1	Experimental observation of transient δ^{18} O interaction between snow and
2	advective airflow under various temperature gradient conditions
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13	Abstract
14	Stable water isotopes (δ^{18} O) obtained from snow and ice samples of polar regions
15	are used to reconstruct past climate variability, but heat and mass transport processes
16	can affect the isotopic composition. Here we present an experimental study on the effect
17	of airflow on the snow isotopic composition through a snow pack in controlled
18	laboratory conditions. The influence of isothermal and controlled temperature gradient
19	conditions on the $\delta^{18}O$ content in the snow and interstitial water vapor is elucidated. The
20	observed disequilibrium between snow and vapor isotopes led to the exchange of

isotopes between snow and vapor under non-equilibrium processes, significantly changing the δ^{18} O content of the snow. The type of metamorphism of the snow had a significant influence on this process. These findings are pertinent to the interpretation of the records of stable isotopes of water from ice cores. These laboratory measurements

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suggest that a highly resolved climate history is relevant for the interpretation of the
snow isotopic composition in the field.

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28 Keywords: snow, isotope, isothermal, metamorphism, advection, tomography, post-depositional process

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30 **1. Introduction**

Water stable isotopes in polar snow and ice have been used for several decades as 31 proxies for global and local temperatures (e.g. Dansgaard, 1964; Lorius et al., 1979; 32 Grootes et al., 1994; Petit et al., 1999; Johnsen et al., 2001; EPICA Members, 2004). 33 However, the processes that influence the isotopic composition of precipitation in high-34 latitude are complex, making direct inference of paleo temperatures from the isotopic 35 record difficult (Cuffey et al., 1994; Jouzel et al., 1997, 2003; Hendricks et al., 2000). 36 Several factors affect the vapor and snow isotopic composition, which give rise to ice 37 core isotopic composition, starting from the process of evaporation in the source region, 38 transportation of the air mass to the top of the ice sheet, and post-depositional processes 39 (Craig and Gordon, 1964; Merlivat and Jouzel, 1979; Johnsen et al. 2001; Ciais and 40 Jouzel, 1994; Jouzel and Merlivat, 1984; Jouzel et al., 2003; Helsen et al., 2005, 2006, 41 2007; Cuffey and Steig, 1998; Krinner and Werner, 2003). Mechanical processes such 42 as mixing, seasonal scouring, or spatial redistribution of snow can alter seasonal and 43 44 annual records (Fisher et al., 1983; Hoshina et al., 2014). Post-depositional processes associated with wind scouring and snow redistribution are known to introduce a "post-45 depositional noise" in the surface snow. Comparisons of isotopic records obtained from 46 closely located shallow ice cores have allowed for estimation of a signal-to-noise ratio 47

and a common climate signal (Fisher and Koerner, 1988, 1994; White et al., 1997;
Steen-Larsen et al., 2011; Sjolte et al., 2011; Masson-Delmotte et al., 2015). After
deposition, interstitial diffusion in the firn and ice affects the water-isotopic signal but
back-diffusion or deconvolution techniques have been used to establish the original
isotope signal (Johnsen, 1977; Johnsen et al., 2000).

53 Snow is a bi-continuous material consisting of fully connected ice crystals and pore space (air) (Löwe et al., 2011). Because of the proximity to the melting point, the high 54 vapor pressure causes a continuous recrystallization of the snow microstructure known 55 as snow metamorphism, even under moderate temperature gradients (Pinzer et al., 56 57 2012). The whole ice matrix is continuously recrystallizing by sublimation and deposition, with vapor diffusion as the dominant transport process. Pinzer et al. (2012) 58 showed that a typical half-life of the ice matrix is a few days. The intensity of the 59 recrystallization is dictated by the temperature gradient and this can occur under mid-60 latitude or polar conditions. Temperature, and geometrical factors (porosity and specific 61 surface area) also play a significant role (Pinzer and Schneebeli, 2009; Pinzer et al., 62 2012). 63

The interpretation of ice core data and the comparison with atmospheric model 64 results implicitly rely on the assumption that the snowfall precipitation signal is 65 preserved in the snow-ice matrix (Werner et al., 2011). Classically, ice-core stable-66 isotope records are interpreted as reflecting precipitation-weighted signals, and 67 compared to observations and atmospheric model results for precipitation, ignoring 68 snow-vapor exchanges between surface snow and atmospheric water vapor (e.g. Persson 69 et al., 2011). However, recent studies carried out on top of the Greenland and Antarctic 70 ice sheets combining continuous atmospheric water vapor isotope observations with 71 daily snow surface sampling document a clear day-to-day variation of isotopic 72

composition of surface snow between precipitation events as well as diurnal change in 73 the snow isotopes (Steen-Larsen et al., 2014a; Ritter et al., 2016; Casado et al., 2016). 74 This effect was interpreted as being caused by the uptake of the synoptic-driven 75 atmospheric water-vapor isotope signal by individual snow crystals undergoing snow 76 metamorphism (Steen-Larsen et al., 2014a) and the diurnal variation in moisture flux 77 78 (Ritter et al., 2016). However, the impact of this process on the isotope-temperature reconstruction is not yet sufficiently understood, but crucial to constrain. This process, 79 compared to interstitial diffusion (Johnsen, 1977; Johnsen et al., 2000), will alter the 80 isotope mean value. The field observations challenge the previous assumption that 81 82 sublimation occurred molecular layer-by-layer with no resulting isotopic fractionation (Dansgaard, 1964; Friedman et al., 1991; Town et al., 2008; Neumann and Waddington, 83 2004). It is assumed that the solid undergoing sublimation would not be unduly enriched 84 in the heavier isotope species due to the preferential loss of lighter isotopic species to 85 the vapor (Dansgaard, 1964; Friedman et al., 1991). Because self-diffusion in the ice is 86 about three orders of magnitude slower than molecular diffusion in the vapor, the 87 amount of isotopic separation in snow is assumed to be negligible. 88

Snow has a high permeability (Calonne et al., 2012; Zermatten et al., 2014), which 89 facilitates diffusion of gases and, under appropriate conditions, airflow (Gjessing, 1977; 90 91 Colbeck, 1989; Sturm and Johnson, 1991; Waddington et al., 1996). In a typical Antarctic and Greenland snow profile, strong interactions between the atmosphere and 92 93 snow occurs, especially in the first 2 m (Neumann and Waddington, 2004; Town et al., 2008), called the convective zone. In the convective zone, air can move relatively freely 94 and therefore exchange between snow and the atmospheric air occurs. Air flowing into 95 the snow reaches saturation vapor pressure nearly instantly through sublimation 96 (Neumann et al., 2008; Ebner et al., 2015a). Models of the influence of the so-called 97

'wind pumping'-effect (Fisher et al., 1983; Neumann and Waddington, 2004), in which 98 99 the interstitial water vapor is replaced by atmospheric air pushed through the upper meters of the snow pack by small-scale high and low pressure areas caused by irregular 100 grooves or ridges formed on the snow surface (dunes and sastrugi), have assumed that 101 the snow grains would equilibrate with the interstitial water vapor on timescales 102 103 governed by ice self-diffusion. However, no experimental data are available to support this assumption. With this in mind the experimental study presented here is specifically 104 105 developed to investigate the effect of ventilation inside the snow pack on the isotopic 106 composition. Only conditions deeper than 1 cm inside a snowpack are considered. 107 Previous work showed that (1) under isothermal conditions, the Kelvin effect leads to a saturation of the pore space in the snow but does not affect the structural change (Ebner 108 109 et al., 2015a); (2) applying a negative temperature gradient along the flow direction leads to a change in the microstructure due to deposition of water molecules on the ice 110 matrix (Ebner et al., 2015b); and (3) a positive temperature gradient along the flow had 111 a negligible total mass change of the ice but a strong reposition effect of water 112 molecules on the ice grains (Ebner et al., 2016). Here, we continuously measured the 113 isotopic composition of an airflow containing water vapor through a snow sample under 114 both isothermal and temperature gradient conditions. Micro computed-tomography 115 116 (μCT) was applied to obtain the 3D microstructure and morphological properties of 117 snow.

118 **2. Experimental setup**

Isothermal and temperature gradient experiments with fully saturated airflow and defined isotopic composition were performed in a cold laboratory at around $T_{\text{lab}} \approx -15$ °C with small fluctuations of ± 0.8 °C (Ebner et al., 2014). Snow produced from deionized tap water in a cold laboratory (water temperature: 30 °C; air temperature: -20

°C) was used for the snow sample preparation (Schleef et al., 2014). The snow was 123 sieved with a mesh size of 1.4 mm into a box, and isothermally sintered for 27 days at -124 5 °C to increase the strength, in order to prevent destruction of the snow sample due to 125 the airflow, and to evaluate the effect of metamorphism of snow. The morphological 126 properties of the snow are listed in Table 1. The sample holder (diameter 53 mm, height 127 128 30 mm, 0.066 liter) was filled by a cylinder cut out from the sintered snow. To prevent air flow between snow sample and the sample holder walls, the undisturbed snow disk 129 was filled in at a higher temperature (about -5 °C) and sintering was allowed for about 1 130 131 h before cooling down and start of the experiment. The setup of Ebner et al. (2014) was modified by additionally inserting a water vapor isotope analyzer (Model: L1102-I 132 Picarro, Inc., Santa Clara, CA, USA) to measure the isotopic ratio δ^{18} O of the water 133 134 vapor contained in the airflow at the inlet and outlet of the sample holder. The experimental setup consisted of three main components (humidifier, sample holder, and 135 the Picarro analyzer) connected with insulated copper tubing and Swagelok fitting (Fig. 136 1). The tubes to the Picarro analyzer were heated to prevent deposition of water vapor 137 and thereby fractionation. The temperature was monitored with thermistors inside the 138 humidifier and at the inlet and outlet of the snow sample. A dry air pressure tank 139 controlled by a mass flow controller (EL-Flow, Bronkhorst) generated the airflow. A 140 humidifier, consisting of a tube (diameter 60 mm, height 150 mm, 0.424 liter volume) 141 filled with crushed ice particles (snow from Antarctica with low δ^{18} O composition), was 142 used to saturate the dry air entering the humidifier with water vapor at an almost 143 constant isotopic composition. The air temperature in the humidifier and at the inlet of 144 the snow sample was maintained at the same value (accuracy ± 0.2 K) to limit the 145 influence of variability in absolute vapor pressure and isotopic composition. We 146 measured the δ^{18} O of the water vapor produced by the humidifier before and after each 147

experimental run ($\delta^{18}O_{hum}$). The outlet flow ($\delta^{18}O_{a}$) of the sample holder was continuously measured during the experiment to analyze the temporal evolution of the isotopic signal. All data from the Picarro analyzer were corrected to the humidity reference level using the established instrument humidity-isotope response (Steen-Larsen et al. 2013; 2014b). In addition, VSMOW-SLAP correction and drift correction were performed. We followed the calibration protocol and used the calibration system described in detail by Steen-Larsen et al. (2013; 2014b).

The sample holder described by Ebner et al. (2014) was used to analyze the snow by 155 156 μ CT. Tomography measurements were performed with a modified μ -CT80 (Scanco Medical). The equipment incorporated a microfocus X-ray source, operated at 70 kV 157 acceleration voltage with a nominal resolution of 18 µm. The samples were scanned 158 with 1000 projections per 180° in high-resolution setting, with typical adjustable 159 integration time of 200 ms per projection. The field of view of the scan area was 36.9 160 mm of the total 53 mm diameter, and subsamples with a dimension of $7.2 \times 7.2 \times 7.2$ 161 mm³ were extracted for further processing. The reconstructed μ CT images were filtered 162 using a $3 \times 3 \times 3$ median filter followed by a Gaussian filter ($\sigma = 1.4$, support = 3). The 163 Otsu method (Otsu, 1979) was used to automatically perform clustering-based image 164 thresholding to segment the grey-level images into ice and void phase. Morphological 165 166 properties in the two-phase system were determined based on the exact geometry obtained by the µCT. Tetrahedrons corresponding to the enclosed volume of the 167 triangulated ice matrix surface were applied on the segmented data to determine 168 porosity (ε) and specific surface area (SSA). The mean pore size distribution was 169 estimated using the opening-size-distribution operation. This operation can be imagined 170 as virtual sieving with different mesh size (Haussener et al., 2012). 171

Three experiments with saturated advective airflow through the snow sample were 172 performed to record the following parameters and analyze their effects: (1) isothermal 173 conditions to analyze the influence of curvature effects (Kaempfer al et., 2007); (2) 174 positive temperature gradient applied to the snow sample where cold air entering the 175 sample is heated while flowing through the sample in order to analyze the influence of 176 177 sublimation; (3) negative temperature gradient applied to the snow sample where warm air entering the sample is cooled while flowing across the sample, to analyze the 178 179 influence of net deposition. During the temperature gradient experiments, a temperature difference of 1.4 °C and 1.8 °C was imposed resulting in a gradient of +47 K m⁻¹ and -60 180 K m⁻¹, respectively. The runs were performed at atmospheric pressure and with a 181 volume flow rate of 3.0 liter min⁻¹ corresponding to an average flow speed in the pores 182 of $u_{\rm D} \approx 30$ mm s⁻¹. We performed the experiments with airflow velocities in the snow 183 sample at $u_{\rm D} \approx 30 \text{ mm s}^{-1}$, which is a factor of three higher than calculated by Neumann 184 (2003) for a natural snow pack. But looking at the Reynolds number, describing the 185 flow regime inside the pores, our experiments (Re ≈ 0.7) were in the feasible flow 186 regime (laminar flow) of a natural snow pack (Re ≈ 0.65). In experiment (2) the outlet 187 temperature and in experiment (3) the inlet and also the humidifier temperature were 188 actively controlled using thermo-electric elements. Variations in temperature of up to \pm 189 0.8 °C were due to temperature fluctuations inside the cold laboratory, leading to 190 191 slightly variable temperature gradients and mean temperature in experiment (2) and (3). 192 Table 1 presents a summary of the experimental conditions and the morphological properties of the snow samples. All snow samples were taken from the same snow block 193 with an average density of $\approx 210 \text{ kg m}^{-3}$. The density given in Table 1 was the density of 194 the snow sample in each experiment measured by μ CT. At the end of each experiment, 195 the snow sample was cut into five layers of 6 mm height and the isotopic composition of 196

each layer was analyzed to examine the spatial δ^{18} O gradient in the isotopic composition of the snow sample.

A slight increase with a maximum of 0.7 ‰ of δ^{18} O in the water vapor produced by the humidifier was observed in experiment (1), with lower increases during experiments (2) and (3) (Table 2). This change of ~0.7‰ is not significant compared to the difference between the isotopic composition of the water vapor and the snow sample in the sample holder of ~53‰ and the temporal change of the water vapor isotopes on the back side of the snow sample.

In the first approximately 30 min, the isotopic composition of the measured outflow air $\delta^{18}O_a$ increased from a low $\delta^{18}O$ to a starting value of around -29‰ in each experiment. This was due to memory effect and one possible effect might be condensed water left in the tubes from a prior experiment which had no further impact on the experiments (Penna et al., 2012).

210 **3. Results**

211 **3.1 Isothermal condition**

The experiment (1) was performed for 24 h at a mean temperature of $T_{\text{mean}} = -15.5$ °C. $\delta^{18}O_a$ decreased exponentially in the outlet flow observed throughout the experimental run as shown in Fig. 2. Initially, the $\delta^{18}O_a$ content in the flow was -27.7 ‰ and exponentially decreased to -47.6 ‰ after 24 h. The small fluctuations in the $\delta^{18}O_a$ signal at $t \approx 7$ h, 17 h and 23 h were due to small temperature changes in the cold laboratory.

We observed a strong interaction between the airflow and the snow as manifest by the isotopic composition of the snow. The $\delta^{18}O_s$ signal in the snow decreased by 4.75 – 7.78 ‰ and an isotopic gradient in the snow was observed after the experimental run, shown in Fig. 3. Initially, the snow had a homogeneous isotopic composition of $\delta^{18}O_s =$ -10.97 ‰ but post-experiment sampling showed a decrease in the snow $\delta^{18}O$ at the inlet side to -17.75 ‰ and at the outlet side to -15.72 ‰. The spatial $\delta^{18}O_s$ gradient of the snow had an approximate slope of 0.68 ‰ mm⁻¹ at the end of the experimental run. Table 2 shows the $\delta^{18}O$ value in snow at the beginning (t = 0) and end (t = 24 h) of the experiment.

227 **3.2** Air warming by a positive temperature gradient along the airflow

The experiment (2) was performed over a period of 24 h with an average 228 temperature gradient of approximatively +47 K m⁻¹ (warmer temperatures at the outlet 229 of the snow) and an average mean temperature of -14.7 °C. As in the isothermal 230 experiment (1), we observed a relaxing exponential decrease of $\delta^{18}O_a$ in the outlet flow 231 throughout the measurement period as shown in Fig. 2, but the decrease was slower 232 compared to the isothermal run. Initially, the $\delta^{18}O_a$ content in the flow coming through 233 the snow disk was -29.8 ‰ and exponentially decreased to -41.9 ‰ after 24 h. The 234 small fluctuations in the $\delta^{18}O_a$ signal at $t \approx 2.7$ h, and 12.7 h were due to small 235 temperature changes in the cold laboratory. 236

The $\delta^{18}O_s$ signal in the snow decreased 4.66 – 7.66 ‰ and a gradient in the isotopic composition of the snow was observed after the experimental run, shown in Fig. 3. Initially, the snow had a homogeneous isotopic composition of $\delta^{18}O_s = -11.94$ ‰, but post-experiment sampling showed a decrease at the inlet side to -19.6 ‰ and at the outlet side to -16.6 ‰. The spatial $\delta^{18}O_s$ gradient of the snow had an approximate slope of 1.0 ‰ mm⁻¹ at the end of the experimental run. Table 2 shows the $\delta^{18}O_s$ value in snow at the beginning (t = 0) and end (t = 24 h) of the experiment.

3.3 Air cooling by a negative temperature gradient along the air flow

The experiment (3) was performed for 84 h instead of 24 h to better estimate the 245 trend in $\delta^{18}O_a$ in the outlet flow. An average temperature gradient of approximately -60 246 K m⁻¹ (colder temperatures at the outlet of the snow) and an average mean temperature 247 of -13.2 °C were observed during the experiment. As in the previous experiments, a 248 relaxing exponential decrease of $\delta^{18}O_a$ in the outlet flow was observed throughout the 249 experimental run as shown in Fig. 2, but the decrease was slower compared to the 250 isothermal run and temperature gradient opposed to the airflow. Initially, the $\delta^{18}O_a$ 251 content in the flow was -29.8 ‰ and exponentially decreased to -37.7 ‰ after 84 h. The 252 small fluctuations in the $\delta^{18}O_a$ signal at $t \approx 7.3$ h, 21.3 h, 31.3 h, 45.3 h, 55.3 h, 69.3 h, 253 and 79.3 h were due to small temperature changes in the cold laboratory. 254

The $\delta^{18}O_s$ signal in the snow decreased 4.46 – 15.09 ‰ and a gradient in the isotopic composition of the snow was observed after the experimental run, shown in Fig. 3. Initially, the snow had an isotopic composition of $\delta^{18}O_s = -10.44$ ‰ but post-experiment sampling showed a decrease at the inlet side to -25.53 ‰ and at the outlet side to -15.00 ‰. The spatial $\delta^{18}O_s$ gradient of the snow had an approximate slope of 3.5 ‰ mm⁻¹ at the end of the experimental run. Table 2 shows the $\delta^{18}O_s$ value in snow at the beginning (t = 0) and end (t = 84 h) of the experiment.

262 **4. Discussion**

All experiments showed a strong exchange in δ^{18} O between the snow and watervapor saturated air resulting in a significant change of the value of the stable isotopes in the snow. The advective conditions in the experiments were comparable with surface snow layers in Antarctica and Greenland, but at higher temperature, especially compared to interior Antarctica.

The results also showed strong interactions in δ^{18} O between snow and air depending on the different temperature gradient conditions. The experiments indicate that

temperature variation and airflow above and through the snow structures (Sturm and 270 Johnson, 1991; Colbeck, 1989; Albert and Hardy, 1995) seem to be dominant processes 271 affecting water stable isotopes of surface snow. The results also support the statement 272 that an interplay between theoretically expected layer-by-layer sublimation and 273 deposition at the ice-matrix surface and the isotopic content evolution of snow cover 274 275 due to mass exchange between the snow cover and the atmosphere occurs (Sokratov and Golubev, 2009). The specific surface area of snow exposed to mass exchange (Horita et 276 al., 2008) and by the depth of the snow layer exposed to the mass exchange with the 277 278 atmosphere (He and Smith, 1999) plays an important role. Our results support the 279 interpretation that changes in surface snow isotopic composition are expected to be significant if large day-to-day surface changes in water vapor occur in between 280 precipitation events, wind pumping is efficient and snow metamorphism is enhanced by 281 temperature gradients in the upper first centimeters of the snow (Steen-Larsen et al., 282 2014a). 283

We expect that our findings will lead to improvement of the interpretation of the 284 water stable isotope records from ice cores. Classically, ice core stable isotope records 285 are interpreted as paleo-temperature reflecting precipitation-weighted signals. When 286 comparing observations and atmospheric model results for precipitation with ice core 287 288 records, such vapor-snow exchanges are normally ignored (e.g. Persson et al., 2011; Fujita and Abe, 2006). However, vapor-snow exchange enhanced by recrystallization 289 rate seems to be an important factor for the high variation in the snow surface $\delta^{18}O$ 290 signal as supported by our experiments. It was hypothesized that the changes in the 291 snow-surface δ^{18} O reported by Steen-Larsen et al. (2014a) are caused by changes in 292 large-scale wind and moisture advection of the atmospheric water vapor signal and 293

snow metamorphism. The strong interaction between atmosphere and near-surface snowcan modify the ice core water stable isotope records.

The rate-limiting step for isotopic exchange in the snow is isotopic equilibration between the pore-space vapor and surrounding ice grains. The relaxing exponential decrease of δ^{18} O in the outflow of our experiments predicted that full isotopic equilibrium between snow and atmospheric vapor will not be reached at any depth (Waddington et al., 2002; Neumann and Waddington, 2004) but changes towards equilibrium with the atmospheric state occurs (Steen-Larsen et al., 2013, 2014a).

302 As snow accumulates, the upper 2 m are advected through the ventilated zone 303 (Neumann and Waddington, 2004; Town et al., 2008). In areas with high accumulation rate (e.g. South Greenland), snow is advected for a short time through the ventilated 304 zone. The snow exposed a relatively short time to vapor snow exchange would result in 305 higher spatial variability compared to long-time exposure. However, the effects of snow 306 ventilation on isotopic composition may become more important as the accumulation 307 rate of the snow decreases ($< 50 \text{ mm a}^{-1}$), such that snow remains in the near-surface 308 ventilated zone for many years (Waddington et al., 2002; Hoshina et al., 2014; Hoshina 309 310 et al., 2016). As the snow remains longer in the near-surface ventilated zone, a larger δ^{18} O exchange between snow and atmospheric vapor will occur. Consequently, the 311 312 isotopic content of layers at sites with high and low accumulation rates can evolve 313 differently, even if the initial snow composition had been equal, and the sites had been 314 subjected to the same histories of air-mass vapor.

Despite a relatively small change in the difference between the isotopic composition of the incoming vapor and the snow, large differences in the isotopic composition of the water vapor at the outlet flow exist for the three different experimental setups. Based on the difference in the outlet water vapor isotopic composition, we hypothesized that

different processes are at play for the different experiments. It is obvious that there is a 319 fast isotopic exchange with the surface of the ice crystals, and a much slower timescale 320 on which the interior of the ice crystals is altered. Due to the low diffusivity of $H_2^{16}O$ 321 and $H_2^{18}O$ in ice $(D_{H_2^{18}O} \approx D_{H_2^{16}O} = \sim 10^{-15} \text{ m}^2 \text{ s}^{-1}$ (Ramseier, 1967; Johnsen et al., 2000), 322 we assumed that the interior of the ice crystals is not altered on the timescale of the 323 experiment. This explained why the net isotopic change of the bulk sample is relatively 324 small compared to the changes in the outlet water vapor isotopes. The effective 'ice-325 diffusion depth' of the isotopic exchange during the experiments is given as $L_{\rm D} = \sqrt{D \cdot t}$, 326 where D is the diffusion coefficient of $H_2^{16}O$ and $H_2^{18}O$ in ice, respectively, and t the 327 experimental time. The calculated 'ice-diffusion depth' L_D , is ~ 9.3 µm for experiments 328 (1) and (2), and ~17.4 µm for experiment (3), respectively, indicating an expected a 329 minimal change of the interior of the ice crystal. However, snow has a large specific 330 surface area and therefore a high exchange area. This has an effect on the $\delta^{18}O$ snow 331 concentration. The fraction of the total volume V_{tot} of ice that is close enough to the ice 332 surface to be affected by diffusion in time t is then $\rho_{ice} \cdot SSA \cdot L_D$, where SSA is the 333 specific surface area (area per unit mass), and $L_{\rm D}$ is the diffusion depth, defined above, 334 for time t. For $t \approx 24$ hours, a large fraction (24 to 43 %) of the total volume V_{tot} of the 335 ice matrix can be accessed through diffusion. It is quite hard to see the total δ^{18} O snow 336 difference between experiments (1) and (2) after the experiment compared to the δ^{18} O 337 of the vapor in the air at the outlet. There is a small, but notable, difference in the total 338 δ^{18} O of the snow between experiment (1) and (2). Due to the higher recrystallization 339 rate of experiment (2) the spatial $\delta^{18}O_s$ gradient of the snow (1.0 % mm⁻¹) is higher than 340 for experiment (1) (0.68 % mm⁻¹). Increasing the experimental time, the δ^{18} O change in 341 the snow increases (experiment (3)). In general, the calculated 'ice-diffusion depth' is 342 realistic under isothermal conditions where diffusion processes are the main factors 343

(Kaempfer and Schneebeli, 2007; Ebner et al., 2015). Applying a temperature gradient, the impact of diffusion is suppressed due to the high recrystallization rate by sublimation and deposition. Due to the low half-life of the ice matrix of a few days, the growth rates are typically on the order of 100 µm per day (Pinzer et al., 2012). Therefore, this redistribution of ice caused by temperature gradient counteracts the diffusion into the solid ice.

By comparing similarities and differences between the outcomes of the three 350 experimental setups we will now discuss the physical processes influencing the 351 interaction and exchange processes within the snowpack between the snow and the 352 advected vapor. We first notice that the final snow isotopic profile of experiment (1) 353 (isothermal) and (2) (positive temperature gradient along the direction of the flow) are 354 comparable to each other. Despite this similarity, the evolution in the outlet water vapor 355 356 of experiment (1) showed a significantly stronger depletion compared to experiment (2). For experiment (3) (negative temperature gradient along the direction of the flow) we 357 observed the smallest change in outlet water vapor isotopes but the largest snow-pack 358 isotope gradient after the experiment. However, this change was caused by 84 hours 359 flow instead of 24 hours. 360

Curvature effects, temperature gradients and therefore the recrystallization rate 361 influence the mass transfer of $H_2^{16}O/H_2^{18}O$ molecules. The higher the recrystallization 362 rate of the snow the slower the adaption of the outlet air concentration to the inlet air 363 concentration (see in experiment (2) and (3)). Under isothermal conditions (experiment 364 (1)) the only effect influencing the recrystallization rate is the curvature effect 365 (Kaempfer and Schneebeli, 2007). However, based on the experimental observations 366 367 (Kaempfer and Schneebeli, 2007) this effect decreases with decreasing temperature and increasing experimental time. Applying an additional temperature gradient on a snow 368

sample causes complex interplays between local sublimation and deposition on surfaces 369 and the interaction of water molecules in the air with the ice matrix due to changing 370 saturation conditions of the airflow. Therefore, the recrystallization rate increases and 371 causes the change in the δ^{18} O of the air. For experiment (2) there is a complex interplay 372 between sublimation and deposition of water molecules into the interstitial flow (Ebner 373 374 et al., 2015c) while for experiment (3) there is deposition of molecules carried by the interstitial flow onto the snow crystals (Ebner et al., 2015b). Furthermore, in the 375 beginning of each experiment there is a tendency to sublimate from edges of the 376 individual snow crystals due to the higher curvature. As the edges were sublimated and 377 378 deposition occurred in the concavities, the individual snow crystals became more rounded, slowing down the transfer of water molecules into the interstitial airflow. We 379 noticed for all three experiments that within the uncertainty of the isotopic composition 380 of the snow, the initial isotopic composition of the vapor was the same and in isotopic 381 equilibrium with the snow. The difference between experiment (1) and (2) lies in the 382 fact that due to the temperature gradient in experiment (2) there is an increased transfer 383 of water vapor with the isotopic composition of the snow into the airflow. Hence the 384 depleted air from the humidifier advected through the snow disk is mixed with a 385 relatively larger vapor flux from the snow crystals. Additionally, we also expected less 386 387 deposition into the concavities in experiment (2) compared to experiment (1). However, it is interesting to note that the final isotopic profile of the snow disk is similar in 388 experiment (1) and experiment (2). We interpreted this as being a result of two 389 processes acting in opposite direction: although relatively isotope-depleted vapor from 390 the humidifier was deposited on the ice matrix there was also a higher amount of 391 sublimation of relatively isotope-enriched vapor from the snow disk in experiment (2). 392 Experiment (3) separates itself from the other two experiments in the way that as the 393

water vapor from the humidifier is advected through the snow disk there is a continuous 394 deposition of very depleted air due the negative temperature gradient. As for the case of 395 experiment (1) and (2) there was also in experiment (3) a constant sublimation of the 396 convexities into the vapor stream. We notice that despite the fact that experiment (3) ran 397 for 84 hours the snow at the outlet side of the snow-disk did not become more 398 399 isotopically depleted compared to experiment (1) and (2). However, the snow on the inlet side became significantly more isotopically depleted. This observation, together 400 401 with the fact that the vapor of the outlet of the snow-disk is less depleted compared to 402 experiment (1) and (2), leads us to hypothesize that there is a relatively larger deposition 403 of isotopically depleted vapor from the humidifier as the vapor is advected through the snow disk. This means that a relatively larger component of the isotopic composition of 404 405 the vapor is originating by sublimation from the convexities of the snow disk and less from the isotopically depleted vapor from the humidifier. 406

Our results and conclusions indicate that there is a need for additional validation. 407 Specifically, it would be crucial to know the mass balance of the snow disk more 408 precisely, which could be done by reconstructing the entire snow disk following the 409 410 change in density and morphological properties over the entire height. Ideally, the entire sample would be tomographically measured with a resolution of $4 \times 4 \times 4 \text{ mm}^3$, each 411 412 cube corresponding to the representative volume. Insights would also be achieved with 413 experiments using snow of the same isotopic composition, but different SSA, as more 414 precise calculation of the different observed exchange rates would be allowed. Additionally, different and colder background temperatures should be tested to better 415 understand inland Antarctic environment and the effect of the quasi-liquid layer, which 416 is necessary for the development of a numerical model. Isotopically different 417 combinations of vapor and snow should be performed. In the present manuscript, vapor 418

with low δ^{18} O isotopic composition was transported through snow with relative high δ^{18} O isotopic composition. It would be interesting to reverse the combination and perform experiments with different combinations to provide more insights on mass and isotope exchanges between vapor and snow. Experiments with longer running time help to understand the change in the ice matrix better under low accumulation conditions.

424

4. Summary and conclusion

Laboratory experimental runs were performed where a transient $\delta^{18}O$ interaction 425 between snow and air was observed. The airflow altered the isotopic composition of the 426 snowpack and supports an improved climatic interpretation of ice core stable water 427 isotope records. The water vapor saturated airflow with an isotopic difference of up to 428 55‰ changed within 24 h and 84 h the original δ^{18} O isotope signal in the snow by up to 429 7.64 ‰ and 15.06 ‰. The disequilibrium between snow and air isotopes led to the 430 observed exchange of isotopes, the rate depending on the temperature gradient 431 conditions. Concluding, increasing the recrystallization rate in the ice matrix causes the 432 temporal change of the δ^{18} O concentration at the outflow to decrease (experiment (2) 433 and (3)). Decreasing the recrystallization rate causes the temporal curve of the outlet 434 concentration to become steeper reaching the δ^{18} O inlet concentration of the air faster 435 436 (experiment (1)).

Additionally, the complex interplay of simultaneous diffusion, sublimation and deposition due to the geometrical complexity of snow has a strong effect on the δ^{18} O signal in the snow and cannot be neglected. A temporal signal can be superimposed on the precipitation signal, (a) if the snow remains near the surface for a long time, i.e. in a low-accumulation area, and (b) is exposed to a history of air masses carrying vapor with a significantly different isotopic signature than the precipitated snow.

These are novel measurements and will therefore be important as the basis for 443 further research and experiments. Our results represent direct experimental observation 444 of the interaction between the water isotopic composition of the snow, the water vapor 445 in the air and recrystallization due to temperature gradients. Our results demonstrate that 446 recrystallization and bulk mass exchange must be incorporated into future models of 447 448 snow and firn evolution. Further studies are required on the influence of temperature and airflow as well as snow microstructure on the mass transfer phenomena for 449 validating the implementation of stable water isotopes in snow models. 450

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Table 1: Morphological properties and flow characteristics of the experimental runs: μ CT measured snow density (ρ), porosity (ε), specific surface area per unit mass (SSA), mean pore space diameter (d_{mean}), superficial velocity in snow (u_D), corresponding Reynolds number (Re = $d_{mean} \cdot u_D/v_{air}$), average inlet temperature of the humidifier and at the inlet ($T_{in,mean}$), average outlet temperature at the outlet ($T_{out,mean}$), and average temperature gradient (∇T_{ave}). Experiment (1) corresponds to the isothermal conditions; Experiment (2) to air warming; and Experiment (3) to air cooling in the snow sample.

	ρ	З	SSA	$d_{\rm mean}$	$u_{\rm D}$	Re	T _{in,mean}	T _{out,mean}	∇T_{ave}
	kg m ⁻³	_	$m^2 kg^{-1}$	mm	m s ⁻¹	_	°C	°C	$\mathrm{K} \mathrm{m}^{-1}$
Experiment (1)	202	0.78	28	0.39	0.03	0.76	-15.5	-15.5	_
Experiment (2)	202	0.78	30	0.36	0.03	0.70	-15.4	-14.0	+47
Experiment (3)	220	0.76	27	0.37	0.031	0.74	-12.3	-14.1	-60

Table 2: δ^{18} O in the vapor in the humidifier (δ^{18} O_{hum}) and of the snow in the sample holder (δ^{18} O_s) at the beginning (t = 0) and end (t = end) of each experiment and the final δ^{18} O content of the snow in the sample holder at the inlet (z = 0 mm) and outlet (z = 30mm). Experiment (1) corresponds to the isothermal conditions; Experiment (2) to air warming; and Experiment (3) to air cooling in the snow sample.

	$\delta^{18}O_{hum}$		$\delta^{18}O_{s,t}=0$	$\delta^{18}O_{s, t} = end$		
	‰ 0		‰	‰		
	t = 0	t = end		z = 0 mm	z = 30 mm	
Experiment (1)	-68.2	-67.5	-10.97	-17.75	-15.72	
Experiment (2)	-66.3	-66.1	-11.94	-19.60	-16.60	
Experiment (3)	-62.8	-62.2	-10.44	-25.53	-15.00	

718 Figure captions

- 719Fig. 1.Schematic of the experimental setup. A thermocouple (TC) and a humidity720sensor (HS) inside the humidifier measured the the mean temperature and721humidity of the airflow. Two thermistors (NTC) close to the snow surface722measured the inlet and outlet temperature of the airflow (Ebner et al., 2014).723The Picarro Analyzer measured the isotopic composition δ^{18} O of the outlet724flow. Inset: 3D structure of 110 × 42 × 110 voxels (2 × 0.75 × 2 mm³)725obtained by the μ CT.
- Temporal isotopic composition of δ^{18} O of the outflow for each of the **Fig. 2**. 726 experimental runs. The spikes in the δ^{18} O were due to small temperature 727 changes in the cold laboratory (Ebner et al., 2014). Exp. (1) corresponds to 728 the isothermal conditions; Exp. (2) to air warming; and Exp. (3) to air 729 cooling in the snow sample. The higher the recrystallization rate of the snow 730 the slower the adaption of δ^{18} O of the outlet air to the inlet air. The 731 illustration in the lower right corner shows the relation between δ^{18} O of the 732 initial snow, inlet, and outlet of the air. 733
- Fig. 3. Spatial isotopic composition of δ^{18} O of the snow sample at the beginning (*t* = 0) and at the end (*t* = end) for each experiment. The air entered at *z* = 0 mm and exited at *z* = 30 mm. Exp. (1) corresponds to the isothermal conditions; Exp. (2) to air warming; and Exp. (3) to air cooling in the snow sample.

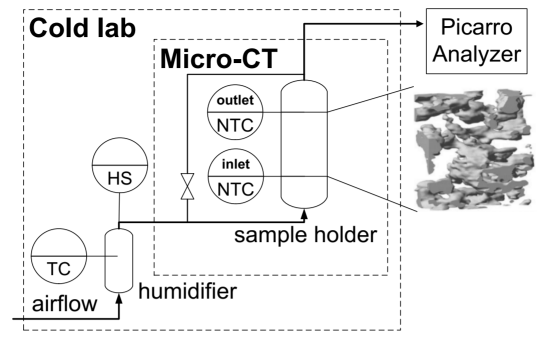
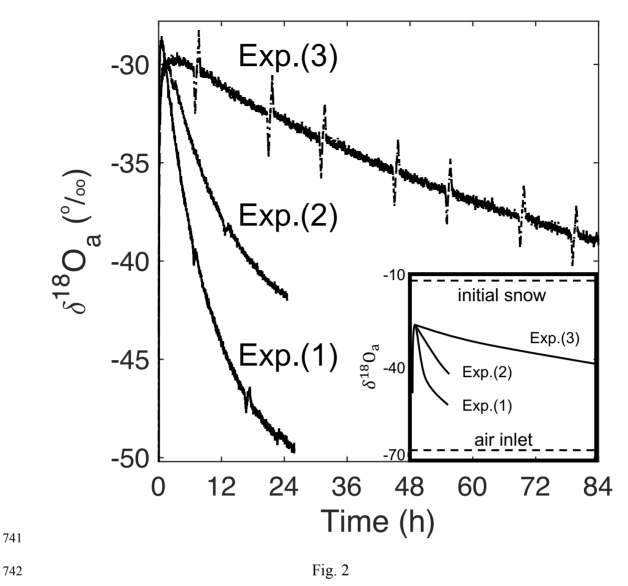




Fig. 1



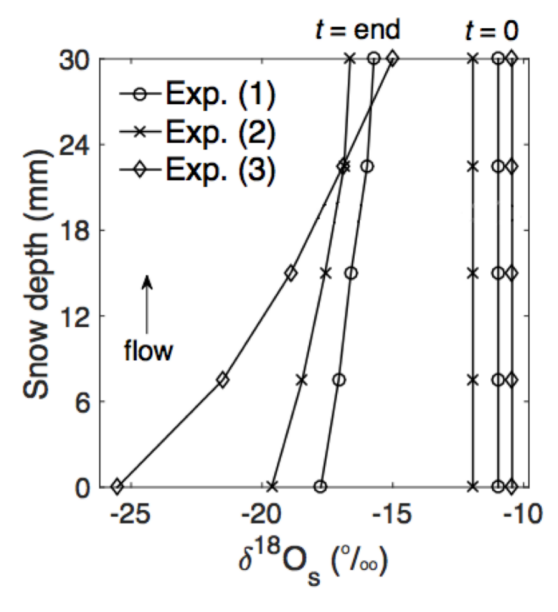


Fig. 3