1	A better understanding of hydroxyl radical photochemical sources in cloud							
2	waters collected at the puy de Dôme station : Experimental versus modeled							
3	3 formation rates							
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21								
22	Abstract. The oxidative capacity of the cloud aqueous phase is investigated during three							
23	field campaigns from 2013 to 2014 at the top of the puy de Dôme station (PUY) in France.							
24	Forty-one cloud samples are collected, and the corresponding air masses are classified as							
25	highly marine, marine and continental. Hydroxyl radical (HO <sup>•</sup> ) formation rates ( $R_{HO^{•}}^{f}$ ) are							
26	determined using a photochemical setup (Xenon lamp that can reproduce the solar spectrum)							
27	and a chemical probe coupled with spectroscopic analysis that can trap all of the generated							
28	radicals for each sample. Using this method, the obtained values correspond to the total							
29	formation of HO <sup>•</sup> without its chemical sinks. These formation rates are correlated with the							
30	concentrations of the naturally occurring sources of HO <sup>•</sup> , including hydrogen peroxide,							
31	nitrite, nitrate and iron. The total hydroxyl radical formation rates are measured as ranging							
32	from approximately $2 \times 10^{-11}$ to $4 \times 10^{-10}$ M s <sup>-1</sup> , and the hydroxyl radical quantum yield							
33	formation ( $\Phi_{HO}$ ) is estimated between 10 <sup>-4</sup> and 10 <sup>-2</sup> . Experimental values are compared							
34	with modeled formation rates calculated by the model of multiphase cloud chemistry							

35 (M2C2), considering only the chemical sources of the hydroxyl radicals. The comparison 36 between the experimental and the modeled results suggests that the photoreactivity of the 37 iron species as a source of  $HO^{\bullet}$  is overestimated by the model, and  $H_2O_2$  photolysis 38 represents the most important source of this radical (between 70 and 99%) for the cloud 39 water sampled at the PUY station (primarily marine and continental).

40

### 41 Keywords

42 cloud water, photochemistry, hydroxyl radical, *in situ* measurements, cloud chemistry model

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### 44 **1** Introduction

In the atmosphere, many trace gases are transformed by the hydroxyl radical (HO<sup>•</sup>), which is 45 considered the most efficient environmental oxidant (e.g., Seinfeld and Pandis, 2006). 46 Evaluating the production of this short-lived species is crucial because it determines the fate 47 of many chemical compounds. In atmospheric water drops and aqueous particles, the 48 hydroxyl radical also controls the fate of inorganic and organic species (Herrmann et al., 49 2010). The HO<sup>•</sup> - mediated oxidation of organic compounds in the aqueous phase can lead to 50 51 the formation of shorter but often multifunctional organic species and, ultimately, to complete mineralization (Charbouillot et al., 2012). Complex chemical reactions catalyzed by HO<sup>•</sup> can 52 also occur in the aqueous phase forming accretion products such as oligomers (Altieri et al., 53 54 2008; Carlton et al., 2007; Perri et al., 2009; Tan et al., 2011; Ervens and Volkamer, 2010; De 55 Haan et al., 2009). These alternative chemical pathways are efficient processes to convert organic compounds into Secondary Organic Aerosols (SOAs) (Ervens et al., 2011). 56

57 The sources of hydroxyl radicals in the aqueous phase strongly differ from those in the gas phase because of the presence of ionic species and metal ions. Aqueous phase reactants that 58 produce HO<sup>•</sup> present high concentrations in water drops and aqueous particles, likely 59 enhancing the HO<sup>•</sup> photochemical production in the condensed phase. This radical can be 60 generated in the aqueous phase by direct photolysis of hydrogen peroxide  $(H_2O_2)$  (Herrmann 61 et al., 2010; Yu and Barker, 2003), iron complexes (Deguillaume et al., 2005), nitrate (NO<sub>3</sub><sup>-</sup>) 62 (Zellner et al., 1990) and nitrite ions (NO<sub>2</sub>) (Zafiriou and Bonneau, 1987). The other 63 significant source of HO<sup>•</sup> in cloud water is the uptake from the gas phase (Arakaki and Faust, 64 1998). The relative importance of the different hydroxyl radical sources depends on the 65 chemical composition of the aqueous phase, which is also strongly variable (Deguillaume et 66 67 al., 2014). HO<sup>•</sup> is further scavenged in the aqueous phase, primarily by dissolved organic

compounds. Evaluation of this sink is difficult because the dissolved organic matter isdiverse, complex and poorly characterized (Herckes et al., 2013).

Uncertainties in HO<sup>•</sup> sinks and sources make its concentrations in atmospheric water highly 70 difficult to estimate. For this estimation, models describing the multiphase cloud chemistry 71 72 have been developed and have considered the reactivity in the gas and aqueous phases along with the mass transfer between the two phases (Ervens et al., 2014; Long et al., 2013; Tilgner 73 74 and Herrmann, 2010). These numerical tools allow the estimation of the steady-state concentration of  $HO^{\bullet}$  ([ $HO^{\bullet}$ ]<sub>ss</sub>), which is a crucial quantity to understand the fate of 75 atmospheric pollutants (Arakaki et al., 2013). The range of the maximal HO<sup>•</sup> concentration 76 varies from 10<sup>-16</sup> to 10<sup>-12</sup> M, depending on the "chemical scenario" (*i.e.*, emission/deposition 77 and the initial chemical conditions) used in the modeling study. The amounts of organic 78 matter and iron are key parameters controlling the [HO<sup>•</sup>]<sub>ss</sub>. These models are expected to 79 underestimate the radical sinks because organic scavengers cannot be exhaustively described 80 81 in the aqueous chemical mechanism (Arakaki et al., 2013).

In this study, we propose the investigation of the hydroxyl radical formation in real cloud 82 83 water sampled at the puy de Dôme mountain (France). The hydroxyl radical formation rate is quantified for 36 cloud water samples collected during 3 field campaigns (2013-2014). 84 85 Because the main photochemical sources (hydrogen peroxide, iron, nitrite and nitrate) are also quantified, we can calculate their relative contributions to the production of the hydroxyl 86 87 radicals. For this purpose, the contribution to the hydroxyl radical formation rate of more concentrated inorganic photochemical sources is investigated separately in synthetic solution. 88 89 In parallel, the model of multiphase cloud chemistry (M2C2) is used to simulate HO<sup>•</sup> 90 formation rates. This model considers explicit aqueous chemical mechanisms, and a "simplified" version of the model is used to reproduce the bulk water irradiation experiments 91 92 (lamp spectrum) under variable physico-chemical conditions (pH, initial concentrations of HO<sup>•</sup> sources) corresponding to the cloud water samples. The comparison between the 93 modeled and experimental HO<sup>•</sup> production rates facilitates quantification of the various HO<sup>•</sup> 94 sources and enables validation of the model to reproduce the oxidative capacity of the 95 96 atmospheric aqueous phase.

- 97
- 98 2 Materials and Methods
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- 100 2.1 Chemicals

Hydrogen peroxide (30% in water, not stabilized), sodium nitrate (purity >99%) and ferrozine
(purity >97%) were obtained from Fluka, while sodium nitrite (purity >98%) and terephthalic
disodium salt (purity >99%) were purchased from ProLabo and Alfa Aesar, respectively. All
of the other chemicals (purity reagent grade) used for the analysis were obtained from SigmaAldrich.

107 Solutions are prepared with deionized ultra-pure aerated milli-Q water from Millipore 108 (resistivity = 18.2 M $\Omega$  cm) under a laminar flux hood. Moreover, glass containers and 109 injection material are washed three times with ultra-pure water before use. If necessary, the 110 pH values are adjusted with perchloric acid (1 N) and NaOH (1 N) using a JENWAY 3310 111 pH-meter within ± 0.01 pH unit. All of the solutions are stored under dark conditions, and the 112 final preparations are performed in a room equipped with a sodium lamp (589 nm emission).

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- 114 2.2 Cloud water sampling
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116 Cloud water is sampled at the puy de Dôme (PUY) station (48°N, 2°E; 1465 m a.s.l.) in the 117 Massif Central region (France). Three campaigns occurred during autumn 2013 from October 118 14<sup>th</sup> to November 6<sup>th</sup>, during spring and autumn 2014 from March 22<sup>nd</sup> to April 5<sup>th</sup> and from 119 November 4<sup>th</sup> to 19<sup>th</sup>. During these periods, the station was primarily located in the free 120 troposphere; thus, the air masses from various origins were not influenced by the local 121 pollution (Hervo, 2013).

122 The cloud droplet sampling is performed by a one stage cloud droplet impactor (Deguillaume 123 et al., 2014). With the air flux used, the lower limit of the aerodynamic diameter is 124 approximately 7  $\mu$ m (Brantner et al., 1994). The impactor used for this study is constructed of 125 stainless steel and aluminum, and cloud droplets are collected by impaction onto a rectangular 126 aluminum plate with an average sampling time of two hours. Cloud water samples are filtered 127 using a 0.45  $\mu$ m PTFE filter within 10 minutes after sampling to eliminate all of the 128 microorganisms and particles that can interfere with the spectroscopic analysis.

Measurements performed immediately after cloud collection are conductivity, redox potential, pH, UV-visible spectroscopy, H<sub>2</sub>O<sub>2</sub> and iron concentrations. Ion chromatography (IC), total organic carbon (TOC), and nitrite analysis are determined less than 24 hours after sampling. At each stage, sampling and analyses are performed with the greatest precaution to minimize all possible external contaminations, and the solutions are stored at 277 K under dark 134 conditions. Hydroxyl radical formation rates  $(R_{HO^*}^f)$  and polychromatic quantum yields 135  $(\Phi_{HO^*})$  are calculated using polychromatic wavelengths.

- 136
- 137 2.3 Physico-chemical measurements
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Different parameters are monitored, including pH, conductivity and redox potential, which 139 140 are measured using a Hanna multiparameter instrument. The UV-Vis spectrum of the collected cloud water is determined with an Agilent Technologies Cary 60 UV-vis 141 spectrophotometer. The TOC concentration is determined with a TOC 5050A analyzer 142 (Shimadzu). Hydrogen peroxide concentration is estimated using p-hydroxyphenilacetic acid 143 (HPAA, purity > 98%) and horseradish peroxidase (POD) (solid containing 150-200 units per 144 mg), according to the spectrofluorimetric quantification method (Miller and Kester, 2002). 145 The formation of the dimer of HPAA is correlated with the concentration of hydrogen 146 peroxide and is detected using a Varian Cary Eclipse Fluorescence Spectrophotometer setting 147 excitation wavelengths at 320 nm, while emission is registered from 340 and 500 nm. The 148 maximum signal is quantified at 408 nm. The scan rate is 600 nm min<sup>-1</sup>, and a bandpass of 10 149 nm is set for excitation and emission. Nitrite ions concentration is determined by 150 derivatization with 2,4-dinitrophenylhydrazine (DNPH) (purity > 97%), in acidic solution 151 (HCl 37%). The UV-absorbing derivative (2,4-dinitrophenilazide) is detected by HPLC. The 152 153 HPLC system (Waters Alliance) equipped with a diode array detector is used with an Eclipse XDB-C18 column (Agilent,  $4.6 \times 150$  mm, 5 µm), and an isocratic method is adopted, using 154 40% acidified water (0.1% phosphoric acid) and 60% methanol. The flow rate is 1 mL min<sup>-1</sup>, 155 and 2,4-dinitrophenilazide is eluted with a retention time of 4.1 min (Kieber and Seaton, 156 1995) and detected at 307 nm. Fe(II) and Fe(III) concentrations are determined by the 157 spectrophotometric method by complexation with ferrozine (purity > 97%), as described by 158 159 Stookey (1970). Fe(II) and Fe(III) represent the oxidative state of the iron species. Adopted complexation method allows us to determine all Fe(II) and Fe(III) species present in solution 160 (i.e., considered as free, aquacomplexes and as complex with other organic molecules). 161 Ascorbic acid (purity reagent grade) is used as the reducing agent to determine total iron. The 162 complex absorption is measured with a Varian Cary 300 Scan Spectrophotometer at 562 nm. 163

164

165 It has been previously demonstrated that filtration does not modify the soluble iron 166 quantification in natural cloud water samples (Parazols et al., 2006; Vaitilingom et al., 2013). It is not possible to measure particulate iron because the ferrozine method cannot solubilize
solid phase iron (the contact time between acidic reagents and particulate iron is too short).
Moreover, the iron particle is expected to be less reactive than the solubilized iron;
consequently, its contribution can be neglected (Arakaki and Faust, 1998).

171 Ion chromatography (IC) analysis is performed employing a DIONEX DX-320 equipped with 172 an IonPac AG11 (guard-column  $4 \times 50$  mm) and an IonPac AS11 (analytical column  $5 \times 250$ 

173 mm) for anions and a DIONEX ICS-1500 equipped with an IonPac CG16 (guard-column  $4 \times$ 174 50 mm) and an IonPac CS16 (analytical column  $5 \times 250$  mm) for cations.

175

### 176 2.4 Statistical analysis

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Principal component analysis (PCA) and hierarchical clustering analysis (HCA) are 178 179 performed with R-3.1.2 software (R Core Team, 2013) using the FactoMineR package (version 1.28, (Lê et al., 2008). This statistical analysis provides a synthetic representation of 180 181 experimental data as a function of the correlations between variables considered and similarities present among the analyzed samples. This technique allows the determination of 182 183 information contained in a set of multivariate data, summarizing it in a few linear combinations of the variables (Deming et al., 1988). HCA data are grouped by similarity, 184 considering all of the information contained in the data set. HCA is a statistical method to 185 qualitatively study the composition of cloud water and can be used to identify the grouping 186 variables that are not well detectable using only PCA. 187

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## 189 2.5 Irradiation experiments

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191 To evaluate the contribution of each possible photochemical source (nitrate, nitrite or 192 hydrogen peroxide) to the hydroxyl radical formation in cloud water, synthetic solutions 193 doped with a single source of oxidant are irradiated to quantify their contribution to the total 194 generation of hydroxyl radicals in a more complex medium.

The photochemical device is composed of a Xenon lamp equipped with a water cooler to avoid the increase of temperature due to the infrared radiations and a mirror to reflect the light vertically. A Pyrex filter was located before the reactor for filtering of light at wavelengths below than 290 nm, corresponding to the lowest wavelengths of the solar emission spectrum. The reactor is a 40 mL cylindrical Pyrex container cooled by water circulation at a temperature of  $278 \pm 2$  K to limit thermal reactions. Samples are continuously stirred with a magnetic stirrer using a Teflon bar to ensure homogeneity.

In Fig. 1, the emission spectrum of the lamp recorded using fiber optics coupled with a charge-coupled device (CCD) spectrophotometer (Ocean Optics USD 2000+UV-VIS) is reported. The energy was normalized with the actinometry results using a paranitroanisole (PNA)/pyridine actinometer (Dulin and Mill, 1982). Over the wavelength range of 290 to 600 nm, a total flux of 157 W m<sup>-2</sup> is measured. The intensity values of the sun emission under clear sky and cloudy conditions at the puy de Dôme mountain in autumn 2013 are also presented in Fig. 1.

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## 210 2.6 Hydroxyl radical formation rate and quantum yield determination

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212 The hydroxyl radical formation rate is determined using terephthalate (TA) (terephthalic disodium salt, purity > 99%) as a probe (Charbouillot et al., 2011). Formation of 213 214 hydroxyterephthalate (TAOH) is quantified using a Varian Cary Eclipse Fluorescence Spectrophotometer, setting excitation wavelengths at 320 nm, while the emission maximum is 215 measured at 420 nm. The scan rate is 600 nm min<sup>-1</sup>, and a bandpass of 10 nm is set for 216 excitation and emission. Terephthalate is a useful probe because it allows the determination of 217 hydroxyl radical formation rates in the presence of fluorescent dissolved organic matter. The 218 concentration of the probe is in a large excess (2 mM) compared with the concentration of 219 organic matter to trap all of the photogenerated hydroxyl radicals and then to estimate a value 220 for the hydroxyl radical formation rate not affected by depletion of HO<sup>•</sup> by other sinks. 221

The reaction between TA and the hydroxyl radical leads to the formation of fluorescent TAOH and non fluorescent secondary products (R1), as follows:

224

$$225 \quad TA + HO^{\bullet} \to TAOH + products \tag{R1}$$

226

The degradation rate of TA ( $R_{TA}^d$ ) and formation rate of hydroxyl radical ( $R_{HO}^f$ ) and TAOH ( $R_{TAOH}^f$ ) can be expressed as follows:

229 
$$R_{HO}^{f} \approx R_{TA}^{d} \approx \frac{R_{TAOH}^{f}}{\gamma}$$
 (1)

230 With

231 
$$R_{TA}^{d} = k_{HO^{\bullet},TA} \Big[ HO^{\bullet} \Big] [TA]$$
(2)

where  $k_{\text{HO}^{\bullet},\text{TA}} = 4.0 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$  is the second order rate constant of the reaction between HO<sup>•</sup> and TA (Charbouillot et al., 2011), and [TA] is the initial concentration of terephthalate and  $\gamma$  is the TAOH formation yield calculated as a function of solution pH. This value is found to be linearly correlated with the pH value and is estimated between the values of 0.15 and 0.25 over the pH range of 4 to 7.

238

Other radicals, such as sulfate  $(SO_4^{\bullet-})$  or chlorine/dichlorine  $(Cl^{\bullet}/Cl_2^{\bullet-})$ , can react with TA, 239 leading to the H-abstraction as first chemical reaction. However, their direct generation is 240 nearly exclusively due to the electron transfer reaction from the corresponding anion (*i.e.*, 241  $SO_4^{2-}$  and Cl<sup>-</sup>, respectively) to the hydroxyl radical. Moreover, considering that the second 242 order rate constant of aromatic compounds with the dichloride radical anion and the sulfate 243 244 radical is expected to be one or two orders of magnitude lower than that with hydroxyl radical (Neta et al., 1988), and considering a relatively low concentration of sulfate and dichlorine 245 radicals in our cloud samples, the TA reactivity can be attributed exclusively to the HO<sup>•</sup>. 246

The quantum yield of hydroxyl radical formation ( $\Phi_{290-400nm}^{HO^{\bullet}}$ ) is defined as the ratio between the formation rate of HO<sup>•</sup> ( $R_{HO^{\bullet}}^{f}$ ) and the number of absorbed photons in Einstein per unit time in the overlap range of 290 to 600 ( $\lambda_1$  and  $\lambda_2$ ) (Eq. 3).

This value evaluates the photochemical process efficiency independent of the experimentalphotochemical conditions.

252 
$$\Phi_{290-400nm}^{HO^{\bullet}} = \frac{R_{HO^{\bullet}}^{f}}{I_{a}}$$
(3)

253 where  $I_a$  can be calculated from the following equation:

254 
$$I_a = \int_{\lambda_1}^{\lambda_2} I_0(\lambda) (1 - 10^{-Abs(\lambda)}) d\lambda$$
(4)

where  $I_0$  (photons m<sup>-2</sup> s<sup>-1</sup>) is the incident photon flux corresponding to the lamp emission and *Abs* is the absorption of cloud water (normalized considering the optical path length of 5 cm inside of the thermostated reactor).

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### 259 2.7 Back-trajectory plots

Backward trajectories of collected air masses are calculated using the HYSPLIT (hybrid
single-particle Lagrangian integrated trajectory) model with the GDAS1 meteorological data
archive and the model of vertical velocity (*http://ready.arl.noaa.gov/HYSPLIT.php*).
Backward trajectories are calculated for 72 hours (Draxler and Rolph, 2012).

265

### 266 2.8 Model description

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The M2C2 combines detailed multiphase chemistry along with the parameterization of the 268 microphysics (Leriche et al., 2001; Leriche et al., 2000; Deguillaume et al., 2004). 269 Particularly, the detailed chemistry of H<sub>x</sub>O<sub>y</sub>, chlorine, carbonates, NO<sub>y</sub>, sulfur, transition 270 271 metal ions (iron, copper, manganese) and the oxidation of volatile organic compounds 272 (VOCs) is included. Photolysis rates are calculated in the gaseous and aqueous phases, and the pH is calculated following the H<sup>+</sup> concentration. Numerical results consist of following 273 the time evolution of the concentrations of each chemical species and calculating at each time 274 275 step the relative contribution of chemical reactions in the production/destruction of chemical compounds. 276

277 In this study, a simplified version of the model is used. The cloud chemical mechanism is restricted to inorganic chemistry (H<sub>x</sub>O<sub>y</sub>, nitrogen, iron) that leads to the HO<sup>•</sup> formation (see 278 Table SM1 for details about the considered reactions). The complexation of iron by oxalate is 279 also considered in the model because it can interfere with the HO<sup>•</sup> formation rates (Long et 280 al., 2013). Laboratory irradiation experiments are simulated with the M2C2 model 281 considering its chemical module and neglecting microphysical processes and mass transfer 282 283 parameterizations. Temperature and pH remained constant during the simulation time. We set the pH for each cloud water sample to the values reported in Table SM2, and the temperature 284 285 is fixed at 278 K, which corresponds to the temperature of the irradiated solutions. The simulated irradiation intensity is held constant and homogenous throughout the experiment. 286 287 The actinic flux of the experimental lamp is discretized in the tropospheric ultraviolet-visible (TUV) model in 156 non-regular intervals over a wavelength range of 120 to 750 nm 288 (Madronich and Flocke, 1999). The photolysis rates of the chemical species are calculated in 289 TUV according to the experimental quantum yields and absorption cross-sections and are 290 indicated in Table SM1. Experimental chemical concentrations (Table SM2) are used to 291 initialize the model (H<sub>2</sub>O<sub>2</sub>, nitrite, nitrate, iron). Moreover, oxalic acid is considered as an 292 organic complexant during the Fe(II)/Fe(III) cycle. The formation rate of HO<sup>•</sup> is calculated by 293

the model considering the modeled contribution of each reaction producing HO<sup>•</sup> during one
hour of experiment.

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### 297 **3** Results and Discussion

**Classification of cloud samples** 

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3.1

300

Recently, physicochemical parameters and concentrations of the major organic and inorganic 301 302 compounds of cloud samples collected over the last ten years at the puy de Dôme are measured and statistically analyzed by PCA (Deguillaume et al., 2014). Along with the 303 304 corresponding back-trajectory plots, 4 different categories of air masses reaching the summit of the PUY could be distinguished, as follows: polluted, continental, marine and highly 305 marine. Highly marine clouds exhibited high concentrations of Na<sup>+</sup> and Cl<sup>-</sup>, and the marine 306 category presented a lower concentration of ions but more elevated pH, while the two 307 remaining clusters, classified as "continental" and "polluted", are characterized by the 308 second-highest and highest levels of  $NH_4^+$ ,  $NO_3^-$ , and  $SO_4^{2-}$ , respectively. 309

310 In Table SM2, the measured physico-chemical composition of the cloud water samples is 311 reported for this study. We use the same statistical analysis to classify these cloud water samples as Deguillaume et al. (2014). PCA is performed using the pH and the concentration 312 of sulfate, nitrate, chloride, sodium and ammonium ions as variables. Fig. 2 reports the scores 313 plot for samples used for the previously reported classification as a function of the attributed 314 class and for the new samples. Three types are identified, as follows: (i) highly marine, (2 315 samples) characterized by pH values of 5.0 and 5.6, respectively, high concentration of 316 chloride and sodium and low concentrations of nitrate, nitrite and ammonium; (ii) marine 317 cloud waters (28 samples), showing pH values between 4.7 and 7.6 and very low 318 concentrations of anions and cations, and (iii) continental samples (11 samples), with pH 319 values from 4.1 to 6.9 and a medium concentration of nitrates, sulfates and ammonium, while 320 321 sodium and chloride concentrations are very low. No sample could be classified as polluted cloud water because polluted cloud waters have been characterized by concentrations of 322 nitrates, sulfates and ammonium higher than 350, 70 and 330 µM, respectively. This 323 statistical analysis confirms that the majority of the collected samples are of marine origin. 324 This statistical analysis is confirmed by the back-trajectory plots from the HYSPLIT model, 325

showing that most of the air masses reaching the puy de Dôme arises from the west sector *i.e.*, 326

- from the Atlantic Ocean. 327
- 328

# 329

#### 3.2 Determination of the hydroxyl radical formation rates and photolysis rates 330

The concentration of the main photochemical sources of HO<sup>•</sup> for each sample is reported in 331 Table 1. Particularly, the Fe(II) and Fe(III) concentrations are below the detection limit (0.01 332  $\mu$ M) for the majority of the collected samples. The highest value found for the Fe(II) 333 concentration is 0.7 µM, while it is 0.6 µM for Fe(III), corresponding to typical values found 334 for marine origin cloud waters (Parazols et al., 2006). The H<sub>2</sub>O<sub>2</sub> concentration values range 335 between 6 and 50 µM, nitrate is evaluated between 2 and 220 µM, while the nitrite 336 concentration is between 0 and  $1.4 \,\mu$ M. 337

The  $R_{Ho}^{f}$  was measured in pure water doped with different concentrations of hydrogen 338 peroxide, nitrate and nitrite on the same order of magnitude as the collected natural samples. 339 The conditions were those used for natural cloud samples (see Sect. 2.6). A linear correlation 340 between  $R_{\mu\sigma}^{f}$  and the concentrations of photochemical precursors is found (Fig. 3). The 341 photolysis rate (J) (s<sup>-1</sup>) is then estimated from the slopes and are reported in Table 4. For 342  $H_2O_2$ , the J value is half of the experimental slope because  $H_2O_2$  provides two  $HO^{\bullet}$  radicals. 343

The hydroxyl radical formation rate  $(R_{\mu\rho}^{f})$  is determined for 36 samples, and its value is 344 estimated between  $3.3 \times 10^{-12}$  and  $4.2 \times 10^{-10}$  M s<sup>-1</sup>. Higher values are obtained for cloud 345 water samples of continental origin, while the values found for marine and highly marine 346 cloud waters are less than  $1.4 \times 10^{-10}$  M s<sup>-1</sup>. The quantum yield of the formation of hydroxyl 347 radicals under polychromatic irradiation between 290 and 400 nm ( $\Phi_{290-400nm}^{HO^{\bullet}}$ , see eq.(3)) is 348 estimated as between  $10^{-5}$  and  $10^{-2}$ . (Table 2). 349

To our knowledge, only a scarce number of data are available in the literature concerning 350 measurements of hydroxyl radical formation rates  $(R_{Ho^{\bullet}}^{f})$  and formation quantum yield 351  $(\Phi^{HO^*})$  in real cloud waters (Table 3). Faust and Allen (1993) measured the photoformation 352 rates of HO<sup>•</sup> (ranging from 1.3 to  $8.3 \times 10^{-10}$  M s<sup>-1</sup>) under monochromatic light (313 nm) and 353 hydroxyl radical quantum yield (between ~  $5 \times 10^{-4}$  and  $10^{-2}$ ) of six continental cloud water 354 samples. Anastasio and McGregor (2001) investigated the photoreactivity of two cloud waters 355 from the Tenerife Islands to compare the obtained values with fog waters. The authors found 356

357  $R_{Ho}^{f}$ , ranging between 3.0 and  $6.9 \times 10^{-10} \,\mathrm{M \, s^{-1}}$ . t that are approximately 1 order of magnitude 358 higher than those reported in this study for marine cloud waters, and the differences can be 359 attributed to the air mass origin, as suggested by Faust and Allen (1993). The authors 360 suggested that long-range terrestrial aerosol and gas transport in continental clouds could 361 provide an additional source of hydroxyl radicals compared with other marine or remote 362 clouds.

363

# **364 3.3 Modeling the hydroxyl radical formation rates**

365

We simulate the hydroxyl formation rate,  $R_{HO^*}^f$  mod, using the model along with the relative 366 contribution (%) of each chemical source (Table SM3). In Fig. 4, the differences between the 367 modeled and experimental HO<sup>•</sup> formation rates are estimated calculating the bias error 368  $((R_{HO^{\bullet}}^{f} \text{ mod } - R_{HO^{\bullet}}^{f} \exp) / R_{HO^{\bullet}}^{f} \exp \text{ in } (\%)).$  Globally, for the whole cloud water samples (black 369 boxplot), the model can reproduce the range of measured hydroxyl radical formation rates 370 with a slight model underestimation (median of the bias error equal to -23%). However, if the 371 cloud samples are discretized as a function of different iron concentration ranges (boxplots in 372 color), then the model tends to overestimate the hydroxyl radicals formation for iron 373 concentrations (Fe(II) + Fe(III)) more than 0.1  $\mu$ M. For concentration of iron between 0.1 and 374 0.4 µM (8 cloud samples), the median of the bias error is 61% whereas for iron concentrations 375 over 0.4 µM (5 cloud samples), the median reaches 260%. For cloud samples in which the 376 iron concentration is 0.4 µM, the modeled contribution to the hydroxyl radical formation of 377 iron (Fenton reaction and photolysis of aqua-complexes) can reach 80% (Table SM3). In the 378 model, Fe(III) is partially complexed with oxalic acid, but the majority of iron for these cloud 379 samples is simulated as aqua-complexes (mainly  $Fe(OH)^{2+}$  and  $Fe(OH)_{2+}^{2+}$ ). Therefore, iron 380 can act as a significant HO<sup>•</sup> source due to its efficient photolysis (Reactions R7, R8, R9, 381 Table SM1) and the Fenton reaction with  $H_2O_2$  (Reactions R11, Table SM1). 382

However, in atmospheric natural water, the chemical composition of organic matter is still not very well characterized (Herckes et al., 2013). Part of this organic matter is expected to efficiently complex metals in cloud water (Okochi and Brimblecombe, 2002). Due to missing information about the iron speciation and complexation in natural cloud water, the model probably overestimates the free Fe(III). Moreover, iron organic complexes are not expected to directly generate hydroxyl radicals but primarily contribute to the oxidative capacity *via*  Fe(II) generation and the Fenton process. To evaluate this hypothesis, we decide to consider the iron as totally complexed by the organic matter present in natural cloud waters. As a first general approximation, we consider iron as not reactive in the model. The new simulations show that the majority of the simulated values (75%) of the hydroxyl radical formation rates are now underestimated by the model (median of the bias error equal to -40%) (Fig. 5).

Based on the laboratory irradiation experiments (sec. 2.6 and Fig. 3), new photolysis rates 394 from nitrite, nitrate and hydrogen peroxide are previously estimated from the hydroxyl radical 395 formation rates (Table 4). For nitrate and H<sub>2</sub>O<sub>2</sub>, the values calculated by the model are lower 396 than the experimental ones; the experimental photolysis rates are higher by a factor ~1.5 for 397  $H_2O_2$  and ~2 for nitrate. For nitrite, the experimental photolysis rate is approximately half of 398 the modeled value. These discrepancies should partially explain the underestimation of HO<sup>•</sup> 399 formation by the model. Therefore, we consider the experimental photolysis rates in the 400 model, and a new comparison of modeled HO<sup>•</sup> formation rates with experimental values is 401 shown in Fig. 5. The median of the bias error is -3% with the 25<sup>th</sup> and 75<sup>th</sup> percentiles at -50 402 and 60%, respectively. Table SM4 reports the distribution of the relative contributions of 403 H<sub>2</sub>O<sub>2</sub>, NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> photolysis to the modeled  $R_{HO^{\bullet}}^{f}$ . While the median value of the nitrite 404 and nitrate contributions is calculated by the model as equal to 1 and 5%, respectively (Fig. 405 6), the main HO<sup>•</sup> contributor is  $H_2O_2$  photolysis (median value of 93%, with the 25<sup>th</sup> and 75<sup>th</sup> 406 percentiles at 85 and 96%, respectively). This result suggests that H<sub>2</sub>O<sub>2</sub> is the key compound 407 that drives the oxidative capacity of our cloud water samples when iron concentrations are 408 409 relatively low or when iron is suggested to be totally complexed by organic matter.

410

### 411 **4** Conclusion

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In this study, we compare experimental and modeled HO<sup>•</sup> formation rates for 41 cloud water 413 samples with contrasted chemical compositions depending on the origin of the air masses. 414 415 This approach helps to elucidate the complex aqueous photoreactivity of natural cloud water 416 that is expected to drive the oxidative capacity. Experimental data are obtained considering only the HO<sup>•</sup> formation sources with addition of an excess of chemical probe leading to the 417 measurements of total generated HO<sup>•</sup> radicals. The first comparison with the M2C2 model 418 shows that the model can reproduce the order of magnitude of measured  $R_{HO^*}^f$  (from  $1.1 \times 10^-$ 419 <sup>11</sup> to  $4.2 \times 10^{-10}$  M s<sup>-1</sup> and from  $1.1 \times 10^{-11}$  to  $2.4 \times 10^{-10}$  M s<sup>-1</sup> for experimental and modeled 420 values, respectively). Some discrepancies appear for samples containing iron concentrations 421

over 0.4 µM in which the model overestimates the contribution of iron(III)-aqua complexes to 422 the HO<sup>•</sup> production rate. Ultimately, the aqueous chemical mechanism in the model was 423 424 modified considering new photolysis rates for H<sub>2</sub>O<sub>2</sub>, nitrite and nitrate estimated by laboratory irradiation experiments. As a sensitivity test, iron reactivity was also suppressed in 425 the model to account for the total complexation of iron. The modeled production rates of  $HO^{\bullet}$ 426 with the updated mechanism are closer to the experimental values. This supports the 427 hypothesis that iron could be strongly complexed by the organic matter in natural cloud water. 428 These complexes could be more stable and less photoreactive, leading to less HO<sup>•</sup> production 429 than that calculated by theoretical models in which only the photochemistry of Fe(III)-430 carboxylate is considered (Weller et al., 2014). In this context, evaluation of the complexation 431 432 of iron by organic compounds in the cloud aqueous phase and the photoreactivity of these complexes should be pursued in the future. Their photoreactivity provides significant data to 433 434 understand the specific role of iron species and, more generally, the oxidant capacity of this medium. 435

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584	Table	Caption
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- Table 1. Concentration of main sources of hydroxyl radical in sampled clouds. 41 samples have been
  analyzed. BDL: below detection limit (0.01 µM for iron and 0.05 µM for NO<sub>2</sub><sup>-</sup>), NM: not measured.
- **Table 2.** Hydroxyl radical formation rate  $(R_{Ho}^{f})$  and hydroxyl radical polychromatic quantum yield formation  $(\Phi_{HO}^{\bullet})^{(290-400 \text{ nm})}$  values estimated from cloud water samples. NM: not measured. Mar: Marine, H-Mar: highly marine and Cont: continental influence. The error on  $R_{Ho}^{f}$  are derived at the 1- $\sigma$  level simply from the scattering of experimental data.
- 593

**Table 3.** Hydroxyl radical formation rates  $(R_{Ho^*}^f)$  and polychromatic quantum yield  $(\Phi_{Ho^*})$  found in literature and in this work.

596

597 Table 4. Modeled photolysis rates calculated by the model *versus* experimental photolysis rates598 obtained from experiments reported in Fig. 3.

Sample	Iron (µM)		$H_2O_2$ $NO_3^-$ (uM) (uM)		$NO_2^-$ (µM)	Sample	Iron (µM)		H <sub>2</sub> O <sub>2</sub> (µM)	NO <sub>3</sub> <sup>-</sup> (μM)	NO <sub>2</sub> <sup>-</sup> (μM)
L	Π	III	(μΜ)	(μ)	(µ.111)	1	Π	III			-
1	NM	NM	12.3	16.7	0.46	22	BDL	BDL	52.3	131.9	0.72
2	NM	NM	9.0	6.1	1.44	23	BDL	BDL	49.4	133.1	0.95
3	NM	NM	15.1	9.9	0.40	24	BDL	BDL	8.1	7.5	BDL
4	NM	NM	14.0	14.2	0.30	25	0.08	0.02	6.7	21.2	0.15
5	NM	NM	13.0	14.7	0.38	26	0.40	0.20	6.8	39.7	BDL
6	NM	NM	7.8	2.6	BDL	27	0.70	0.20	6.6	75.6	BDL
7	NM	NM	6.2	1.7	BDL	28	0.70	0.30	7.2	73.8	BDL
8	NM	NM	9.7	6.9	BDL	29	0.01	0.10	8.0	24.7	0.27
9	NM	NM	8.2	8.2	BDL	30	BDL	0.16	8.8	19.7	0.52
10	NM	NM	10.2	2.3	BDL	31	BDL	0.45	9.1	20.7	0.61
11	NM	NM	17.2	5.6	BDL	32	BDL	BDL	13.1	21.4	0.07
12	BDL	0.57	18.0	24.7	0.28	33	BDL	0.10	2.1	6.1	BDL
13	BDL	0.12	24.6	23.7	1.10	34	0.09	BDL	8.4	10.3	0.47
14	BDL	0.11	12.0	19.0	BDL	35	BDL	BDL	2.2	15.1	0.51
15	BDL	BDL	14.5	19.0	0.23	36	BDL	0.03	2.1	20.3	BDL
16	BDL	BDL	9.1	21.3	0.10	37	BDL	0.03	2.1	18.5	BDL
17	BDL	0.11	16.2	219.6	0.05	38	0.07	BDL	2.4	13.5	0.34
18	0.10	0.01	16.2	205.6	0.07	39	0.04	BDL	3.1	20.8	BDL
19	BDL	BDL	14.9	20.0	0.12	40	BDL	0.01	5.7	39.1	BDL
20	BDL	BDL	15.7	37.4	0.19	41	BDL	0.02	5.3	46.5	0.16
21	BDL	BDL	22.2	72.6	0.42						

**Table 1.** Concentration of main sources of hydroxyl radical in sampled clouds. 41 samples have been analyzed. BDL: below detection limit (0.01  $\mu$ M for iron and 0.05  $\mu$ M for NO<sub>2</sub><sup>-</sup>), NM: not measured.

Table 2. Hydroxyl radical formation rate  $(R_{HO}^{f}, (M \text{ s}^{-1}))$  and hydroxyl radical polychromatic quantum yield formation  $(\Phi_{HO}^{\bullet}, (290-400 \text{ nm}))$  values estimated from cloud water samples. NM: not measured. Mar: Marine, H-Mar: highly marine and Cont: continental influence. The error on  $R_{HO}^{f}$  are derived at the 1-σ level simply from the scattering of experimental data.

Sample	$R^{f}_{HO^{\bullet}}(Ms^{\text{-}1})$	$\Phi_{\mathrm{HO}^{\bullet}}^{290-400nm}$	Class	Sample	$R^{\rm f}_{\rm HO^{\bullet}}(\rm M~s^{-1})$	$\Phi_{\mathrm{HO}^{\bullet}}^{290-400nm}$	Class
1	(3.30±0.23)×10 <sup>-11</sup>	1.27×10 <sup>-3</sup>	Mar	22	(3.37±0.01)×10 <sup>-10</sup>	2.5×10 <sup>-3</sup>	Cont
2	NM	NM	Mar	23	(4.16±0.01)×10 <sup>-10</sup>	1.8×10 <sup>-3</sup>	Cont
3	NM	NM	Mar	24	(5.10±0.01)×10 <sup>-10</sup>	NM	Mar
4	(1.40±0.01)×10 <sup>-10</sup>	1.0×10 <sup>-2</sup>	H-Mar	25	(2.42±0.08)×10 <sup>-11</sup>	1.9×10 <sup>-4</sup>	Cont
5	(1.24±0.02)×10 <sup>-10</sup>	6.0×10 <sup>-3</sup>	H-Mar	26	(1.41±0.01)×10 <sup>-10</sup>	3.4×10 <sup>-3</sup>	Cont
6	(2.77±0.01)×10 <sup>-11</sup>	1.5×10 <sup>-4</sup>	Mar	27	(4.95±0.01)×10 <sup>-11</sup>	1.5×10 <sup>-4</sup>	Cont
7	(5.60±0.06)×10 <sup>-11</sup>	9.0×10 <sup>-3</sup>	Mar	28	NM	NM	Cont
8	(2.48±0.01)×10 <sup>-11</sup>	2.9×10 <sup>-3</sup>	Mar	29	(8.48±0.04)×10 <sup>-11</sup>	1.3×10 <sup>-4</sup>	Mar
9	(2.20±0.02)×10 <sup>-11</sup>	1.8×10 <sup>-3</sup>	Mar	30	(8.43±0.02)×10 <sup>-11</sup>	1.2×10 <sup>-4</sup>	Mar
10	(2.93±0.02)×10 <sup>-11</sup>	2.4×10 <sup>-3</sup>	Mar	31	(6.11±0.21)×10 <sup>-11</sup>	5.1×10 <sup>-5</sup>	Mar
11	(6.77±0.02)×10 <sup>-11</sup>	2.7×10 <sup>-3</sup>	Mar	32	NM	NM	Mar
12	(6.10±0.19)×10 <sup>-11</sup>	1.7×10 <sup>-4</sup>	Mar	33	(3.27±0.23)×10 <sup>-12</sup>	1.3×10 <sup>-5</sup>	Mar
13	(4.66±0.01)×10 <sup>-11</sup>	1.5×10 <sup>-4</sup>	Mar	34	(2.73±0.01)×10 <sup>-11</sup>	8.5×10 <sup>-4</sup>	Mar
14	(2.81±0.01)×10 <sup>-11</sup>	2.0×10 <sup>-4</sup>	Mar	35	(3.60±0.30)×10 <sup>-11</sup>	6.0×10 <sup>-4</sup>	Mar
15	(1.09±0.04)×10 <sup>-11</sup>	9.0×10 <sup>-5</sup>	Mar	36	(5.97±0.12)×10 <sup>-11</sup>	9.6×10 <sup>-5</sup>	Mar
16	NM	NM	Mar	37	(2.41±0.04)×10 <sup>-11</sup>	8.4×10 <sup>-5</sup>	Mar
17	(6.05±0.44)×10 <sup>-11</sup>	4.1×10 <sup>-5</sup>	Cont	38	(5.76±0.13)×10 <sup>-11</sup>	2.7×10 <sup>-4</sup>	Mar
18	(3.39±0.20)×10 <sup>-11</sup>	3.2×10 <sup>-5</sup>	Cont	39	(2.69±0.04)×10 <sup>-11</sup>	1.1×10 <sup>-4</sup>	Mar
19	(8.11±0.02)×10 <sup>-11</sup>	1.9×10 <sup>-4</sup>	Mar	40	(1.27±0.01)×10 <sup>-10</sup>	2.6×10 <sup>-4</sup>	Cont
20	(8.46±0.01)×10 <sup>-11</sup>	8.4×10 <sup>-3</sup>	Mar	41	(1.09±0.01)×10 <sup>-10</sup>	5.1×10 <sup>-4</sup>	Cont
21	(1.54±0.01)×10 <sup>-10</sup>	1.5×10 <sup>-3</sup>	Cont				

**Table 3.** Hydroxyl radical formation rates  $(R_{HO^{\bullet}}^{f}, M \text{ s}^{-1})$  and polychromatic quantum yield  $(\Phi_{HO^{\bullet}})$ 

612 found in literature and in this work.

	$R^f_{HO^{\bullet}}$ (M s <sup>-1</sup> )	$\Phi_{\cdot_{OH}}$	References
Rain water	2.0 - 6.5 × 10 <sup>-11</sup>		(Albinet et al., 2010)
Cloud water	$1.3 - 8.3 \times 10^{-10}$	$4.6 \times 10^{-4} - 1.0 \times 10^{-2}$ at 313 nm	(Faust and
Fog	$0.9 - 6.9 \times 10^{-10}$		Allen, 1993)
Aqueous extracted aerosol particles	$0.4 - 3.8 \times 10^{-10}$	$3.0  imes 10^{-4}$ - $1.7  imes 10^{-3}$	(Arakaki et al., 2006)
Cloud water	3.1 - 6.9 × 10 <sup>-10</sup>		(Anastasio and McGregor, 2001)
Cloud water	$0.3 - 5.9 \times 10^{-10}$	$5.1\times10^{\text{-4}}$ - $3.0\times10^{\text{-3}}$	(Arakaki and Faust, 1998)
Cloud water at the PUY station	$0.2 - 4.2 \times 10^{-10}$	$1.3 \times 10^{-5} - 1.0 \times 10^{-2}$ Polychromatic	This work

	Modeled photolysis rates J (s <sup>-1</sup> )	Experimental photolysis rates $J(s^{-1})$
$H_2O_2 \xrightarrow{h\nu} 2HO^{\bullet}$	$1.52  imes 10^{-6}$	$(2.50\pm 0.11)\times 10^{-6}$
$HNO_2 \xrightarrow{hv} HO^{\bullet} + NO^{\bullet}$	$6.16\times10^{\text{-5}}$	
$NO_2^- + H_2O \xrightarrow{hv} HO^{\bullet} + NO^{\bullet} + HO^-$	$9.98  imes 10^{-6}$	$(5.15\pm 0.30)\times 10^{\text{-6}}$
$NO_3^- + H_2O \xrightarrow{hv} HO^{\bullet} + NO_2^{\bullet} + HO^-$	$6.71 imes10^{-8}$	$(1.23 \pm 0.04) \times 10^{-7}$
$Fe^{3+} + H_2O \xrightarrow{hv} HO^{\bullet} + Fe^{2+} + H^+$	$1.24  imes 10^{-6}$	
$Fe(OH)^{2+} \xrightarrow{h\nu} HO^{\bullet} + Fe^{2+}$	$2.81 imes10^{-4}$	
$Fe(OH)_2^+ \xrightarrow{h\nu} HO^{\bullet} + Fe^{2+} + HO^-$	$3.53 \times 10^{-4}$	

**Table 4.** Modeled photolysis rates calculated by the model *versus* experimental photolysis ratesobtained from experiments reported in Fig. 3.

### 619 **Figure Caption**

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**Fig. 1.** Absorption spectra of different cloud water samples (left axis) and the right-hand axis shows the emission spectrum of the adopted Xenon lamp reaching solutions over the range of 290 to600 nm (total flux intensity =  $157 \text{ W m}^{-2}$ ) compared with the sun emission spectrum (dashed line) for a sunny ( $353 \text{ W m}^{-2}$ ) and a cloudy day ( $90 \text{ W m}^{-2}$ ) in October 2013.

625

**Fig. 2.** Scores plot obtained by PCA analysis of 137 samples (corresponding to 73 cloud events) collected before 2013 and grouped in four classes in function of previously described classification (Deguillaume et al., 2014) and 41 samples (numbered from 80 to 120) collected during 2013 and 2014. These new data correspond to 15 cloud events and are indicated in black triangles. Statistical analysis is performed using 6 selected variables (pH,  $[Na^+]$ ,  $[Cl^-]$ ,  $[SO_4^{2^-}]$ ,  $[NO_3^{-}]$ , and  $[NH_4^+]$ ). The scree plot obtained from autoscaled data shows that two selected principal components (PC) containing a total variance of about 81%.

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Fig. 3. Scatter plot of hydroxyl radical formation rates *vs.* hydrogen peroxide, nitrate and
nitrite concentrations using 2 mM of TA at pH 5.0 under Xenon lamp irradiation. The solid
line is the linear fit, and dashed lines denote the 90% confidence of the linear fit.

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**Fig. 4.** Distribution of the bias error for the whole cloud water samples (black) and for cloud samples discretized as a function of different iron concentration ranges (in color). The bias error is defined by the ratio  $(R_{HO}^{f}, \text{mod} - R_{HO}^{f}, \exp) / R_{HO}^{f}, \exp$  in (%). The number of samples analyzed is indicated above each box plot. The bottom and top lines correspond to the 25<sup>th</sup> and 75<sup>th</sup> percentiles, respectively. The full line represents the median values. The ends of the whiskers are the 10<sup>th</sup> and 90<sup>th</sup> percentiles.

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**Fig. 5.** Distribution of the bias error for the whole cloud water samples for the reference case and for two sensitivity tests performed with the model: (i) the iron chemistry (photolysis of Fe(III) and the Fenton reaction) is neglected in the model; (ii) the iron chemistry is neglected, and the new photolysis rate constants obtained from experimental measurements are implemented in the model. The bias error is defined by the ratio  $(R_{Ho}^{f}, \text{mod} - R_{Ho}^{f}, \exp) / R_{Ho}^{f}, \exp$  in (%). The number of samples analyzed is indicated above each box plot. The bottom

- and top lines correspond to the  $25^{\text{th}}$  and  $75^{\text{th}}$  percentiles, respectively. The full line represents the median values. The ends of the whiskers are the  $10^{\text{th}}$  and  $90^{\text{th}}$  percentiles.
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**Fig. 6.** Distribution of relative contributions of modeled HO<sup>•</sup> formations rates for each photochemical source ( $H_2O_2$ ,  $NO_3^-$  and  $NO_2^-$  photolysis) for the whole cloud water samples. Model outputs are obtained from the sensitivity test in which the iron chemistry is not considered, and new photolysis rate constants from experimental measurements are implemented in the model. The number of samples analyzed is indicated above each box plot. The bottom and top lines correspond to the 25<sup>th</sup> and 75<sup>th</sup> percentiles, respectively. The full line represents the median values. The ends of the whiskers are the 10<sup>th</sup> and 90<sup>th</sup> percentiles.



662

**Fig. 1.** Absorption spectra of different cloud water samples (left axis) and the right-hand axis shows the emission spectrum of the adopted Xenon lamp reaching solutions over the range of 290 to600 nm (total flux intensity =  $157 \text{ W m}^{-2}$ ) compared with the sun emission spectrum (dashed line) for a sunny (353 W m<sup>-2</sup>) and a cloudy day (90 W m<sup>-2</sup>) in October 2013.



**Fig. 2.** Scores plot obtained by PCA analysis of 137 samples collected before 2013 and grouped in four classes as a function of the previously described classification (Deguillaume et al., 2014) and 41 samples collected during 2013 and 2014. These new data correspond to 15 cloud events and are indicated in black triangles. Statistical analysis is performed using 6 selected variables (pH,  $[Na^+]$ , [Cl<sup>-</sup>],  $[SO_4^{2-}]$ ,  $[NO_3^{-}]$ , and  $[NH_4^{+}]$ ). The scree plot obtained from autoscaled data shows that two selected principal components (PC) containing a total variance of about 81%.



Fig. 3. Scatter plot of hydroxyl radical formation rates *vs.* hydrogen peroxide, nitrate and nitrite
concentrations using 2 mM of TA at pH 5.0 under Xenon lamp irradiation. The solid line is the linear
fit, and dashed lines denote the 90% confidence of the linear fit.





**Fig. 4.** Distribution of the bias error for the whole cloud water samples (black) and for cloud samples discretized as a function of different iron concentration ranges (in color). The bias error is defined by the ratio  $\binom{R_{H0}^{f}}{H0}$  mod -  $\binom{R_{H0}^{f}}{H0}$  exp in (%). The number of samples analyzed is indicated above each box plot. The bottom and top lines correspond to the 25<sup>th</sup> and 75<sup>th</sup> percentiles, respectively. The full line represents the median values. The ends of the whiskers are the 10<sup>th</sup> and 90<sup>th</sup> percentiles.



690 Fig. 5. Distribution of the bias error for the whole cloud water samples for the reference case and for two sensitivity tests performed with the model: (i) the iron chemistry (photolysis of Fe(III) and the 691 692 Fenton reaction) is neglected in the model; (ii) the iron chemistry is neglected, and the new 693 photolysis rate constants obtained from experimental measurements are implemented in the model. The bias error is defined by the ratio  $(R_{HO}^{f}, \text{mod} - R_{HO}^{f}, \exp) / R_{HO}^{f}, \exp (\%)$ . The number of samples 694 analyzed is indicated above each box plot. The bottom and top lines correspond to the 25<sup>th</sup> and 75<sup>th</sup> 695 percentiles, respectively. The full line represents the median values. The ends of the whiskers are the 696 10<sup>th</sup> and 90<sup>th</sup> percentiles. 697



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**Fig. 6.** Distribution of relative contributions of modeled HO<sup>•</sup> formations rates for each photochemical source  $(H_2O_2, NO_3^- \text{ and } NO_2^- \text{ photolysis})$  for the whole cloud water samples. Model outputs are obtained from the sensitivity test in which the iron chemistry is not considered, and new photolysis rate constants from experimental measurements are implemented in the model. The number of samples analyzed is indicated above each box plot. The bottom and top lines correspond to the 25<sup>th</sup> and 75<sup>th</sup> percentiles, respectively. The full line represents the median values. The ends of the whiskers are the 10<sup>th</sup> and 90<sup>th</sup> percentiles.