Radiocarbon dating of alpine ice cores with the dissolved organic carbon (DOC) fraction

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15 Abstract

High-alpine glaciers are valuable archives of past climatic and environmental conditions. The 17 interpretation of the preserved signal requires a precise chronology. Radiocarbon (¹⁴C) dating 18 of the water-insoluble organic carbon (WIOC) fraction has become an important dating tool to 19 constrain the age of ice cores from mid-latitude and low-latitude glaciers. However, in some 20 cases this method is restricted by the low WIOC concentration in the ice. In this work, we 21 report first ¹⁴C dating results using the dissolved organic carbon (DOC) fraction, which is 22 present at concentrations of at least a factor of two higher than the WIOC fraction. We 23 evaluated this new approach by comparison to the established WIO¹⁴C dating based on parallel 24 ice core sample sections from four different Eurasian glaciers covering an age range of several 25 hundred to around 20'000 years. ¹⁴C dating of the two fractions yielded comparable ages with 26 WIO¹⁴C revealing a slight, barely significant, systematic offset towards older ages. Our data 27 suggests comparable in magnitude with the analytical uncertainty. We attribute this to be 28 29 caused by offset to two effects of about equal size, but opposite in direction: (i) in-situ produced ¹⁴C contributing to the DOC resulting in a bias towards younger ages and (ii) incompletely 30 removed carbonate from particulate mineral dust (¹⁴C depleted) contributing to the 31 WIOC fraction- with a bias towards older ages. The estimated amount of in-situ produced ^{14}C 32 in the DOC fraction is smaller than the analytical uncertainty for most samples. Nevertheless, 33 34 under extreme conditions, such as very high altitude and/or low snow accumulation rates, DO¹⁴C dating results need to be interpreted cautiously. While in the during DOC extraction 35 36 procedure-the removal of inorganic carbon is monitored to ensure complete removal, the for completeness, the removal for WIOC samples was so far only assumed to be quantitative, at 37 38 least for ice samples containing average levels of mineral dust. Here we estimated an average removal efficiency for WIOC samples was here estimated to be ~96%. We did not find any 39 indication of in-situ production systematically contributing to DO¹⁴C as suggested in a 40 previous of 98±2 %, resulting in a small offset in the order of the current analytical uncertainty. 41 42 Future optimization of the removal procedure has the potential to improve the accuracy and precision of WIO¹⁴C dating. With this study. By we demonstrate, that using the DOC instead 43 of the fraction for ¹⁴C dating is not <u>only a valuable alternative to the use of WIOC fraction for</u> 44 ⁴⁴C dating, the , but also benefits from a reduced required ice mass can be reduced to of 45 typically ~250 g, yielding a to achieve comparable precision of around ±200 years or even 46 better if sample sizes typically required for WIO¹⁴C dating are used. This study shows approach 47 thus has the potential of pushing radiocarbon dating of ice forward even to remote and Polar 48 Regions, regions where the carbon content in the ice is particularly low, when applying the 49 DOC fraction for ¹⁴C dating. 50

52 **1 Introduction**

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For a meaningful interpretation of the recorded paleoclimate signals in ice cores from glacier 54 archives, an accurate chronology is essential. Annual layer counting, supported and tied to 55 independent time markers such as the 1963 nuclear fallout horizon evident by a peak maximum 56 in tritium or other radioisotopes, or distinct signals from known volcanic eruptions in the past 57 is the fundamental and most accurate technique used for ice core dating. However, for ice cores 58 from high-alpine glaciers this approach is limited to a few centuries only, because of the 59 exceptional strong thinning of annual layers in the vicinity of the bedrock. Most of the current 60 analytical techniques do not allow high enough sampling resolution for resolving seasonal 61 62 fluctuations or detecting distinct single events in this depth range. Ice flow models, which are widely used to retrieve full depth age scales (e.g. Nye, 1963; Bolzan, 1985; Thompson et al., 63 64 2006), also fail in the deepest part of high-alpine glaciers due to the complex bedrock 65 geometry.assumption of steady state conditions and the complexity of glacial flow and bedrock 66 geometry limiting realistic modeling of strain rates. Even with 3D models, which require extensive geometrical data, it is highly challenging to simulate a reasonable bottom age (e.g. 67 68 Licciulli et al., 2020). This emphasizes the need for an absolute dating tool applicable to the oldest, bottom parts of cores from these sites. 69

70 Radioactive isotopes contained in the ice offer the opportunity to obtain absolute ages of an ice sample. For millennial scale ice cores, ${}^{14}C$ dating is the technique of choice. With a 71 half-life of 5370 years, dating in the age range from ~250 years to up to ten half-life times is 72 73 theoretically possible, covering the time range accessible by alpine glaciers in the vast majority of cases (Uglietti et al., 2016). The ¹⁴C dating ofapproach using water insoluble organic carbon 74 (WIOC) from glacier ice has become a well-established technique for ice core dating. Samples 75 76 of >10 μ g WIOC can be dated with reasonable uncertainty (10-20%), requiring less than 1 kg of ice from typical mid-latitude and low-latitude glaciers and its accuracy was recently 77 78 validated (Uglietti et al., 2016). Ice samples from mid- and low-latitude glaciers can now be dated with a reasonable uncertainty of 10-20%. Ice sample masses of 200-800 g are usually 79 selected to aim for >10 μ g carbon for $\frac{14}{C}$ analysis with accelerator mass spectrometry (AMS), 80 whereby the respective mass depends on sample age and organic carbon concentrations (Jenk 81 82 et al., 2007; Jenk et al., 2009; Sigl et al., 2009; Uglietti et al., 2016). However; Hoffmann et al., 2018). Accordingly, the low WIOC concentration in some glaciers and in Polar Regions, with 83 84 and the correspondingrelated large demands of ice mass puts a limit to this application.

Concentrations of dissolved organic carbon (DOC) in glacier ice are a factor of 2-8 higher 85 compared to typical WIOC concentrations (Legrand et al., 2007; Legrand et al., 2013; May et 86 al., 2013, Fang et al., in prep.). Using the DOC fraction for ¹⁴C dating could therefore reduce 87 the required amount of ice or, for sample sizes similar to what would be needed for ¹⁴C dating 88 by WIOC, improve the achievable analytical (dating) precision which strongly depends on the 89 absolute carbon mass even for state-of-the-art micro-radiocarbon dating. The underlying 90 hypothesis of applying the DOC fraction for ¹⁴C dating is the same as for the WIO¹⁴C dating 91 approach (Jenk et al., 2006; Jenk et al., 2007; Jenk et al., 2009). DOC in ice is composed of 92 93 atmospheric water soluble organic carbon (WSOC) contained in carbonaceous aerosol particles and organic gases taken up during precipitation (Legrand et al., 2013). WSOC is formed in the 94 atmosphere by oxidation of gases emitted from the biosphere or from anthropogenic sources 95 (Legrand et al., 2013; Fang et al. in prep.) -and subsequent condensation of the less volatile 96 products. Carbonaceous aerosols transported in the atmosphere can be deposited on a glacier 97 by wet and dry deposition. Before the industrial revolution, these organic carbon species, then 98 entirely of non-fossil origin, contain the contemporary atmospheric ¹⁴C signal of the time when 99 the snow deposited on the glacier (Jenk et al., 2006). In view of the analytical precision 100 101 achievable with this method, the turn over time from atmospheric CO2 to deposited aerosol is 102 negligible (Fang et al., in prep.). For both WIOC and WSOC, carbon from biomass burning and oceanic organic matter can potentially introduce a reservoir effect (sources of aged carbon). 103 104 The mixed age of trees in Swiss forests today is estimated to be slightly less than 40 years (Mohn et al., 2008). Back in time, prior to extensive human forest management, the mixed age 105 106 of trees in Europe was likely older and the mean age of old-growth forest wood ranged from around 70 to 300 years depending on the region, i.e. the tree species present (Gavin, 2001, 107 Zhang et al., 2017). Prior to the use of fossil fuels about 50% of WIOC is estimated to originate 108 from biomass burning (Minguillon et al., 2011). For biogenic DOC, May et al. (2013) estimated 109 a turnover-time of around 3 to 5 years, corresponding to a 20% contribution from biomass 110 burning. With a mean age of burned material (aged wood plus grass and bushes) of 150±100 111 years, this results in a potential in-built age from biomass burning for WIOC and DOC of 75±50 112 and 30±20 years, respectively. Such an in-built age is negligible considering the analytical 113 uncertainty, which is similarly the case for a bias from oceanic sources, since concentrations 114 of marine organic tracers are more than one order of magnitude lower than terrestrial tracers 115 for the vast majority of glacier sites. This conclusion is supported by the fact that Uglietti et al. 116 (2016) did not identify such a bias, when comparing $WIO_{14}^{14}C$ ages with ages derived by 117 independent methods. 118

For analyzing DO¹⁴C in ice cores, one of the major limitations is the relatively low 119 extraction efficiency ranging from 64 % (Steier et al., 2013) to 96 % (May et al., 2013; Fang 120 et al., 2019) and the high risk of sample contamination (Legrand et al., 2013) potentially 121 introduced during drilling, storage, and sample processing. A first attempt to use DOC for ¹⁴C 122 dating of ice samples was conducted by May (2009) using a set-up for a combined analysis of 123 both, the DOC and WIOC fraction with subsequent radiocarbon micro-analysis. However, 124 these first results suggested a potential in-situ production of ¹⁴C in the DOC fraction based on 125 the obtained super modern $F^{14}C$ values (i.e. $F^{14}C$ values higher than ever observed in the recent 126 or past ambient atmosphere). Building on these initial findings, May (2009) questioned the 127 applicability of the DOC fraction for radiocarbon dating. Although the in-situ ¹⁴C production 128 of ¹⁴CO and ¹⁴CO₂ in air bubbles contained in polar ice has been studied thoroughly and is 129 rather-well understood (Van de Wal et al., 1994; Lal et al., 1997; Smith et al., 2000), possible 130 mechanisms of ¹⁴C in-situ production followed by formation inof organic compounds seem far 131 less likely and have are not been investigated and only few studies exist to date (Woon, 2002).; 132 Hoffmann, 2016). To further explore the potential of DO¹⁴C for dating ice, a DOC extraction 133 setup for radiocarbon analyses was designed and built at the Paul Scherrer Institut (PSI). In 134 order to minimize potential contamination, the entire system is protected from ambient air by 135 136 inert gas (helium) flow or vacuum. To maximize the oxidation efficiency, the PSI DOC methodology applies an ultraviolet (UV) photochemical oxidation step supported by addition 137 of Fenton's reagent. The setup has been characterized by a high extraction efficiency of 96% 138 and a low overall process blank being superior in the resulting blank to sample ratio compared 139 140 to other systems (Fang et al., 2019). The system can handle samples with volumes of up to ~350 mL-allowing.¹⁴C analysis on. With this volume, samples with DOC concentrations as 141 low as 25-30 µg/kg can be analyzed, yielding the minimal carbon mass required for reliable 142 $\frac{14}{C}$ analysis (~10 µg C). Pooling samples from several subsequent extractions would be 143 feasible, allowing dating of samples with lower DOC concentration. In this study, we evaluate 144 ¹⁴C dating with the DOC fraction by comparing to results from the well- established and 145 validated WIO¹⁴C dating method. This is not only analytically highly challenging, but also 146 because of the very limited availability of the precious sampling material needed in a rather 147 large quantity $(\rightarrow (\text{total for both fractions} > 500 \text{ g}) - \text{and})$, ideally covering a wide range of ages 148 from a few hundred to several thousands of years. Here, we succeeded to analyze such parallel 149 samples from four different Eurasian glaciers-were analyzed in parallel. 150

152 **2** Sample preparation and ¹⁴C analysis

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To validate the DOC ¹⁴C dating technique, a total of 17 samplesice sections from the deep parts 154 of ice cores from the four glaciers Colle Gnifetti, Belukha, Chongce (Core 1), and Shule 155 156 NanshanShu Le Nan Shan (SLNS) were selected (Figure 1). They were sampled in parallel to directly compare DOC and WIOC concentrations and $\frac{14}{C}$ dating results. The high-alpine 157 glacier Colle Gnifetti is located in the Monte Rosa massif of the Swiss Alps, close to the Italian 158 159 border. A 76 m long core was retrieved from the glacier saddle in September 2015 at an altitude of 4450 m asl. (45°55'45.7''N, 7°52'30.5''E); Sigl et al., 2018), only 16 m away from the 160 161 location of a previously dated core obtained in 2003 (Jenk et al., 2009). The low annual net accumulation rate at this site (~0.45 m w.e. yr^{-1}) provides access to old ice covering the 162 Holocene (Jenk et al., 2009). Four samples were selected from the bottom 4 m (72-76 m) closest 163 164 to bedrock- (72-76 m depth). The Belukha core was drilled in May/June 2018 from the saddle between the two summits of Belukha (49°48'27.7"N, 86°34'46.5"E, 4055 m asl.), the highest 165 166 mountain in the Altai mountain range. The bedrock was reached and the total length of the core is 160 m. Three samples were analyzed from the deepest part (158-160 m). Seven, and three 167 samples were analyzed from the deep parts of SLNS and Chongce, respectively. The SLNS ice 168 169 core was retrieved in May 2010 from the south slope of the ShulenanshanShu Le Nan Shan Mountain (38°42'19.35"N, 97°15'59.70"E, 5337 m asl.). The bedrock was reached and the total 170 171 length of the ice core is 81.05 m (Hou et al., submitted). The Chongce ice cap is located in the western Kunlun Mountains on the northwestern Tibetan Plateau, covering an area of 163.06 172 km² with a volume of 38.16 km³- (Hou et al., 2018). The ice analyzed in this study was sampled 173 from Chongce Core 1, one of three ice cores drilled in October 2012 (35°14'5.77"N, 174 81°7'15.34"E, 6010 m asl.). Two of those cores reached bedrock with lengths of 133.8 m (Core 175 1) and 135.8 m (Core 2). In 2013, two more ice cores were recovered tofrom a higher altitude 176 177 of 6100 m asl., reaching bedrock with lengths of 216.6 m (Core 4) and 208.6 m (Core 5) at a higher altitude of 6100 m asl. (Hou et al., 2018). Find The annual net accumulation rate is about 178 0.14 m w.e. yr⁻¹ for Core 3, located less than 2 km away from Core 1. A summary of the 179 180 metadata for the study sites and ice cores can be found in the supplement (Table S1) and details 181 about all samples sampling depths and sample sizes in Table 1. No results from any of the cores analyzed in this study have been published previously. 182

All <u>sample sampled ice sections</u> were decontaminated in a cold room (-20°C) by cutting off the surface layer (~3 mm). <u>Each sample was) and each section</u> split into two parallel

sections samples to perform both WIOC and DOC ¹⁴C analysis. Samples for WIO¹⁴C-dating 185 were prepared following the protocol described in Uglietti et al. (2016) with a brief summary 186 provided in the following. In order to remove potential contamination in the outer layer of the 187 ice core, pre-cut samples from the inner part of the core were additionally rinsed with ultra-188 pure water (Sartorius, 18.2 M Ω ×cm, TOC < 5ppb), resulting in samples masses ranging from 189 ~300 to 600 g (Table 1). To dissolve carbonate potentially present in the ice, melted samples 190 were acidified with HCl to pH < 2, before being sonicated for 5 min. Subsequently, the 191 contained particles were filtered onto pre-baked (heated at 800 °C for 5 h) quartz fiber filters 192 193 (Pallflex Tissueqtz-2500QAT-UP). In a second carbonate removal step, the filters were acidified 3 times with a total amount of 50 µL 0.2M HCl, left for 1 h, rinsed with 5 mL ultra-194 pure water and finally left again for drying. These initial steps were performed in a laminar 195 flow box to ensure clean conditions. At the Laboratory for the Analysis of Radiocarbon with 196 AMS (LARA) of the University of Bern the particle samples were then combusted in a thermo-197 optical OC/EC analyzer (Model4L, Sunset Laboratory Inc, USA) equipped with a non-198 dispersive infrared (NDIR) cell to quantify the CO₂ produced, using the well-established Swiss 199 4S protocol for OC/EC separation (Zhang et al., 2012). Being coupled to a 200 kV compact 200 accelerator mass spectrometer (AMS, MIni CArbon DAting System MICADAS) equipped 201 202 with a gas ion source via a Gas Interface System (GIS, Ruff et al., 2007; Synal et al., 2007, Szidat et al., 2014), the LARA Sunset-GIS-AMS system (Agrios et al., 2015; Agrios et al., 203 2017) allowed for final, direct online ¹⁴C measurements of the CO₂ produced from the WIOC 204 fraction. 205

For DO¹⁴C analysis, sample preparation follows the procedure described in Fang et al. 206 207 (2019). After transfer of pre-cut samples to the laboratory and before being melted, samples were further decontaminated in the pre-cleaned melting vessel of the extraction setup by rinsing 208 209 with ultrapure water before being melted, (sample mass loss of about 20-30 %), all performed under helium atmosphere. Simultaneously, a pre-cleaning step was applied to remove potential 210 contamination in the system. For this, 50 mL ultra-pure water was injected into the reactor and 211 212 acidified with 1 mL of 85% H_3PO_4 . To enhance the oxidation efficiency, 2 mL of 100 ppm FeSO₄ and 1 mL of 50 mM H₂O₂ (Fenton's reagent) was also injected into the base water 213 before turning on the UV lights for ~20 min, thereby monitoring the process via the online 214 NDIR CO₂ analyzer. After the ice melted, the meltwater was filtrated under helium atmosphere, 215 using a pre-baked in-line quartz fiber filter. The sample volume was determined by measuring 216 the reactor fill level. The filtrate was acidified by mixing with the pre-treated base water. After 217

degassing of CO₂ from inorganic carbon was completed as monitored by the CO₂- detector, 1 218 mL of 50 mM H₂O₂ was injected into the reactor right before the irradiation started. During 219 UV oxidation, water vapor was removed by cryogenic trapping at -60 °C and produced CO₂ 220 was trapped in liquid nitrogen. All steps were carried out under a constant flow of helium. The 221 sample CO₂ was further cleaned from residual water vapor and quantified manometrically 222 before being sealed into a glass vial for offline ¹⁴C analyses. The CO₂ gas from DOC in the 223 glass vial was directly injected into the MICADAS using a cracking system for glass vials 224 under vacuum, allowing to then carry the CO₂ gas in a helium flow to the AMS ion source 225 226 (Wacker et al., 2013). Procedural blanks were determined and continuously monitored by processing and analyzing frozen ultra-pure water (Sartorius, 18.2 M Ω cm, TOC< 5ppb) similar 227 to natural ice samples. They were prepared every time when cutting ice and then 228 processed/analyzed along with the samples at least twice a week. Procedural blanks are 1.3±0.6 229 μ g C with an F¹⁴C of 0.69±0.15 (n=76) and 1.9±1.6 μ g C with an F¹⁴C value of 0.68±0.13 230 (n=30) for WIOC and DOC, respectively. 231

All ${}^{14}C$ results are expressed as fraction modern (F ${}^{14}C$), which is the ${}^{14}C/{}^{12}C$ ratio of the 232 sample divided by the same ratio of the modern standard referenced to the year 1950 (NIST, 233 SRM 4990C, oxalic acid II), both being normalized to -25‰ in δ^{13} C to account for isotopic 234 fractionation. All AMS F¹⁴C values presented here are finally corrected for the system and 235 236 method characteristic contributions as reported previously (e.g. Uglietti et al., 2016 and Fang et al., 2019). For WIOC analysis using the Sunset-GIS-AMS system this includes a correction 237 for the system background, i.e. constant contamination $(0.91\pm0.18 \ \mu gC$ with $F^{14}C$ of 238 239 0.72±0.11). For the cracking system applied for DOC samples the constant contamination is 240 $0.06\pm0.18 \ \mu gC$ with $F^{14}C$ of 0.50 ±0.11). Further corrections applied account for the AMS cross contamination (0.2% of the previous sample), and procedure blanks ($1.26\pm0.59 \mu gC$ with 241 $F^{14}C$ of 0.69±0.15 for WIOC samples and 1.9±1.6 µgC with a $F^{14}C$ value of 0.68±0.13 for 242 DOC samples)procedural blanks (see above). All uncertainties were propagated throughout 243 data processing until final.¹⁴C calibration. These corrections, have a larger effect on low carbon 244 mass samples (higher noise-to-sample ratio), resulting in a larger dating uncertainty. Therefore, 245 we only discuss samples with a carbon mass larger than 10 µg as recommended in Uglietti et 246 al. (2016). Radiocarbon ages are calculated following the law of radioactive decay using 5570 247 years as the half-life of radiocarbon, thus age equals $-8033 * \ln (F^{14}C)$ with -8033 years being 248 Libby's mean lifetime of radiocarbon. Radiocarbon ages are given in years before present (BP) 249 with the year of reference being 1950 (Stuiver and Polach, 1977). To obtain calibrated ¹⁴C ages, 250

the online program OxCal v4.3.2 with the IntCal13 radiocarbon calibration curve was used (Reimer et al., 2013; Ramsey, 2017). Calibrated ages, also given in years before present, are indicated with (cal BP) and denote the 1σ range unless stated otherwise.

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255 **3 Results**

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257 **3.1 DOC and WIOC concentrations**

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DOC concentrations are generally higher compared to the corresponding WIOC concentrations 259 (Figure 2). For all samples from the four glaciers, the DOC/WIOC concentration ratio ranges 260 261 from 1.2 to 4.0 with an average of 1.9 ± 0.6 (Table 2). This is at the lower end of previously reported average DOC/WIOC ratios of 2-8 (Legrand et al., 2007; Legrand et al., 2013, Fang et 262 al., in prep.). This is likely explained by temporal variability because most samples in this study 263 are several thousand years old, whereas the literature data only covers the last few centuries, 264 265 including values from the industrial period in which additional anthropogenic sources exist (e.g. fossil DOC precursors). It is interesting to note that the average DOC/WIOC ratio at Belukha 266 267 (2.5) is higher compared to the other sites (Colle Gnifetti, SLNS and Chongce is 1.8, 1.7 and 1.6, respectively). Because the Belukha glacier is surround by extensive Siberian Forests, the 268 higher ratio may be explained by particularly high emissions of biogenic volatile organic 269 270 compounds. This is corroborated by the observation that DOC concentrations are highest at this site $(241 \pm 82 \,\mu\text{g/kg})$ (Figure 2). Absolute concentrations of DOC and WIOC are slightly 271 lower at Colle Gnifetti ($112 \pm 12 \,\mu$ g/kg and $63 \pm 13 \,\mu$ g/kg, respectively) compared to the other 272 three glaciers (Table 1 and 2). Mean DOC and WIOC concentrations in the ice from the Tibetan 273 Plateau are 211 \pm 28 µg/kg and 123 \pm 19 µg/kg for SLNS and 156 \pm 40 µg/kg and 99 \pm 37 µg/kg 274 for Chongce, respectively. These values are higher compared to the pre-industrial (PI) average 275 values found in European Alpine glaciers, not only compared to the few samples from Colle 276 Gnifetti of this study, but also to previously reported values from the Fiescherhorn glacier with 277 PI-DOC of ~95 µg/kg (Fang et al., in prep.) and PI-WIOC of ~30 µg/kg (Jenk et al., 2006), 278 respectively; and from Colle Gnifetti with PI-WIOC of ~30 µg/kg (Legrand et al., 2007; Jenk 279 280 et al., 2006).

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282 **3.2 Radiocarbon results**

For all four sites, F¹⁴C of both fractions (WIOC and DOC) decreases with depth, indicating the 284 expected increase in age (Figure 2, Table 1 and 2). For three of the sites (Colle Gnifetti, Belukha 285 and SLNS), the corresponding DOC and WIOC fractions yielded comparable F¹⁴C values, with 286 no statistical evidence for a significant difference (Mann-Whitney U-test, U=79.5, n=14, 287 p=0.41>0.05). They scatter along the 1:1 ratio line-and, are significantly correlated (Pearson 288 correlation coefficient r=0.987986, p < .01, n=14) and both intercept (0.021025 ± 0.033034) 289 290 and slope (1.036034 ± 0.048050) are not significantly different from 0 and 1, respectively (Figure 3a3). Nevertheless, Figure 2 and Figure 3a suggest a slight systematic offset towards 291 lower F¹⁴C values for WIOC compared to DOC, seems evident if looking at least for higher 292 $E^{14}C$ values. Figures 2 and 3. This is particularly obvious for the samples from Chongce 293 samples, characterized by a high mineral dust load (see and from a site of very high elevation 294 with low net accumulation. For these samples, the $F^{14}C$ DOC-WIOC offset is significant 295 (discussion in Sect. 4.2) and for which $F^{14}C$ values of DOC and WIOC differ significantly.4.3). 296

For all sites, the calibrated ¹⁴C ages from both fraction fractions show an increase in age 297 with depth (Table 3). The ages range from ~0.2 to 20.3 kyr cal BP for DOC and ~0.98 to 22.4298 kyr cal BP for WIOC, respectively. In both fractions, the oldest age was derived for the sample 299 300 from the deepest part of the Belukha ice core. Samples from Colle Gnifetti generally showed younger ages (< 2 kyr cal BP). The two ice cores from the Tibetan Plateau (SLNS and Chongce) 301 302 cover a similar age span from ~ 0.2 ± 0.1 to 5.5 ± 0.3 kyr cal BP in the DOC fraction. WIO¹⁴C gaveresulted in a similar ages age range for the samples from SLNS (0.98 ± 0.4 to 6.6 ± 0.8 kyr 303 304 cal BP), but was considerably older for Chongce they resulted much older (3.1±0.7 to 11.0±1.7 305 kyr cal BP, see-discussion in SectionSect. 4.2 and 4.3).

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307 4 Discussion

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309 4.1 Radiocarbon dating with the DOC fraction

In Table 3, we present the first radiocarbon dating results of ice using the DOC fraction, with the exception of one dating point being part of the dataset to establish the very recent, first complete chronology of the Mt. Hunter ice core, Alaska (same setup and methodology as described here, Fang et al., submitted)... The DOC calibrated ¹⁴C age of ice increases with depth for all four sites, as expected for undisturbed glacier archives from the accumulation zone. The fact that none of the samples analyzed in this study (n=17) resulted in super modern $F^{14}C$

values (>1) and the obtained significant correlation between the F¹⁴C of WIOC and DOC (Sect. 316 3.2) and the resulting calibrated.¹⁴C ages (Pearson r = 0.988, p < .01, n=14, Figure S1) represent 317 strong evidence against the previously suggested ¹⁴C in situ production in the DOC fraction 318 (May, 2009). For samples from three out of the four sites, our results (Sect. 3) indicate no 319 significant difference in $F^{14}C$ between DOC and WIOC, with the latter fraction being validated 320 for allowing accurate dating of the surrounding ice (Uglietti et al., 2016). With the new DO¹⁴C 321 dating method an average dating uncertainty of around ± 200 years was achieved for samples 322 with an absolute carbon mass of 20-60 μ g, if the and ice is younger than ~6 kakyrs (Table 2). 323 This and 3). The analytical uncertainty mainly arises from the correction offor the procedure 324 blank introduced during sample treatment prior to AMS analysis (see Sect. 2 for details about 325 other corrections), contributing with 20 to 70 % to the final overall dating uncertainty. The 326 contribution thereby depends on carbon mass (larger for small samples) and sample age (the 327 larger the bigger the difference between sample and blank $F^{14}C$). See howHow the overall 328 analytical uncertainty of $F^{14}C$ decreases with higher carbon mass is shown in Figure S2 (see 329 Sect. 2 for details on blank corrections). S1. For DOC concentrations observed in this study, 330 an initial ice mass of about 250 g was required, with about 20-30 % of the ice being removed 331 during the decontamination processes inside the DOC set-up, yielding ~200 g of ice available 332 333 for final analysis. Our Expected based on previously reported DOC/WIOC concentration ratios (Sect. 3.1), the results here confirmed that with this new technique, the required ice mass can 334 thus be reduced by more than a factor of two compared to the mass needed for ${}^{14}C$ dating using 335 the WIOC fraction (expected based on the previously reported DOC/WIOC concentration ratio, 336 337 Sect. 3.1). Consequently, using the DOC instead of the WIOC fraction for ¹⁴C dating, a higher dating precision can be achieved for ice samples of similar mass. An additional benefit is that 338 the DOC extraction procedure allows monitoring to ensure complete the removal of inorganic 339 carbon (for completeness (see Sect. 2), i.e. interfering carbonate which is important to avoid a 340 potential age bias (see Sect. 4.2), leading to an improved accuracy.3). 341

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4.2 Potential contribution of carbonates to ¹⁴C of WIOC in situ production to DO¹⁴C 344

As described in Section 3.2, no significant difference between F¹⁴C of DOC and WIOC was observed
 for the ice samples from Colle Gnifetti, Belukha and SLNS (Figure 3). However, the observed offset
 for the samples from Chongce glacier needs to be discussed. Previously published WIOC ¹⁴C ages
 from the upper parts of the Chongce Cores 2 and

Previous studies suggested that $\frac{14}{C}$ of the DOC fraction may be influenced by in-situ 349 production of ¹⁴/₋C in the ice matrix (May 2009; Hoffman 2016). Induced by cosmic radiation, 350 the production of $\frac{14}{C}$ atoms within the ice matrix, i.e. by spallation of oxygen within the water 351 molecule, is a well-known process (Lal et al., 1987; Van de Wal et al., 1994). Earlier studies 352 indicated that in-situ produced ¹⁴C atoms mostly form CO, CO₂ and CH₄ (Petrenko et al., 353 2013), but also can form methanol and formic acid (Yankwich et al., 1946, Woon, 2002). The 354 mechanism of incorporation of in-situ produced $\frac{14}{10}$ C incorporation into organic molecules is 355 not well understood (Woon, 2002; Hoffman, 2016). Hoffmann (2016) performed neutron 356 357 irradiation experiments on Alpine glacier ice, showing that about 11-25 % of the initially produced $\frac{14}{C}$ atoms entered into the DOC fraction. The resulting effect on $F^{14}C$ of DOC 358 consequently depends on (i) the number of 14C atoms produced in the ice (14C in-situ 359 production), (ii) the fraction of these atoms incorporated into DOC, and because F^{14} C is based 360 on a $\frac{14}{C}/\frac{12}{C}$ ratio, (iii) the DOC concentration in the ice (the higher the smaller the resulting 361 shift in F¹⁴C-DOC). 362

The natural neutron flux, relevant for the $\frac{14}{14}$ C production rate, strongly depends on 363 altitude and latitude with a generally uniform energy distribution of the incoming neutrons 364 (Gordon et al., 2004). The ¹⁴C in-situ production in natural ice further depends on the depth in 365 the glacier and the snow accumulation rate of the site (Lal et al., 1987), determining the totally 366 received neutron radiation. Following Lal et al. (1987), the number of in-situ produced $\frac{14}{10}$ C 367 atoms in each of our ice samples was estimated, assuming an average incorporation into DOC 368 of 18±7 % (Hoffmann, 2016) (Table 4, equations and input parameters in the Supplementary 369 Material). The average F^{14} C-DOC shift for all samples is 0.044±0.033. We find a good 370 correlation between the measured F¹⁴C DOC-WIOC offset and the ¹⁴C in-situ caused F¹⁴C-371 DOC shift which explains about 50 % of the offset (Pearson r=0.82, Figure 4) and after 372 correcting for it improves the overall agreement between $F^{14}_{--}C$ of DOC and WIOC (Figure 5). 373 The shift is largest for the Chongce samples (0.109 ± 0.048) as a result of the high production 374 375 rate at 6 km altitude in combination with the low annual net accumulation rate at this site (0.14 m w.e. yr^{-1}). The calculated shift for samples from the SLNS core, from similar latitude but 376 from a site lower in altitude (5 km) and experiencing higher net accumulation (0.21 m w.e. yr-377 $\frac{1}{1}$, is significant lower with 0.038±0.016. The samples from Belukha and Colle Gnifetti are 378 least affected (0.013±0.006 and 0.033±0.013, respectively). 379

380 We find that while the effect of in-situ ${}^{14}C$ production causes only a negligible shift in 381 $\underline{F}^{14}C$ -DOC for most samples (masked by the analytical uncertainty), it can become significant for ice samples from sites of exceptional high altitude and experiencing low annual net
 accumulation rates in addition, such as the Chongce ice cap (6010 m asl., 0.14 m w.e. yr⁻¹;
 Figure 4). Note that for any site, the size of this effect gets reduced the higher the DOC
 concentration of the sample.

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387 <u>4.3 Potential contribution of carbonates to ¹⁴C of WIOC</u>

Under the basic assumption that the initially emitted fractions of DOC and WIOC are of similar 389 age, an additional contribution from $\frac{14}{C}$ -depleted carbonate (low $F^{14}C$) to the WIOC would 390 cause an $F^{14}C$ offset between the two fractions. Previously published WIOC ^{14}C ages from 391 the upper parts of the Chongce Core 2 and Core 4, less than 2 and ~6 km away from Core 1, 392 did show large scatter with no clear increase in age with depth for samples younger than 2 393 394 kakyrs. It was speculated that this was at least partly caused by the visible, exceptionally high loading of mineral dust on the WIOC filters (Hou et al., 2018). Such high mineral dust loading 395 396 was also observed during filtration of the Chongce Core 1 samples analyzed in this studypresented here. High mineral dust content in the ice can influence.¹⁴C dating with WIOC 397 in two ways, by affecting filtration through clogging of the filter and by potentially contributing 398 with ¹⁴C-depleted carbon from carbonate, as has been discussed in most previous studies. They 399 all concluded, that although for dust levels typically observed in ice cores from high elevation 400 glaciers, no significant bias is detectable for ¹⁴C of WIOC, but it wasis of concern for the 401 elemental carbon (EC) fraction combusted at higher temperatures during OC/EC separation. 402 403 This fraction EC – as well as total carbon (TC), the sum of OC and EC) – is thus not recommended to be used for radiocarbon dating (Jenk et al., 2006; Jenk et al., 2007; Jenk et al., 404 2009; Sigl et al., 2009; Uglietti et al., 2016). In any case of similar age of, the DOC 405 andcarbonate removal efficiency during WIOC fraction, a contribution of ¹⁴C-depleted 406 carbonate (low F¹⁴C) to WIOC would result in an F¹⁴C offset between the two fractions and 407 yield a slope value different from 1, since it most strongly affects high F¹⁴C values (i.e. younger 408 samples). This can easily be understood by the concept of isotopic mass balance. Therefore, to 409 testsample preparation was never quantified. 410

Here, the hypothesis that incomplete removal of carbonate resulted inmay have caused
 the F¹⁴C DOC-WIOC offset observed inremaining after accounting for DO¹⁴C in-situ
 production (Sect. 4.2) was tested. Applying an isotopic mass balance based model to our dataset,

414 wethe carbonate removal efficiency in WIO¹⁴C samples was estimated the carbonate removal
415 efficiency of our procedure by applying an isotopic mass balance model. We thereby used the .
416 The Ca²⁺ concentration in the <u>ice</u> samples was thereby used as a tracer for calcium carbonate
417 (see Supplement for details see supplement). The obtained).

We find a carbonate removal procedure incomplete by around 2 % (i.e. an average 418 removal efficiency of -96% would amount to a $-98\pm2\%$) to be sufficient for explaining the 419 remaining part of the observed $F^{14}_{--}C$ DOC-WIOC offset Figure 5). In terms of residual 420 carbonate carbon mass on the filter of -4, this equals to $< 2 \mu gC$ on average (Table 4). The 421 effect of this not entirely complete removal, can easily explain the F¹⁴C DOC-WIOC offset 422 observed for the Chongce ice core samples. By accounting for the estimated carbonate 423 contribution, the offset for these samples S2). On the one hand, this is significantly reduced to 424 the level of scatter observed for the three other sites (Figure 3b; for an according Figure in 425 agreement with calibrated ¹⁴C ages instead of F¹⁴C see S1). Based on these model results, we 426 are also confident to conclude that residual carbonate carbon is the most likely cause to explain 427 the slight systematic F¹⁴C DOC-WIOC shift we observed for the other three sites. The old 428 samples from Belukha contain the highest Ca²⁺ concentrations in this study. With their origin 429 in the Late Glacial or Glacial to Holocene transition period, these samples might not be 430 comparable to the other samples in terms of mineral dust sources and transport. Their Ca²⁺ to-431 432 carbonate ratio or the abundance of the more soluble form of bicarbonate can thus be assumed to be at least slightly different compared to the rest of the samples with the determined average 433 434 removal efficiency thus yielding a slight underestimation for the samples from Belukha (estimated to rather be ~99%, see Supplement). In any case, our results confirm the findings of 435 previous studies, indicating that the potential carbonate related bias for ¹⁴C dating using WIOC 436 is hardly detectable for ice samples with normal dust loading, typically being less than the 437 analytical uncertainty (around 10-20% of the determined age, see supplement Figure S3). For 438 example, Uglietti et al. (2016) did not detect such an effect when comparing dating results from 439 WIO¹⁴C with ages from independent methods. (effect masked by the analytical uncertainty, 440 see Figure S2). For example, Uglietti et al. (2016) did not detect such an effect when 441 successfully validating WIO¹⁴C dating results with ages from independent methods. On the 442 other hand, it demonstrates that a slightly below average removal efficiency for ice samples 443 444 containing visibly high loading of mineral dust can already cause a notable offset (93-97 % for 445 Chongce). The likely bigger particle size in such samples will affect their solubility, i.e. 446 increase the dissolution time required in the acid treatment step. In the current procedure, this

time is not adjusted accordingly (Sect. 2). Based on these results, we consider a small offset from incomplete carbonate removal to be a very likely reason contributing to the measured $F^{14}C$ DOC-WIOC, i.e. resulting dating offset (Figure 5). Instead of a correction, which does not seem feasible for this effect because of large uncertainties and likely substantial site-to-site (sample-to-sample) variations, we suggest future improvement in the analytical procedure of the carbonate removal step (e.g. a slight increase in acid concentration and an increase of the reaction time).

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455 **4.34** DO¹⁴C ages in the context of published chronologies

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In the following we will discuss our new DO¹⁴C results in the context of ages from previous 457 studies (Table 5, Figure 4). For final calibration of ${}^{14}C$ ages, most of those earlier studies took 458 advantage of the assumption of sequential deposition in the archive, which seems very 459 460 reasonable considering thei.e. a continuous, undisturbed and preserved sequential deposition of annual snow layers on top of each other on the glacier surface. Particularly in case of 461 462 relatively large analytical uncertainties compared to the age difference of the samples, the sequential deposition model can moderately constrain the probability distribution of the 463 464 calibrated age range in each sample of the dataset. For consistency we applied the same 465 calibration approach here by using the in-built OxCal sequence model (Ramsey, 2008). For all DO¹⁴C data presented in this study, there is While the underlying assumption may not generally 466 be valid for all sites, and individually needs to be carefully assessed, we find no difference in 467 the calibrated ages using the sequence model and the ages from the conventional calibration 468 approach (Table 3). for all DO¹⁴C data presented in this study (Table 3). Note, that no 469 correction for a potential in-situ $\frac{14}{C}$ bias was applied to the DO $\frac{14}{C}$ data used here (Section 470 471 4.2).

We obtained the oldest age of ~21 kyr cal BP for the bedrock ice at Belukha, indicating this glacier to be oldest and of Pleistocene origin (Figure 4)... This is older than the previously reported age of ~11 kyr cal BP, which (Table 5, Figure 6). The latter age was retrieved obtained for thean ice core from the nearby Belukha West Plateau glacier extracted in 2003 (B03) and not from the saddle as the 2018 core (B18) analyzed in this study (Aizen et al., 2016; Uglietti et al., 2016). However,) opposed to the 2018 core extracted from the saddle (B18) analyzed in this study. Also, the according sample from B03 was from a slightly shallower depth (0.6-0.3 479 m above bedrock-) than the sample analyzed from B18 in this study. The age range modeled for B03 atfor the same depth above bedrock as sampled in this study (0.5-0 m) is in better 480 agreement with ~28 kyr cal BP withand a very large uncertainty of ~15 kyr (Uglietti et al., 481 482 2016). Thus Overall, our new age for the oldest ice at Belukha thus reasonably agrees well-with this the previous result, but yields a much better constrained age with a strongly reduced 483 uncertainty of ± 4 kyr. The two glaciers from the Tibetan Plateau (SLNS and Chongce) show 484 485 very similar bottom ages of ~5-6 kyr cal BP₇ (Figure 6), which is in agreement with the previously reported age range of Tibetan Plateau glaciers (Hou et al., 2018). The bottom age 486 of Chongce Core 1 determined here <u>based on DO¹⁴C</u> (5.6 \pm 0.3 kyr cal BP) is slightly younger 487 than the previously reported bottom age in Core 2 based on WIO¹⁴C (6.3 ± 0.3 kyr cal BP, Hou 488 et al., 2018), which is likely related toin agreement with the influence of residual carbonates 489 on the latter (findings discussed in Section 4.2 and 4.3. Nevertheless, our new age is still in the 490 range of the previously estimated bottom age (Table 5, Figure 6). The bottom most sample of 491 the Colle Gnifetti 2015 (CG15) core could not be dated because the small amount of ice 492 available yielded an insufficient carbon mass of $<10 \ \mu g$ for ^{14}C analysis. Previous WIO¹⁴C 493 dating of a core obtained at Colle Gnifetti in 2003 (CG03) also revealed ice of Pleistocene 494 495 origin with the ice at bedrock being older than 15 kyr cal BP (Jenk et al., 2009). TheAs expected, 496 the age obtained in this study from a shallower depth was much younger with 1.2 kyr cal BP, but. This is in excellent agreement with the age of CG03 atfor a similar depth (~74 m below 497 498 surface). This is; Table 5, Figure 6). We consider this as a clear indication that the CG15 ice 499 core-drilling did not reach bedrock.

500 <u>Overall, the dating with DO¹⁴C results in ages which are in good agreement with the</u> 501 age ranges reported in earlier studies. Even though a contribution from in-situ ¹⁴C to DO¹⁴C 502 was not considered in the comparison here, we find that the dating by the DOC fraction does 503 not lead to significantly different results compared to dating by WIO¹⁴C or cause a different 504 interpretation about the oldest ice still present for any of the sites.

505

506 **5 Conclusion**

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In this study, we evaluated and successfully validated the DO¹⁴C dating technique by directly comparing with <u>direct comparison of dating</u> results from with the well-established WIO¹⁴C dating method onusing parallel <u>ice</u> samples. Achieving this goal was not only analytically

demanding but also highly challenging because of due to the very limited availability of the 511 sampling material, requiring ice in rather large quantities and spanning a wide range of ages. 512 The obtained DO¹⁴C ages for four different Eurasian glaciers, ranging from 0.2 ± 0.2 to 20.3 513 \pm 4.1 kyrs cal BP, agreed well with the respective WIO¹⁴C ages (0.98 \pm 0.4 to 22.54 \pm 1.1 kyrs 514 cal BP) and with previously published chronologies from these ice core sites. This underlines 515 the great potential for applying $DO^{14}C$ analysis for ice core dating. Effects of in-situ ¹⁴C 516 production on DO¹⁴C ages, as suspected in previous studies, were not observed. Our data 517 confirmed that pre-industrial DOC concentrations are higher by a factor of about two than 518 WIOC concentrations in high alpine ice cores. With the new DO¹⁴C dating methodWith this 519 new method, an average dating uncertainty of around ± 200 years was achieved for samples 520 with an absolute carbon mass $> 20 \mu g$ and ages up to ~ 6 ka. For DOC concentrations observed 521 in this study, an initial ice mass of about 250 g was required. The sample mass can thus be 522 reduced by more than a factor of two compared to the mass needed for.¹⁴C dating using the 523 WIOC fraction. Furthermore, the new DO¹⁴C dating revealed a slight age bias of the WIOC 524 fraction towards older ages, which however was found to be significant only for ice containing 525 high concentrations of mineral dust and in particular if younger than ~1000 years, in agreement 526 with findings of previous studies. In any case, for this age range, other independent dating 527 methods exist and the use of these in combination with radiocarbon ages in a glaciological flow 528 model to derive a final ice core age-depth relationship provides additional constraint, thus 529 reducing potential bias, for these younger ice sections (less weight given to the potentially 530 biased radiocarbon ages in this range). Nevertheless, future work for improving the carbonate 531 removal procedure is encouraged to further increase the accuracy of ¹⁴C dating with the WIOC 532 fraction of $> 20 \mu g$ and ages up to ~ 6 kyrs. For DOC concentrations observed in this study, an 533 534 initial ice mass of about 250 g was required. Our data confirmed previous results that concentrations of pre-industrial DOC are higher by about a factor two compared to WIOC 535 concentrations in high alpine ice cores. This shows that the required ice mass to achieve similar 536 precision is reduced by at least a factor of two for ¹⁴/₂C dating when using the DOC instead of 537 the WIOC fraction. Accordingly, an improvement in precision can be achieved for same sample 538 mass. Compared to WIOC, a downside of using the DOC fraction for $\frac{14}{14}$ C dating is a more 539 demanding and time consuming extraction procedure. In addition, because of its higher 540 solubility and a related higher mobility of DOC in case of meltwater formation, this fraction is 541 only applicable for dating ice which had been cold throughout its "lifetime". Beneficial 542 compared to WIOC, there is no potential for a dating bias by carbonates of mineral dust for 543 DO¹⁴C. However, our results confirm previously suggested potential dating biases from in-situ 544

¹⁴C causing DO¹⁴C dates to shift towards younger ages. While we find the effect to be small 545 (at the level of analytical uncertainty), it may become significant for $DO^{14}C$ dating of ice 546 547 samples from sites of e.g. exceptional high altitude, experiencing low annual net accumulation 548 rates in addition. For such sites, a reasonably accurate correction to account for the age bias 549 seems feasible according to our results, although at the cost of an increase in the final dating uncertainty. Nevertheless, we think this new dating method has a great potential to open up 550 551 new fields for radiocarbon dating of ice for example from remote regions, where concentrations of organic impurities in the ice are particularly low. 552

In summary the main benefits of DO¹⁴C dating of glacier ice compared to WIO¹⁴C dating are 553 the reduced ice amount required, or higher precision for the same sample mass. The DOC 554 fraction is not affected from potential carbonate bias from mineral dust with the applied 555 methodology and thus ensures high dating accuracy. This new dating method opens up new 556 fields for radiocarbon dating of ice for example from remote or Polar Regions, where 557 concentrations of organic impurities in the ice are particularly low. Compared to the WIO¹⁴C 558 559 dating, a downside is the more demanding and time consuming DOC extraction procedure and because of its higher solubility, the higher mobility of DOC in case of meltwater, thus only 560 561 being applicable for dating ice which had been cold throughout its "lifetime".

562

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574 Data availability

575 The data is provided in the Tables.

576 Author contributions

577 LF and TS performed ¹⁴C analysis. LF, TS, TMJ, and MS wrote the manuscript while all 578 authors contributed to the discussion of the results. MS designed the study.

579 Competing interests

580 The authors declare that they have no conflict of interest.

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Core section	Depth (m)	Ice mass (kg)	WIOC (µg)	Concentration (µg/kg)	Bern AMS Nr.	F ¹⁴ C (±1σ)	14 C age (BP, ±1 σ)
CG110	72.1-72.7	0.570	35.2	61.9±3.3	11770.1.1	0.875±0.011	1073±105
CG111	72.7-73.4	0.539	38.7	71.8±3.8	11771.1.1	0.848±0.011	1321±101
CG112	73.4-73.9	0.536	23.7	44.1±2.4	11772.1.1	0.852±0.015	1284±143
CG113	73.9-74.6	0.549	39.8	72.4±3.8	11773.1.1	0.786±0.011	1937±109
Belukha412	158.3-159.0	0.443	37.8	85.2±4.5	11766.1.1	0.367±0.010	8055±211
Belukha414	159.5-160.3	0.336	27.8	82.6±4.4	11768.1.1	0.212±0.014	12473±535
Belukha415	160.3-160.9	0.319	39.3	123.3±6.5	11769.1.1	0.100±0.011	18462±899
SLNS101	56.8-57.5	0.420	41.5	98.9±2.1	12325.1.1	0.902±0.047	825±420
SLNS113	64.7-65.4	0.427	45.3	106.1±2.5	12324.1.1	0.852±0.046	1284±438
SLNS122	68.9-69.7	0.424	58.5	138.0±3.6	12323.1.1	0.807±0.046	1727±459
SLNS127	71.8-72.5	0.483	50.9	105.3±2.5	12322.1.1	0.695±0.046	2921±532
SLNS136	76.7-77.5	0.374	50.6	135.2±3.0	12321.1.1	0.521±0.046	5235±706
SLNS139	78.9-79.6	0.485	61.2	126.3±3.6	12320.1.1	0.521±0.045	5232±703
SLNS141-142	80.3-81.0	0.413	61.7	149.5±3.8	12319.1.1	0.489±0.046	5754±750
CC237	126.0-126.7	0.352	22.4	63.7±1.8	12328.1.1	0.704±0.049	2815±555
CC244	130.2-130.8	0.311	29.8	95.9±2.2	12327.1.1	0.639±0.048	3602±600
CC252	133.4-133.8	0.174	23.8	136.7±4.3	12326.1.1	0.316±0.049	9256±1250

Table 1 WIOC samples analyzed from Colle Gnifetti, Belukha, SLNS and Chongce ice cores.

Core section	Depth (m)	Ice mass (kg)	DOC (µg)	Concentration (µg/kg)	Bern AMS Nr.	$F^{14}C(\pm 1\sigma)$	¹⁴ C age (BP, ±1σ)	DOC/WIOC
CG110	72.1-72.7	0.171	18.9	110.0±2.7	11575.1.1	0.943±0.030	474±259	1.8
CG111	72.7-73.4	0.207	25.5	122.9±3.0	11576.1.1	0.901±0.021	836±190	1.7
CG112	73.4-73.9	0.248	23.6	95.0±2.3	11577.1.1	0.889±0.021	943±192	2.2
CG113	73.9-74.6	0.246	29.5	119.4±2.9	11578.1.1	0.849±0.016	1312±151	1.7
Belukha412	158.3-159.0	0.172	28.5	165.0±4.0	11581.1.1	0.315±0.024	9284±624	1.9
Belukha414	159.5-160.3	0.128	41.9	327.4±7.9	11584.1.1	0.239±0.019	11505±648	4.0
Belukha415	160.3-160.9	0.102	23.7	231.0±5.6	11585.1.1	0.144±0.041	15584±2365	1.9
SLNS101	56.8-57.5	0.238	44.0	184.9±4.5	12458.1.1	0.972±0.016	227±131	1.9
SLNS113	64.7-65.4	0.213	39.4	185.2±4.5	12459.1.1	0.942±0.016	484±137	1.7
SLNS122	68.9-69.7	0.234	57.9	248.0±6.0	12460.1.1	0.773±0.010	2073±101	1.8
SLNS127	71.8-72.5	0.252	57.8	229.7±5.5	12461.1.1	0.730±0.009	2527±101	2.2
SLNS136	76.7-77.5	0.220	48.3	219.1±5.3	12462.1.1	0.657±0.009	3380±112	1.6
SLNS139	78.9-79.6	0.208	48.1	230.8±5.6	12463.1.1	0.580±0.009	4381±131	1.8
SLNS141-142	80.3-81.0	0.246	43.8	177.5±4.3	12464.1.1	0.550±0.010	4809±151	1.2
CC237	126.0-126.7	0.208	28.5	136.6±3.3	12454.1.1	0.980±0.023	161±185	2.1
CC244	130.2-130.8	0.167	21.7	129.8±3.1	12455.1.1	0.800±0.018	1789±185	1.4
CC252	133.4-133.8	0.120	24.3	202.5±4.9	12456.1.1	0.546±0.016	4854±239	1.5

 Table 2 DOC samples analyzed for Colle Gnifetti, Belukha, SLNS and Chongce ice cores.

Table 3 Calibrated WIO¹⁴C and DO¹⁴C ages using OxCal v4.3.2 with the Intcal13 radiocarbon calibration curve. Ages are given as the OxCal provided μ -age $\pm 1\sigma$, which is the calibrated mean age accounting for the age probability distribution. In addition, calibrated ages derived when applying the OxCal sequence deposition model for further constraint are shown.

Core section	WIOC	WIOC Cal age	DOC	DOC Cal age
	Cal age	with sequence	Cal age	with sequence
	(cal BP)	(cal BP)	(cal BP)	(cal BP)
CG110	1024±110 1004±119	1003±99 968±1049	464±235	403±196
CG111	1238±96 <u>1224±103</u>	1198±81<u>1174±86</u>	810±169	749±123
CG112	1209±130<u>1190</u>±142	1310±98 1292±103	901±176	947±139
CG113	1898±130<u>1889</u>±138	1890±123 <u>1869±143</u>	1222±153	1248±144
Belukha412	8957±265 8960±266	8953±247 8954±268	10695±867	10701±861 10686±865
Belukha414	14780±781<u>14796</u>±782	14741±691<u>14802</u>±774	13646±893	15063±737<u>13670±880</u>
Belukha415	22485±1112 22441±1107	22343±949 22497±1107	20264±4073	20605±3936 20393±4033
SLNS101	851±395 848±396	707±317 701±315	250±145	226±137
SLNS113	1298±452<u>1297</u>±453	1264±333 1255±331	480±131	505±111
SLNS122	1775±513 1769±514	1902±427<u>1901±4301</u>	2057±129	2056±129
SLNS127	3178±677 <u>3175±679</u>	3223±625 3221±629	2585±125	2585±125
SLNS136	6033±829 <u>6030±824</u>	5424±617 5426±620	3635±138	3636±137
SLNS139	6032±811<u>6026</u>±820	6171<u>6177</u>±567	5014±191	5007±187
SLNS141-142	6619±841<u>6626±831</u>	7069±676 7081±689	5519±188	5531±176
	3057±704	2956±56 4		233±153
	4050±764	4214±699		1738±212
CC237	10998±1689<u>3051</u>±703	10948±1656 2886±617	237±151	5580±295
CC244	<u>4057±769</u>	<u>4210±713</u>	1737±211	
CC252	<u>11000±1697</u>	<u>11017±1716</u>	5580±294	

Core section	Ca ²⁺ -concentration	ice sample mass	residual carbonate C
	(ppb)	(kg)	(µgC)
CG110	100	0.570	0.3
CG111	110	0.539	0.3
CG112	61	0.536	0.2
CG113	59	0.549	0.2
Belukha412	4191	0.443	10.2
Belukha414	7566	0.336	13.9
Belukha415	3737	0.319	6.5
SLNS101	1400*	0.420	3.2
SLNS113	same	0.427	3.3
SLNS122	same	0.424	3.3
SLNS127	same	0.483	3.7
SLNS136	same	0.374	2.9
SLNS139	same	0.485	3.7
SLNS141-142	same	0.413	3.2
CC237	2170 [#]	0.352	4.2
CC244	same	0.311	3.7
CC252	same	0.174	2.1

Table 4 Estimated residual carbonate carbon on the analyzed WIOC filters. Ca²⁺ concentrations, used here as a tracer for carbonates, are average values for the sampled ice core sections (or site if data not available) of which the mass is also indicated.

*No Ca²⁺ concentrations are available for SLNS, instead the average Ca²⁺ concentration over the last 7000 years measured on the nearby Puruogangri ice cap on the central Tibetan Plateau are used here(Thompson et al., 2006).

[#]-Ca²⁺ concentration over the period of 1903-1992 from another core drilled on the Chongce ice cap by a different group (ChongYi et al., 2016).

Table 4 Estimate of the effect from in-situ ${}^{14}C$ production on $F^{14}C$ -DOC. For comparison, the measured $F^{14}C$ offset between DOC and WIOC is also shown.

Core section	Ice mass	Carbon mass	Depth	<u>P</u> o	<u>In-situ ¹⁴C</u>	<u>In-situ F¹⁴C-DOC</u>	Observed F ¹⁴ C	<u>In-situ</u>	<u>In-situ</u>
	<u>(g)</u>	<u>(µg)</u>	<u>(m w.e.)</u>	<u>(¹⁴C atom g⁻¹ ice yr⁻</u>	(atoms)	offset	DOC-WIOC	<u>corrected F¹⁴C-</u>	corrected DOC
				<u>1</u>)			<u>offset</u>	DOC	Cal age
CC110	171	19.0	EE 0	220	1107	0 022±0 012	0.00010.000	0 010+0 022	<u>(Cal BP)</u>
	1/1	<u>10.9</u>	55.0	<u>520</u>	1197	<u>0.055±0.015</u>	<u>0.068±0.032</u>	<u>0.910±0.055</u>	<u>/52±2/3</u>
<u>CG111</u>	207	25.5	<u>56.3</u>	328	<u>1197</u>	<u>0.030±0.012</u>	<u>0.053±0.024</u>	<u>0.901±0.024</u>	<u>1045±207</u>
<u>CG112</u>	<u>248</u>	<u>23.6</u>	<u>56.7</u>	<u>328</u>	<u>1197</u>	<u>0.038±0.015</u>	<u>0.037±0.026</u>	<u>0.889±0.026</u>	<u>1225±250</u>
<u>CG113</u>	<u>246</u>	<u>29.5</u>	<u>57.0</u>	<u>328</u>	<u>1197</u>	0.030±0.012	0.064±0.019	<u>0.849±0.020</u>	<u>1546±208</u>
<u>Belukha412</u>	<u>172</u>	<u>28.5</u>	<u>142.7</u>	<u>286</u>	<u>921</u>	<u>0.017±0.007</u>	<u>-0.052±0.026</u>	<u>0.315±0.025</u>	<u>11271±902</u>
<u>Belukha414</u>	<u>128</u>	<u>41.9</u>	<u>143.9</u>	<u>286</u>	<u>921</u>	0.009±0.003	<u>0.027±0.024</u>	<u>0.239±0.020</u>	<u>14096±964</u>
<u>Belukha415</u>	<u>102</u>	<u>23.7</u>	<u>144.5</u>	<u>286</u>	<u>921</u>	0.012±0.005	<u>0.043±0.043</u>	<u>0.144±0.041</u>	<u>21571±4753</u>
<u>\$LNS101</u>	<u>238</u>	<u>44</u>	<u>47.9</u>	<u>345</u>	<u>2666</u>	<u>0.044±0.017</u>	<u>0.070±0.050</u>	<u>0.972±0.023</u>	<u>587±187</u>
<u>\$LNS113</u>	<u>213</u>	<u>39.4</u>	<u>54.4</u>	<u>345</u>	<u>2656</u>	0.044±0.017	<u>0.089±0.050</u>	<u>0.942±0.023</u>	<u>837±184</u>
<u>\$LNS122</u>	<u>234</u>	<u>57.9</u>	<u>58.1</u>	<u>345</u>	<u>2651</u>	0.033±0.013	-0.034±0.047	<u>0.773±0.016</u>	<u>2483±210</u>
<u>\$LNS127</u>	<u>183</u>	<u>57.8</u>	<u>60.5</u>	<u>345</u>	<u>2647</u>	<u>0.026±0.010</u>	0.029±0.047	<u>0.730±0.014</u>	<u>2967±197</u>
<u>\$LNS136</u>	<u>220</u>	<u>48.3</u>	<u>64.7</u>	<u>345</u>	<u>2641</u>	<u>0.037±0.014</u>	<u>0.135±0.047</u>	<u>0.657±0.017</u>	<u>4264±304</u>
<u>\$LNS139</u>	<u>208</u>	<u>48.1</u>	<u>66.5</u>	<u>345</u>	<u>2638</u>	0.035±0.014	<u>0.058±0.046</u>	<u>0.580±0.016</u>	<u>5600±290</u>
<u>SLNS141-142</u>	<u>246</u>	<u>43.8</u>	<u>67.7</u>	<u>345</u>	<u>2636</u>	<u>0.045±0.018</u>	<u>0.061±0.047</u>	<u>0.550±0.020</u>	<u>6323±363</u>
<u>CC237</u>	<u>208</u>	<u>28.5</u>	<u>113.7</u>	<u>497</u>	<u>5371</u>	<u>0.120±0.046</u>	<u>0.275±0.054</u>	0.980±0.052	<u>1240±498</u>
<u>CC244</u>	<u>167</u>	<u>21.7</u>	<u>117.6</u>	<u>497</u>	<u>5353</u>	<u>0.126±0.049</u>	<u>0.161±0.051</u>	0.800±0.052	<u>3509±799</u>
<u>CC252</u>	<u>120</u>	<u>24.3</u>	<u>120.2</u>	<u>497</u>	<u>5341</u>	<u>0.080±0.031</u>	<u>0.231±0.051</u>	<u>0.546±0.035</u>	<u>7007±635</u>

Site	Study	Core	Dating method	Depth above bedrock (m)	Age (cal BP)
Colle Gnifetti	this study	CG15	DO ¹⁴ C	(74.3 m below surface).*	1248 ± 144
	Jenk et al., 2009	CG03	WIO ¹⁴ C	(73.5 m below surface) [#]	1152 ± 235
	Jenk et al., 2009	CG03	Model	(74.3 m below surface). ^{&}	$1160 \pm ^{140}_{170}$
	Jenk et al., 2009	CG03	WIO ¹⁴ C	0.6-0	>15000
	Jenk et al., 2009	CG03	Model	oldest ice estimate	$19100 \pm ^{4800}_{4500}$
Belukha	this study	B18 (saddle)	DO ¹⁴ C	0.5-0	20605 ± 3936 20393 ±
	Aizen et al., 2016	B03 (west plateau)	WIO ¹⁴ C	0.6-0.3	$\frac{4033}{11015 \pm 1221}$
	Uglietti et al., 2016	B03 (west plateau)	Model	0.6-0	28500 ± 16200
SLNS	this study	SLNS	DO ¹⁴ C	0.4-0	5531 ± 176
	no previous results				
Chongce	this study	Core 1	$DO^{14}C$	0.2-0	5580 ± 295
	Hou et al., 2018	Core 2	WIO ¹⁴ C	1.2-0.8	6253 ± 277
	Hou et al., 2018	Core 2	Model	oldest ice estimate	$9000 \pm ^{7900}_{3600}$

Table 5 $DO^{14}C$ dating results for near bedrock ice compared to results from previous studies (visualized in Figure 4<u>6</u>).

*precise bedrock depth unknown at this coring site, *sampled depth being closest to depth sampled in this study; (CG03 and CG15 drill sites only <u>16 m apart</u>), *modeled age at same depth as sampled in this study.



Figure 1: Location of the four glaciers Colle Gnifetti, Belukha, Chongce, and Shu Le Nan Shan (SLNS). Map made from Matlab R2019b geobasemap. Colle Gnifetti is located in the Monte Rosa massif in the Swiss Alps, Belukha glacier in the Altai mountain range, Russia, the Chongce ice cap on the northwestern Tibetan Plateau, China, and the SLNS at the south slope of the Shulenanshan Mountain, China.



Figure 2: Comparison of results from the WIOC and DOC fractions for the studied four sites. (a) concentrations (b) $E^{14}C$. The error bars denote the overall analytical 1 σ uncertainty.





Figure 3: Scatter plot showing the correlation between \mathbb{F}^{14} <u>C of WIOCWIO¹⁴</u><u>C</u> and DOC. (a) <u>Measured values DO¹⁴<u>C</u> results for the four sites (see legend). In terms of \mathbb{F}^{14} <u>C (top) and</u> <u>calibrated ages (bottom).</u> For the linear fit shownin both panels, the data from Chongce (open symbols) was not-excluded. Shaded areas indicate the 95% confidence band.</u>



Figure 4 Estimated in-situ ${}^{14}C$ offset to $F{}^{14}C$ -DOC plotted against the measured offset between $F{}^{14}C$ of the DOC and WIOC fraction.



Figure 5 Scatter plots showing the correlation between WIO¹⁴C and DO¹⁴C results for all samples. In terms of $F^{14}C$ (top) and calibrated ages (bottom). (a) Measured values as shown in Figure 3 but with the linear fit applied to all data (Chongce included). (b) Same as panel (a), but DOC ¹⁴C results corrected for in-situ ¹⁴C contribution. (c) Same as panel (a), but DOC and WIOC ¹⁴C results corrected for in-situ ¹⁴C and accounting for potentially incompletely removed carbonate, respectively. An estimated average carbonate removal efficiency of 98±2 % was used here. Error bars in panel (a) and (b) reflect the propagated uncertainty of analysis and correction. In panel (b) and (c), measured values are shown as gray crosses.



Figure 4 Age of ice at the very bottom of the four glaciers (close to bedrock). DO¹⁴C dates. are shown



Figure 6 Comparison of our DO¹⁴C ages (not corrected for in-situ) with filled squares and dating results from previous studies if available. For the four sites of Belukha (BEL), Colle Gnifetti (CG), Shu Le Nan Shan (SLNS) and Chongce (CC), DO¹⁴C ages (green) and previously reported WIO¹⁴C ages indicated by open circles. The (open circles) for similar sampling depths are shown. Gray bars indicate previously modeled, ¹⁴C based bedrock age estimates (additionally for CG the modeled estimate of age for the Belukha bottom age (gray bar) issampling depth of this study). Previously published data are from Uglietti et al. (2016). The modeled Colle Gnifetti bottom age from) (BEL, West Plateau), Jenk et al. (2009) and the indicated WIO¹⁴C value is from ice sampled from the CG03 core at similar depth as in this study. For Chongce, the result of the deepest WIOC sample from core 2 is shown together with the modelled bottom age ((CG), and Hou et al., (2018). Find all-) (CC). See Table 5 for underlying data and information summarized in Table 5. details.