

# Influence of particle size distribution on carbonbased flowable electrode viscosity and desalination efficiency in flow electrode capacitive deionization

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# <sup>1</sup> Influence of particle size distribution on carbon-

# <sup>2</sup> based flowable electrode viscosity and <sup>3</sup> desalination efficiency in flow electrode <sup>4</sup> capacitive deionization.

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# 15 Abstract

16 Flow-electrode capacitive deionization (FCDI) is a potential energy-efficient electrochemical water desalination technology that combines ion-exchange membranes and flowable 17 electrodes (FE). The performance of activated carbon (AC)-based FEs in FCDI systems is 18 influenced by various factors, including particle size. Here, particle size commercial AC was 19 20 reduced by dry ball milling to produce FEs for FCDI. Physical analysis by scanning electron microscopy and particle size analyzer confirmed the particle size reduction after dry ball milling. 21 Then, the effect of particle size distribution on the rheological behavior and desalination 22 23 performances of AC-based FEs was investigated by desalination performance indicators, such 24 as desalination efficiency, average salt adsorption rate and charge efficiency. FE samples were prepared using different mixtures of fine-size range (0.65-0.92 µm; fine AC, FAC) and 25 large-size range (1.5-2.3 µm; AC) particles. The AC to FAC particle ratio influenced the 26 rheological properties of AC-based FEs in FCDI. Bimodal mixtures with 75% of FAC and 25% 27 of AC (0.75:0.25 FAC:AC) particles displayed the highest desalination efficiency compared 28 with pure AC and pure FAC FEs (72% versus 34% and 71%, respectively). Moreover, in our 29 FCDI set-up, 0.75:0.25 FAC:AC bimodal mixtures showed better flowing properties than the 30 pure FAC FEs, characterized by high viscosity. 31

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Keywords: Flow electrode capacitive deionization, activated carbon flow electrode, particle size
 distributions, rheology, stability and desalination.

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# 36 Graphical abstract



37 38

# 39 **1. Introduction**

40 Capacitive deionization (CDI) is used to eliminate charged species from water with various applications, such as brackish and sea water desalination, <sup>1, 2</sup> water softening,<sup>3</sup> and wastewater 41 remediation.<sup>4</sup> CDI is an energy-efficient, cost-effective, and ecofriendly technology.<sup>5</sup> The first 42 43 and most widely-used CDI set-up requires two porous carbon electrodes separated by a 44 spacer through which water flows perpendicularly to the applied electric field.<sup>5</sup> In membrane CDI (MCDI), a key variation of the basic CDI set-up<sup>6</sup>, the feed water flows between an anion 45 46 exchange membrane at the anode and a cation exchange membrane at the cathode.<sup>7</sup> The 47 membrane main role is to reduce co-ion adsorption. This improves the charge efficiency and can increase the charge storage in the electrode porous structure.<sup>8</sup> In the last decade, a new 48 MCDI-based approach was developed to introduce carbon-based flowable electrodes.<sup>9</sup> The 49 basis of this technology, called flow-electrode CDI (FCDI), is similar to that of electrochemical 50 flow capacitors<sup>10</sup>. The process is based on electrical double layers that are formed on 51 52 interfaces between porous activated carbon (AC)-based electrodes and an electrolyte (e.g. NaCl solution).<sup>11</sup> lons are adsorbed and stored in the electrical double layers when a cell 53 voltage in the range of 1–1.2 V is applied to the cell<sup>12</sup>. FCDI has three major advantages 54 55 compared with other CDI systems. First, the continuous desalination of the feed water flowing through a single cell does not require an in situ discharge step for regenerating the ion-56 saturated AC-based electrodes, which is done in a separate process. This allows the easy 57 scaling-up of FCDI<sup>13</sup>. Second, the continuous inflow of uncharged carbon particles into the 58 59 charging cell increases the capacitance, which is needed for desalination, above that of staticelectrode CDI systems.<sup>14</sup> This offers more flexibility because the desalination rate and kinetics 60 61 can be easily controlled by adjusting the saline solution, flow electrode retention time, and the applied current charge. Third, in AC-based flowable electrodes, pores are more exposed and 62 easily accessible for ion adsorption. 63

The choice of electrode material is a crucial factor in CDI system performance. The ideal CDI electrode material should meet the following criteria: 1) large specific surface area, high ion

adsorption sites; 2) high conductivity and ion mobility; 3) hydrophilicity to ensure full contact 66 67 between the electrode porous structure and water; 4) electrochemical stability at different pH and voltages to ensure the system reliability and sustainability; 5) easily shapeable in function 68 of the design requirements and economically feasible (i.e. availability, cost, and recyclability) 69 for industrial development. Carbon-based materials are mostly used in this field because they 70 71 meet these requirements. Moreover, to improve the properties of the carbon-based materials used as CDI electrodes, many methods have been tested and many carbon forms have been 72 exploited, for instance AC, AC fibers, carbon aerogels, AC cloths,<sup>15,16</sup> carbon nanotubes,<sup>17</sup> 73 graphene,<sup>18</sup> carbon nanofibers,<sup>19</sup> carbon sheet<sup>20</sup>. 74

75 Porous AC, a cheap, easily available and reusable material, is widely used to prepare flowable electrodes for FCDI.<sup>21</sup> The electrochemical performance of AC-based flowable electrodes is 76 influenced by several parameters, including mass loading, particle size and morphology, 77 specific surface area, surface charge, viscosity, and hydrophilicity.<sup>22</sup> However, they display low 78 wettability, pore inaccessibility for ions, and low conductivity. Therefore, to improve its features 79 and overcome some of its limitations, AC can undergo physical or chemical treatments. For 80 instance, pores can be expanded using physical treatments, such as thermal modification 81 using inert gases (e.g. N<sub>2</sub>, Ar, CO<sub>2</sub>).<sup>23</sup> Surface area can be improved by exposure to KOH or 82 NaOH followed by thermal treatment.<sup>24</sup> Much research has been done on hierarchical porous 83 carbons with interconnected micro-mesopores for increasing ion adsorption.<sup>25</sup> Recent studies 84 85 have investigated the particle size effect on the desalination performance of AC-based electrodes in CDI.<sup>26</sup> Cohen et al. combined dilute slurry electrodes and dense fluidized bed 86 electrodes for better electric conductivity and lower resistance.<sup>27</sup> Zhang et al. found that AC-87 based electrodes with different particle sizes provide different CDI performances. Particularly, 88 a mixture of large and small particles gave the best desalination performance in CDI.<sup>26</sup> 89 However, the effect of particle size distribution on the viscosity and desalination performance 90 of carbon slurry electrodes in FCDI has not been investigated yet. In a previous study, we 91 92 found that reducing AC particle size in flowable electrodes can double the desalination rate in FCDI.<sup>28</sup> This was attributed to the enhanced connectivity between particles that increases 93 94 charge transport and storage, thus improving conductivity through the AC percolating network. Moreover, in AC-based electrodes with finer particles, obtained by grinding, pores were more 95 exposed, leading to faster ionic diffusion.<sup>28</sup> However, viscosity was increased upon particle 96 size reduction. As the flowable electrode must flow along a narrow channel in the FCDI set-97 up, its rheological features are a crucial factor for limiting pressure drops and ensuring the 98 absence of clogging. Indeed, the higher viscosity of the flowable electrode with fine AC 99 100 particles, compared with normal AC particles, led to pumping and flowing difficulties inside the 101 FCDI system. Therefore, in the present study, we wanted to decrease the viscosity of AC-102 based flowable electrodes, while maintaining the good desalination performance obtained with 103 fine AC particles. To this aim, we mixed two AC particle size ranges to produce flowable 104 electrodes for FCDI. We prepared bimodal slurries with different ratios of fine AC (FAC) to AC particles with the same carbon loading (10%) and characterized their rheological properties 105 and desalination performance to find the best compromise between these parameters. 106

Our AC modification strategy for desalination improvement in FCDI is very simple, compared
 with the complex physical and chemical AC modifications that have been recently used in CDI.
 This is important, especially due to the major improvements in FCDI desalination efficiency

and viscosity reduction observed with these bimodal flowable electrodes compared with pure

111 AC and FAC flowable electrodes.

112

# 113 2. Experimental

# 114 **2.1. Material preparation**

Commercial AC (Darco; CAS no: 7440-44-0) and sodium chloride (NaCl, CAS no: 7647-14-5,
 MW 58.44 g mol-1, 99%) were purchased from Sigma Aldrich. Cationic and anionic exchange
 membranes were purchased from Membranes International Inc. (Ringwood, New Jersey,
 USA). Deionized water (18 MΩ cm<sup>-2</sup>) was prepared using a purification system.

AC powder was dry ball-milled using 50-mm diameter balls (350 rpm rotation speed) at room temperature for 90 minutes to obtain finer particles (

				121
FE (10 % wt)	AC (% wt)	FAC (% wt)	[NaC (V=7	<b>i] (g/Ĺ)</b> <b>0 mĽ)<sup>2</sup></b> 123
FE1	10	0	1	124
FE2	8	2	1	125
FE3	5	5	1	126
FE4	2.5	7.5	1	127
FE5	0	10	1	128

)Erreur ! Source du renvoi introuvable..<sup>28</sup> Ball-milled AC, called fine particle AC (FAC), and pristine AC with different particle size ranges (0.3-0.6 μm and 0.8-1.6 μm, respectively) were used for this study.

AC suspensions, used as flowable electrodes, were prepared by mixing the desired amounts of AC and FAC (10% wt as total 129 weight percentage in each electrode) in 1 g/L NaCl solution (Erreur ! Source du renvoi 130 introuvable.). Two different packing characteristics were chosen; i) monomodal distribution, 131 where slurry electrodes contained only one particle size range (AC or FAC), and ii) bimodal 132 distribution where slurry electrodes were mixtures of AC and FAC at different ratios. Then, the 133 mixtures were sonicated for 2 h and stirred for 1 h to obtain a uniform particle distribution in 134 suspension. The slurry electrode in a beaker was continuously stirred and was fed into the cell 135 throughout the FCDI experiments (repeated three times for each sample). 136

137



140 **Figure 1**: Planetary dry ball milling of activated carbon (AC) into fine activated carbon (FAC).

FE (10 % wt)	AC (% wt)	FAC (% wt)	[NaC (V=70	l] (g/Ľ) ) mL) 143
FE1	10	0	1	144
FE2	8	2	1	± 1 1
FE3	5	5	1	145
FE4	2.5	7.5	1	146
FE5	0	10	1	147

141 **Table 1**: Composition of the flowable electrodes (FE) used in this study

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# 2.2. AC Material Characterization

For all sample characterizations, carbon suspensions prepared in 1 g/L NaCl solution 150 (experimental conditions) were used. Our previous study included a detailed physical 151 152 characterization of AC and FAC (after grinding) samples<sup>28</sup>. AC and FAC particle size distribution and their zeta potentials were measured by dynamic light scattering using a 153 Litesizer 500 particle size analyzer (Anton Paar, France). The particle structure was analyzed 154 by scanning electron microscopy (SEM) (SEM, Hitachi S4800, Tokyo, Japan), and the atomic 155 composition (chemical moieties) of the material was studied before and after ball milling 156 (analyzed surface area =  $300 \times 700 \mu$ m) by X-ray photonelectron spectroscopy (XPS) 157 (Spectrometre XPS/UPS KRATOS AXIS Ultra DLD). The surface area was measured using 158 159 N<sub>2</sub> adsorption/desorption at 77 K. The specific surface area was calculated using the Brunauer-160 Emmett-Teller method (Micromeritics 2020 ASAP, Merignac France). The total pore volume (V<sub>t</sub>) was calculated from the amount of N<sub>2</sub> adsorbed at 0.99 relative pressure (P/P<sub>0</sub>). The 161

mesopore volume (V<sub>meso</sub>) was calculated with the Barrett-Joyner-Halenda model. The dynamic 162 viscosity of the different carbon suspensions (flow electrodes) was measured using a Paar 163 Physica UDS 200 Rheometer, a concentric cylinder adapted for heterogeneous solutions or 164 water suspensions, as a function of the shear rate at 25 °C. Dynamic viscosity describes the 165 resistance of a liquid solution to flow, whereas the shear rate is the deformation speed of the 166 167 suspension under an applied force. The physical stability or sedimentation rate of the different AC slurries over time was measured with a Formulaction TURBISCAN® in which static multiple 168 light scattering is used to monitor particle migration in liquids. A measurement head with two 169 detectors moves along the cell height to measure the transmission (T) and backscattering (BS) 170 signals that are related to particle size and concentration in the sample. The changes over time 171 are a sign of destabilization that is quantified using the TURBISCAN<sup>®</sup> stability index (TSI). 172 Each experiment was repeated three times for each electrode type. The electrochemical 173 properties of AC and FAC were tested by cyclic voltammetry (CV) method. CV tests were 174 performed using Origalys OrigaStat OGS080 potentiostat and 3 electrodes system with a 5 175 176 mm carbon glass rotating disk electrode (RDE) as working electrode while a platinum mesh and a saturated 3 M KCl, Ag/AgCl electrode served as counter and reference electrodes 177 respectively. 10% wt. activated carbon suspensions were prepared, 100 µl were deposited on 178 the surface of the RDE then dried at 80°C before adding one drop of nation ink to hold the 179 material on the electrode. Voltammetry measurements were performed at a -0.2 V-0.4 V vs ref 180 operating potential window in 5 g/L NaCl solution as electrolyte. The electrical double laver 181 capacitance (EDLC) was determined using cyclic voltammetry cycles at 0.02 V. s<sup>-1</sup> scan rate. 182

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# 184 **2.3. FCDI experiments**

Figure 2 shows the FCDI experimental set-up and its operating principle. The system was 185 composed of a homemade FCDI cell, a potentiostat (Origalys Electrochem, Origaflex), a 186 187 conducti-meter (HANNA Instruments) to monitor the saline solution conductivity during the desalination experiment, and two beakers (one for the flowable electrode and the other for the 188 saline solution) with two peristaltic pumps to recirculate both solutions through the 189 190 electrochemical cell. Experiments were performed in a closed loop reactor at 1V constant voltage, using 5g/L NaCl as feed solution (FS), and lasted 30 min. The FDCl cell (Erreur! 191 Source du renvoi introuvable.(a)), specifically designed for these experiments, was 192 193 composed of two current collectors (0.9mm depth, 8.5 cm width, and 14.5 cm length) with two channels each as flow electrode inflow and outflow of 7.8 cm depth, two ion exchange 194 membranes (IEM) of 142 cm<sup>2</sup> effective area, a spacer (0.9 mm thick), two stainless steel plates, 195 196 and nails to hold all parts together.

197 The electrical field created inside the FCDI cell drives ions present in the FS (5 g/L NaCl) 198 through a spacer, sandwiched between two IEM, to be electrostatically attracted to the charged 199 flowable electrodes that circulate through the flow paths carved on the current collectors, as 200 shown in Erreur! Source du renvoi introuvable.(a). When the flowable electrodes exit the channels, they are fed back into the reservoir, stirred, and then re-circulated inside the cell. 201 The resulting current at the electrodes was monitored throughout the experiment by 202 203 chronoamperometry. The FS and flowable electrodes were flowing at a constant flow rate of 3 ml.min<sup>-1</sup> and 40 ml.min<sup>-1</sup> respectively. Throughout the process, the conductivity of the FS that 204 205 was recirculated in a closed loop system was monitored at room temperature with a

206 conductimeter (Hannah Instrument), as depicted in Erreur ! Source du renvoi
 207 introuvable.(b). To mention that no significant water transmembrane migration was observed
 208 during the desalination experiments.



Figure 2: (a) FCDI cell. (b) Experimental set-up and operating principle. CEM: cationic exchange membrane; AEM: anionic exchange membrane

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213 FCDI desalinating performance indicators were calculated for each experiment.

Salt adsorption capacity (SAC) indicating the mass of adsorbed salt as function of the adsorbent electrode weight is calculated according to equation (1) and expressed in mg of salt adsorbed per g of electrode material:

217 
$$SAC = \frac{(C_0 - C_f)V}{m_{electrode}}$$
(1)

218 Where  $C_0$  is the initial solute concentration of the feed solution,  $C_f$  the final solute concentration,

- 219 V is the volume of the feed solution and m<sub>electrode</sub> is the mass of the electrode material.<sup>29</sup>
- 220

The salt removal efficiency or desalination efficiency (DE) was calculated using equation (2):

223 
$$DE = \frac{(C_0 - C_f)}{C_0}$$
 (2)

- where  $C_0$  and  $C_f$  are the initial and final concentrations (mol. L<sup>-1</sup>) of the FS.<sup>1</sup>
- The salt adsorption rate (SAR), which represents the amount of salt adsorbed ( $\mu$ g) per
- electrode surface (cm<sup>2</sup>) per unit of time (min), was calculated using equation (3):

228 
$$SAR (\mu g. cm^{-2}. min^{-1}) = \frac{(C_0 - C_f) V}{A t}$$
 (3)  
229

where V is the volume of the feed solution (L), A is the contact area between the feed electrode
 and the current collector, and t is the charging time.<sup>30</sup>

The charge efficiency (CE) or faradaic efficiency is the ratio of the electrical charges related specifically to ion removal over the total electrical charge applied by the potentiostat. It represents the energy efficiency of the system and was calculated using equation (4):

235

236 
$$CE = \frac{z (C_0 - C_f) V F}{\int I dt}$$
 (4)

237

where z is the equivalent charge of ions, V is the FS volume, F is the Faradaic constant, and  $\int I dt$  is the integrated quantity of charge passed to the system over time.<sup>31</sup>

The Reynolds number shows the electrode flowing nature in the channel and was calculated with equation (5):

$$242 \quad Re = \frac{\rho \, v \, D}{\eta} \tag{5}$$

where  $\rho$  and  $\nu$  are the flowable electrode density and velocity, respectively, in the flow channel calculated from the flow rate of the flowable electrode (0.0024 m<sup>3</sup> h<sup>-1</sup>) to the area of the flow cell (5.4 x 10<sup>-6</sup> m<sup>2</sup>), D is the hydraulic diameter of the flow channel (6 mm), and  $\eta$  is the mean coefficient of dynamic viscosity of the flowable electrode in function of the shear rate at 25 °C.

Stokes's law of sedimentation shows the drag force resisting the fall of small particles througha fluid medium under the influence of gravity, and is described by equation (6):

249 
$$V = \frac{2(\rho_p - \rho_f)}{9\eta} g R^2$$
(6)

where  $\rho_p$  is the particle density,  $\rho_f$  is the fluid density, *g* is the acceleration of gravity, *R* is the particle radius, and  $\eta$  is the viscosity.<sup>32</sup>

The linear pressure drop is the loss of mechanical energy caused by friction on the common walls of the network. It is directly affected by the fluid viscosity, and is described by equation (7):

255 
$$\Delta P = \mu \frac{L}{D} \frac{1}{2} \rho v^2$$
 (7)

where  $\mu$  is the friction coefficient with  $\mu = \frac{64}{Re}$ , L and D are the canal length and diameter, 256 257  $\rho$  is the fluid apparent density, and  $\nu$  its velocity.<sup>33</sup>

The electrical double-layer capacitance (EDLC) was calculated according to equation (8): 258

$$259 \quad C_{EDL} = \frac{\int I dV}{2 \, u \, A \, \Delta V} \tag{8}$$

where  $C_{EDL}$  is the double layer capacitance, *I* is the current response,  $\Delta V$  is the potential 260 window, A is the effective area, and u is the scan rate.<sup>34</sup> 261

262

### 3. Results and discussion 263

### Particle size reduction 264 3.1.

AC and FAC morphological and structural properties were investigated in our previous study.<sup>28</sup> 265 The dynamic light scattering (DLS) data (Erreur ! Source du renvoi introuvable. (a)) showed 266 the difference in particle size distribution between pristine AC and FAC samples: 0.8-1.6 µm 267 268 versus 0.3 to 0.6 µm respectively.



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### 3.2. Structural and textural characteristics 272

The structural and textural characteristics as well as the chemical moieties and composition of 273 AC and FAC samples were described in our previous study and are summarized in Figure S1, 274

Table S1 and Figure S2 in supplementary information. Briefly, all AC characteristics were maintained in FAC after dry ball grinding, without any significant difference in texture (pores structure and surface area).<sup>28</sup>

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# **3.3.** Physical and chemical properties

# 280 *3.3.1. Atomic composition*

The atomic composition of the carbon material before and after ball milling was investigated 281 using XPS. No band reduction was observed after ball milling. However, according to XPS 282 283 analysis, the O1s atomic percentage of the carbon material slightly increased after ball milling from 5.10 % (AC) to 5.25 % (FAC), which can be due to the generation of new oxygenated 284 surface functional groups after ball milling. Figure 4 a shows the complete spectra of both AC 285 and FAC, while c and d present the resolved C 1s spectrum of AC and FAC respectively into 286 their individual peaks. 284.00, 284.80 and 286.10 eV binding energies correspond to C 1s sp2, 287 C 1s sp3 and C 1s C-O respectively.<sup>35</sup> As well Figure 4 b and c show a slight increase in the 288 C 1s C-O band area from 3.92% to 5.34% after ball milling (FAC) proving the formation of new 289 290 C-O functional groups on the surface of the activated carbon material.



Figure 4: XPS complete survey of AC and FAC (a), deconvoluted XPS C 1s spectra of AC (b) and FAC (c)

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## 295

# 5 **3.3.2.** Rheological properties and zeta potential: effect of particle distribution

To improve the flowing properties of carbon-based flowable electrodes in FCDI systems, 296 297 bimodal distributions were used by mixing (different ratios) of FAC (size range: 0.3-0.6 µm) 298 and AC (size range: 0.8-1.6 µm). This was based on the hypothesis that fine particles can fill 299 the gaps between larger particles (packing effect), increasing the connectivity within the carbon network, while keeping enough free spaces for water to flow through (i.e. acceptable viscosity). 300 301 The packing effect on the dynamic viscosity and surface charge (zeta potential) was studied first by measuring the viscosity in function of the shear rate and zeta potential of carbon 302 suspensions with monomodal and bimodal distributions (different FAC to AC ratios) (Figure ). 303 The apparent viscosities of all carbon suspensions decreased when the applied shear rate 304 305 was increased, showing a flow behavior typical of a pseudo-plastic fluid, this phenomenon is typical of non-Newtonian fluids.<sup>36</sup> Moreover, their viscosity (Figure 5(a)) increased 306 307 proportionally with the FAC percentage. Upon dry ball grinding of AC, each particle is divided 308 into many particles, thus increasing the particle number for the same weight percentage and flowable electrode volume. Consequently, the higher specific surface area of particles 309 increases interactions with water, and dissipates more energy due to friction. This also 310

promotes particle-particle interactions, notably repulsion forces, leading to higher resistance to 311 flow or higher viscosity. This is observed particularly at low shear rates.<sup>37</sup> This effect becomes 312 less significant at higher shear rates because the interactive forces between particles are less 313 effective. Moreover, wide and narrow particle size ranges are associated with low and high 314 viscosity, respectively.<sup>38</sup> This explains the lower viscosity of AC samples because their particle 315 size range was wider than that of FAC samples. Suspensions with wide particle size 316 317 distribution (large polydispersity) aggregate more easily than suspensions with narrow polydispersity. This gives more of free space that facilitates the movement (flow) of individual 318 particles, and thus lower viscosity. These results are in agreement with those by Harmadi et 319 al.<sup>39</sup> and Buranasrisak et al.<sup>40</sup> who studied the particle size effect on the rheology and stability 320 321 of carbon-water mixtures.

322 Zeta potentials represent the surface charge of particles in suspension. The particle size effects on zeta potential and apparent viscosity was studied. The electrostatic repulsive forces 323 of particles in suspension under an applied voltage depend on their surface charge. It was 324 found that zeta potential increases after ball-milling, which can be attributed to the increase in 325 oxygenated functional groups on the surface of the material as shown previously in the XPS 326 study. The addition of large particles with low zeta potential to the flow electrodes leads to 327 decreased repulsive behavior, and decrease the suspension viscosity because the smaller 328 specific surface area leads to less friction, as explained previously. However, suspensions of 329 330 finer particles, which have strong surface charge (high zeta potential) increase particle-particle repulsion and the resistance to flow or viscosity. Similarly, zeta potential measurements 331 (Figure (b)) showed the interdependence between viscosity, surface charge, and FAC 332 percentage. As finer particles in suspension have stronger surface charge (higher zeta 333 potential), particle-particle repulsion is promoted, decreasing aggregation. Consequently, the 334 whole space is occupied by well-dispersed particles, and less space is available for free water 335 flow, leading to higher viscosity. Upon reducing FAC percentage in the suspension, repulsion 336 forces between particles were decreased as well as resistance to flow, and therefore, flowing 337 338 behavior in the FCDI system was improved. These results are fully in line with those by 339 Buranasrisak *et al.*<sup>40</sup>.





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shear rate). All experiments were carried out with carbon suspensions with 10 % wt carbon loading,
fine particle size range of 0.3-0.6 μm, and large particle size range of 0.8-1.6 μm.

# 345 *3.3.3. Reynolds number*

Reynolds number (i.e. the inertial forces to viscous forces ratio) is used to categorize the fluid 346 systems where viscosity has an important effect in controlling the fluid velocity or flow pattern, 347 348 a critical parameter in our study. For all mixtures, the Reynolds number (calculated using 349 equation (5)) was <2100, indicating a laminar flow (i.e. the flow type needed for flowable electrodes in FCDI systems) (Table 2). As flow velocity is directly proportional to the Reynolds 350 351 number, this explains the decrease of the Reynolds number in carbon suspensions with higher FAC percentages because they decrease the flow velocity. Consequently, when the viscosity 352 increased with higher FAC: AC ratios, the Reynolds number decreased. 353

 354
 Table 2: Reynolds number of the indicated carbon suspensions

 FAC : AC ratio
 Reynolds n°

 0.0 : 1.0  $57.787 \pm 0.006$  

 0.2 : 0.8  $45.033 \pm 0.006$  

 0.5 : 0.5  $30.620 \pm 0.001$  

 0.75 : 0.25  $21.808 \pm 0.008$  

 1.0 : 0.0  $13.106 \pm 0.030$ 

# 355

# 356 **3.3.4**. *Physical stability*

357 AC particles suspended in water tend to settle due to gravitational force. In FCDI systems, this can cause clogging inside the channels, thus reducing the particle availability for ion adsorption 358 359 during the process. The destabilization kinetics of suspensions with different fine to large particle ratios (at 10% wt) were evaluated by measuring the TSI (TURBISCAN® technology) 360 over 30 minutes (Erreur ! Source du renvoi introuvable.). The TSI (i.e. destabilization) 361 progressively increased for samples with lower FAC percentages. This indicates that the 362 stability of the carbon suspensions increased with higher FAC percentages, although no 363 difference was observed for samples with fine particles to large particle ratios of 0.75:0.25 and 364 1.0:0.0. Particle size reduction is associated with slower sedimentation rates. This is explained 365 by the sedimentation velocity law shown in equation (6). Indeed, the particle size has an effect 366 on the sedimentation velocity. According to Stokes' law, the particle sedimentation velocity is 367 368 proportional to the density difference between the solid and liquid phase, inversely proportional to the fluid viscosity, and proportional to the particle diameter square. This explains the 369 important decrease in sedimentation velocity with FAC percentage increase in the 370 suspensions: smaller particle size is associated with higher viscosity. FAC particles remained 371 in suspension and well dispersed for a longer period of time, thus increasing the intra-particle 372 interactions in solution. These results are in agreement with the rheology and zeta potential 373 data, and explain the behavior of the different carbon particle size suspensions as flowable 374 electrodes. 375



Figure 6: TURBISCAN<sup>®</sup> stability index (TSI) of the carbon suspensions at different ratios of fine particles
 to large particles. Below level A (lowest TSI dash line) on the TSI visual scale, no destabilization is
 observed. Level A marks the beginning of the very early destabilization stage. Level B indicates the
 beginning of destabilization, and levels C and D describe high sedimentation stages.

# **381 3.4. Electrochemical properties**

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To evaluate the electrochemical behavior of activated carbon electrodes with different 382 FAC:AC ratios, cyclic voltammetry (CV) measurements were carried out using a 1 g/L NaCI 383 384 electrolyte solution at 2 mV. s<sup>-1</sup> scan rate with a -0.2 – 0.4 V voltage window. The CV results 385 are shown in Figure 7 (a). All the electrodes displayed a symmetrical box-like shape, which reflects good capacitor characteristic. The larger area obtained under the CV curves implies 386 the higher specific capacitance of the materials. The 1.0: 0.0 FAC: AC ratio (pure FAC) 387 electrode showed the highest area obtained under the CV curves, indicating the highest 388 389 specific capacitance compared to the pure AC electrode. As well, based on the cyclic 390 voltammetry cycles, the EDL capacitance of the different electrodes of different FAC: AC ratios was calculated according to equation 8 and presented in figure 7 (b).<sup>41</sup> Thus, in all 391 392 cases, the addition of FAC to the AC electrodes enhances their double layer capacitance due 393 to the increased conductivity of the flow electrodes caused by the improved interconnectivity 394 between the carbon particles in the case of FAC as explained in details before. Thus pure 395 FAC electrode showed the highest specific capacitance of 35 F.cm<sup>-2</sup> compared to 25.5 F.cm<sup>-2</sup> in the case of pure AC marking a higher conductivity of FAC compared to AC. 396



**Figure 7**: Cyclic voltammetry cycles (a) and electrical double layer capacitance (b) for activated carbon electrodes with different FAC:AC ratios.

# 400 **3.5. Desalination performance**

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The electrosorption characteristics of the different carbon suspensions (10% wt carbon 401 loading) as flow electrodes were investigated without clogging in our FCDI set-up. FCDI 402 performance indicators (DE, SAC, SAR and CE) were calculated for the different mixtures 403 (Table 3 and Figure (b and c)). In all cases, FAC addition strongly improved their salt adsorption 404 capacity and desalination performance in FCDI. The higher specific surface area, more 405 accessible pores, better conductivity observed upon particle size reduction explain the 406 improved DE, SAC, and SAR values (Table 3, Figure 8 (b)). Higher surface area promotes 407 mass transfer (ion adsorption), which is the limiting mass transfer phenomenon in FCDI. This 408 409 strongly influences the SAR values, as seen after FAC addition in the flowable electrodes.

Sharper drop in ionic conductivity of the FS when using bimodal mixtures or pure FAC 410 compared with pure AC was observed (Figure 7 (a)). Flowable electrodes with 0.75:0.25 and 411 1.0:0.0 FAC:AC ratios exhibited the best DE, the highest SAC and SAR among all samples 412 (Table 3 Figure 8 (b and c)). DE was 72%, SAC was 32.3 mg.g<sup>-1</sup>, and SAR was 1772 µg.cm<sup>-</sup> 413 <sup>2</sup>.min<sup>-1</sup> for the 0.75:0.25 sample compared with 34%, 15.3 mg.g<sup>-1</sup> and 839 µg.cm<sup>-2</sup>.min<sup>-1</sup> for the 414 415 pure AC flow electrode respectively. This can be due to the improved ion diffusion pathways 416 within the carbon network because fine particles filled the void between larger particles, thus increasing contact areas within the carbon network, and consequently conductivity in the 417 418 flowable electrode. The lower desalination performance of the flowable electrode with AC particles might be due to the lower repulsion forces between particles that aggregate to each 419 420 other, thus masking the pores that become less accessible to ions. Furthermore, the higher resistance and lower conductivity caused by water occupying the large gaps between large 421 AC particles create long ion diffusion pathways. These parameters explain the lower 422 423 performance metrics of these flowable electrodes. The chronoamperometry curves (Figure S3 424 in SI) for the different carbon slurries showed the current density in function of time at a 425 constant applied voltage (1V). Current density was lower for pure AC suspensions than for the samples with different FAC percentages. Therefore, the total charge transferred (which is the 426 427 integration of the chronoamperometry curves) was lower for large AC particles, resulting in lower DE. The current density progressively increased with the percentage of FAC added, and 428

429 DE improved. Conversely, CE was comparable among samples (Figure 8 (c)), likely because 430 this parameter is mostly influenced by the ion transfer through the selective IEM. For all samples, the same FCDI system with the same IEM was used, the conductivity of the saline 431 432 solution passing between the electrodes was the same (5 g/L NaCl), and the quantity of carbon in the electrode compartments was largely enough to receive ions from the central 433 434 compartment. The high selectivity of cationic and anionic membranes in such conditions explains the comparable CE values among samples.<sup>42</sup> Analysis of the relationship between 435 the apparent viscosity and DE of the different flowable electrodes (Figure 8 (d)) showed that 436 with the 0.75:0.25 FAC:AC ratio bimodal mixture, viscosity decreased by 30%, overcoming the 437 438 major limitation of FAC-based flowable electrodes in FCDI (i.e. high viscosity). The challenge is always to find a compromise between these parameters to improve desalination 439 performance in FCDI, while maintaining good flowing properties of the electrodes inside the 440 system. In this study a good compromise between these parameters was achieved with the 441 0.75:0.25 FAC:AC ratio bimodal mixture. 442

443	Table	<b>3</b> : sa	lt ad	Isorption	ca	pac	ity (	(SAC)	and	laverage	salt	adsorption	rate	(SAR)	of	carbon	flow
						-											

FAC : AC ratio	SAC (mg.g <sup>-1</sup> )	SAR (µg.cm <sup>-2</sup> .min <sup>-1</sup> )				
0.0 : 1.0	15.3	0.51				
0.2 : 0.8	27.6	0.92				
0.5 : 0.5	28.9	0.96				
0.75 : 0.25	32.3	1.08				
1.0 : 0.0	31.9	1.06				

444 electrodes with different FAC : AC ratio



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Figure 8: Feed solution conductivity (a), desalination efficiency and salt adsorption rate (SAR) (b), and charge efficiency (c) of carbon-based flowable electrodes with different FAC: AC ratios, during 30 min of FCDI experiment time. Relationship between desalination efficiency and apparent viscosity (d).

449 The rheological characteristics of the carbon slurries were studied to optimize their flowability inside the FCDI system because this has a direct effect on the pumping energy and cost. To 450 this aim, viscosity effect on the pressure drop was assessed when comparing the desalination 451 performances of the different flowable electrodes. In every FCDI process, there is always a 452 compromise between particle size and viscosity. Smaller carbon particles increase 453 conductivity, and consequently the process efficiency, for instance in terms of charge transfer, 454 maximum salt capacity, mass transfer. However, adding small particles to the carbon slurry 455 increases viscosity and also pumping costs due to the pressure drop increase. In our set-up, 456 pressure drop (calculated with equation (7)) increased with the viscosity increase (Figure 9). 457 With higher FAC percentages in the slurry, DE improved but viscosity increased, and also the 458 pressure drop, leading to higher pumping energy and costs. Therefore, it is important to find 459 the best compromise between particle size distribution and viscosity in order to obtain an 460 energy-efficient process with good performance without clogging problems inside the FCDI 461 462 system.



Figure 9: Relationship between desalination efficiency, apparent viscosity, and pressure drop increase
 of carbon-based flowable electrodes with the indicated FAC: AC ratios, during 30 min of FCDI
 experiment time.

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# 468 **4. Conclusion**

This work investigated the effect of particle size distribution on the viscosity and desalination 469 performance of AC-based flowable electrodes in FCDI. Mixtures of two different AC particle 470 sizes were used: commercial AC and FAC obtained after ball-milling commercial AC. This 471 experimental work shows that particle size distribution has a considerable effect on ion 472 473 diffusion length and on the ion transport pathway that is determined by the packing density of 474 flowable electrodes. Particularly, with pure AC suspensions, intraparticle and interparticle ion 475 diffusion limited the desalination capacity (34% of DE). By mixing two different particle sizes, particle size distribution became broader, leading to better desalination performance, as 476 indicated by the 72% of DE and improved flowing properties of the flowable electrode in FCDI 477 (decrease by 33% of viscosity compared with the pure FAC suspension that achieved 70% of 478 479 DE) in FCDI. These results highlighted the importance of particle size distribution and packing density on the behavior of flowable electrodes optimized for FCDI-based desalination. Fine 480 particles and combined (0.75: 0.25 FAC: AC ratio) slurries showed better desalination 481 482 performance compared to large particle slurries. However, large particle slurries showed better 483 fluidity and easier flow in our water desalination system leading to lower pressure drop. Thus, good compromise must be done between high desalination capacity and acceptable pressure 484 drop (viscosity) to have an optimal electrochemical desalination process. To find the best 485 balance, more research is needed, including molecular dynamic simulations that can improve 486 experimental procedures and save time. 487

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492 and M.C.; Visualization: F.Z., M.T., M.B. and M.C.; Supervision: F.Z., M.B., P.S. and M.C.;
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