

1 Article

2 **Dispersion-Free Extraction of In(III) from HCl Solutions Using a** 3 **Supported Liquid Membrane Containing HA324H+Cl⁻ Ionic Liquid as** 4 **Carrier**

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9

10 **Abstract:** The transport of indium(III), from HCl solutions, across a supported liquid membrane in
11 flat-sheet configuration was investigated, being the carrier the ionic liquid HA324H+Cl⁻ (derived
12 from the tertiary amine Hostarex A324 and hydrochloric acid). Different variables affecting the
13 metal transport: hydrodynamic conditions in the source and receiving phases, metal and HCl
14 concentrations in the source phase, and carrier concentration in the membrane phase, were
15 investigated. Also the transport of indium(III) using carriers of various nature: ionic liquids,
16 alcohol, ketone, phosphine oxide and phosphoric ester, was compared. The metal transport was
17 modelled describing the transport mechanism as: diffusion across the source diffusion layer, a fast
18 interfacial chemical reaction, and diffusion of the InCl₄--carrier complex through the membrane
19 support. Diffusional parameters for the transport of indium(III), from the experimental data and
20 the model, were estimated.

21 **Keywords:** indium(III); ionic liquid; supported, liquid membrane; Hostarex A324; transport.

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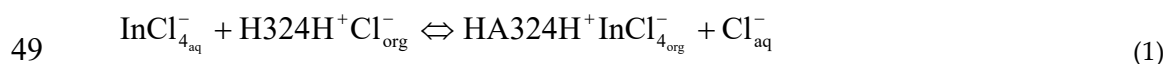
23 1. Introduction

24 Indium is classified by the European Commission as a critical raw material both for its
25 economics and high supply risk, which an expected increased in its demand over the next years. The
26 above being a consequence of its special properties and wide industrial uses, i.e., liquid-crystal
27 displays (LCDs), electronics, catalysts, etc. Thus, the recovery of this element from the above sources
28 is an important target, and several technologies had been proposed to resolve such a situation [1-13].
29 Besides the above, indium is considered a hazardous element due to its carcinogenic character [14],
30 being its removal from residual aqueous solutions i.e., resulted from a number of the above
31 processes, is of the foremost importance, and here it is when liquid membranes must be considered
32 as a technology suitable for the recovery of metals or other solutes present, in the few mg/L
33 concentration order, in the wastewaters. Included in these liquid membranes operations, supported
34 liquid membranes in their various configurations are useful for this task. Before scaling up the
35 technology in the form of hollow fiber modules, the investigation of a given system, using supported
36 liquid membrane in flat-sheet configuration, is necessary in order to obtain information about the
37 mass transfer process involved in the membrane operation. One important part, in real terms the
38 key-of-the-operation, is the carrier used to transport the metals, and included in them, ionic liquids
39 are a group of chemicals, which properties make of them to be considered as *green solvents*, and
40 among other uses [15-18] suitable for the removal of metals from aqueous solutions [19-28].

41 As a part of the investigations carried out from our group related from the removal of
42 indium(III) using liquid membrane technologies [5], this work presented results about the removal
43 of indium(III), from HCl solutions, using a flat-sheet supported liquid membrane (FSSLM)
44 containing the ionic liquid HA324H+Cl⁻, as carrier. Different hydrodynamic and chemical variables
45 affecting the indium(III) transport process were investigated.

46 2. Results

47 It was shown [10], that the extraction of indium(III), from hydrochloric acid media, by the ionic
48 liquid HA324H⁺Cl⁻ dissolved in Solvesso 100, occurred via the next equilibrium:



50 where the subscripts org and aq referred to the organic and aqueous phases, respectively. Thus, the
51 extraction process occurred when the equilibrium was shifted to the right, and metal strip occurred
52 shifting the equilibrium to the left. It was also determined that the value of the equilibrium constant
53 value, for the above equilibrium, was 10.96 in 1 M HCl medium.

54 2.1. Influence of the stirring speed in the source phase on indium(III) permeation

55 In every separation technology, i.e., liquid membranes, ion exchange or adsorption, and
56 working in batch conditions, it is of a necessity to find, in experimentally form, the best
57 hydrodynamic conditions to ensure maximum solute transport, adsorption, etc., thus, in the present
58 system, the influence of the stirring speed applied to the source phase on metal permeation was
59 investigated by the use of source phases containing 0.01 g/L In(III) in 2 M HCl, and a receiving
60 solution of 0.1 M HCl. The membrane phase was a 0.23 M ionic liquid in Solvesso 100 solution
61 supported in a Durapore GVHP4700 solid support. Table 1 shown the results derived from these
62 experiments.

63 **Table 1.** Influence of the stirring speed on permeability of indium(III)

Stirring speed (min ⁻¹)	Px10 ³ (cm/s)
375	1.5
500	1.9
750	2.9
1000	1.5
1500	1.0

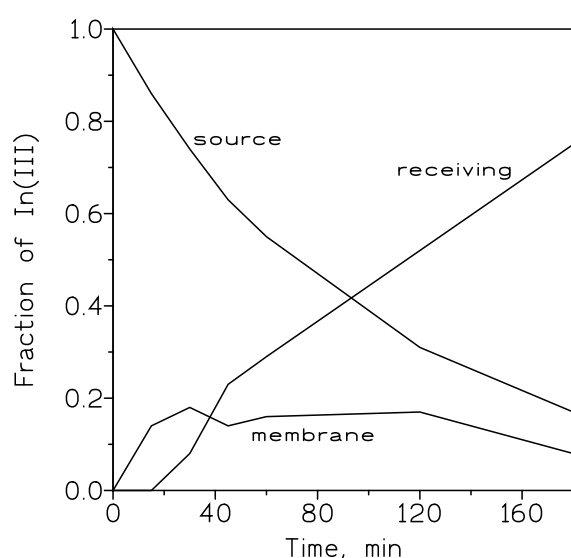
64 It is shown that the indium(III) permeation increased from 375 to 750 min⁻¹ and then decreased,
65 this is attributable to a continuous decrease of the source phase boundary layer with the increase of
66 the stirring speed on this phase, and that a limiting permeability value is obtained at 750 min⁻¹, at
67 this point, the system reached a minimum in the thickness of the aqueous source layer and
68 indium(III) transport maximized, thus:

69

$$70 \quad P_{\text{lim}} = \frac{D_{\text{aq}}}{d_{\text{aq}}} \quad (2)$$

71 being D_{aq} the metal diffusion coefficient in the aqueous source phase (averaging a value of 10^{-5} cm/s),
 72 and d_{aq} the thickness of the aqueous source layer, being P_{lim} 2.9×10^{-3} , the value of d_{aq} is estimated to
 73 be of 3.4×10^{-3} cm. Thus, this value represented the minimum thickness of the stagnant source phase
 74 layer, considering, the experimental conditions used in this work. It is also shown in Table 4 that at
 75 stirring speeds above 750 min^{-1} , the metal permeation decreased, being this caused to the turbulence
 76 caused by the stirring speed and the progressive instability of the liquid membrane.

77 In the limiting conditions and after 3 hours, the percentage of metal recovered in the receiving
 78 solution is of 75% with respect to the initial In(III) concentration of the source solution, and 90% with
 79 respect to the solute transported from the source to the membrane phases. Figure 1 shown the metal
 80 distribution between the three phases, there is a lag time, of near 30 min between this time and time
 81 zero, before indium(III) begins to appear in the receiving solution; from 120 min, indium(III) is
 82 transported against its concentration gradient driven by the difference in acidity between the source
 83 and receiving phases.



84

85 **Figure 1.** Fraction of indium(III) between the source, membrane and receiving phases. Data from the
 86 experiment at 750 min^{-1} .

87 2.2. Influence of the stirring speed in the receiving phase on In(III) permeation

88 In the present work, the receiving phase was of 0.1 M HCl, since previous data [10] indicated
 89 that this solution was a good strippant for this system. Experiments carried out with this solution, as
 90 the receiving phase, the same source and membrane phases than above, and 750 min^{-1} as stirring
 91 speed of the source phase, indicated that in the $500\text{-}750 \text{ min}^{-1}$ range of stirring speeds applied on the
 92 receiving phase, there was no appreciable difference in the transport of the metal. Thus, stirring
 93 speeds of 750 and 500 min^{-1} in the source and receiving phase were selected for further experiments.

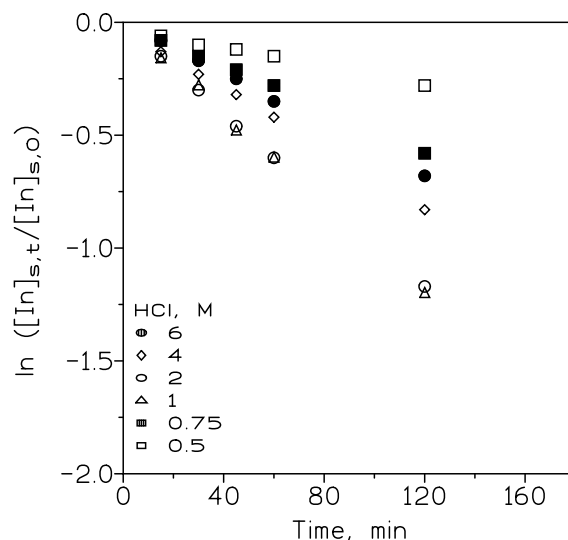
94 2.3. Influence of HCl concentration in the source phase on indium(III) transport

95 The variation of the HCl concentration in the source phase on In(III) permeation was
 96 investigated using the same receiving and membrane phases than above, being the source phase of
 97 0.01 g/L In(III) in different HCl media. The results of the experiments were showed in Figure 2, by
 98 plotting $\ln([In]_{s,t}/[In]_{s,0})$ versus time, and being $[In]_{s,t}$ and $[In]_{s,0}$ the metal concentrations in the source
 99 phase at an elapsed and initial times, respectively.

100 It was showed, that the indium permeation increased with the increase of the HCl concentration in
 101 the source solution up to 1-2 M, and then further decreased at higher HCl concentration in this

102 phase, this was probably attributable to the increase of the aqueous ionic strength or/and at an
 103 increment of the ions population in this phase (population growth effect), which often decreased
 104 metals permeation.

105



106 **Figure 2.** Influence of HCl concentration in the source phase on In(III) transport. Stirring speeds of
 107 the source and receiving phases: 750 and 500 min^{-1} , respectively.

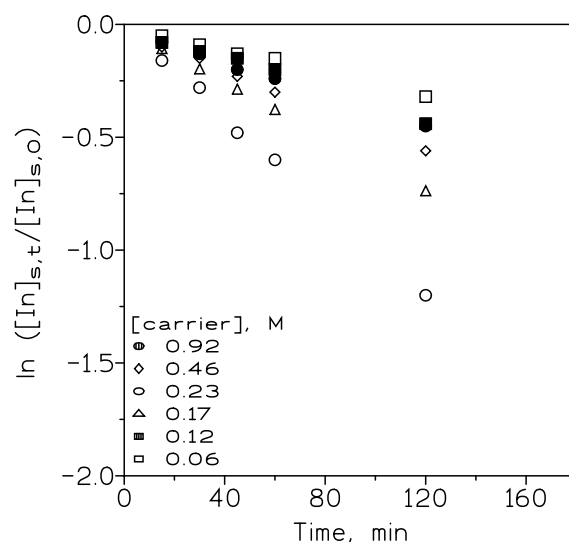
108 Moreover, the percentage of indium(III) recovered in the receiving phase was, after three hours,
 109 around 90% when using 1 or 2 M HCl solutions.

110 2.4. Influence of the carrier concentration on In(III) permeation

111 It was mentioned above that the carrier solution is the key for the liquid membrane operation,
 112 since a liquid membrane with no carrier phase resulted in no metal transport. Figure 3 showed the
 113 results derived when a 0.01 g/L in 1 M HCl source phase was transported across a membrane,
 114 containing organic phases of different concentrations of the ionic liquid in Solvesso 100, and using a
 115 0.1 M HCl solution as receiving phase. As it can be observed, a maximum in metal permeability
 116 ($3.0 \times 10^{-3} \text{ cm/s}$) was reached when a carrier concentration of 0.23 M was used in the membrane phase.
 117 The above was attributable that, at low carrier concentrations diffusion of the indium(III)-carrier
 118 complex across the liquid membrane governed the step, but at the maximum indium(III)
 119 permeation, diffusion of indium(III) across the source phase boundary layer is the rate-determining
 120 process. Accordingly to eq. (2), d_{aq} was estimated as $3.3 \times 10^{-3} \text{ cm}$, value which matched perfectly with
 121 the previously obtained at 2 M HCl concentration in the source phase.

122 When higher carrier concentrations, i.e. 0.46 and 0.92 M, were used in the membrane phase, the
 123 increase of the organic phase viscosity decreased metal transport, being the above attributable to an
 124 increase of the membrane resistance.

125



126 **Figure 3.** Influence of the carrier concentration on In(III) transport. Source phase: 0.01 g/L In(III) in 1
 127 M HCl. Membrane phase: different concentrations of the ionic liquid, in Solvesso 100, imbibed in
 128 GVHP4700 supports. Receiving phase: 0.1 M HCl. Stirring speeds as in Figure 2.

129 *2.5. Influence of the initial indium(III) concentration in the source phase on the metal permeation and flux*

130 These results were derived from membrane phases containing 0.23 M ionic liquid, in Solvesso
 131 100, supported in a GVHP4700 support, and receiving phases of 0.1 M HCl. The source phases were
 132 of various In(III) concentrations (0.01-0.1 g/L) in 1 M HCl. Table 2 resumed these results. Firstly, it
 133 can be seen that the permeation coefficient, P , decreased as the initial metal concentration in the
 134 source phase increased, and secondly, the initial metal flux increased with the increase of the initial
 135 indium(III) concentration in this phase.

136 **Table 2.** Variation of the indium(III) permeation coefficients (P) and fluxes (J) with the variation of
 137 the initial metal concentration in the source phase

Initial In(III) concentration (g/L)	$P \times 10^3$ (cm/s)	$J \times 10^{10}$ (mol/cm ² ·s)
0.01	3.0	2.6
0.05	0.82	3.6
0.1	0.72	6.3

138 Stirring speed in the source and receiving phases: 750 and 500 min⁻¹, respectively

139 The decrease in metal permeation, with the increase of the indium(III) concentration, may be
 140 due to that the membranes pores become saturated with the increasingly metal concentrations,
 141 whereas in the case of the flux, these results were logical since the flux varies with metal
 142 concentrations accordingly to:

143

144
$$J = P[\text{In}]_{s,0} \quad (3)$$

145 and thus, there was an increase of the flux value with the increase of the metal concentration in the
146 source phase. Accordingly with the flux values showed in Table 2, it was seemed that in the range of
147 indium(III) concentrations used in this work, the permeation process was controlled by diffusion of
148 metal species.

149 With respect to the metal recovered in the receiving phase, these percentages varied between
150 90-99% (after 3 h), with respect to the metal transported from the source to the membrane phase, for
151 the three initial indium(III) concentrations used in this work.

152 2.6. Influence of the support characteristics on indium(III) transport

153 Results derived with GVHP4700 support, were compared when HVHP4700 support was used
154 as solid support in this system, with all the experimental variables as in Figure 3, except that in this
155 case, the membrane phase was of a 0.23 M carrier, in Solvesso 100, solution supported in the HPVP
156 solid support.

157 The initial fluxes, see eq. (3), obtained from these experiments were of 2.6×10^{-10} mol/cm² s and
158 1.7×10^{-10} mol/cm² s for the GVHP and HVHP supports, respectively; thus, being all the
159 characteristics similar to both supports but the pore size (2.2×10^{-3} cm for GVHP versus 4.5×10^{-3} cm
160 for HVHP), this last characteristic dominated the metal transport, and apparently, the less the pore
161 size the best the transport, and also with respect to the indium(III) recovery in the receiving phase:
162 (91% for GVHP and 59% for HVHP supports, after 3 hours and, as above, with respect to the metal
163 transported from the source to the membrane phases).

164 2.7. Indium(III) transport using different carriers

165 The transport of indium(III) was compared when different carriers were used. In this case, the
166 source phase was of 0.01 g/L in 1 M HCl and the receiving phase of 0.1 M HCl. The membrane
167 phases were of 0.17 M carrier in Solvesso 100 solutions supported in the GVHP4700 solid support,
168 and the stirring speeds applied to the source and receiving phase were, as usual, 750 and 500 min⁻¹,
169 respectively.

170 These transport results, together with the indium(III) recovery in the receiving phase, were
171 summarized in Table 3.

172 Overall, and under the present experimental conditions, with the ionic liquid HA324H⁺Cl⁻ the
173 best transport results were obtained, followed by the two ionic liquids Cyphis IL101 and Cyphos
174 IL102, derived from a phosphonium salt. The worst transport result was obtained when
175 2-ethyl-hexanol was the carrier for indium transport. For each type of carrier used here, in the case of
176 carriers derived from quaternary ammonium, the apparent order found was: tertiary amine
177 (Hostarex A324) > Akiquat 336 > primary amine (Primene JMT); in the case of phosphorous
178 derivatives: phosphonium salts > phosphine oxides > phosphoric ester, and for organics containing
179 C-O bonds: ketone > alcohol.

180 With respect to the indium recovered in the receiving phase, the sequence was not as clear,
181 however near quantitative In(III) recovery was achieved with Cyphos IL101, Cyphos IL102 and
182 Cyanex 923, being 0.1 M HCl solutions not good receiving medium in the case of the primary amine
183 (acting as the corresponding ammonium derivative), the ketone and the alcohol.

184

185

186

187 **Table 3.** Results of In(III) transport, and metal recoveries in the receiving phase, using different
 188 carriers.

Carrier	P (cm/s)	^b In recovery (%)
Cyphos IL102	1.2x10 ⁻³	99
Cyphos IL101	1.4x10 ⁻³	99
Cyanex 923	8.5x 10 ⁻⁴	99
Tributylphosphate	4.1x10 ⁻⁴	6
^a Primene JMT	3.1x10 ⁻⁴	nil
Aliquat 336	2.9x10 ⁻⁴	50
^a Hostarex A324	1.8x10 ⁻³	83
2-ethyl-1-hexanol	4.2x10 ⁻⁷	nil
Isopenty-methylketone	3.9x10 ⁻⁴	nil

189 ^aActing as the corresponding quaternary ammonium salt. ^bAfter 3 hours, and with respect to the
 190 metal transported from the source to the membrane phases

191 With respect to the indium recovered in the receiving phase, the sequence was not as clear,
 192 however near quantitative In(III) recovery was achieved with Cyphos IL101, Cyphos IL102 and
 193 Cyanex 923, being 0.1 M HCl solutions not good receiving medium in the case of the primary amine
 194 (acting as the corresponding ammonium derivative), the ketone and the alcohol.

195 2.8. Estimation of the diffusional parameters

196 Accordingly to eq.(1), the equilibrium constant of the reaction can be written as:

197

$$198 \quad K = \frac{[\text{HA324H}^+\text{InCl}_4^-]_{\text{org}} [\text{Cl}^-]_{\text{aq}}}{[\text{InCl}_4^-]_{\text{aq}} [\text{HA324H}^+\text{Cl}^-]_{\text{org}}} \quad (4)$$

199 Following the same reasoning published elsewhere [28], an expression for the permeability
 200 coefficient can be written as:

201

$$202 \quad P = \frac{K[\text{HA324H}^+\text{Cl}^-]_{\text{org}} [\text{Cl}^-]_{\text{aq}}^{-1}}{\Delta_{\text{org}} + \Delta_{\text{aq}} \left(K[\text{Cl}^-]_{\text{aq}}^{-1} [\text{HA324H}^+\text{Cl}^-]_{\text{org}} \right)} \quad (5)$$

203

204 where Δ_{aq} and Δ_{org} are the transport resistance related to the diffusion by the source and membrane
 205 phases, respectively. This expression combines the diffusional and equilibrium parameters involved
 206 in the transport of indium(II), from HCl solutions, across a membrane supporting the ionic liquid.
 207 Then:

208

$$209 \quad \frac{1}{P} = \Delta_{\text{aq}} + \Delta_{\text{org}} \frac{1}{K[\text{Cl}^-]_{\text{aq}}^{-1}[\text{HA}324\text{H}^+\text{Cl}^-]_{\text{org}}} = \Delta_{\text{aq}} + \Delta_{\text{org}} \frac{1}{b} \quad (6)$$

210 thus, a plot of $1/P$ versus $1/b$ for various carrier concentrations in the membrane phase and 1 M Cl-
 211 concentration in the source solution, may resulted in a straight line with intercept and slope to
 212 estimate the transport resistance in the source phase ($\Delta_{\text{aq}}= 166$ s/cm) and in the membrane
 213 phase($\Delta_{\text{org}}= 0.84$ s/cm), respectively. The estimate value of the membrane diffusion coefficient:

214

$$215 \quad D_{\text{org}} = \frac{d_{\text{org}}}{\Delta_{\text{org}}} \quad (7)$$

216

217 was 1.5×10^{-2} cm²/s, whereas the transport resistance in the source phase was estimated as 166 s/cm.

218 The diffusion coefficient of the indium(III)-ionic liquid complex in the bulk organic phase can
 219 be estimated by:

220

$$221 \quad D_{\text{b,org}} = D_{\text{org}} \frac{\tau^2}{\varepsilon} \quad (8)$$

222

223 and $D_{\text{b,org}}$ being of 5.6×10^{-2} cm²/s. It can be observed, that D_{org} had a lower value than $D_{\text{b,org}}$, the reason
 224 may be caused to the diffusional resistance due to the membrane thickness located between the
 225 source and receiving phases.

226 Moreover, an apparent diffusion coefficient for indium(III) can be estimated as:

227

$$228 \quad D_{\text{org}}^{\text{a}} = \frac{Jd_{\text{org}}}{[\text{carrier}]_{\text{org}}} \quad (10)$$

229

230 by assumption of a constant carrier concentration of 0.23 M, then the value of this coefficient is
 231 1.3×10^{-8} cm²/s.

232

233 3. Materials and Methods

234 The amine tertiary Hostarex A324 (Hoechst), had as active group: tri-isooctyl amine, was used
 235 as the organic precursor of the ionic liquid, the inorganic moiety was hydrochloric acid, and the ionic
 236 liquid was generated by reaction of both [10]. Other carriers used in this investigation had the
 237 composition showed in Table 5. All were used without further purification. Other chemicals used in
 238 the work were of G.R. quality, except the organic diluent Solvesso 100 (99% aromatics) which was
 239 obtained from Exxon Chem. Iberia, and was also used without further purification. Though many
 240 authors claimed that ionic liquids were used without dilution, the real fact was that in most, if not all
 241 the cases, an organic diluent was needed, and its use facilitated the liquid-liquid extraction
 242 operation because, and among others:

243 i) decreased the high viscosity of the ionic liquid, and, thus, facilitated phase disengagement in the
 244 settler,

245 ii) adequate the range of carrier concentrations to any particular use, avoiding the use of an excess of
 246 carrier in the process, with the benefits of decreasing the economic input of it, and favouring the
 247 metal transport, as one can see, i.e. in this investigation, that an excess of carrier decreased
 248 indium(III) permeability.

249 **Table 5. Chemical composition and source for the carriers**

Carrier	Chemical composition	Source
Cyphos IL102	Trihexyl tetradecylphosphonium bromide	Cytec Ind.
Cyphos IL101	Trihexyl tetradecylphosphonium chloride	Cytec Ind.
TBP	Tri-n-butyl phosphate	Fluka
Primene JMT	mixture of t-alkylprimary amines	Rohm and Haas
Cyanex 923	mixture of tri-n-alkyl phosphine oxides	Cytec Ind.
Aliquat 336	Tri-octyl methylammonium chloride	Fluka
MIPK	Isopentyl-methylketone	Fluka
2-Ethyl-1-hexanol	Alcohol	Fluka

250 Transport experiments were carried out in a two-compartment cell, one compartment each for
 251 the (200 cm³) source and (200 cm³) receiving phases, with the membrane support separating both
 252 aqueous phases. The source and receiving solutions were mechanically stirred by means of four
 253 blades impellers (11.5 cm diameter). Indium(III) permeability (P) was estimated by the use of the
 254 common relationship:

255

$$256 \ln \frac{[\text{In}]_{s,t}}{[\text{In}]_{s,0}} = -\frac{A}{V} Pt \quad (11)$$

257 where $[In]_{s,t}$ and $[In]_{s,0}$ were the indium concentrations in the source phase at an elapsed time and
258 time zero, respectively, A the effective membrane area (11.3 cm^2), V the volume of the source phase,
259 and t the elapsed time. Indium was analyzed in the source and receiving phases by AAS.

260 The membrane support used in the investigation was Durapore GVHP4700, with 0.75%
261 (porosity, ϵ), $12.5 \times 10^{-3} \text{ cm}$ (thickness, d_{org}), 1.67 (tortuosity, τ) and $2.2 \times 10^{-5} \text{ cm}$ (pore size). Durapore
262 HVHP4700 support had the same specifications than above but $4.5 \times 10^{-5} \text{ cm}$ (pore size). Both were
263 composed of polyvinylidene difluoride. The liquid membranes were prepared by immersion of the
264 support in a carrier solution, during 24 hours, and drip it during 20 seconds before placed it in the
265 cell.

266 4. Conclusions

267 This investigation demonstrated the ionic liquid $HA324H^+Cl^-$ is efficient for the transport of
268 indium(III) from hydrochloric acid solutions. It was experimentally found that the best conditions
269 for indium(III) transport were: 750 and 500 min^{-1} stirring speeds in the source and receiving phases,
270 respectively, 1-2 M HCl concentration in the source phase and carrier concentration of 0.23 M. In
271 terms of metal transport, this carrier compares well with other carriers of various types, though
272 indium(II) recovery in the receiving solution was not the best in comparison with results derived by
273 the use of other carriers, i.e. phosphonium salts and the phosphine oxide Cyanex 923. The kinetic
274 model for indium(III) permeation showed that the permeation process is controlled by diffusion of
275 the $InCl_4^-$ species across the source phase layer and diffusion of the $In(III)$ -carrier complex across
276 the liquid membrane, being the former dominant when the carrier concentration in the membrane
277 phase is low.

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279 and F.A.L. investigation; F.J.A. writing-original draft; F.J.A. and F.A.L. writing-review & editing.

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284

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