# Aqueous-interfacial and on-water molecular reactions across diverse chemistries

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#### 44 Abstract

This review aims to critically analyze the current state of knowledge in the emerging field of 45 chemical reactivity at aqueous interfaces. The area has evolved rapidly since the discovery of the 46 so-called "on-water catalysis" effect, alluding to the fact that many chemical reactions 47 experience a dramatic acceleration at the surface of water or different aqueous interfaces with 48 hydrophobic media. The immense importance of this phenomenon is discussed first by reviewing 49 some critical experimental studies in the fields of atmospheric and synthetic organic chemistry, 50 as well as related research exploring the origins of life. The physicochemical aspects of the topic 51 are analyzed afterwards. First, with a concise analysis of issues such as the structure, the 52 dynamics, and the thermodynamics of adsorption/solvation processes at aqueous interfaces. 53 Then, presenting the basic theories intended to explain interface catalysis, followed by the results 54 of advanced ab initio molecular dynamics simulations. Though some topics addressed here have 55 already been the focus of previous reviews, their interconnection across diverse disciplines has 56 not been sufficiently highlighted in the literature. For this reason, this manuscript seeks to 57 provide a common perspective by trying to identify the most fundamental issues still 58 incompletely understood in this fast-moving domain. 59

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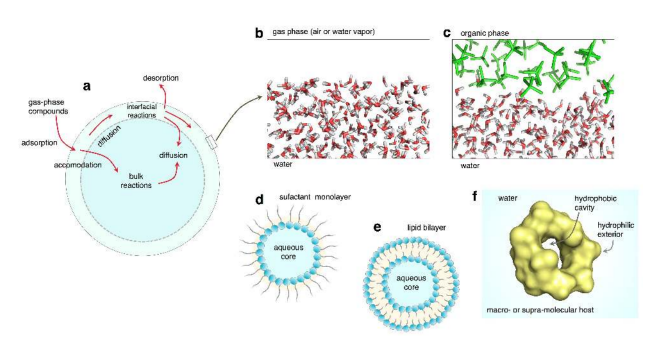
### 62 Introduction

Chemistry at aqueous interfaces is a vast subject that encompasses processes from quite 63 different domains. Specifically, we deal here with processes occurring at liquid water-vapor 64 interfaces (or air-water interfaces) and the interfaces of liquid water with hydrophobic 65 environments. Figure 1 illustrates a few examples of systems that are covered by this review. 66 Many chemical and photochemical reactions are dramatically accelerated when they occur at 67 these interfaces, in comparison with gas-phase or bulk water, and this phenomenon is now 68 designated as "on-water" catalysis.<sup>1</sup> The term chemistry "on-water" must be understood here in a 69 broad sense, i. e. the chemistry that occurs at, or near aqueous interfaces in oil-water emulsions 70 and other dispersed systems, aerosols, sprays, nano and micro water droplets, as well as extended 71 air-water interfaces. It goes without saying that such a variety of systems may involve 72 phenomena implying quite different mechanisms, which makes the implementation and 73 interpretation of experimental measurements often a complex task. The potential implications of 74 interfacial reactions are widespread because they are omnipresent in atmospheric, environmental, 75 biological, prebiotic, or synthetic organic chemistry, to cite the most relevant domains. Interfaces 76 of liquid water with either solids or biomolecules, as well as the surface of ice, share many 77 similarities with the former but are not directly concerned by the subject of the present review. 78 The reasons underlying rate acceleration at aqueous interfaces remain unclear however. In 79 contrast to bulk solvation, the theory of interfacial solvation is still in its early stages. The 80 formation of hydrogen-bonds with dangling protons at the interface was first proposed to explain 81 the catalytic role of the interface,<sup>2</sup> but many other causes can be invoked: confinement of 82 reagents, partial solvation, preferential orientations, curvature in nanodroplets, water surface pH, 83 etc. It is worth pointing out that physicochemical concepts from the bulk are not always 84 applicable at interfaces, as the latter are disordered systems of nanometric thickness displaying 85 sharp configurational fluctuations. Experimental studies based on macroscopic properties such as 86 surface tension<sup>3,4</sup> have provided invaluable data on interfacial thermodynamics and structural 87

properties. However, only with the progress of non-linear second-harmonic generation (SHG)
 and sum-frequency generation (SFG) spectroscopies,<sup>5</sup> and other interface-sensitive molecular
 techniques, the microscopic details of interfacial phenomena are being elucidated. In parallel, *ab initio* Molecular Dynamics (MD) simulations and related approaches have provided priceless
 information on these issues.<sup>6-8</sup>

Nevertheless, the literature remains scattered across various fields. In fact, despite the similarities between all these chemistries and the existence of some reviews on restricted aspects of the topic, a general discussion on the available experimental and theoretical studies, placing them in a shared perspective, is still lacking. In this review, we will provide such a perspective through a comprehensive and critical survey of the recent literature aiming to highlight the main challenges that need to be addressed in order to advance the state-of-the-art in the field.

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**Figure 1.** Aqueous interfaces contemplated in this review: a) the air-water interface at the surface of a water droplet with indications of the different processes that can take place, b) detail of the liquid water-vapor interface, c) interface of liquid water with a non-miscible organic solvent ( $CCl_4$  here), d) inverted micelle in aqueous organic aerosols, e) vesicles, f) macro- or supra-molecular systems with a hydrophobic cavity that can host hydrophobic guest molecules.

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## 112 Chemical reactions at aqueous interfaces

Many contributions in this field come from the atmospheric chemistry community because 113 reactions at the air-water interface of cloud water droplets and aqueous aerosols may proceed at 114 higher rates than gas-phase reactions, influencing the atmospheric budget of trace gases.<sup>9-12</sup> 115 Further interest comes from the field of synthetic green chemistry. The need to develop green 116 processes for the synthesis of organic compounds that decrease the negative environmental 117 impact of current industrial practices pleads for the use of non-organic solvents such as water. 118 Experiments have shown that reactions in water microdroplets generated by electrospray 119 ionization undergo remarkable acceleration with respect to bulk-phase processes, and due to 120 large surface to volume ratio, the air-water interface is thought to play a key role.<sup>13-16</sup> Moreover, 121 122 dispersed systems such as polyelectrolyte solutions, micellar solutions, oil-in-water microemulsions or vesicle dispersions, have been proposed to overcome water solubility 123 limitations and develop biomimetic reactors within which reactions can proceed.<sup>17,18</sup> Aqueous 124 interfaces have also been evoked as possible environments in which prebiotic processes could 125 have taken place and led to the origin of life. For instance, orientation, alignment and proximity 126 of functional groups is essential to the synthesis of peptides by the ribosome, and air-water 127 interfaces in inverted micelle atmospheric aerosols or in the surface of oceans and lakes could 128 have been a rudimentary prebiotic system mimicking this functioning.<sup>19,20</sup> 129

We have selected some illustrative experimental works, and organized them in four specific (and to some extent, arbitrary) areas that are not disjointed, but rather overlap in many ways.

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Atmospheric and environmental chemistry. The role of condensed matter in the Earth's
 atmosphere is widespread. Aerosols scatter sunlight and serve as seeds for the formation of

clouds, which has significant consequences in climate regulation.<sup>21</sup> Condensed matter in its 135 different forms also supplies a medium for chemical transformations. Well-known examples are 136 the oxidation of SO<sub>2</sub> to sulfate in water droplets, which leads to acid-rain formation in the 137 troposphere,<sup>22</sup> or the heterogeneous reactions that lead to ozone depletion in the stratosphere.<sup>23</sup> 138 Indeed, despite a small volume fraction of atmospheric condensed matter (about 7% of the total 139 volume of the troposphere contains clouds, and a moderately dense cloud contains about  $5 \times 10^{-7}$ 140 cm<sup>3</sup> of water per cm<sup>3</sup> of air),<sup>9</sup> its relevance is now recognized.<sup>9,24,25</sup> It influences the atmospheric 141 budget of trace gases through the modification of the cycles of nitrogen, sulfur, and various 142 atmospheric oxidants such as ozone.<sup>26,27</sup> In addition, some reaction pathways that are unfeasible 143 in the gas-phase (e.g. ionic dissociations) may be quite favorable in the condensed phase, 144 producing new species.<sup>28</sup> In line with the subject of this review, we put the focus on liquid water 145 interfaces (water droplets, aqueous aerosols) even though the heterogeneous reactions at the 146 surface of solid matter such as carbonaceous particles or mineral dust have comparable 147 importance. 148

When atmospheric trace gases interact with a water droplet, several phenomena can take place 149 (Figure 1) including uptake, diffusion and reaction at the surface, desorption, mass-150 accommodation, diffusion and reaction in the bulk.<sup>29</sup> Bulk reactions are relatively well-151 understood<sup>30</sup> but not those occurring at the air-water interface. Several studies have confirmed 152 that the efficiency of interfacial processes in the atmosphere may be quite significant, compared 153 to bulk processes.<sup>7,11,31-33</sup> This is due in part to the high surface to volume ratio characterizing 154 atmospheric droplets and aerosols. However, there is evidence of specific effects that accelerate 155 chemical and photochemical reactions at aqueous interfaces; some particular examples using 156 different experimental platforms are outlined below. 157

Adsorption of trace organic molecules on water film surfaces enhance their reaction with atmospheric oxidants.<sup>33</sup> Electrospray-mass-spectrometry studies<sup>34</sup> showed that when benzoate is allowed to react with OH radicals at the air-water interface, H-abstraction from the aromatic ring

is mainly observed. At the same time, this reaction is negligible both in the gas-phase and bulk 161 water. The rationalization of this results goes in terms of the higher polarity of the transition state 162 for the OH-radical addition compared to H-abstraction.<sup>34</sup> A similar experimental technique was 163 used by Enami et al<sup>35</sup> to study the reactivity of isoprene at mildly acidic water surfaces showing 164 that it can undergo cationic oligomerization. The authors suggested a superacidity behavior of 165 the air-water interface with pH < 4 water, a result that has raised some debate (see below). Fatty 166 acids, which are generally not sensitive to actinic radiation, produce aldehydes and other 167 oxygenated species when a monolayer at the water surface is irradiated in the 280-330 nm 168 region.<sup>12</sup> The process seems to involve UV-absorption to a triplet state followed by the homolytic 169 OH dissociation or by reaction with an adjacent fatty acid molecule at the air-water interface. 170 The conclusions of these experiments, however, have been challenged by subsequent studies by 171 Shrestha et al<sup>36</sup> and Rapf et al,<sup>38</sup> who have emphasized the need of photoinitiators for reactions of 172 this type to take place, as fatty acids are not themselves photoactive. Upon irradiation of 173 nebulized pyruvic acid, zymonic acid has been observed among the products formed,<sup>39</sup> as 174 opposed to other conditions, suggesting that it could be generated by reactions at the droplet 175 surface. Other interesting interface-assisted photochemical processes can be found in the review 176 by George et al.<sup>11</sup> Colussi and coworkers<sup>40-46</sup> have devoted considerable effort to the study of the 177 ozonolysis reaction and the chemistry of the Criegee intermediate at the air-water interface, 178 which are chemical processes with broad implications in the atmosphere, as they represent a 179 major sink for unsaturated volatile organic compounds produced by plants, particularly isoprene 180 and monoterpenes. The oxidation reaction of anthracene by ozone on aqueous surfaces was 181 studied by Mmereki et al,<sup>47</sup> who showed that it may be of comparable importance to gas-phase 182 oxidation by OH in the atmosphere. Chemistry at the surface of sea-salt aerosols and its 183 atmospheric implications have been extensively studied by Finlayson-Pitts and coworkers, who 184 have emphasized the role played by the air-water interface. For instance, the main sources of Cl<sub>2</sub> 185 and Br<sub>2</sub> gases from sea-salt aerosols under dark conditions are the interfacial reactions of the 186

corresponding halide anion with OH and O<sub>3</sub>, respectively.<sup>48</sup> When concentrated NaCl aerosols 187 are irradiated at 254 nm in the presence of O<sub>3</sub> to generate OH radicals, the observed amount of 188 Cl<sub>2</sub> gas produced is in good agreement with estimates based on field measurements in the marine 189 boundary layer.<sup>10</sup> At the surface of aqueous aerosols, halide ions (and also some cations) 190 influence other interfacial reactions such as the production of NO<sub>2</sub> from photolysis of nitrate.<sup>49-51</sup> 191 Some fundamental knowledge about the water effects on reactions at the air-water interface of 192 water droplets and aqueous aerosols comes from the study of small water clusters and further 193 details on this topic can be found in the review by Vaida.<sup>52</sup> 194

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Microdroplets as synthetic chemical reactors. Reactivity in microdroplets is emerging as an up-196 and-coming tool in synthetic organic chemistry. Acceleration of many organic reactions in 197 aqueous media has been known for decades, <sup>53-56</sup> especially after the seminal work by Breslow 198 and coworkers on the Diels-Alder reaction.<sup>57,58</sup> This is rather good news because one could 199 consider water as the ideal green solvent. Studies by Sharpless and coworkers<sup>1</sup> pointed out that 200 some reactions proceed optimally in pure water when insoluble reactants are stirred in the 201 aqueous medium, and denoted such processes "reactions on-water". Such processes occur in 202 aqueous suspension and thus, hydrophobic effects might be claimed to provide the driving-force 203 for rate acceleration. Yet, experimental results showed that observed rates are not the sole 204 consequence of an effective concentration increase,<sup>1</sup> and since the pioneer work of Sharpless and 205 coworkers,<sup>1</sup> "on-water" chemistry has been steadily expanding (see for instance the reviews by 206 Butler and Coyne<sup>59</sup> and by Butler et al<sup>60</sup>). 207

In recent years, synthesis in small volume microreactors has been the subject of intense research.<sup>14</sup> This includes studies in microdroplets generated by a variety of electrospray and other spray mass-spectrometry methods,<sup>13,15,20,61-66</sup> levitated droplets,<sup>67</sup> thin films on surfaces<sup>68,69</sup> or microfluidic systems.<sup>70-73</sup> In many cases, the reaction rates are higher with respect to the reference bulk reaction (see a counterexample here<sup>74</sup>) but the effects responsible for such rate

accelerations are still unclear. Confinement of reagents and increased concentration (due to 213 solvent evaporation) are probably important factors,<sup>14</sup> but the large surface to volume ratio 214 characterizing these systems also points at specific interface effects.<sup>13-16</sup> Experimental data 215 supporting this statement were reported by Cooks and coworkers<sup>75</sup> in the study of competitive 216 substituent effects in Claisen–Schmidt reactions. Other experiments by Mellouli et al<sup>72</sup> using a 217 biphasic microfluidic approach, which allows getting better control of the generated interfaces 218 and water surface area, concluded that stabilizing hydrogen-bonds play a role in decreasing the 219 activation energy, as previously suggested by Jung and Marcus<sup>2</sup> (see below). The observed rate 220 increase is sometimes very large. For instance, the Pomeranz-Fritsch synthesis of isoquinoline in 221 charged microdroplets generated by electrospray has been reported to be at least 10<sup>6</sup> times faster 222 than in bulk.<sup>76</sup> Likewise, Enami et al<sup>77</sup> showed that Fenton ( $Fe^{2+} + H_2O_2$ ) and Fenton-like ( $Fe^{2+} + H_2O_2$ ) 223  $O_3$ ) reactions proceed  $10^3$ - $10^4$  faster at aqueous interfaces than in bulk aqueous media due to a 224 modified geometry of the hydration shell of Fe<sup>2+</sup>, which may have implications not only for 225 advanced oxidation processes but also in atmospheric and biological chemistries. Other exciting 226 results have been obtained by Lee et al,<sup>78</sup> who have observed spontaneous formation of hydrogen 227 peroxide in sprayed water microdroplets. The authors have considered and analyzed several 228 possible mechanisms and concluded that the process occurs at or near the interface, where the 229 strong intrinsic electric field is enough to ionize hydroxyl anions, generating hydroxyl radicals 230 that then recombine to form  $H_2O_2$ . Although the mechanism is not fully understood, the result is 231 quite significant because H<sub>2</sub>O<sub>2</sub> has great importance in biomedical and industrial applications, 232 and it is also a key compound in the atmosphere due to its oxidative capacity.<sup>79</sup> The results of 233 Lee et al<sup>78</sup> have been supported by the work of Gao et al<sup>80</sup> showing that Dakin and Baeyer-234 Villiger oxidation reactions proceed in water microdroplets without the addition of any peroxides 235 and acid or base catalysts, as usually required.<sup>80</sup> 236

It is worth reminding that reactions in microdroplets and electrospray-mass spectrometry techniques are not only interesting from the point of view of analysis and synthesis, as

<sup>239</sup> mentioned above, but also to study a wide variety of problems in atmospheric,<sup>35,43,45</sup> biomedical<sup>81-</sup> <sup>83</sup> or other domains in which aqueous interfaces play a central role. Moreover, possible scale-up <sup>241</sup> of microdroplet chemical synthesis by heated ultrasonic nebulization opens interesting industrial <sup>242</sup> perspectives.<sup>84</sup> Finally, it must be noted here that experiments with electrospray techniques and <sup>243</sup> their interpretation as purely interfacial reactions have raised certain controversy in the literature <sup>244</sup> because of the possible influence of ions<sup>85</sup> and gas phase chemistry.<sup>86</sup> Two illustrative examples <sup>245</sup> of the controversy will be commented on below.

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**Prebiotic chemistry.** The role played by aqueous interfaces on the chemical mechanisms that led 247 to the origin of life on Earth has received considerable interest in recent years. It is explained by 248 the fact that compartmentalization, as well as the emergence of genetic materials, is considered 249 to be a key prerequisite in the long journey towards protocells capable of growth, division and 250 Darwinian evolution.<sup>87</sup> Colloidal systems, resulting from self-organization of amphiphilic 251 molecules in aqueous environments, provide such suitable compartments in which complex 252 chemical reactions could have taken place in the prebiotic era. In particular, vesicles formed in 253 bulk waters (lakes, rivers) have attracted considerable attention because the amphiphilic bilayer 254 that separates the aqueous interior from the exterior media in these structures bears a 255 resemblance to cell membranes.<sup>17,18,88-90</sup> In such confined volumes, molecular crowding increases 256 the probability of reactive encounters between chemical species, and at the same time, limits the 257 diffusion of products. Hence, the synthesis of complex biomolecules required for the 258 development of primitive living organisms is strongly favored compared to similar reactions in 259 bulk.<sup>73</sup> Though molecular crowding is not the only important feature controlling the chemical 260 reactivity inside the vesicle, and several works have emphasized the importance of the interface 261 in terms of molecular alignment, electric charge, pH, etc. For instance, experiments have shown 262 that the polycondensation of aminoacids and peptides is assisted by the lipidic bilayer, not only 263

as a favorable environment for the reaction to take place,<sup>91,92</sup> but also as an active acid-base
 catalyst.<sup>93</sup>

Other possible prebiotic chemical reactors are the inverted micelles structures of atmospheric 266 organic aerosols.<sup>94-96</sup> In these systems, the organic content may be quite high (up to 50%), and 267 there is direct evidence that palmitic and other fatty acids form the organic film on the exterior of 268 marine aerosols.<sup>97,98</sup> Interestingly, it has been shown that the size of bacteria and viruses can be 269 predicted from atmospheric aerosols by combining atmospheric aerodynamics and gravity 270 equations.<sup>95,96</sup> Here too, the role of the interface has been emphasized and supported by different 271 studies. Using infrared reflection absorption spectroscopy (IRRAS) and Langmuir trough 272 methods, Griffith and Vaida<sup>99</sup> have observed peptide-bond formation in the Leucine ethyl ester 273 condensation process in presence of Cu<sup>2+</sup> ions at the air-water interface. Such condensation 274 reactions are thermodynamically and kinetically unfavorable in aqueous environments, but at the 275 air-water interface, there's evidence suggesting the spontaneous peptide bond formation. The 276 interaction of Cu<sup>2+</sup> ions with the amine group of the Leucine ester might play a role by inducing 277 an orientational change. Note that the probe depth of the IRRAS technique can be as large as 1-2 278  $\mu$  m, i. e. much larger than other interface sensitive techniques such as SFG, for instance, but 279 this probe depth was considered suitable for the reactive region of interest.<sup>99</sup> 280

Experiments in microdroplets have been reported as well. Lee et al<sup>100</sup> have observed 281 spontaneous reduction of several organic molecules without assistance of reducing agents, 282 catalysts of external charges, which could represent an essential reduction route in prebiotic 283 conditions. The mechanism is unclear but might involve the oxidation of OH<sup>-</sup> at the droplet 284 surface, likewise in the spontaneous formation of  $H_2O_2$  described above.<sup>78</sup> Nam et al<sup>101</sup> have 285 studied aqueous microdroplets containing a mixture of sugars and phosphoric acid, and observed 286 287 that phosphorylation proceeds spontaneously in such conditions. The effect has been ascribed to a cancellation of the entropic barrier when the process occurs at the surface of the microdroplet, 288 whereas such barrier prevents the uncatalyzed reaction to proceed in bulk solution. Nam et al 289

have also reported the synthesis of uridine<sup>101</sup> and other ribonucleosides<sup>20</sup> in an aqueous 290 microdroplet containing D-ribose, phosphoric acid, and a nucleobase. As an example of the 291 controversy surrounding some results obtained with electrospray techniques, Jacobs et al<sup>86</sup> have 292 reported different conclusions for the reaction between sugars and phosphoric acid. The authors 293 have used an experimental setup in which droplet generation is separated from ionization, so that 294 they have been able to analyze different possible sources of rate acceleration. They have 295 concluded that part of the products could originate from gas-phase chemistry, which in some 296 cases may complicate the interpretation of rate acceleration in droplets generated by 297 electrosprays or its variants. 298

The preceding results exemplify the role aerosols and microdroplets could have played for the generation of chemical complexity in prebiotic chemistry,<sup>19</sup> which could have also involved sunlight-driven processes.<sup>102</sup>

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**Reactions at organized molecular interfaces.** Quite a diverse variety of processes can be placed 303 in this category that includes systems possessing an organized amphiphilic interface with ability 304 for molecular recognition, possibly including a binding site, and compartments that can host 305 chemical reactions. Of course, some systems described above belong to this category, such as the 306 vesicles hosting prebiotic chemical reactions or the atmospheric organic aerosols structured as 307 inverted-micelles. Chemical reactions in biological membranes could be included in this class of 308 interfacial processes too.<sup>103</sup> Nevertheless, the focus here is on synthetic reactions in water that 309 mimic the functioning of enzymatic catalysis in biology, which have particular interest in the 310 field of Green Chemistry. The term "artificial enzymes" was coined by Breslow,<sup>104,105</sup> who 311 introduced the use of functionalized macromolecules, mainly cyclodextrins, as water-soluble 312 catalysts that can host a non-polar reactant guest in a hydrophobic cavity. The design of enzyme 313 mimics or "chemzymes" is a field of intense research,<sup>106-109</sup> which has turned into the more 314 general one of "molecular reaction vessels". Antibody catalysts or "abzymes",<sup>110</sup> functionalized 315

nanomaterials or "nanozymes",<sup>111,112</sup> dendrimers,<sup>113,114</sup> micellar<sup>115</sup> and other disperse interface-rich
structures (polyelectrolyte solutions, microemulsions, vesicles, ..),<sup>18</sup> as well as enzymes confined
in small-volume environments<sup>17</sup> have been considered in detail previously. Therefore they will
not be further described here.

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## **Solvation at the water surface**

The hydrogen-bond network formed by water molecules in the liquid state, and its 322 cooperative character, confer this environment its unique properties. At the surface of water, the 323 network is inevitably disrupted and the physical and chemical properties of molecules lying there 324 (hydrogen-bonds, dipole moment, acidity, etc) differ from those in the bulk. To address how 325 these changes affect chemical reactivity is a complex issue that requires a close examination of 326 the structure and properties of the water surface. This section reviews some theoretical and 327 experimental aspects on structural (hydrogen-bonding), chemical (acid/base) and solvation 328 (dynamics, thermodynamics) properties of the liquid water-vapor interface. 329

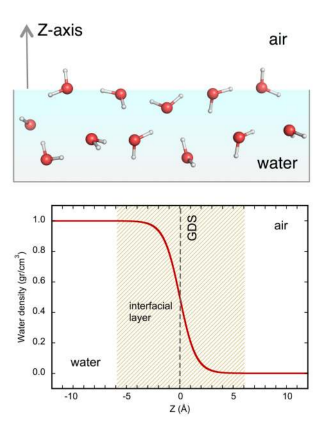
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Chemical properties of the water surface. The structure of the water surface has been a subject 331 of intense debate for many years.<sup>116</sup> Most of the current knowledge comes from SFG vibrational 332 spectroscopy and from calculations. Du et al<sup>117</sup> reported the first SFG spectrum of the liquid 333 water-vapor interface and the authors concluded that about 20% of water molecules display a 334 dangling bond, the free OH bond that is projected into the vapor phase. This result predicted by 335 pioneer MD simulations<sup>118,119</sup> was subsequently confirmed and rationalized by classical SFG and 336 ab initio simulations.<sup>120-122</sup> Further theoretical studies support a 2D H-bond network of interfacial 337 waters (the water "skin") with oscillating OH bonds around a plane parallel to the instantaneous 338 surface.<sup>123</sup> A schematic view of the water surface is displayed in Figure 2, which also shows a 339 typical density profile from classical MD simulations. The thickness  $\delta$  of the air-water interface 340 is usually deduced from the density profile  $\rho(z)$  by fitting a function: 341

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$$\rho(z) = \frac{\rho_0}{2} \left(1 + \tanh\left(\frac{Z - Z_G}{\delta}\right)\right) \tag{1}$$

where  $\rho_o$  is the bulk density,  $Z_G$  is the position of the Gibbs-dividing-surface (Z at which  $\rho(z) = \frac{\rho_o}{2}$ ). Values of  $\delta$  can change significantly with the theoretical model<sup>124-127</sup> but common values are 10-15 Å at 300K.

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Figure 2. Schematic structure of the water surface showing free OH groups pointing towards the air layer, and
 typical density profile of water at the air-water interface from MD simulations. The vertical dashed-line indicates the
 Gibbs-dividing-surface (GDS) where the density is half of the bulk density. The width of the interface layer depends
 on simulation models but is typically in the range 10-15 Å.

353 The dynamics of water reorientation has been a broadly studied subject, both in bulk water

<sup>354</sup> (see for instance Laage and Hynes<sup>128</sup>) and at interfaces.<sup>127,129-136</sup> Simulations<sup>130</sup> and experiments

using femtosecond pump/probe vibrational sum-frequency spectroscopy<sup>129</sup> have shown that

reorientation of free OH groups in the liquid-vapor interface takes place on a subpicosecond time

scale, i. e. several times faster than in bulk. In contrast, simulations by Verde et al<sup>127</sup> have shown

that reorientation of bonded OH groups happens at a rate similar to that of bulk water.

Particularly relevant to chemical reactivity is the acid/base character of the water surface, i.e. 359 its ability to donate or accept protons, an issue that remains incompletely elucidated and has 360 raised intense controversies in the literature. This issue is connected, on the other hand, to the 361 properties of water in nanoconfined environments such as inversed micelles, a topic that we will 362 not develop here but which has attracted a lot of attention, while it remains incompletely 363 understood (see for instance the works by Levinger and coworkers<sup>137,138</sup>). Interestingly, 364 experiments and calculations reveal unforeseen acid/base behavior of aqueous interfaces. For 365 instance, HCl is fully dissociated at the interface but HNO<sub>3</sub> is essentially in its molecular 366 form,<sup>139-141</sup> unless ions are present,<sup>142</sup> and HCOOH dissociates faster at the interface than in the 367 bulk.<sup>143</sup> Vibrational spectroscopic studies of the ionization state of the L-phenylalanine 368 aminoacid indicated a decrease of the pK<sub>a</sub> of its polar groups at the ait-water interface.<sup>144</sup> 369 Depending on experiments and calculations (see for instance<sup>145-157</sup>), apparent opposite 370 conclusions have been deduced for the interface affinity of hydronium and hydroxide ions and 371 their spatial distribution, though most recent SFG experiments on D<sub>2</sub>O-air interface indicate that 372 the hydrated proton is much more surface-active than the hydroxide anions.<sup>158</sup> Discordant results 373 are probably explained by inherent difficulties in interpreting experiments, and by the limited 374 accuracy of numerical simulations, besides the fact that results from different methods may 375 correspond to different probing depths. Electrospray mass spectrometry experiments by Colussi 376 and coworkers<sup>35,148,149,159</sup> led them to conclude that (in their own words):<sup>160</sup> "(1) water is more 377 extensively self-ionized at the surface than in the bulk, and (2) interfacial  $H_3O^+$  is a stronger acid 378 (a "superacid") and interfacial OH a stronger base than their bulk counterparts likely due to 379 limited hydration". According to these authors, the acidic or basic behavior of the surface of 380 water would rather be interpreted in terms of the availability of proton or hydroxide ions at a 381 given pH, with pH~3 being neutral (instead of 7 as in bulk).<sup>148,149,161</sup> An enhanced autolysis of 382 water at hydrophobic interfaces due to the strong local electric-field gradient was already 383 reported by Beattie<sup>162</sup> (with an isoelectric point around pH 4) trying to explain the contrasting 384

observed electro-osmotic properties of microfluidic channels. On the theoretical side, water self-385 ionization has been found to be more favorable in water clusters of 20<sup>163</sup> or 21<sup>164</sup> water 386 molecules, compared to bulk solution. This unexpected result is probably a consequence of the 387 topology of the hydrogen-bond network, and could serve as a clue for elucidating the acid/base 388 properties of water in extended aqueous interfaces. A complete survey and a comparative 389 analysis of experimental and theoretical data before 2016 can be found in the review of Agmon 390 et al<sup>165</sup> and in the paper by Saykally.<sup>166</sup> As an example of the ongoing discussion, one can refer to 391 the experiments on isoprene oligomerization in aqueous electrosprays and mildly acidic water by 392 Enami et al<sup>35</sup> that we have mentioned above. Gallo et al<sup>85</sup> have carried out another study of this 393 system by comparing the reactivity in electrosprays and isoprene-water emulsions with adjusted 394 pH, in an attempt to differentiate between pure interfacial effects and effects due to the 395 conditions characterizing the electrosprays experiments (charge separation, concentration of 396 reactants). According to these authors, the absence of chemical reactions in emulsions suggests 397 that the high-voltages in the electrosprays play a key role, leading to charge-separation that 398 facilitates the formation of partially hydrated, highly-reactive hydronium ions, that then catalyze 399 the process. The author's conclusion was supported by theoretical calculations comparing the 400 reactivity of  $(H_3O^+)(H_2O)_n$  clusters of different size. Further works by Colussi and Enami<sup>161</sup> and 401 Gallo et al<sup>167</sup> have discussed the effects that the partial solubility (milimolar level) of isoprene in 402 water might have on the fate of reactions in the case of isoprene-water emulsions. It is worth 403 mentioning in this respect the work by Butler et al,<sup>60</sup> who used the *endo/exo* preference in 404 Huisgen cycloadditions to classify reactions (in-water vs on-water) as a function of the 405 hydrophobicity of one of the reactants, i.e. its water solubility: on-water reactions do not display 406 increased *endo*-effects relative to organic solvents, in contrast to in-water reactions. To sum up, 407 the case of isoprene oligomerization emphasizes the difficulties to derive definite conclusions 408 about interface effects on reactivity and the presence of on-water catalysis. The combined use of 409

multiple analytical platforms and of elaborated numerical simulations will be most useful to test
 different hypotheses and get more insights in this field.

Finally, one should note that ab initio MD simulations of the water liquid-vapor interface have highlighted the augmented reactivity with respect to excess protons and electrons by an analysis of the HOMO and LUMO energies at interfacial layers.<sup>122</sup>

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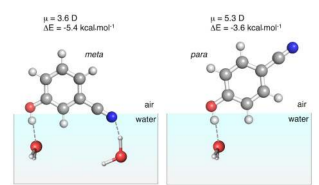
The "polarity" of the water surface. Solvent polarity has been a widely used concept in 416 Chemistry to rationalize solvation phenomena.<sup>28</sup> Following the « like dissolves like » 417 principle, polar solvents are likely to dissolve polar compounds and favor their most polar 418 conformations. Non-polar solvents, on their side, are likely to dissolve non-polar compounds. 419 Though a precise definition of "solvent polarity" is not straightforward, the use of empirical 420 parameters derived from linear Gibbs energy relationships has been very popular in Organic 421 Chemistry.<sup>28</sup> In Computational Chemistry, polarizable continuum models based (essentially) on 422 the static dielectric constant of the solvent have been very successful to study processes in bulk 423 solution,<sup>168</sup> and more recently at interfaces as well.<sup>169-171</sup> 424

However, with regard to the solvation power of aqueous interfaces, the use of the concept of 425 "solvent polarity" entails some difficulties. In fact, experimental attempts to characterize the 426 polarity of aqueous interfaces using second-harmonic spectroscopy have led to conflicting 427 results. On one hand, Eisenthal and coworkers<sup>172</sup> tried to derive an interface polarity scale 428  $(E_{\tau}(30))$  for a betaine dye at several water interfaces. They deduced a simple relationship 429 according to which the polarity of a liquid interface is the arithmetic average of the polarity of 430 the two bulk phases, pointing at a dominant effect of long-range solute-solvent interactions. For 431 the air-water interface, the polarity would be close to that of a low polar solvent. On the other 432 hand, further measurements with coumarin derivatives<sup>173</sup> and other dyes<sup>174,175</sup> have demonstrated 433 the limitations of the "arithmetic average" rule, claiming that the polarity of aqueous interfaces is 434 not a well-defined concept. The apparent "polarity" of the interface strongly depends on solute's 435

structure since subtle modifications of the later (stereochemistry, hydrophobic groups) can
produce significant changes on the former. This is because the solute's position and orientation
relative to the interfacial boundary rely on its structure, and so does the water response.

Theoretical studies can clarify the issues in the definition of interface polarity. Classical and 439 first-principles MD simulations of glyoxal (O=CH-HC=O) have shown that water interfaces 440 selectively stabilize the polar cis-conformer (the two polar C=O bonds pointing in the same 441 direction) over the apolar trans-conformer (C=O bonds pointing in opposite directions).<sup>176,177</sup> 442 This result can be explained by the fact that both, stereochemistry and polarity favor the 443 interaction of the cis-isomer with the interface. Stereochemistry and polarity, however, do not 444 always go in the same direction, as in the case of *meta*- and *para*-cyanophenol isomers. Ab initio 445 calculations using a dielectric model<sup>171</sup> show that despite its lower polarity, the *meta*-isomer has 446 a higher interface affinity because, in this case, but not in the case of the para-isomer, the -CN 447 and -OH groups can simultaneously interact with the aqueous layer (Figure 3). 448

Finally, in Figure 4, we illustrate the differences between the bulk-water and air-water 449 interface reaction-field potentials, i. e. the electrostatic potentials created by the polarized water 450 medium, in the case of methanol obtained by MD simulations.<sup>178</sup> Methanol is an important 451 atmospheric compound and its air-water interface affinity and structure have been thoroughly 452 described by SFG spectroscopy measurements and theoretical simulations.<sup>5,178</sup> As shown in 453 Figure 4, , there are topological differences between the two potentials that do not correspond to 454 those that would be expected for two media differing simply by their "polarity" gradation. The 455 potentials around the OH groups are indeed quite similar, while a large difference appears 456 around the CH<sub>3</sub>-group, which is of course the consequence of a preferred orientation of methanol 457 at the interface. Roughly, the CH<sub>3</sub>-group points towards the air layer (and is basically not 458 solvated), and the OH-group points towards the water layer (and has an almost complete 459 hydration shell), although the details of the solvation dynamics discussed below draw a slightly 460 more complicated picture. 461



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**Figure 3**. Schematic view of the *meta-* and *para-*isomers of cyanophenol adsorbed at the air-water interface showing the expected hydrogen-bonds with water molecules. The plotted values for the gas-phase dipole moment ( $\mu$ ) and for the electrostatic interaction energy with the interface ( $\Delta$ E) have been obtained using quantum chemistry calculations and a simple dielectric model of the air-water interface.<sup>171</sup> The values reveal that despite a lower dipole moment of the *meta-*isomer with respect to the *para-*isomer, its electrostatic interaction energy is higher (in absolute value) owing to the possible simultaneous contact of both polar groups with the water surface.

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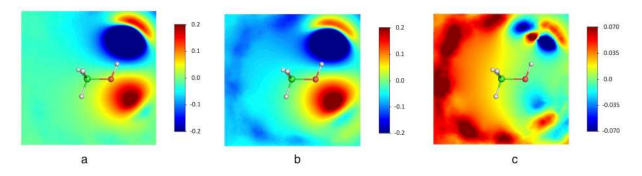
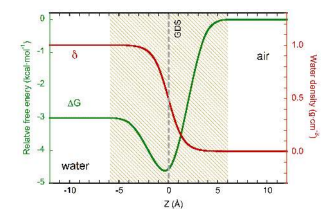


Figure 4. Calculated electrostatic potential (atomic units) created by the solvent water molecules surrounding 473 methanol. The graphs correspond to time averages of the potential obtained from QM/MM Molecular Dynamics 474 simulations.<sup>178</sup> The surfaces are displayed in an arbitrary methanol-fixed coordinate system but in the simulation 475 there are no constraints imposed to the methanol or water molecules, which are flexible to vibrate, rotate and 476 translate. Graph (a) corresponds to the calculation at the air-water interface: the left green part reveals that at the 477 478 interface, the average potential around the methyl group is close to zero, as corresponds to the fact that this group is most of the time pointing towards the air layer. Graph (b) corresponds to the calculation in bulk water: as shown, the 479 potential in the right part (around the OH group) is very similar to the potential obtained at the interface, but the 480 potential around the methyl group (left part, blue-green surface) is significantly different. In the bulk, water 481 molecules around the methyl group undergo orientational polarization, and may form weak hydrogen-bonds with the 482 methyl H atoms. Graph (c) displays the difference between the two potentials (interface - bulk): it confirms that the 483 most relevant disparity holds for the region around the CH<sub>3</sub> hydrophobic group. 484

486	Thermodynamics and dynamics of solvation. The energetics of solvation at aqueous interfaces
487	is a vast subject with extensive literature and a multitude of facets. The specific topics dealt with
488	here are those that could be considered most relevant for understanding chemical reactivity of
489	organic compounds at air-water interfaces. In the field of atmospheric chemistry, several reviews
490	have already been published describing the uptake and accommodation processes, the energetics

of interface adsorption, and the experimental techniques.<sup>4,29,33</sup> MD simulations have allowed, on 491 the other hand, to obtain the potential of mean force for the adsorption and accommodation 492 processes of many chemical species. One of the most remarkable findings of these studies (see 493 for instance<sup>32,33,124,125,179-186</sup>) has been the significant interface affinity, not only of hydrophobic or 494 amphiphilic organic molecules, which is an expected result, but of small polar systems and even 495 ions as well. An archetypal free energy profile for moving a neutral water-soluble compound 496 from the gas-phase to bulk water across the air-water interface is shown in Figure 5 (the 497 solvation of ions is considered in deeper detail below). The free energy decreases from air to 498 bulk with a minimum at the interfacial layer. These profiles are useful to obtain Henry's 499 constants and surface excess properties.<sup>182</sup> Theoretical analysis<sup>181</sup> of the solvation of organic 500 molecules in water droplets has revealed that the surface preference is principally due to 501 enthalpic effects. Namely, the total water-water interaction energy is more negative when the 502 solute is at the surface of the droplet because, when it is in the bulk, some water-water hydrogen-503 bonds are disrupted. Entropic effects further enhance the surface preference when the system 504 bears large apolar groups (e.g. 4-5 carbon atoms or longer hydrophobic chains). 505

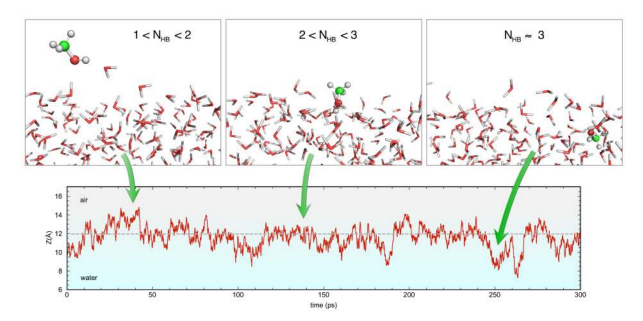
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**Figure 5**. Schematic plot for the relative free-energy profile ( $\Delta G$ , green) for a neutral (water soluble) solute crossing the air-liquid water interface. The density profile of water ( $\delta$ , red) is also shown. The free energy decreases from the air layer (right part of the figure) to the interface, where it displays a minimum (hatched area) close to the Gibbsdividing-surface (GDS, Z=0 here), then increases from the interface to the bulk. Note that the width of the interface layer (hatched area) is about 1nm. Depending on the solute's structure and on its hydrophilicity/hydrophobicity character, the free-energy profile can display substantial differences, e.g. a free-energy maximum can occur between the interfacial layer and the bulk water, and the sign of the relative air-bulk water free-energy can be reversed.

The molecular dynamics of solutes adsorbed at the air-water interface displays two 516 fundamental differences with respect to the bulk. First, axial oscillations of the solute's position 517 across the average interface plane may be quite large and this implies concomitant fluctuations 518 of the instantaneous hydration shell. This is illustrated in Figure 6 for methanol at the air-water 519 interface.<sup>178</sup> Second, due to the asymmetry of the interface and the existence of preferred 520 orientations of the solute, the interface orientational dynamics differs in general from the bulk. 521 Reorientational relaxation at the interface can be characterized by time- and polarization-522 resolved pump-probe SFG spectroscopy<sup>187</sup> or by MD simulations. Calculation of the rotational 523 autocorrelation functions of the methane derivatives MeCl, MeCN, and MeOH, which are 524 important organic compounds in the troposphere, shows that the reorientation decay times 525 increase with their hydrogen-bonding capability, i. e. with the strength of their interface 526 anchoring.178 527

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**Figure 6.** QM/MM MD simulation of methanol at the air-water interface.<sup>178</sup> The lower panel shows the fluctuations of the solute's axial position (Z-axis) with respect to the average interface plane (Z=12Å). The snapshots in the upper panel illustrate different situations in which the methanol molecule, depending on its relative position with respect to the interface (air or water layers) is more or less hydrated; the average number of methanol-water hydrogen-bonds (N<sub>HB</sub>) in different cases is indicated.

Interface affinity of ions. Ions in the outermost interface layers are more easily available to 536 catalyze chemical reactions (e.g. on sea salt-aerosols) and it is therefore crucial to set-up a scale 537 of interface affinity values. Beyond that, interface affinities are valuable to establish 538 kosmotropic/chaotropic scales, predict the surface tension of electrolyte solutions or explain the 539 Hofmeister series.<sup>188,189</sup> In the classical view of electrolytes that considers the interface as an 540 abrupt discontinuity between two dielectric continuum media,<sup>190</sup> the air-water interface is devoid 541 of ions. In such models, the point-charge q in a dielectric with dielectric constant  $\epsilon_1$  (water) 542 interacts with its image charge  $q' = q(\epsilon_1 - \epsilon_2)/(\epsilon_1 + \epsilon_2)$  in the dielectric with constant  $\epsilon_2$  (air), 543 and therefore is repelled from the interface ( $\epsilon_2 < \epsilon_1$ ). The divergence found in this model for 544 ions approaching the interface can be untangled for finite radii ions.<sup>191</sup> Although a full 545 understanding of the topic is still lacking, many endeavors have been made to get beyond the 546 classical view. The macroscopic view from surface tension and electrostatic potential 547 measurements has been supplemented by data from interface-sensitive spectroscopic techniques 548 such as SFG, by elaborated dielectric continuum theories and MD simulations, providing new 549 insights.<sup>179,188,189,192-200</sup> 550

Hard non-polarizable ions (such as F<sup>-</sup> or the alkali cations), and multiply charged ions (such as 551 sulfate) behave classically and are repelled from the interface, but large polarizable anions (such 552 as I or Br and to a lower extent Cl), display a propensity for the air-water interface. The case of 553 hydronium discussed above is an exception, and its interface affinity results from specific 554 hydrogen-bonding properties. Though it goes beyond the limits of the present review, the 555 stability of the solvated electron at aqueous interfaces has also been studied<sup>201-203</sup> because of 556 potential implications in numerous chemical processes (radiation chemistry, electron-transfer, 557 redox and electrochemical reactions, etc). 558

The interface affinity of ions has been explained (at least qualitatively) by a favorable balance between electrostatic and cavitation energies.<sup>199,204,205</sup> The later represents the energy cost required to disrupt water-water interactions in the medium in order to create a hole where the ion

is placed. The cavitation energy drops when the ion moves from bulk to the interface, and for 562 bulky soft ions it can overcompensate the loss of favorable ion-water electrostatic interactions. In 563 such a case, the ion stabilizes at the interface. The role of anion polarizability has been 564 emphasized<sup>179,195</sup> and though correlation with interface affinity is not always apparent,<sup>206</sup> this term 565 must be taken into account for a quantitative description of the adsorption energetics, as recently 566 reported for aqueous solutions of ClO<sub>4</sub><sup>-</sup> using SFG spectroscopy.<sup>205</sup> Under the effect of the local 567 electric field (the permanent field due to the asymmetry of the interface and the ion-induced 568 reaction-field), the electronic cloud of large polarizable anions is distorted and the induced 569 dipole moment contributes to enhancing the solvation of the ion at the interface.<sup>192</sup> Solvation 570 dispersion-forces may also influence interfacial adsorption,<sup>199,204</sup> specially at oil-water 571 interfaces.188 572

Cations are dragged to the interface from the bulk through the electrostatic interactions with 573 the anions and cumulate in nearby inner layers,<sup>192</sup> although their distribution is quite sensitive to 574 the type of counterions present.<sup>189</sup> Anions and cations interact differently with water, and 575 according to Levin and dos Santos,<sup>188</sup> alkali cations are repelled from the interface because they 576 are strongly hydrated, while anions may behave either as kosmotropes or chaotropes. The 577 distribution of anions and cations near the air-water interface is also influenced by the 578 electrostatic potential originated by the orientation of water molecules at the interface, although 579 the role of this surface potential still remains unclear.<sup>207</sup> Indeed, classical calculations using 580 point-charge force-fields predict the air layer to be more electropositive than water (in 581 congruence with the image of dangling protons pointing towards the air layer), while explicit 582 treatment of the electronic cloud in ab initio simulations predicts the opposite trend.<sup>179,188,208</sup> Thus, 583 the anionic adsorption predicted with polarizable force-fields is probably overestimated.<sup>188</sup> The 584 adsorption energy of ions has been decomposed in entalphic and entropic terms in some 585 cases,<sup>179</sup> and the simulations by Caleman et al<sup>209</sup> and Otten et al<sup>198</sup> concluded that adsorption of 586