1	Quantification of ice nuclei active at near 0°C temperatures
2	in low altitude clouds at the puy de Dôme atmospheric
3	station (1465 m a.s.l.)
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20 Abstract

21 The distribution, abundance and nature of ice nucleation active particles in the atmosphere are 22 major sources of uncertainty in the prediction of cloud coverage, precipitation patterns and 23 climate. Some biological ice nuclei (IN) induce freezing at temperatures at which most other 24 atmospheric particles exhibit no detectable activity (>-10°C). Their actual contribution to the 25 pool of IN in clouds remains poorly known, but numerical studies suggested their likely significance in atmospheric processes. In this study, cloud water was collected aseptically 26 27 from the summit of puy de Dôme (1465 m a.s.l., France) within contrasted meteorological and 28 physico-chemical situations. Total and biological (i.e. heat-sensitive) IN were quantified by 29 droplet-freezing assay between -5°C and -14°C. We observed that freezing was systematically 30 induced by biological material, between -6°C and -8°C in 92% of the samples. Its removal by 31 heat treatment consistently led to a decrease of the onset freezing temperature, by 3°C or more in most samples. At -10°C, 0 to ~220 biological IN mL⁻¹ of cloud water were measured (i.e. 0 32 to $\sim 22 \text{ m}^{-3}$ of cloud air based on cloud liquid water content estimates) and these represented 33 65% to 100% of the total IN. Based on back-trajectories and on physico-chemical analyses, 34 35 the high variability observed resulted probably from a source effect, with IN originating mostly from continental sources. Assuming that biological IN were all bacteria, at maximum 36 37 0.6% of the bacterial cells present in cloud water samples could have acted as IN at -8°C, 38 1.5% at -10°C, and 3.1% at -12°C. The dataset generated here will help elucidating the role of 39 biological and bacterial IN on cloud microphysics by numeric modelling, and their impact on 40 precipitation at local scale.

41

42 **1. Introduction**

43 The formation of clouds and their evolution have global impacts on Earth's climate. Within 44 the last decade, considerable efforts have been made in order to identify and quantify the 45 particles acting as ice nuclei (IN) in the atmosphere. Those particles are responsible for the 46 heterogeneous nucleation of ice in supercooled clouds, leading to modifications of their 47 radiative properties and initiating precipitation. At temperatures colder than about -15°C, 48 feldspar particles were recently demonstrated to account for a great part to the pool of IN in 49 mixed-phase clouds at a global scale (Atkinson et al., 2013). However, at warmer 50 temperatures, most of the mineral aerosols as well as metallic and soot particles exhibit only 51 very low or undetectable IN activity (INA), and the best ice nuclei candidates are biological 52 (bacteria, fungi), or biogenic (macromolecules derived from living organisms, such as 53 proteins) (Conen et al., 2011; DeMott and Prenni, 2010). Hence, biological IN are thought 54 to largely influence clouds' evolution within the upper range of temperatures around freezing 55 (e.g. Möhler et al., 2007). Among those, the most efficient natural IN described so far are 56 bacteria, with representatives active at temperatures as "warm" as -2°C (Maki et al., 1974); 57 other very active biological IN different from bacteria were also detected in the air, but their 58 exact nature remains unknown (Garcia et al., 2012). Specimens of INA bacteria have been 59 recovered from all the compartments of the water cycle: freshwaters (Maki and Willoughby, 60 1978; Morris et al., 2008), clouds (Joly et al., 2013) and precipitation at high altitude (Sands 61 et al., 1982) or closer to the ground (Constantinidou et al., 1990; Maki and Willoughby, 62 1978; Šantl-Temkiv et al., 2009; Stephanie and Waturangi, 2011). This supports the hypothetical concept termed "bioprecipitation" that such bacteria could participate to 63 64 hydrological cycles by triggering precipitation (Morris et al., 2004).

Fig. 1 summarizes our current quantitative knowledge about high-negative-temperature 65 (>-15°C) IN in the atmosphere and in the environmental compartments of the water cycle. 66 67 The main results of the present study are also indicated. Most plants harbor relatively large 68 populations of epiphytic ice nucleation active (INA) bacteria (Constantinidou et al., 1990; Lindemann et al., 1982; Lindow et al., 1978; Maki and Willoughby, 1978; Morris et al., 69 70 2008), so the main source of atmospheric biological IN is probably vegetation (Pöschl et al., 71 2010). Recently, oceans were also cited as possible emitters of biogenic IN into the 72 atmosphere (Burrows et al., 2013).

In the air at low altitude, Garcia et al. (2012) observed concentrations of 90 to 460 IN m⁻³ 73 74 active at -10°C over vegetated agricultural areas, most of which were classified as biological 75 based on their sensitivity to heat. In the latter study, INA bacterial cells were estimated to 76 represent only a small fraction of the total airborne bacteria (~0.002%). Nevertheless, some 77 specimens of INA bacterial strains have been recovered by culture from atmospheric samples 78 (e.g. Stephanie and Waturangi, 2011). At high altitude, notwithstanding their suspected 79 importance in atmospheric processes, much less quantitative data of high-temperature IN is available. Their concentration there is in general much below 25 m⁻³, but it can vary 80 drastically between <1 and $\sim100 \text{ m}^{-3}$ within very short timeframes (Bowers et al., 2009; 81 82 Conen et al., 2012; Xia et al., 2013). Interestingly, the highest concentrations were observed 83 at high relative humidity.

Airborne IN can be transported to regions very distant from the source of emission and affect rain patterns after being incorporated into clouds (**Creamean et al., 2013**). In the single 86 orographic cirrus cloud event studied by **Pratt et al. (2009)**, about half of the 46 ice crystals 87 residues (140-700 nm in diameter; -31°C ambient temperature) had a mass spectrometry 88 signature typical of mineral dust, while about 33% were biological particles. More recent and 89 more extensive *in situ* observations of cirrus clouds at temperatures $< -30^{\circ}$ C showed that 90 biological particles are probably much more scarce among the solid residues of ice crystals 91 (*i.e.* less than 1%), but that rather mineral dust and metallic particles dominate (Cziczo et al., 92 2013). However, observing ice crystal residues does not guarantee identifying the actual IN, 93 and cirrus are high-altitude, very low temperature and non-precipitating clouds, so probably 94 not the most appropriate environments for investigating high-temperature and biological IN. 95 A quantitative study of high-temperature atmospheric IN at lower altitude was led at the Jungfraujoch summit in the Alps (3450 m a.s.l.); concentrations of 0 to 3.8 m⁻³ were measured 96 97 when clouds were present on the site (Xia et al., 2013). Albeit, as emphasized by authors, their "precision was low" due to a limited air sample volume of less than 3 m³. 98

99 Fresh snow and rain collected at different locations over the planet, from poles to subequatorial regions, carried ~1 to ~100 IN active at -7°C per liter of water. Most were altered by heat treatment and were thus categorized as biological, and about half of these were probably bacteria (Christner et al., 2008a, 2008b). INA bacteria were reported to be relatively more abundant in rainfall than in the air at a given site (Stephanie and Waturangi, 2011), which may indicate that INA bacteria are preferentially incorporated into rainfall than other bacteria.

106 Based on these studies, biological IN are undoubtedly present throughout the water cycle. 107 They represent an important fraction of the pool of high subzero temperature IN where they 108 were unambiguously quantified: in the air at low altitude, and in precipitation. However, our 109 knowledge about their relative abundance in clouds is still scarce, which limits the evaluation 110 of their impact on hydrological cycles using modeling approaches (Hoose et al., 2010; 111 Phillips et al., 2008). As stressed by DeMott and Prenni (2010), it is technically not possible 112 to provide any realistic concentration of airborne IN particles at the altitude of a cloud from 113 measurements in precipitation, due to possible dilution/concentration effects and to non-114 nucleation particle scavenging. With the objective to provide quantitative data of IN 115 concentration in clouds that could be utilized for modeling purposes, cloud water samples 116 were collected throughout the year and under various meteorological situations from the 117 summit of puy de Dôme mountain in France (1465 m a.s.l). Total and biological IN 118 concentrations were measured by the droplet-freezing method (immersion freezing mode) between -5°C and -14°C. Data were then analyzed against meteorological, chemical and
biological variables, and maximum possible values of INA bacteria concentration were
inferred.

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123 **2.** Materials and methods

124 **2.1. Cloud water sampling and meteorological measurements**

125 Twelve random cloud events non-orographic and non-precipitating at the sampling site during collection were sampled at the puy de Dôme station (45° 46' 20" North, 2° 57' 57" East, 1465 126 127 m above sea level; see Figure S1 for localization) between June 2011 and October 2012. 128 These were identified as samples #76 to #87 following the numbering of cloud events 129 sampled at puy de Dôme since 2001, for which chemical and microbiological datasets are 130 http://wwwobs.univ-bpclermont.fr/SO/beam/data.php.Sampling publically available at 131 operations were decided after visual estimation of cloud optical thickness at the sampling site. 132 Cloud droplets were selectively collected using single-stage aluminum droplet impactors (cut-133 off diameter: ~7 µm, Kruisz et al., 1992) sterilized by autoclave, as in Vaïtilingom et al., 134 (2012) and previous studies from our group. Meteorological data was recorded continuously 135 during sampling by the Observatory of the Globe of Clermont-Ferrand (OPGC)'s atmospheric 136 station. The following parameters were considered in our analysis: temperature (T; Vaisala), 137 relative humidity (RH; Vaisala), liquid water content (LWC; Gerber PVM-100), and 138 cumulated precipitation downwind the sampling site (Figure S1). For most samples, LWC 139 was not available. Sample collection rates were not judge reliable enough for estimating it due 140 to variations of collection efficiency with droplet size distribution or ice formation (Kruisz et 141 al., 1992). So, in those cases, LWC was approximated from archive data by assigning the 142 minimum, average or maximum value observed in clouds at puy de Dôme (0.1, 0.3 or 0.6 g 143 m⁻³, respectively; **Deguillaume et al., 2014**) depending on the sample collection rate.

The global meteorological context was examined through 72-hour back-trajectories of the air masses sampled using the HYSPLIT model (HYbrid Single-Particle Lagrangian Integrated Trajectory) with GDAS1 meteorological data archive and default settings (**Draxler and Rolph, 2010**) and satellite visible images of Europe and France from Eumetsat, available for academic purpose at http://www.woksat.info/wwp.html.

149 **2.2.** Physico-chemical characterization and total cell counts

150 Cloud water samples were recovered either liquid or frozen onto the impaction plate 151 depending on ambient temperature during sampling. For each sample, pH was measured 152 (Consort multiparameters C830) and major inorganic and organic ions were examined by ion 153 chromatography (Dionex DX320 for anions and Dionex ICS1500 for cations).

Total bacteria were counted by epifluorescence microscopy on DAPI stained samples as in Vaïtilingom et al., (2012). Directly after collection, samples were fixed by the addition of 2% formaldehyde (final concentration; from 20% stock solution prepared in phosphate buffer 0.1 M, pH 7.0), and incubated in the presence of 2.5 µg mL⁻¹ of DAPI (4',6-diamino-2phenylindol) in the dark for at least 20 min before filtration on GTBP black filters (0.22 µm porosity; Millipore). Filters were then mounted on microscope slides and observed under UVepifluorescence microscopy (λ_{exc} = 365 nm; λ_{em} = 420 nm) (Leica DM-IRB).

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162 **2.3. Droplet-freezing assays**

163 The ice nucleation activity (INA) of the cloud water samples in the immersion freezing mode 164 was determined within 2 hours after collection following the well-tried droplet-freezing 165 method (Vali, 1971). Thirty-two to 160 drops (Table 2) of 20 µL were distributed in 0.2 mL 166 microtubes designed for high thermal conductivity and preventing aerial contamination and 167 evaporation (Stopelli et al., 2013). These were placed in a cooling bath (Julabo F34-ED) at 168 decreasing temperatures from -5°C to -14°C, with 1°C intervals for 8 min. The tubes were 169 visually inspected at the end of each temperature step and those still liquid were counted. The concentration (mL^{-1}) of ice nuclei C_{IN} at the temperature T in the suspensions was calculated 170 171 using the equation in Vali (1971): $C_{IN} = [\ln (N_{total}) - \ln (N_{tiouid})]_T / V \times (1 / D_f)$

where N_{total} is the total number of droplets, N_{liquid} the number of droplets still liquid after 8 minutes at the temperature T, V the volume of the droplets assayed (mL) and D_f the dilution factor of the suspension. Under our experimental conditions, the quantification limits ranged from 1.6 to 173.3 IN mL⁻¹ in the case where 32 droplets were assayed, and from 0.3 to 253.8 IN mL⁻¹ in the case where 160 droplets were assayed. Negative controls consisted of ultrapure sterile water droplets and these remained liquid over all the range of temperatures investigated.

180 **2.4. Biological IN quantification**

181 For each sample, the concentration of biological IN (INA_{bio}) was calculated as the difference 182 between the concentration of IN measured in untreated sample (INAtotal) and the concentration 183 of IN measured after heating for 10 minutes at 95°C (INA_{heated}), as in Christner et al. (2008a) 184 and in Garcia et al. (2012). Heat denatures protein structures, so it eliminates at least a 185 certain fraction of biological IN without altering non-biological material. When [(INA_{heated})_{T-1} 186 - $(INA_{heated})_T$] exceeded $[(INA_{total})_{T-1} - (INA_{total})_T]$, this calculation artificially led to a 187 decrease in the concentration of INAbio at T-1 compared to T and values of INAheated were 188 corrected for being consistent with the values of INAtotal. Following this rule, three values of 189 INA_{heated} were corrected: -12°C in sample #79, -10°C in sample #82 and -11°C in sample #86.

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191 **2.5. Statistical analyses**

Principal Component Analysis was made using R software version 2.12.2 (R Core Team,
2011), and non-parametric tests (Pearson's rank correlation test, Mann-Whitney test) were
performed as most data were not normally distributed and the number of samples was quite
low (< 30), using PAST version 2.04 (Hammer et al., 2001).

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197 **3. Results and discussion**

3.1. Main characteristics of the cloud water samples

199 Twelve cloud water samples were collected between 29 June 2011 and 10 October 2012 from 200 cloud events lasting in total approximately 20 to 180 hours at the puy de Dôme, based on 201 relative humidity measurements (Table 1; Figure S1). The meteorological context associated 202 with each sampling period is presented in Fig S1. Most of the air masses sampled originated 203 from West (Atlantic Ocean) and travelled over different continental areas in Europe before 204 reaching the puy de Dôme, following different trajectories. Sampling operations were started 205 about 5 to 110 hours after clouds arrived at the sampling site. Cloud water was then collected 206 for 1h10 to 5h15, and after sampling, the sampling site remained embedded in cloud for 5 to 207 more than 160 additional hours.

208 Only clouds that were non-precipitating at the sampling site were collected, but at some 209 occasions rainfalls occurred in the vicinity. The amount of precipitation that felt at 5 sites 210 downwind the sampling site around the sampling period of time was measured (see Fig S1 and Table 1); most cloud events were not or slightly precipitating in this area, with less than 1 mm of rain accumulated considering the 5 rain gauges together. In contrast, it reached 1.6 and 7 mm for samples #76 and #77, respectively. Ambient temperature during sampling ranged from -1.5°C to 13.3°C, so some samples consisted of ice formed upon impaction on the collectors (samples #80 through #84); other samples were collected as liquid.

Bacteria concentration in the samples ranged between 1.65×10^3 to 3.37×10^4 mL⁻¹. The 216 chemical composition varied greatly from one sample to another (Table S1): pH ranged from 217 218 4.6 to 6.2, which are typical values for cloud water (e.g. Deguillaume et al., 2014). 219 Ammonium (16.8 to 531.1 µM), sodium (0.6 to 145.7 µM), nitrate (1.0 to 126.0 µM) and 220 sulfate (0.5 to 52.2 µM) dominated among inorganic ions, and formate was the most abundant 221 dissolved carboxylic acid (3.2 to 109.6 µM). The chemical signature of the samples attested 222 of mixed influences from oceanic and continental sources, the respective contributions to the 223 global chemical composition of which were more or less marked depending on the origin of 224 the air mass.

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3.2. Quantification of total and biological ice nuclei

227 The total concentration of IN active between -5°C and -14°C was determined by droplet 228 freezing assays. In 11 of the 12 cloud samples (92%), the onset temperature of freezing (i.e. 229 temperature at which the first droplet froze) was -8°C or warmer. Only sample #87 started to 230 freeze at colder temperature (-11°C) (Table 2; Fig. 2). Ice initially formed due to the presence of 0.6 to 8.5 IN mL⁻¹ (Table 2; Fig. 3a). Two samples (#81 and #83) were clearly outlying 231 with much higher IN concentrations ($\sim 70 \text{ mL}^{-1}$ at -8° C). Overall, the onset freezing 232 temperature was significantly correlated with the concentration of IN in the sample at the 233 234 warmest temperatures (Table S2; p < 0.03, $0.66 < \rho < 0.79$ with IN concentrations at -8°C and -9°C). After correction for LWC (Fig. 3b), the concentration of IN per volume of cloud air 235 ranged from 0.06 to more than 71.1 m⁻³ between -6°C and -14°C. This is in the range of 236 237 concentrations typically observed in the air at high altitude (Fig. 1) (Bowers et al., 2009; Xia 238 et al., 2013), and one order of magnitude lower than the concentrations measured at low 239 altitude (Garcia et al., 2012). Rain and surface snow samples analyzed using similar methods 240 by Christner et al. (2008a, 2008b) had total IN concentrations of about ~1 to ~300 per liter 241 of water at -8°C, i.e. 2 orders of magnitude fewer than in our cloud water samples. This 242 probably resulted from the relative dilution of insoluble particles in precipitation compared to 243 cloud water (Flossmann and Wobrock, 2010), and from differences in sample handling: Christner et al. filtered samples for concentrating particles larger than 0.22 μm, so smaller
IN particles were missed, among which some could have originated from bacteria (Phelps et
al., 1986). In addition, it is possible that a fraction of IN particles was not recovered from the
filters.

248 Heating samples for 10 min at 95°C invariably decreased the highest temperature of freezing 249 (Fig. 2), in general by 3°C to 4°C, and by 1°C (sample #81) to more than 4°C in samples #77 250 and #79, respectively (Table 2). This indicated that heat sensitive IN (thereafter termed 251 biological IN) were systematically responsible for freezing at the warmest temperatures. The 252 proportion of biological IN in samples did not depend on the absolute total IN concentration 253 (Table S2; p > 0.05). As other IN were activated at lower temperature, the relative 254 contribution of biological IN decreased with decreasing temperature, from 97% to 100% of 255 the total number of IN active at -8°C to as low as 77% at -12°C (Table 2). These are in 256 accordance with observations of IN in the air (Garcia et al., 2012) and in precipitation 257 (Christner et al., 2008a, 2008b).

258 The average absolute concentrations of biological and non-biological IN are represented on 259 Fig. 4. Since heat treatment does probably not inactivate every IN site of biological material 260 such as fungi or pollen (Pummer et al., 2012), the concentrations of biological IN reported 261 here should be seen as conservative (i.e. lowest possible) values. Clearly, non-biological (i.e. 262 heat resistant) particles contribution became significant only around -12°C and colder. We 263 examined the influence of the different variables measured on the IN content of our samples. 264 Table S2 shows Spearman's correlation matrices (p-values, p and n) linking the variables 265 together. Among noticeable correlations, coldest sampling temperatures were linked with highest onset freezing temperatures (Table S2; Spearman's rank correlation test; n = 12, p =266 0.0361, $\rho = -0.61$). Consistently, IN activity was higher in samples collected frozen than in 267 268 samples collected liquid: higher IN concentrations at the highest temperatures and warmer 269 onset freezing temperature (Mann-Whitney's test; medians = -8° C and -6° C, respectively). 270 This result is quite surprising if, logically, one considers that the most active IN should be 271 activated and precipitated first, and so that cold air masses should be depleted in highest 272 temperatures IN compared to warmer air masses. Bigg (1996)'s observations of airborne IN in 273 the Artic indeed suggested that such selection process occurs in the atmosphere. In our case, 274 the minimum temperature during sampling was $> -2^{\circ}C$ (Table 1), so likely still too warm for 275 leading to any temperature partitioning of IN in the clouds sampled. So, despite the fact that 276 the influence of freezing on further IN concentration measurements in our samples cannot be totally excluded, it is possible that the relationship observed results from a higher expression
level of IN proteins by bacteria in the coldest clouds (Nemecek-Marshall et al., 1993).

Clouds which precipitated downwind the puy de Dôme had globally a shorter lifespan at the
sampling site (Table S2). Despite the potential influence of IN on precipitation, no correlation
was found here between IN concentrations and local rainfalls.

- 282 Principal component analysis (PCA) revealed 2 different groups of IN depending on their 283 temperature of activity, with a net separation between -10°C and -11°C (Fig. S2). This 284 demonstrated differences in the origin of the two sets of IN and so probably in their nature as well. The clear positive correlation existing between $IN_{T\leq-11^{\circ}C}$ and soluble inorganic ions 285 286 concentrations supports their inorganic composition (Fig. S2). The concentrations of IN_{T>-11°C}, i.e. biological IN, and Ca²⁺ were positively correlated, while the trend toward Chloride, 287 288 which mostly originates from marine environment (Warneck, 1999) was negative (Table S2). 289 These tend to situate the sources of biological IN on the continent, at the puy de Dôme site, 290 probably including both regional and more distant areas.
- 291 Considering cloud droplets as spherical, we propose an extrapolation of IN concentration per 292 droplet based on the total IN concentration measured. Thus, for a population of cloud droplets 293 distributed as a single mode of 20 μ m in diameter, at the temperature of -8°C there was a 294 maximum of 1 IN every ~3×10⁶ droplets, and the median value corresponded to 1 IN every 295 ~5×10⁷ droplets.
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297 **3.3. Estimation of the contribution of bacteria to biological IN**

298 Joly et al. (2013) proposed an estimation of the concentration of INA bacteria in clouds based on laboratory results. It was proposed that between 0 and \sim 500 bacterial cells mL⁻¹ could act 299 300 as IN in cloud water at -10°C. This very wide range needed clarification. In order to 301 discriminate bacterial IN from other biological IN, Christner et al. (2008a, 2008b) suggested 302 treating samples with lysozyme. This was intended to alter bacterial cell wall and selectively 303 eliminate bacterial IN. Lysozyme is indeed responsible for the lysis of peptidoglycans by 304 hydrolyzing the 1,4-β-linkages between N-acetylmuramic acid and N-acetylglucosamine, so 305 it is particularly active towards Gram-positive bacteria; its efficiency towards Gram-negative species is much less marked and it requires additional treatments incompatible with droplet 306 307 freezing assays (Masschalck and Michiels, 2003; Repaske, 1956). So far, all INA bacteria 308 described in literature including those encountered in clouds were Gram-negative species 309 (Cochet and Widehem, 2000; Joly et al., 2013). We verified lysozyme efficiency in altering
310 INA of bacteria on 2 of our cloud samples and on laboratory cultures of INA Gamma311 Proteobacteria (Gram-negative) isolated from cloud water (those reported in Joly et al.,
312 2013): lysozyme had no effect on the freezing profiles (not shown). So, this treatment was
313 finally judged not reliable enough here for suppressing specifically bacterial INA and it was
314 not applied further.

In our samples, bacteria concentration ranged from 1.6×10^3 to 3.4×10^4 mL⁻¹, which is within 315 316 the range of concentrations typically observed in cloud water at the puy de Dôme site 317 (Vaïtilingom et al., 2012) (Table 1). As expected, since only a small proportion of bacteria is 318 actually IN active and that this can be very variable even within INA+ bacterial strains 319 cultures (e.g. Joly et al., 2013), IN concentration did not vary with bacteria concentration. 320 Rather, it was significantly correlated with most ion concentrations, particularly strongly with 321 K^+ and NO₃⁻ (Table S2), suggesting similar source, i.e. continental origin. In order to provide 322 an estimation of the possible proportion of INA bacteria in our samples, biological IN 323 concentration was normalized to bacteria concentration (Table 3). This has to be considered 324 as an upper estimate as it obviously assumes only one IN site per cell, which is the most likely 325 (Hartmann et al., 2013), and it ignores the fact that a certain but unknown fraction of 326 biological materials other than bacteria could also have been inactivated by heat and 327 contributed to the population of biological IN, such as cell fragments for example (Hartmann 328 et al., 2013). At the temperature of -6°C, a maximum of 0.1% of the bacteria could have been 329 responsible for freezing (sample #82). This proportion reached maxima of 1.24% at -9°C and 3.06% at -12°C (in samples #83 and #85, respectively), or about 200 INA cells mL⁻¹. In the 330 331 air over vegetated areas, INA bacteria were estimated to contribute only ~0.002% of the total 332 cells (Garcia et al., 2012), and this proportion falls to less than 0.001% at high (Xia et al., 333 **2013**). In snowfall, comparable estimations gave a very similar fraction of 0.4% of bacterial cells acting as IN between -4°C and -7°C (Christner et al., 2008a) (Fig. 1). In laboratory 334 335 cultures of INA bacteria, the proportion of individual cells actually acting as IN largely 336 depends on the strain. Except in some exceptionally efficient microorganisms for which this 337 can reach up to more than 4%, this is often around 1% at -9°C, and in general well below 0.1% at -6°C (Joly et al., 2013; Šantl-Temkiv et al., 2009; Yankofsky et al., 1981). So, at 338 339 temperatures below -6°C, the proportion of INA bacterial cells in clouds basically matched 340 laboratory cultures of INA+ strains.

Low pH (i.e. pH ~4) was shown to negatively impact bacterial INA (**Turner et al., 1990**). This suggested attenuation of bacterial IN efficiency in polluted clouds due to anthropogenic emissions responsible for acidification (**Attard et al., 2012**). Among the set of clouds investigated here, only sample #79, with a pH of 4.6, was clearly under influence of Human emissions. Yet its freezing profile was not different from others, and on the whole we found no significant relationship between pH and total or biological IN concentrations (Spearman's correlation test; the p-values ranged between 0.46 and 1 between -6 and -13°C).

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349 **4. Conclusion**

350 To our knowledge, this study constitutes the first quantitative dataset of biological IN 351 measured directly in cloud water. A basic but straightforward experimental set up allowed to determine that the concentration of total IN varies in general between ~ 1 and $\sim 200 \text{ mL}^{-1}$ 352 353 at -10°C. As previously observed in the air (Garcia et al., 2012) and in precipitation 354 (Christner et al., 2008a), heat-sensitive material, i.e. biological particles, were systematically 355 responsible for freezing at the warmest temperatures and largely dominated the population of 356 IN particles at temperatures down to -11°C. These data support the possibility that biological 357 material could contribute to clouds evolution by triggering precipitation at temperatures close 358 to 0°C.

A certain proportion of the biological IN detected in the cloud water samples were likely bacterial cells. Some specimens were indeed previously recovered by culture from several clouds collected at that site (**Joly et al., 2013**). Assuming that the biological IN observed were all bacterial cells, between 0% and about 1.5% of the total bacteria were IN at -10°C. This extends to much higher values than the proportion of around 0.001% and 0.4% proposed for air (**Garcia et al., 2012; Xia et al., 2013**) and precipitation, respectively (**Christner et al., 2008a**).

Our experimental procedure by conventional droplet freezing assay only allowed processing a limited number of samples, which also limited our conclusions. In addition, seasonality was not approached here. The development of online measurements is opening new perspectives in the prospection for atmospheric IN, and, in the near future, it should greatly help elucidating their role and environmental drivers (e.g., **Bundke et al., 2010; Huffman et al., 2013**).Such estimates of in-cloud biological IN concentrations will allow the community of atmospheric scientists to explore, e.g. using cloud-resolving models, the extent to which these particles can contribute to cloud glaciation, to modification of cloud radiative properties andto regional precipitation patterns.

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530 Table 1. Main characteristics of the cloud events sampled. Samples recovered as ice formed

531 upon impaction in the sampler are indicated in italic. See detailed ion composition in Table

532 SM1.

		Sampling period (UTC)				Cloud period (UTC) ^a								
Sample	Date	From	To	Sampling duration (h)	Volume sampled (mL)	From	To	Cloud event duration (h) ^a	Time in cloud before sampling (h) ^a	Time in cloud after sampling (h) ^a	Precipitation accumulated in the vicinity (mL) ^b	Mean sampling temperature (°C)	Mean LWC during sampling (g m ^{.3})	Bacteria concentration (mL ⁻¹)
# 76	29-Jun-11	6:30 AM	11:45 AM	5.25	> 200	6/28/11 10:00 PM	6/30/11 0:00 AM	26	8.5	12.3	1.6	11.5	0.6	n.d.*
# 77	7-Jul-11	1:50 PM	3:00 PM	1.17	15	7/7/11 9:00 AM	7/8/11 6:00 AM	21	4.8	15	7	12.0	0.1	n.d.*
# 78	20-Jul-11	7:30 AM	9:10 AM	1.67	47	7/19/11 3:00 PM	7/23/11 4:00 PM	97	16.5	78.8	0.2	8.3	0.3°	12355
# 79	7-Nov-11	1:00 PM	2:30 PM	1.50	193	11/6/11 8:00 AM	11/8/11 11:00 AM	51	29	20.5	0.4	7.0	0.6 ^c	10825
# 80	20-Jan-12	12:45 PM	3:00 PM	2.25	55	01/18/12 11:00 PM	01/26/12 0:00 AM	169	37.7	129	0	-0.4	0.3^{c}	9980
# 81	23-Jan-12	1:00 PM	4:00 PM	3.00	53	01/18/12 11:00 PM	01/26/12 0:00 AM	169	110	56	0	-1.2	0.1^{c}	33724
# 82	19-Mar-12	12:10 PM	4:10 PM	4.00	45	3/17/12 11:00 PM	3/21/12 11:00 AM	84	37.2	42.8	0.2	-1.5	0.1^{c}	1648
# 83	4-Apr-12	6:10 AM	9:20 AM	3.17	29	4/3/12 11:00 PM	4/6/12 12:00 PM	61	7.2	50.7	0.25	-0.4	0.1^c	14914
# 84	18-Apr-12	8:10 AM	12:15 PM	4.08	31	4/17/12 6:00 PM	4/25/12 6:00 AM	180	14.2	161.8	0	0.2	0.1^{c}	3902
# 85	25-Jun-12	1:35 PM	5:00 PM	3.42	66	6/25/12 01:00 AM	6/26/12 12:00 AM	35	12.6	19	0	13.3	0.3°	4474
# 86	13-Sep-12	7:50 AM	9:50 AM	2.00	75	9/12/12 7:00 PM	9/13/12 3:00 PM	20	12.8	5.2	0.8	6.0	0.6 ^c	5199
# 87	10-Oct-12	8:40 AM	9:50 AM	1.17	70	10/08/12 9:00 PM	10/11/12 0:00 AM	63	35.7	26.2	0	9.4	0.6 ^c	19658

a: Defined as RH > 95% based on hourly average (see Fig. S1).

b: Sum of precipitation accumulated at 5 rain gauge stations in the vicinity of puy de Dôme (Royat, Farnette, Sayat, Trois Ponts and Blanzat) (see Fig. S1).

c: Estimation from sample collection rate and puy-de-Dôme data archive.

533 * n.d.: not determined.

Table 2. Total IN concentration and proportion of heat-sensitive IN in the cloud water
samples between -5°C and -14°C. Values below the detection limit are presented as '0' for
visual clarity, and '>' indicates values higher than our quantification limit.

Sample	n =*	Onset freezing temperature (°C)	Onset freezing temperature after heat treatment (°C)	Decrease of onset freezing temperature by heat treatment (°C)	IN mL ⁻¹ [total (% heat sensitive)]										
				ase 1pei trea		Temperature (°C)									
				Decre. tem	-5°C	-6°C	-7°C	-8°C	-9°C	-10°C	-11°C	-12°C	-13°C	-14°C	
# 76	32	-8	-12	4	0 (- %)	0 (- %)	0 (- %)	4.9 (100%)	18.7 (100%)	31.6 (100%)	45.0 (100%)	118.4 (99%)	n.d.	n.d.	
# 77	32	-8	<-12	>4	0(-%)	0 (- %)	0(-%)	4.9 (100%)	12.3 (100%)	14.4 (100%)	92.8 (100%)	138.6 (100%)	n.d.	n.d.	
# 78	32	-8	-11	3	0(-%)	0 (- %)	0 (- %)	1.6 (100%)	8.5 (100%)	16.5 (100%)	26.1 (94%)	69.3 (88%)	n.d.	n.d.	
# 79	32	-8	<-12	>4	0(-%)	0 (- %)	0 (- %)	3.2 (100%)	8.5 (100%)	12.3 (100%)	16.5 (100%)	16.5 (100%)	n.d.	n.d.	
# 80	32	-8	-11	3	0(-%)	0 (- %)	0 (- %)	4.2 (100%)	12.3 (100%)	16.5 (100%)	18.7 (92%)	53.4 (91%)	n.d.	n.d.	
# 81	32	-7	-8	1	0(-%)	0 (- %)	8.5 (100%)	63.4 (97%)	>173.3 (<99 %)	>173.3 (<97%)	>173.3 (<95%)	>173.3 (<92 %)	n.d.	n.d.	
# 82	32	-6	-9	3	0(-%)	1.6 (100%)	6.7 (100%)	10.4 (100%)	18.7 (74%)	21.1 (66%)	28.8 (70%)	53.4 (77%)	n.d.	n.d.	
# 83	160	-6	-10	4	0(-%)	0.6 (100%)	7.4 (100%)	73.2 (100%)	184.4 (100%)	219.1 (99%)	>253.8 (<97%)	>253.8 (<93%)	n.d.	n.d.	
# 84	64	-6	-9	3	0(-%)	1.6 (100%)	2.4 (100%)	11.4 (100%)	16.5 (90%)	18.7 (92%)	30.2 (95%)	41.3 (88%)	110.6 (66%)	>207.9	
# 85	160	-8	-12	4	0(-%)	0 (- %)	0 (- %)	2.2 (100%)	3.2 (100%)	5.3 (100%)	34.7 (100%)	138.6 (99%)	>253.8 (<91%)	>253.8 (<45%	
# 86	32	-7	-10	3	0 (- %)	0 (- %)	1.6 (100%)	3.2 (100%)	8.5 (100%)	14.4 (89%)	14.4 (89%)	18.7 (83%)	49.0 (93%)	>173.3 (<56 %	
# 87	32	-11	-13	2	0(-%)	0 (- %)	0 (- %)	0 (- %)	0 (- %)	0 (- %)	3.2 (100%)	28.8 (100%)	83.7 (72%)	>173.3 (<46 %	
Med	ian	-8	-10,5	3				4.9	>12.3	>16.5	>29.5	>61.4	>97.2	>190.6	
11100		Ŭ	- 0,0	5	0 (- %)	0 (- %)	0 (- %)	(100%)	(100%)	(100%)	(<96 %)	(<92%)	(<82%)	(<51%)	
Mi	in	-11	-13	1	0 (- %)	0 (- %)	0 (- %)	0 (- %)	0 (- %)	0 (- %)	3.2 (70%)	16.5 (91%)	49.0 (66%)	>173.3 (<45%)	
Ma	ax	-6	-8	>4		1.6	8.5	73.2	184.4	219.1	>253.8	>253.8	>253.8	>253.8	
			0	~ 7	0 (- %)	(100%)	(100%)	(100%)	(100%)	(100%)	(100%)	(100%)	(93%)	(<56%	

*Number of 20 µL droplets assayed by immersion freezing assays

n.d.: not determined.

539

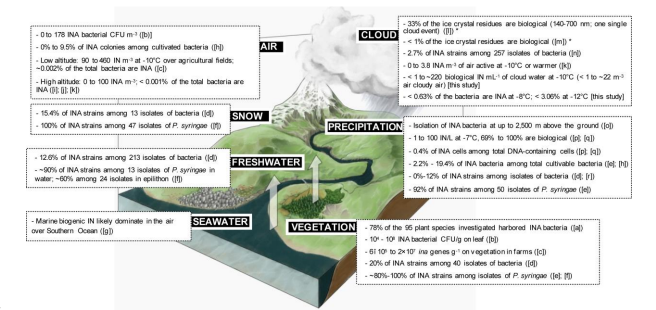
540 Table 3. Inferred maximum possible fraction of INA bacteria among total bacteria in the samples based on heat-sensitive IN concentrations and on total bacteria counts. A '>' indicate 541 values higher than experimental quantification limit for heat-sensitive IN. 542

	Temperature (°C)										
Sample	-6°C	-7°C	-8°C	-9°C	-10°C	-11°C	-12°C	-13°C	-14°C		
# 78	0.00%	0.00%	0.01%	0.07%	0.13%	0.20%	0.49%	>0.49%	>0.49%		
# 79	0.00%	0.00%	0.03%	0.08%	0.11%	0.15%	0.15%	>0.15%	>0.15%		
# 80	0.00%	0.00%	0.05%	0.12%	0.17%	0.17%	0.49%	>0.49%	>0.49%		
# 81	0.00%	0.03%	0.18%	>0.51%	>0.51%	>0.51%	>0.51%	>0.51%	>0.51%		
# 82	0.10%	0.41%	0.63%	0.84%	0.84%	1.23%	2.49%	>2.49%	>2.49%		
# 83	0.00%	0.05%	0.49%	1.24%	1.46%	>1.66%	>1.66%	>1.66%	>1.66%		
# 84	0.04%	0.06%	0.29%	0.38%	0.44%	0.73%	0.93%	1.86%	>2.95%		
# 85	0.00%	0.00%	0.05%	0.07%	0.12%	0.77%	3.06%	>5.15%	>5.15%		
# 86	0.00%	0.03%	0.06%	0.16%	0.25%	0.25%	0.30%	0.88%	>1.87%		
# 87	0.00%	0.00%	0.00%	0.00%	0.00%	0.02%	0.15%	0.31%	>0.41%		
Mean	0.01%	0.06%	0.18%	>0.35%	>0.40%	>0.57%	>1.02%	>1.40%	>1.62%		

IN/ bacterial cell

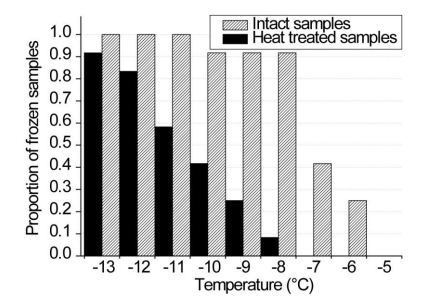
Temperature (°C)

543



544

Figure 1. Schematic summarizing our current knowledge about the abundance of biological 545 546 IN active at temperatures $\geq -10^{\circ}$ C in the different environmental links of the water cycle. A 547 "' indicates data relative to ice crystal residues in clouds at much colder temperatures. [a] 548 Lindow et al. (1978); [b] Lindemann et al. (1982); [c] Garcia et al. (2012); [d] Maki and 549 Willoughby (1978); [e] Constantinidou et al. (1990); [f] Morris et al. (2008); [g] Burrows et 550 al. (2013); [h] Stephanie and Waturangi (2011); [i] Bowers et al. (2009); [j] Conen et al. (2012); [k] Xia et al. (2013); [l] Pratt et al. (2009); [m] Cziczo et al. (2013); [n] Joly et al. 551 552 (2013); [o] Sands et al. (1982); [p] Christner et al. (2008a); [q] Christner et al. (2008b); [r] 553 Šantl-Temkiv et al. (2009).



555

557

556 Figure 2. Cumulative proportion of cloud samples for which at least one freezing event was

observed during IN assays, in the absence of treatment (shaded bars) or after heating at 95°C

558 for 10 minutes (black bars).

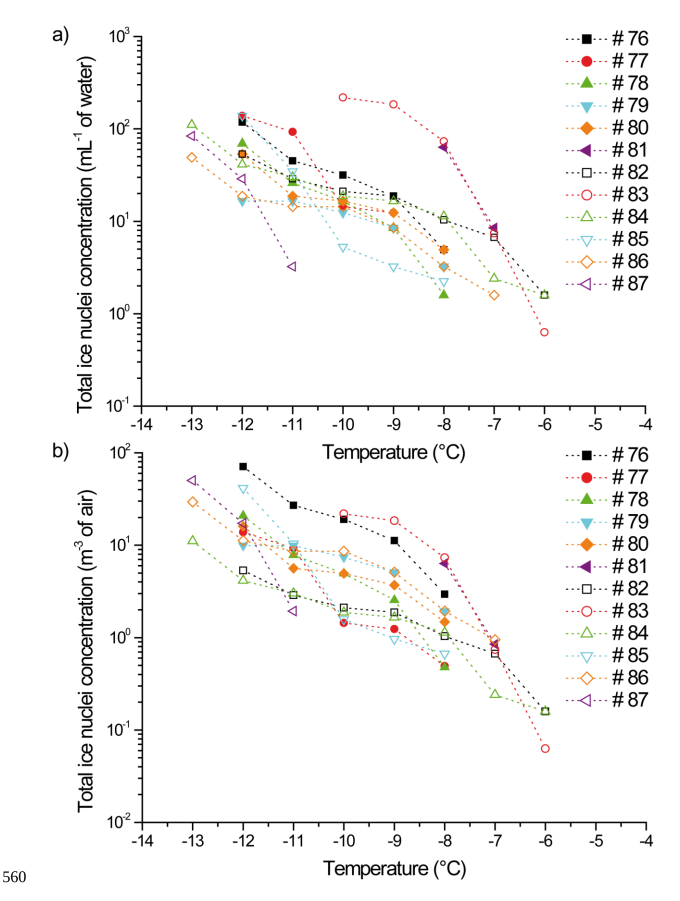
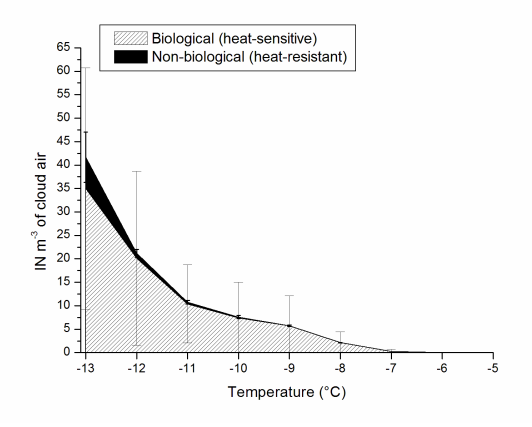


Figure 3. Cumulative concentration of total IN in the cloud samples. (a) per volume of water sample (mL⁻¹) and (b) per corresponding volume of cloud air (m⁻³).



563

Figure 4. Mean cumulative concentrations of biological (heat-sensitive. shaded area) and nonbiological (heat-resistant, black area) IN in clouds (n=12) per volume of air. The sum of the two categories corresponds to the mean concentration of total IN. The lower bound was considered for values below the detection limit.