of the filter's through pores after polishing and etching is still too large. For this reason, the introduction of the α -Al₂O₃ filler into the superficial openings of the pre-treated support is expected to prove crucial for the reduction of their size, ensuring gas tightness (and therefore H₂ selectivity) of the completed membrane. Table 3 shows the results of the Two-way ANOVA performed on the outcome variables listed in Table 1.

The results of the two-way ANOVA highlight a statistically significant contribution to Δ of the interaction between the number of aspiration cycles and the filler size. When an interaction is statistically significant, analyzing solely the main effects can be misleading. Therefore, the interaction effect is observed in Fig. 3. In particular, all filler sizes underperform with the lowest number of cycles, meaning that more

Table 3 – ANOVA test results on outcome variables Ra, Rz and Δ . The relevant factor effect is assumed significant if p-value<0.05, with a confidence interval of 95%.							
Laser confocal microscopy, Δ							
Factor	Factor levels	p-value	Significance				
Filling cycles [—]	10/15/20	0,581	_				
Filler size [µm]	1,5/5/18	2,3•10 ⁻⁸	***				
Interaction [-,µm]	-	0,002	**				
Profilometry, Ra							
Factor	Factor levels	p-value	Significance				
Filling cycles [—]	10/15/20	0,001	**				
Filler size [µm]	1,5/5/18	0,123	_				
Interaction [-,µm]	-	0,512	-				
Profilometry, Rz							
Factor	Factor levels	p-value	Significance				
Filling cycles [—]	10/15/20	0,001	***				
Filler size [µm]	1,5/5/18	0,025	*				
Interaction [-,µm]	_	0,782	-				



Fig. 3 – Interaction effect plot of dependent variable Δ against factor "Filling cycles", grouped by "Filler size". The points indicate the average value for each factor levels combination.



Fig. 4 – Main independent effects plots of factor "Filling cycles" and factor "Filler size" on dependent variable Rz. The points indicate the average Rz decrease values for each independent factor level.



Fig. 5 – Sole main effect plot of factor "Filler size" on dependent variable Ra. The points indicate average Ra decrease for each level of the sole factor "Filler size".

than 10 cycles are required for a low Δ ; at 15 cycles, the best performing size is 5 μ m, while at 20 cycles the largest size (18 μ m) and the smallest size (1.5 μ m) also contribute to a Δ reduction. This result suggests that 5 μ m is a filler that most suits the shape of the superficial pore mouths of the support, requiring less cycles to reach a performance plateau with respect to the largest and smaller filler, which will require more cycles. This result is confirmed by the ANOVA performed on the roughness parameter Rz. For Rz a statistically significant contribution of both independent factors is observed, while the interaction effect is lost. This can be explained by the nature of the outcome variable Rz, which averages roughness profile extremes along a measuring length, rather than purposefully imaged pore mouths, yielding to a loss of information with respect to microscopy. Nevertheless, in Fig. 4b it is noticeable how a 5 µm filler promotes the largest reduction in Rz, independently from the number of cycles performed. However, a large number of



Fig. 6 – (a)Mean flow pore diameter precent decrease with increasing filling cycles, for each filler size, measured via CFP. (b) N_2 permeance percentage decrease with increasing filling cycles, for each filler size. (c) Pore flow distribution measured via CFP of a filter filled 20x with 18 μ m α -Al₂O₃, measured via CFP between 10% and 90% of total dry flow.



Fig. 7 – Dual optical-laser confocal imaging of a Hastelloy filter's selected pore mouth filled with 1.5 μ m α -Al₂O₃, 5 μ m α -Al₂O₃, and 18 μ m α -Al₂O₃.

cycles is preferred independently from the chosen filler size (Fig. 4b). For Ra, a further loss of information is introduced within the ANOVA results. This loss is attributed to the averaging nature of Ra, which does not consider any extremes of the roughness profile, but rather averages all the deviations from the profile's mean line. A statistically significant contribution is attributed solely to the filler size, highlighting how 5 μ m is the best performing in terms of Ra reduction independently from the number of applied cycles, which, in turn, proves to be insignificant (Fig. 5).

All three analyses conclude that 5 μ m reaches a performance plateau quicker than the other sizes, making it the most suitable size for all surface morphology parameters improvement, preferably in combination with an amount of aspiration cycles greater than 15. However, microscopically, as the number of cycles grows larger, the 1.5 and 18 μ m size will contribute to the improving of the pore morphology.

The evolution of surface morphology parameters is not the only phenomenon to be considered while applying a filler to the selected support. An optimal filler design should in fact guarantee sufficient support's average pore diameter reduction and pore distribution sharpening, while preserving gas permeance through the porous media.

From Fig. 6c it is clear how an 18 µm filler sharpens the filter's pore flow distribution around an average pore diameter of about 200 nm, promoting a reduction of the mean flow pore by 80% (Fig. 6a). This shows the filler ability to place itself in large pore necks with respect to the smaller sizes, which only contribute to morphology improvement but require more packing to influence the PSD. Similarly, as observed in Fig. 6b, the 18 µm filler is the most influential in the decrease of permeance in the sample filter, meaning it is affecting the pore necks rather than the surface openings. This conclusion is further confirmed by the dual optical-laser confocal imaging in Fig. 7, in which the 18 μ m filler is observed to close the largest defects, while the smaller fillers tend to assume a packed configuration, leaving larger defects open. This renders the 18 µm filler suitable for a first reduction in large support's openings, while fillers of smaller size can be layered to promote a subsequential improvement of the superficial pore morphology. These observations, paired with the results obtained by Chi et al. who observed the same behavior for α-Al₂O₃ fillers on lower media grade stainless steel tubes, allow to speculate that fillers with about half of the size of the superficial pore openings are the most suitable for pore flow distribution modification, while fillers of lower dimensions can be used for surface morphology improvement [35].

For this reason, the 18 μ m filler is selected as base for the asymmetric support. To then improve the morphology of the support's surface, a 5 μ m and 1.5 μ m filler are introduced subsequentially. The microscopy in Fig. 8 clearly highlights the presence of large α -Al₂O₃ filler underneath the smaller particles, promoting uniform leveling of the pore mouths. Fig. 9 shows the cumulative flow distribution through the pores of a sample filter filled with the selected asymmetric design. The mean flow pore is sufficiently reduced around 100 nm, a comparable value to the one of the α -Al₂O₃ supports commonly used in ceramic membranes preparation. At this stage, most of the largest pore necks are reduced in size by the fillers. However, about 10% of the inert gas flow is still

measured through pores larger than 500 nm. This pore distribution tail depends on both the initial filters themselves and any leftover large openings after pre-treatment. The control of this tail, although representing the minority of the distribution, proves in fact crucial to minimize membrane's N_2 permeation and thus selectivity, ensuring reproducibility of performance. For this reason, further optimization on the filling procedure is the subject of planned studies.

Pd-Ag deposition

In Fig. 10a the surface morphology of a membrane obtained via deposition of Pd–Ag when carried out on a 0.5 μ m media grade filter solely treated with the boehmite based smoothening layer is shown. The boehmite based layer allows for uniform metal deposition on the support's surface. However, the sole presence of this layer is not enough to promote full pore closure and obtain a defect-free Pd membrane. In Fig. 10b the same deposition is carried out on a support in which a 5 μ m α -Al₂O₃ filler has been introduced. The leveling of the pore surface promoted by the filling of the large support's mouths allows for the Pd–Ag to fully close the superficial openings, obtaining the desired defect-free dense layer even on the large superficial pore mouths. For this reason, the



Fig. 8 – Dual optical-laser confocal imaging of a Hastelloy filter's selected pore mouth, asymmetrically filled with α -Al₂O₃ of decreasing size.



Fig. 9 – Cumulative percentage flow distribution through the pores of an asymmetrically filled sample filter before and after boehmite layer deposition, measured via CFP.



Fig. 10 – Dual optical-laser confocal imaging and height distribution of a Pd–Ag layer deposited on a large superficial opening of (a) an unfilled Hastelloy filter, (b) a Hastelloy filter filled with α -Al₂O₃. Both samples are coated with a boehmite-based smoothening layer.

Table 4 $-$ H ₂ /N ₂ perm-selectivity comparison of Pd-based membranes deposited via electroless plating onto steel-based supports pre-treated with similar methods.						
Support Material/media grade/pre- treatment	Filler Material/particle size	Interdiffusion barrier Material	H ₂ permeance [mol/m²/Pa/s]	H ₂ /N ₂ [-]	Ref	
Hastelloy X/0.1 μm/pre- treated by Mott Corp.	Al_2O_3+YSZ powder/-	No additional layer	1,0·10 ⁻⁶	200 000	[34]	
Ceramic/–/double skin	_/_	_	4,6·10 ⁻⁶	26 000	[12]	
Inconel 600/-/-	YSZ powder + boehmite -based sol/–	Blow coated YSZ	3,4·10 ⁻⁶	8050	[38]	
PSS/0.2 μm/-	CeO/1-4 µm	No additional layer	1,2·10 ⁻⁶	infinite	[44]	
PSS/0.5 μm/polished and etched in alkaline sol and HCl	_/_	none	5,0·10 ⁻⁷	5000	[45]	
Inconel 600/0.5 μm/—	YSZ/50 nm	Blow coated YSZ	7,4·10 ⁻⁷	335	[46]	
PSS/-/-	YSZ/50 nm	Blow coated YSZ	3,0·10 ⁻⁶	240	[47]	
Hastelloy X/0.5 μm/6 h polish	_/_	Boehmite	2,0·10 ⁻⁷	512	[24]	
Hastelloy X/0.5 μm/6 h polish, 30s aqua regia etch	α-Al ₂ O ₃ /300 nm	boehmite-based layer 1.2 wt%	1,1.10 ⁻⁶	~6300	#MS, This work	
Hastelloy X/0.5 μm/6 h polish, 30s aqua regia etch	α-Al ₂ O ₃ /300 nm	boehmite-based layer 1.2 wt%	1,1.10 ⁻⁶	~830	#MS2 This work	
Hastelloy X/0.5 μm/6 h polish, 30s aqua regia etch	α-Al ₂ O ₃ /300 nm	boehmite-based layer 1.2 wt%	9,2·10 ⁻⁷	~3600	#MS3 This work	
Hastelloy X/0.5 µm/6 h	α -Al ₂ O ₃ /18 μ m +	boehmite-based layer 0.9 wt%	7,0·10 ⁻⁷	~43 200	#MA, This work	
polish, 30s aqua regia etch	$5\ \mu m\ +1.5\ \mu m$					

choice of a suitable filling procedure proves essential to impact final membrane performance and reproducibility. In Table 4 membranes prepared with different filling procedures are compared in terms of perm-selectivity at 400 °C (see Fig. 11).

Pd–Ag membranes prepared with a small size filler present poor selectivity compared to the membrane prepared with the asymmetric design. In particular, symmetrically filled membrane #MS has been selected due to its high H₂/N₂ selectivity, and its performance is compared to #MAS, prepared on a support filled asymmetrically. Both membranes have been characterized in terms of activation energy via linear regression through the Arrhenius plot in Fig. 12, where #MS shows an activation energy of ~6 kJ·mol⁻¹ and #MA of 9,3 kJ mol⁻¹. While #MA is well in agreement with the activation energy range for thin Pd layer membranes, #MS has a lower value, possibly explained by the presence of scattered larger defects. This deduction is confirmed by the n-values of each membrane, amounting to 0,71 and 0,51, respectively. #MA rate determining step for hydrogen transport is given by the Pd layer, while for #MS the value suggests an influence of the metallic support [34].



Fig. 11 – H_2 permeating flux vs H_2 partial pressure at 400 °C for #MA and #MS.



Fig. 12 – Linear regression performed on Arrhenius plot to determine membrane's activation energy as slope (Δ Ea, in J·mol⁻¹) and pre-exponential factor as intercept.



Fig. 13 – Ideal H_2/N_2 selectivity of #MA, measured at 400 °C,450 °C,500 °C with a trans-membrane pressure of 1,2,3 bar. The measurements are carried out right after membrane annealing.

#MA displays an outstanding H_2/N_2 selectivity (Fig. 13), accounting that this membrane is prepared on a filter with 0.5 µm media grade and 50 µm large superficial pore mouths, the largest in literature obtaining membranes with selectivity >10000. This result proves that it is indeed possible to achieve high-performing membranes using unrefined metallic filters. Pd–Ag membranes prepared with the same filler aspirationbarrier deposition technique reported in literature show promising performances on steel-based supports with larger media grades (Table 4). This two-step procedure therefore classifies itself as a promising standard method for composite PSS/ceramic/Pd based membranes on cheaper support options.

Conclusion

Hastelloy porous filters with large media grade (0.5 µm), 2 µm mean flow pore, 50 µm superficial pore mouths and high surface roughness (6 µm Ra) have been successfully modified to improve their superficial characteristics to be used as support of thin Pd-Ag membranes with high H₂ permeation and selectivity. Surface roughness can be preliminarily reduced via wet polishing method, producing a decrease in the porosity and gas permeation which can be reverted by subsequential chemical etching in aqua regia. The wide pore flow distribution of the filters can be reduced for the most part up to 200 nm via introduction of α -Al₂O₃ filler particles. The 18 μ m filler is the most suitable to reduce the pore size distribution applying at least 20 filling cycles, while fillers of lower size (5 µm, 1.5 µm) prove more suitable for further surface morphology improvement. By combining all the studied pretreatments, a highly selective membrane (~40000) was obtained on a support filled with α -Al₂O₃ particles with decreasing particle size. However, further reproducibility studies are needed to ensure complete elimination of leftover unfilled pore mouths on the support filters.

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.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ijhydene.2023.03.306.

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