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1	Direct contact membrane distillation of refining waste
2	stream from precious metal recovery: chemistry of silica and
3	chromium (III) in membrane scaling
4	
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### 20 Abstract

Precious metals such as platinum group metals (PGMs) with distinct catalytic 21 22 activity are widely used as active components in various industrial catalysts. It is, therefore, highly desirable to recover these valuable components from end-of-life 23 24 products. We explored the treatment of refining wastewater from precious metals recovery using direct contact membrane distillation (DCMD). The role of various 25 initial pH of refining wastewater on DCMD performance was assessed. Results 26 suggest that hydrochloride acid (HCl) and high-quality water can be reclaimed from 27 28 the real refining wastewater by adjusting initial pH. Furthermore, DCMD water flux decline was mainly caused by silica and chromium (III) scaling, which was dependent 29 on initial pH of refining wastewater. Silica scaling was responsible for the decrease of 30 31 DCMD performance when the initial pH of refining wastewater shifted from original 0.03 to 5 and 7. Silica oligomers in the concentrated feed with various initial pH were 32 identified using mass spectra. Whereas chromium (III) scaling was discovered, 33 34 resulting in the used polytetrafluoroethylene (PTFE) membrane surface in green when the initial pH of refining wastewater was 3. Dichlorotetraaquochromiun, 35 [Cr(H<sub>2</sub>O)<sub>4</sub>Cl<sub>2</sub>]Cl·2H<sub>2</sub>O was identified by X-ray photoelectron spectroscopy and 36 ultraviolet and visible absorbance spectra as the main species contributing to the green 37 colour of the scaled PTFE membrane surface. Our results suggest that DCMD can be 38 used as a promising and feasible solution for resource recovery from acidic refining 39 40 waste stream.

- 41 Keywords: precious metal recovery; refining wastewater; membrane distillation;
- 42 silica and chromium (III) scaling

### 44 **1. Introduction**

Precious metals such as platinum group metals (PGMs) are widely used as the 45 active components of various industrial catalysts due to their distinct catalytic activity, 46 chemical inertness, corrosion resistance and thermoelectric stability, which were 47 considered as "Vitamin of modern industry" [1-5]. The loading of precious metals in 48 catalysts ranged from 0.02 to 100% [1]. The most significant applications of precious 49 metals are electronics and catalystic industry, consuming over 90% of precious metals 50 [2]. It was reported that about 65% of palladium (Pd, 182.65 tons), 45% of platinum 51 52 (Pt, 98 tons) and 84% of rhodium (Rh, 25.6 tons) were used in catalytic converters [1, 2]. 53

However, the global reserve of PGMs is only 66,000 tons and their reserve in 54 55 Earth's crust is extremely low. For instance, In China, natural PGMs resources are extremely limited with total reserve of only about 350 tons. Only less than three tons 56 of PGMs were mined in China annually, but demands for Pt and Pd were over 141 57 58 tons. Therefore, there is urgent demand to recycle these precious metals from end-of-life products to realize the sustainable development of precious metals [6, 7]. 59 By 2016, about 30% of PGMs were recovered from spent catalysts, namely 34 tons of 60 Pt, 61 tons of Pd and 7.2 tons of Rh. Precious metals in the spent catalysts are often 61 leached in hydrochloric acid medium with oxidizing agents like HNO<sub>3</sub>, Cl<sub>2</sub>, NaClO, 62 NaClO<sub>3</sub>, H<sub>2</sub>O<sub>2</sub>, etc. through hydrometallurgical process [1, 8-10]. Besides precious 63 metals, spent catalysts also contain many heavy metals [11]. Therefore, wastewater 64 simultaneously generated from precious metals recovery process often contained high 65

66 concentrations of acids and various kinds of heavy metal ions, which has been 67 regarded as huge challenge [12]. From the perspective of resource recovery and 68 environmental protection, waste stream from precious recovery is crucial to be treated 69 before discharging into environment.

70 It has been reported that wastewater containing precious metals can be treated by membrane filtration [13], electrochemical approach [14] and biosorption [15, 16]. For 71 example, forward osmosis (FO), an osmotically driven membrane technology, was 72 widely used in treatment of various industry wastewater [17-23] and was also capable 73 74 of Pd accumulation from printed circuit board (PCB) plant wastewater using an electroless nickel plating solution as draw solution [13]. In addition, microbial fuel 75 cell (MFC) as one kind of bioelectrochemical systems was also employed for 76 77 recovery of precious metals such as silver. High silver removal rate  $(83 \pm 0.7\%)$  and recovery  $(67.8 \pm 1\%)$  efficiencies were achieved from MFC fed with silver laden 78 artificial wastewater (MFC-Ag) after 72 h operation. COD removal rate of MFC-Ag 79 80 was up to  $82.7 \pm 1.5\%$  [14]. Furthermore, biosorption with advantages of low cost and high effectiveness at low concentrations and environmentally friendly nature has been 81 82 widely developed for the recovery of metals ions from aqueous and waste solutions. A range of bioadsorbents, such as Escherichia coli [24, 25], Shewanella oneidensis 83 MR-1[26], Enterococcus faecalis [27], Phomopsis sp. XP-8 [28], Enterobacter 84 cloacae SgZ-5T [29], Galdieria sulphurariaetc [30] etc., have been reported towards 85 86 biorecover Pd (II), Pt and Au from synthetic solutions. However, it is noteworthy that the majority of the studies above were focused on precious metals recovery or 87

accumulation from synthetic solutions. Till now, treatment of real wastewater derived 88 from precious metals recovery has been not reported. Therefore, it is necessary to 89 90 investigate the treatment of real wastewater generated from process of precious metals recovery. However, in economically undeveloped and remote area in China, the actual 91 92 solution for removing heavy metal ions in the real wastewater derived from precious recovery was precipitation in alkali environment. This traditional physicochemical 93 method not only increases cost of wastewater treatment but also produces large 94 amount of physical and chemical sludge. Therefore, the dewatered sludge always 95 96 contains various kinds of heavy metal ions and is a potential big threat to environment. In addition, China is a serious water shortage country, which is one of the 13 countries 97 that lack of water all over the world [31]. From the perspective of water reclamation 98 99 and environmental protection, if water reuse and heavy metal ions removal to an acceptable level is realized in application including industrial reuse, municipal green, 100 and agricultural irrigation, which would have a great significant impact on cost 101 reduction and decrease the negative influence on environment. An alternative 102 technology that promises to achieve this objective is membrane distillation (MD). 103

Membrane distillation (MD) emerged as an advanced membrane technology was used to recover valuable salts [32, 33] as well as treatment of high salinity solution [34-36]. Compared with other membrane technologies such as reverse osmosis, MD possesses several advantages for brine minimization, such as low operating pressure, high water recovery, potential for 100% rejection of non-volatile solutes and small footprint [37]. However, it is well known that MD is a very energy intensive process.

To reduce or replace extra energy input, industrial waste heat and solar energy were 110 used in MD process [38-40]. However, in the field of treatment of industrial waste 111 112 stream via MD process, there was no report on the application of MD technology in treatment of the real wastewater generated from process of precious metals recovery. 113 The aim of this work was to assess the treatment of the real refining wastewater with 114 strong acidity from precious metals recovery through DCMD process. Effects of 115 various initial pH of wastewater on DCMD performance and membrane scaling were 116 systematically assessed. The changes of membrane surface color and morphology 117 118 caused by silica and chromium (III) scaling under various pH values were recorded and interpreted, respectively. Various characterization techniques were employed to 119 elucidate the underlying mechanisms of membrane scaling during the DCMD 120 121 filtration.

### 122 **2. Material and methods**

#### 123 **2.1. MD membrane and wastewater**

Commercial flat sheet MD membrane (PTFE-PVDF/PET) with mean pore size
of 0.45 μm was purchased from Shanghai Minglie New Material Co., Ltd. The detail
properties of commercial PTFE membrane were listed in Table S1. The structures of
brine side facing feed solution and permeate side were shown in Fig. S1.

The real refining wastewater produced from the recovery of precious metals in spent catalysts was kindly provided by Sino-Platinum metals resources (Yimen) Co., Ltd. The key characteristic of the refining wastewater used is shown in Table 1. The refining wastewater is a clear yellow solution with a conductivity of 199.2 mS/cm and initial pH of 0.03. Sodium and potassium ions are the major cations with
concentrations of 11.36 g/L and 10.06 g/L, respectively. The major anion is chloride
ion with concentration of 48.99 g/L.

1	2	5
4		J

Table 1 Key characteristic of refining wastewater used in this study

Analyzed items	Unit	Concentration
Conductivity	mS/cm	$199.2 \pm 1.4$
рН	-	$0.03\pm0.02$
COD	mg/L	$3620\pm200$
Phosphate	mg/L	$400 \pm 80$
Chloride	mg/L	$48993.8 \pm 1200.7$
Nitrate	mg/L	$1107.32 \pm 50.82$
Silicon	mg/L	$11.92 \pm 3.50$
Sodium	mg/L	$11364 \pm 200$
Potassium	mg/L	$10058\pm300$
Zinc	mg/L	$536.71 \pm 8.75$
Aluminum	mg/L	$196.22\pm3.05$
Calcium	mg/L	$64.07 \pm 1.56$
Iron	mg/L	$46.50 \pm 0.37$
Magnesium	mg/L	$36.80\pm1.28$
Copper	mg/L	$14.46\pm0.62$
Chromium	mg/L	$9.48 \pm 1.13$
Nickel	mg/L	$6.27\pm0.52$

Manganese	mg/L	$4.82\pm0.21$
Silver	mg/L	$0.99\pm0.12$
Barium	mg/L	$0.79\pm0.15$
Platinum	mg/L	$0.57\pm0.08$

137 **2.2. DCMD setup and experiments** 

A DCMD module made of transparent Perspex with a channel depth of 4 mm 138 and effective area of 30 cm<sup>2</sup> (i.e. length 10 cm, width 3 cm) was used for experiments. 139 Two same polyester spacers with diamond mesh were used to support the membrane 140 and to promote the flow turbulence in both sides. The detail information of the 141 142 module and spacer can be available in the previous report [34]. The schematic diagram and photo of the lab-scaled DCMD system used in this work is presented in 143 144 Fig. S2 and Fig. S3 as reported previously [41-43]. The real refining wastewater with 145 various initial pH was used as feed. Before experiments, feed and ultrapure water were put in the jacketed glass bottles with total volume of 1.5 liters, respectively. The 146 effective volume of feed and ultrapure water was 1.2 and 1.5 liters for all experiments, 147 respectively. Temperature of permeate side and brine side was maintained at 20±1°C 148 and 60±1°C by circulating water cooler and thermostat circulating water bath, 149 respectively. The flow rate for both sides was controlled at 300 mL/min 150 151 (corresponding to the cross flow velocity of 8.3 cm/s) for all experiments by two same peristaltic pumps (Langer, BT600). 152

To investigate the influence of various initial pH of wastewater on MD 153 performance, the initial pH of wastewater as the feed was adjusted from original 0.03 154 to 3, 5 and 7 using sodium hydroxide (23 wt%). Before experiments, the wastewater 155 with various initial pH was first filtered by filter paper with mean pore size of 40 µm 156 to remove visible species and then filtered again using 0.45 µm filter membrane to 157 remove the small sized particles to maximally reduce membrane fouling during 158 DCMD process. The permeate flux  $(J, \text{ kg/m}^2\text{h})$  was calculated by measuring the 159 weight changes of permeate ( $\Delta m$ , kg)) with a precision balance (OHAUS, AR4202CN) 160 divided by time intervals ( $\Delta t$ , h)) and membrane area (A, m<sup>2</sup>), which was described in 161 equation (1). Additionally, the conductivity of the accumulated permeate was online 162 monitored by a conductivity meter (SUNTEX, EC-4110RS). Both the data of balance 163 164 and conductivity meter was recorded by a computer equipped with a data logging system. 165

$$J = \frac{\Delta m}{\Delta t \cdot A} \tag{1}$$

167

166

## 168 **2.3. Analytical methods**

Key element concentration of refining wastewater was determined with inductively coupled plasma-atomic emission spectroscopy (ICP-AES). The concentrations of chloride, phosphate and nitrate in refining wastewater were measured by ion chromatography (IC, LC20AT, Shimadzu, Japan). Field-emission scanning electron microscopy (FESEM, Hitachi S-4800) equipped with energy dispersive X-ray fluorescence spectrometer was used for analyzing the morphologies

of membranes and elementary composition of scalants. X-ray photoelectron 175 spectroscopy (XPS) (Escalab 250Xi) was employed to analyze element chemical 176 177 bonding. Fourier Transform Infrared Spectrometer (Nicoletln 10MX/Nicolet6700) was employed to analysis the ultraviolet and visible absorbance of the blank and 178 scaled membrane surface. Electrospray ionization mass spectrometry (ESI-MS, 179 Agilent 11000) was used to identify the silica oligomers formed in the concentrated 180 solution with various initial pH. ESI negative ionization mode was used and the direct 181 injection flow of the sample was 0.15 mL/min. 182

183

### 184 **3. Results and discussion**

### **3.1 DCMD performance**

186 Fig. 1(A) shows the permeate flux patterns under various initial pH of refining wastewater. The curves of permeate flux displayed the similar trend, gradually 187 decreased with the increasing of recovery. The initial flux for the original pH of 0.03 188 and 7 of refining wastewater were the same of 13 kg/m<sup>2</sup>h, which is a little higher than 189 that for pH of 3 and 5 also with same flux of 11.5 kg/m<sup>2</sup>h. Then, the corresponding 190 permeate flux for the initial pH of refining wastewater adjusted from original pH of 191 0.03 to 3, 5 and 7 declined to 8.4, 4.7, 4.6 and 5.9 kg/m<sup>2</sup>h, respectively, when 192 recovery reached to 60%. One reason for the decrease of fluxes is that feed solution 193 concentrated and the partial vapor pressure of water in the feed solution decreased as 194 195 increasing of recovery. Another reason is possibly that membrane scaling occurred under various initial pH of wastewater, which resulted in membrane scaling in varying 196

degree, thus caused various initial flux and flux at recovery of 60%. The changes of 197 permeate conductivity were depicted in Fig. 1(B). The permeate conductivity for 198 original pH of 0.03 and 3 increased exponentially, while the curves of permeate 199 conductivity for pH of 5 and 7 were much smoother. It was found amazingly that the 200 permeate conductivity increased from initial 7.6 µS/cm to 18.3 mS/cm when recovery 201 reached to 67% for the wastewater with original pH of 0.03. However, for the 202 wastewater with initial pH of 3, 5 and 7, the corresponding final permeate 203 conductivity was 338, 57.5 and 31.5  $\mu$ S/cm, when the recovery went up to 66%, 60% 204 205 and 60%, respectively. These results indicated that high-quality permeate can be reclaimed by controlling initial pH of feed solution. However, it is noteworthy that the 206 permeate conductivity for the pH of 3 spiked after recovery of 60%, indicating that 207 208 the PTFE membrane was possibly wet.

The pH values of the solution in permeate and feed tanks were also tested before 209 and after experiments. As shown in Fig. 2, compared with ultrapure water with 210 original pH of 5.46 in the permeate tank, the pH of the solution in the permeate tank 211 after experiment first decreased from 5.46 to 1.31 and 3.41, then increased to 6.45 and 212 7.2 as the initial pH of wastewater as feed shifted from original 0.03 to 3, 5 and 7, 213 respectively. In order to reveal the reason for the pH changes, water quality of the 214 solution in the permeate tank after experiment was characterized when original 215 refining wastewater was used as feed. The majority of anion is chloride ions (Cl<sup>-</sup>) with 216 concentration of 545 mg/L and only small amount of cations such as  $Ca^{2+}$  (4.78 mg/L), 217 K<sup>+</sup> (0.38 mg/L), Na<sup>+</sup> (0.36 mg/L), Mg<sup>2+</sup> (0.56 mg/L) coexists in the permeate (Table 218

219 2). It can be concluded from the data in Table 2 that the permeate for wastewater with original pH of 0.03 mainly consisted of hydrochloric acid (HCl). The reason for 220 collection of HCl from waste stream with pH of 0.03 via DCMD process is possibly 221 that under feed temperature of 60°C, the partial pressure of HCl over aqueous solution 222 increased, resulting in Henry's law constant of HCl increase and solubility of HCl 223 decline. Therefore, during DCMD filtration of wastewater with original pH of 0.03, 224 HCl fast volatilized from feed solution with temperature of 60°C and passed through 225 PTFE membrane into permeate, which resulted in permeate conductivity fast 226 increased (Fig 1 B). These results above indicated that refining wastewater can be 227 well treated as well as recovery of hydrochloric acid and high-quality water via 228 DCMD process through adjusting initial pH of wastewater. 229





Fig. 1. DCMD experiment performance, permeate flux (A) and conductivity (B) as a function of recovery for refining wastewater (Temperature of feed side and permeate side was maintained at  $60\pm1^{\circ}$ C and  $20\pm1^{\circ}$ C, respectively; both the volumetric flow rates of feed and permeate were controlled at 300 mL/min; the sodium hydroxide with concentration of 23 wt% was used to adjust the feed pH).





**Fig. 2** pH changes of concentrated feed solution and permeate after experiments under

Table 2 Water quality of permeate from DCMD process using original refining

242

wastewater as feed.

Analytes	Concentration (mg/L)
Chloride	$545 \pm 10$
Potassium	$0.38\pm0.08$
Sodium	$0.36\pm0.05$
Calcium	$4.78\pm0.22$
Magnesium	$0.56\pm0.13$

243

## 244 **3.2 Membrane surface color and contact angle**

<sup>239</sup> various initial pH of wastewater as feed.

The colour of membrane surface facing feed has not changed significantly before 245 and after experiments and was still white when the initial pH of the feed was adjusted 246 247 from original 0.03 to 5 and 7 (Fig. 3). However, significant derivation of membrane surface colour was observed under the condition of wastewater as feed with initial pH 248 of 3. As shown in Fig. 3, the membrane surface facing the feed and the permeate side 249 of membrane facing cold side were both green, which was unexpected. Furthermore, 250 the permeate side of membrane was more green than that of the brine side of 251 membrane, meant that the greenish scalant possibly penetrated into membrane surface. 252 This phenomenon was interpreted and the greenish scalant was identified in the 253 following sections. 254

The contact angles of the brine side of PTFE membrane after experiments were 255 256 shown in Fig. 4. For the refining wastewater with original pH of 0.03, the contact angle of PTFE membrane surface decreased from 112° to 72°, which provided clear 257 evidence that the PTFE membrane surface after usage was partially wet. In contrary, 258 the contact angle increased a little to 116° when the initial pH of the feed solution 259 shifted to 7. However, for the refining wastewater with initial pH of 3 and 5, it was 260 found that the contact angles of PTFE membrane surface were not obtained due to 261 water droplet spreading, resulted from membrane wetting. These results indicated that 262 the brine side of PTFE membrane surface after treatment of the real refining 263 wastewater with initial pH of 3 and 5 were thoroughly wet. The phenomenon of 264 contact angle changes was possibly related to the change of membrane surface 265 morphologies and scalants deposited on membrane surfaces, which was observed as 266

267 below.





Fig. 3. The photos of brine and permeate sides of PTFE membrane surfaces after experiment under various initial pH of feed solution





Fig. 4. Contact angle of brine side of PTFE membrane after treatment of refiningwastewater with various initial pH.

### 276 **3.3 Membrane surface morphology**

To elucidate the microscopic changes of membrane surface, SEM-EDX has been 277 conducted on each scaled brine side of PTFE membrane surface. As shown in Fig. 278 5(A), the morphologies of various membrane surfaces were distinct. There is 279 considerable amount of small white particles on the surface fibers of PTFE membrane 280 after treatment of refining wastewater with original pH of 0.03. The corresponding 281 result of EDX (Table 2) showed that the main metallic element was K with ratio of 282 1.66 wt%, followed by Al (1.32 wt%) and Na (0.94 wt%). Apart from inherent C 283 284 element of the PTFE membrane, the main non-metallic element was Si with ratio of 42.41 wt%, followed by O (37.53 wt%) and Cl (1.70 wt%). It was reported that Si 285

species in solution with pH less than 9 was mainly in the form of H<sub>4</sub>SiO<sub>4</sub> in previous 286 work [44-46]. The polymerization of silica takes place via condensation mechanism in 287 288 the presence of hydroxide ions. It started with a dimerization reaction that is typically considered to involve a non-ionic silicic acid molecule and an ionic silicic acid 289 molecule (Eqs. (2) - (3)). The dimerization also occurs in an acidic solution but in a 290 much slower rate via the reaction scheme in Eq. (4). Additionally, it was also reported 291 that in the presence of metal ions such as  $Na^+$ ,  $Mg^{2+}$ ,  $Al^{3+}$ ,  $Ca^{2+}$  and  $Fe^{3+}$ , dimerization 292 was facilitated by neutralizing the surface charge of silica and allow aggregation of 293 294 particles [47-50]. Based on the analysis from the perspective of elementary composition of scalant and silica polymerization, it can be concluded that the white 295 small scalants deposited on the fibers were possible mixture of mono-silicic acid and 296 297 silica oligomers.

298 
$$H_4SiO_4 + OH^- = H_3SiO_4^- + H_2O$$
 (2)

299 
$$H_3SiO_4^- + H_3SiO_4^- = (OH)_3Si - O - Si(OH)_3 + OH^-$$
 (3)

$$300 H_4SiO_4 + H_4SiO_4 = (OH)_3Si - O - Si(OH)_3 + H_2O (4)$$

However, compared with surface morphology of the PTFE membrane after treatment of refining wastewater with original pH of 0.03, the surface of the PTFE membrane after treatment of refining wastewater with initial pH of 3 was seriously fouled by the evidence that a lot of small spherical particles not only deposited on the surface fibers (Fig. 5(B)), but also aggregated in the gap between the fibers. The corresponding EDX result (Table 2) showed that beside the C element the major non-metallic elements were P, O and Cl with ratio of 26.26 wt%, 24.37 wt% and 5.48 wt%, respectively. The highest content metallic element was Cr with ratio of 17.57
wt%, followed by K (7.66 wt%), Na (5.33 wt%) and Al (5.08 wt%). Based on this
result, it can be explained reasonably that the greenish scaled membrane surface
showed in Fig. 4 was caused by trivalent chromium (Cr (III)) as the Cr (III) ions in
water solution was green [51, 52].

Surface fibers of membrane after treating refining wastewater with initial pH of 5 313 were relatively clear, large massive scalants were mainly found at the crossing of 314 surface fibers beside few small white particles deposited on the surface fibers (Fig. 315 316 5(c)). According to the corresponding EDX results (Table 2), the major non-metallic elements were O (29.69 wt%), Cl (16.77 wt%), P (12.40 wt%), and Si (0.88 wt%). Na 317 was the main metallic element with a ratio of 11.47 wt% followed by Cr (8.76 wt%), 318 319 K (7.69 wt%) and Al (3.88 wt%). Based on the scalants morphologies and EDX elementary composition analysis, the main scalants was possible a mixture of 320 amorphous silica and silica oligomers. 321

322 The surface morphology of PTFE membrane after treatment of neutral (pH=7) refining wastewater looked similar to that of membrane after treatment of original 323 refining wastewater (Fig. 5(D)). EDX result (Table 2) showed that the major 324 non-metallic element was Cl with ratio of 28.49 wt%, followed by O (16.23 wt%), P 325 (5.61 wt%), Si (1.62 wt%) and S (1.06 wt%). The metal elements in the order of 326 content from high to low were Na (13.48 wt%), K (9.23 wt)%, Cr (5.1 wt%), Fe (2.55 327 wt%) and Al (0.62 wt%). Compared with Si ratio on the membrane surface after 328 treating original refining wastewater, the Si ratio on the membrane surface after 329

treatment of neutral (pH=7) refining wastewater was much lower. The membrane
surface morphology was in line with Si ratio on the scaled membrane surface. Only
few small white particles were found on surface fibers, resulting in the maximum
contact angle the PTFE membrane after experiment.

Overall, based on the shape of scalants and their EDX elementary composition 334 analysis, it can be concluded that silica and chromium scaling were occurred on 335 membrane surface. The membrane scaling caused by silica and chromium (III) of the 336 PTFE membrane after treatment of refining wastewater with various initial pH values 337 was in varying degree by evidence that the surface morphologies were distinct, which 338 was possibly the main reason for the changes of DCMD fluxes and contact angles. 339 However, it was noteworthy that silica and chromium scaling occurred and their 340 341 impacts on system performance were dependent on initial pH of refining wastewater.



343

Fig. 5. SEM images of PTFE membrane surface at various initial pH of refining
wastewater; (A), (B), (C) and (D) is SEM of scaled membrane treatment of refining
wastewater with initial pH of original 0.03, 3, 5 and 7, respectively.

348 Table 3 The corresponding element ratio of the scaled PTFE membrane surfaces349 characterized by SEM in Fig. 5

рН	Element ratio (wt%)										
	С	0	Cl	Si	S	Р	K	Na	Al	Cr	Fe
0.03	14.44	37.53	1.70	42.41	/	/	1.66	0.94	1.32	/	/
3	8.15	24.37	5.48	/	/	26.26	7.66	5.33	5.08	17.57	/
5	8.46	29.69	16.77	0.88	/	12.4	7.69	11.47	3.88	8.76	/
7	16	16.2	28.5	1.62	1.06	5.61	9.23	13.5	0.62	5.1	2.55

#### 350 **3.4 Mass spectra identified silica oligomers**

Mass spectra provided important information of silica oligomers during silica 351 related scaling. The mass spectra of concentrated refining wastewater with various 352 initial pH (Fig. 6) were compared and the mass/charge ration (m/z) and possible 353 354 structure of silica oligomers were tabulated (Table 4). For the concentrated refining wastewater with original pH of 0.03 and 3, the major species in the concentrated 355 solution were mono-silica acid with m/z of 113 and dimmer-linear silica acid with m/z356 of 136.9, 155 and 172.9 (Fig. 6 and Table 4). This result indicated that oligomerisation 357 358 of monomer silica proceeded via formation of dimmer-linear silicates [53-55]. Therefore, the saclants deposited on the PTFE membrane surface was possibly 359 mixture of mono- and dimmer-linear silica acid. The deposition of silica on PTFE 360 361 membrane is likely to occur via a homogeneous nucleation process, with silica aggregates formed in refining wastewater prior to PTFE membrane surface. The 362 explanation was also consistent with the silica scaling morphology, as distinct silica 363 crystals, especially for original pH of 0.03 (Fig. 5). However, this situation was not 364 for the concentrated refining wastewater with initial pH of 5 and 7. Besides the 365 mono-silica acid with m/z of 113 and dimmer-linear silica acid with m/z of 155 and 366 172.9, few trimer-cyclic or -linear and tetramer-cyclic or -linear were also found in 367 the concentrated solution. This phenomenon could be interpreted that mono-silica 368 acid firstly deposited on the PTFE membrane surface and then initiate silica 369 polymerization on the membrane surface, resulting in an amorphous silica scaling 370 morphology, especially for pH of 5. 371





Fig. 6. Mass spectra of the concentrated feed solution with various initial pH after DCMD treatment (A, original pH of 0.03; B, pH=3; C, pH=5;



m/z	Possible molecular formula	Intensity ( $\times 10^2$ )	Possible structure	Ref
113.1	H4SiO4·(OH) <sup>-</sup>	30 (original pH) 19 (pH=3) 18 (pH=5) 37 (pH=7)	•	[53]
136.9	Si <sub>2</sub> O <sub>4</sub> (OH) <sup>-</sup>	13 (pH=3) 15 (pH=5)	●●-2H₂O	[55]
155.1	Si <sub>2</sub> O <sub>3(</sub> OH) <sub>3</sub> -	26 (pH=3)	●●-1H <sub>2</sub> O	[54]
172.9	Si <sub>2</sub> O <sub>2</sub> (OH) <sub>5</sub> -	90 (original pH) 78 (pH=3) 67 (pH=7)	••	[53-55]
252.8	Si <sub>3</sub> O <sub>5</sub> (OK)(OH) <sub>2</sub> -	6 (pH=7)		[54]
270.7	Si <sub>3</sub> O <sub>5</sub> (OK)(OH) <sub>4</sub> -	23 (pH=5)	▲ -1H₂O	[54]
292.8	Si <sub>4</sub> O <sub>6</sub> (OH)5 <sup>-</sup>	7 (pH=5) 7 (pH=7)	-1H <sub>2</sub> O -2H <sub>2</sub> O -2H <sub>2</sub> O	[53, 54]

**Table 4** Possible structures of silica oligomers determined by electrospray ionization mass spectrometry under various initial pH of wastewater.

### 377 3.5 Chromium (III) scalant identification

Based on the detailed analysis of membrane morphologies and elementary composition from section 3.3, the reason for the scaled PTFE membrane in green under initial pH of 3 was caused by chromium (III). To identify the green species on the scaled membrane surface, the brine side of the scaled PTFE membrane was examined by XPS and UV-Vis spectrophotometer to elucidate the interaction between chromium (III) and membrane and to identify the specific green species.

### 384 3.5.1 Chromium (III)-membrane interaction

385 Based on the results in sections 3.3 chromium (III) scaling was found on the surface of membrane after treating real refining wastewater with initial pH of 3. To 386 elucidate chemical origin of chromium (III)-membrane interaction, high solution XPS 387 388 was conducted to examine the chemical bond of Cr 2p of the scaled membrane surfaces. The Cr 2p spectra ranging from 568 to 592 eV is shown in Fig. 7. The peaks 389 of Cr 2p binding energy were found at 578.5 eV and 577.4 eV. Based on previous 390 report [56, 57], the peak of Cr 2p binding energy at 578.5 eV and 577.4 eV was the 391 characteristic bond of Cr-Cl and Cr-O, respectively. Further, the intensity of Cr-O was 392 much higher than that of Cr-Cl, indicating that the number of Cr-O was more than that 393 of Cr-Cl in the compounds. According to the analysis above and quality of the real 394 refining wastewater (Table 1), it can be considered that the greenish compounds 395 deposited on the brine side of membrane was chloroaquochromium (III) complexes 396 that was possibly derived from the formation of hydrated isomers of chromic chloride 397 hexahydrate (CrCl<sub>3</sub>·6H<sub>2</sub>O) [51, 52]. According to the color of the isomers, the scaled 398

membrane surface in green was caused by either dichloropentraaquochromiun,
[Cr(H<sub>2</sub>O)<sub>5</sub>Cl]Cl<sub>2</sub>·H<sub>2</sub>O or dichlorotetraaquochromiun, [Cr(H<sub>2</sub>O)<sub>4</sub>Cl<sub>2</sub>]Cl·2H<sub>2</sub>O.

401



402

Fig. 7. High solution Cr 2p scan by X-ray photoelectron spectroscopy of scaled
membrane after treatment of refining wastewater with various initial pH of 3.

405

406 3.5.2 UV-vis absorbance spectra identified dichloroaquochromium (III)
407 complexes

To identify the primary species responsible for the scaled membrane surface in green under condition of feed with initial pH of 3, the test of UV-Vis absorbance of the original and scaled membrane was carried out. In the visible region (380-780 nm), the spectrum of UV-Vis absorbance of the blank membrane was linear (Fig. 8).

However, two characteristic peaks at 450 nm and 635 nm with the maximum 412 absorbance value of 0.088 and 0.096, respectively, appeared in the UV-Vis 413 414 absorbance spectrum of the brine side of the scaled membrane in green. Since previous studies [51, 52, 58] demonstrated that the UV-vis absorption spectrum of 415  $[CrCl_2(H_2O)_4]^+$  showed the characteristic peaks at 450 and 635 nm in the region 416 between 200 and 800 nm, the dichlorotetraaquochromiun, [Cr(H<sub>2</sub>O)<sub>4</sub>Cl<sub>2</sub>]Cl·2H<sub>2</sub>O, 417 was the main species responsible for the appeared greenish color. It was reported that 418 the structure of dichlorotetraaquochromiun was octahedral [59]. However, it should be 419 pointed out that the dichlorotetraaquochromiun has two isomeric structures, namely 420 cis and trans (Fig. 9). Therefore, green specie was possibly mixture of 421 dichlorotetraaquochromium with two isomeric structures. 422



423

424 Fig. 8. UV-Vis absorbance spectra of the blank membrane and scaled membrane after

treatment of refining wastewater with initial pH of 3.



Fig. 9. The molecular space structure of isomeric dichlorotetraaquochromiun (a, *cis*; b, *trans*)

430

### 431 **4.** Conclusion

In this work, DCMD process was employed to treat real refining wastewater with strong acidity from recovery of precious metals in spent catalysts. System performance was assessed under condition of various initial pH of the real refining wastewater. The major findings and conclusions drawn from this work were summarized as follows.

- High initial water flux ranged from 11.5 to 13 kg/m<sup>2</sup>h were obtained under
  various initial pH of refining wastewater as feed through DCMD process.
- 439 Relative pure hydrochloride acid was reclaimed from original refining
  440 wastewater through DCMD process.

30

High-quality water was available from refining wastewater by adjusting pH
from acidic to neutral.

• Silica scaling was the main reason for the decrease of system performance over time when the pH of refining wastewater was adjusted from original 0.03 to 5 and 7.

Chromium (III) scaling was detected, which resulted in the greenish surfaces
of PTFE membrane when the initial pH of refining wastewater was 3. The identified
dichlorotetraaquochromiun, [Cr(H<sub>2</sub>O)<sub>4</sub>Cl<sub>2</sub>]Cl·2H<sub>2</sub>O, was the main species responsible
for the appeared greenish colour.

450

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