New evidence on the microstructural localization of sulfur & chlorine in polar ice cores with implications for impurity diffusion

Pascal Bohleber^{1,2,3}, Nicolas Stoll³, Piers Larkman^{1,3}, Rachael H. Rhodes⁴ and David Clases⁵

Alfred Wegener Institute Helmholtz Center for Polar and Marine Research, Bremerhaven, Germany.

5 ²Goethe University Frankfurt am Main, Frankfurt am Main, Germany

³Ca' Foscari University of Venice, Department of Environmental Sciences, Informatics and Statistics, Italy,

⁴Department of Earth Sciences, University of Cambridge, Cambridge, UK.

⁵Nano Micro LAB, Institute of Chemistry, University of Graz, Graz, Austria

Correspondence to: Pascal Bohleber (pascal.bohleber@awi.de)

Abstract. Aerosol-related impurities play an important part in the set of paleoclimate proxies obtained from polar ice cores. However, in order to avoid misinterpretation, post-depositional changes need to be carefully assessed, especially in deep ice. Na, S and Cl are among the relatively abundant impurity species in polar ice (albeit still at the low ppb level in bulk samples), with important applications to paleoclimate reconstructions and dating, e.g. via identification of volcanic eruptions. Especially S has been studied intensely with respect to peak broadening with depth/age related to diffusion, but the precise physical mechanisms remain unclear. Mapping the two-dimensional impurity distribution in ice with laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) has shown great potential for studying ice-impurity interactions, but the analytically more challenging elements S and Cl have not been targeted thus far. We show here that signals of S and Cl can be detected in Greenland and Antarctic ice by LA-ICP-MS mapping. In ice without evidence of volcanic activity, and unenhanced impurity concentrations, we obtain multi-elemental maps for Na, Cl and S at high resolution up to 10 µm and also include some exemplarily high resolution maps with a spot size down to 1 µm, We use Na as a previously investigated reference element and find a high level of localization of Na, S and Cl at grain boundaries but also some dispersed occurrence within grain interiors in dust-rich ice. The new maps support a view on diffusive transport not only through ice veins but also along grain boundaries. In the EPICA Dome C ice core samples we do not find any clear differences in impurity localization, between samples from the Holocene and last glacial period. These results extend early studies targeting the localization of impurities, in particular through measuring S and Cl, and highlight the benefit of integrating such direct measurements with modelling efforts to determine the physical processes behind impurity diffusion.

1 Introduction

Aerosol-related atmospheric impurity records are an important part of the unique combination of climate proxies archived in polar ice cores (e.g. Fischer et al., 2021; Legrand & Mayewski 1997). However, it is known that the original impurity distribution at the foundation of interpreted climatic signals can be disturbed by various post-depositional processes, and their

hat gelöscht: .

hat formatiert: Englisch (USA)

hat gelöscht: & sodium

hat formatiert: Englisch (USA)

hat gelöscht: not containing enhanced concentrations resulting from a volcanic eruption,

hat gelöscht: using

hat gelöscht: to

hat formatiert: Englisch (USA)

hat gelöscht: and, exemplarilydown even 1µm

hat gelöscht: (and Na)

hat formatiert: Englisch (USA)

hat gelöscht:

hat gelöscht: but

hat gelöscht: show

hat gelöscht: this regard

hat gelöscht: in the EDC ice core

hat formatiert: Englisch (USA)

.

impact needs to be detected and constrained to avoid misinterpretation (e.g. Traversi et al., 2009, Stoll et al. 2021).

45 Understanding the variability of impurities in the deepest and highly thinned ice core layers is particularly urgent for new "Oldest Ice" efforts to retrieve climatic records older than 1 million years from Antarctica — driven by the unsolved challenge to understand the cause of the "Mid-Pleistocene Transition" (Brook et al., 2006; Fischer et al., 2013; Yan et al., 2019; Wolff et al., 2022). Impurity (re-)localization is relevant for understanding climate record preservation, but it is also critical for the study of deformation processes on the microscale, and thus a topic of wide significance for ice core scientists and glaciologists

50 alike.

Post-depositional relocation processes are expected to depend crucially on the physical and chemical properties of the individual impurity species. One very important process of post-depositional impurity relocation is the diffusion of ionic impurities through the ice matrix. Sodium, chloride and sulfate provide an important reference ensemble in this regard, motivating the focus of the work presented here. For the upper 350 m of the deep EPICA Dome C (EDC) ice core drilled in central Antarctica, Barnes et al. (2003) investigated the diffusion of ionic sodium, chloride and sulfate and found evidence for the diffusion of only chloride and sulfate. Evidence for diffusion came from the distinct broadening and damping of sulfuric acid peaks in meltwater analysis, associated with volcanic eruptions that provide crucial tie points for ice core chronologies (e.g., Severi et al., 2007; Svensson et al., 2013; Fujita et al., 2015; Sigl et al., 2022). The rate of sulfate diffusion is typically described by the term "effective diffusivity" (units: m² yr⁻¹) - "effective" because it represents a time-weighted value that is the cumulative result of all diffusion experienced since deposition. Notably, the effective diffusivity metric lacks a foundational connection to a physical process (or processes) driving the diffusion in the ice. The localization of sulfur in the ice matrix is a key issue for describing diffusion processes in models, however. In a prominent modelling study, Rempel et al. (2001) suggested that a process termed "anomalous diffusion", happening within the connected network of veins, can lead to advection of sulfate peaks away from their original position. Ng (2021) achieved an important extension of the modelling efforts to include the "Gibbs Thompson effect", finding that if this effect is included, significant dampening of the peaks will prevent any detectable migration away from their original position. Regarding the implications of the findings, Ng distinguishes two scenarios, one with impurities completely localized at the ice veins (scenario 1) and another with significant contributions from grain boundaries and/or grain interiors (scenario 2). It becomes clear that the localization of impurities within the ice plays a key role in conceptualizing potential diffusion mechanisms. Yet, existing data do not provide sufficient constraint, e.g. to generally distinguish between the above two scenarios by Ng (2021). Notably the localization of the impurities in the ice can already provide significant new insight by itself and does not necessarily need to include precise concentration estimates.

Despite early attempts, observational evidence has not provided a fully comprehensive picture on the matter thus far. We

75 describe a few key studies here; detailed summaries with additional information are given elsewhere (e.g. Stoll et al., 2021).

Due to the low eutectic point of sulfuric acid, it was hypothesized that it is mostly located within a network formed by connected triple junctions (the intersection of grain boundaries of three grains) where it can diffuse along the veins

hat gelöscht: a

hat gelöscht: challenge

hat verschoben (Einfügung) [2]

hat gelöscht: 1

hat gelöscht: therefore

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

hat gelöscht:

One very important process in this framework is the potential diffusion of ionic impurities through the ice matrix, which has been detected in deep EPICA Dome C (EDC) ice through

hat formatiert: Englisch (USA)

hat gelöscht: (Barnes et al., 2003)

hat gelöscht: , which

hat verschoben (Einfügung) [4]

hat verschoben (Einfügung) [5]

hat verschoben (Einfügung) [3]

hat gelöscht: also from a diffusion modelling point of view

hat gelöscht: E

hat gelöscht: es

hat nach oben verschoben [4]: The rate of sulfate diffusion is typically described by the term "effective diffusivity" (units: m² yr¹) – "effective" because it represents a time-weighted value that is the cumulative result of all diffusion experienced since deposition. Effective diffusivity is typically calculated as the diffusion coefficient in a 1D model fitted to match the peak broadening or peak damping observed in empirical measurements (Barnes et al., 2003; Rhodes et al. 2024, Fudge et al. 2024). ¶

hat nach oben verschoben [5]: Notably, the effective diffusivity metric lacks a foundational connection to a physical process (or processes) driving the diffusion in the ice.

hat gelöscht: Based on meltwater analysis, it has been shown that while these peaks last typically only a few years in the upper ice sections (consistent with the duration of volcanic acid deposition on the ice sheet), their apparent duration at significant depth can stretch to a few decades (Rhodes et al., 2024). The rate of sulfate diffusion is typically described by the term "effective diffusivity" (units: m" yr") – "effective" because it represents a time-weighted value that is the cumulative result of all diffusion experienced since deposition. Effective diffusivity is typically calculated as the diffusion coefficient in a 1D model fitted to match the peak broadening or peak damping observed in empirical measurements (Barnes et al., 2003; Rhodes et al. 2024, Fudge et al. 2024). ¶

hat gelöscht: The localization of impurities within the ice plays a key role in conceptualizing potential diffusion mechanisms.

subsequently (Mulvanev et al., 1988). Barnes et al. (2003) discussed two different mechanisms distinguishing a "connected" versus "disconnected" ice vein or grain boundary network, both implying an indirect control on the effectivity diffusivity through the grain growth rate. Two recent studies extended this approach and focused on determining the effective diffusivity for sulfate at greater depths in the EDC core. Using the same input dataset, the determined values for the effective diffusivity differ beyond the Holocene, which is attributed mainly to different methodologies (Fudge et al., 2024; Rhodes et al., 2024). Importantly, the effective diffusivity of sulfate appears to decrease rapidly following deposition, with more than one order of magnitude difference between the early Holocene and the last glacial period (Rhodes et al., 2024). Rhodes et al. (2024) speculate that the decrease could be due to change in the localization of the impurities and therefore a change in diffusion mechanism.

Using cryo-scanning electron microscope (SEM) X-ray analysis or Raman spectroscopy on ice samples from shallow depths, S signals were detected primarily at a few investigated triple junctions, where very high concentrations were found but no trace of S and Cl was measurable elsewhere (Wolff et al., 1988; Mulvaney et al., 1988; Fukazawa et al., 1998). Later studies with cryo-SEM X-ray analysis found S mainly in inclusions but NaCl at grain boundaries (Cullen & Baker, 2000) and soluble impurities at grain boundaries (Barnes et al., 2002). Methodological differences, the low number of investigated triple junctions, the young age of the samples, and especially limits of detection have to be taken into account when attempting to put this work in a general view. At face value, Na and Cl are predominantly found at grain boundaries and triple junctions, while S may also be found in inclusions. Salts containing S (e.g. CaSO₄) are primarily found in inclusions and the grain interior, respectively (Table 2 in Stoll et al., 2021), suggesting that their contribution to impurity diffusion is likely limited unless they become displaced to grain boundaries and possibly dissociate into soluble impurities.

130

145

150

Exploring the localization of impurities thoroughly over larger numbers of grains has so far been limited primarily due to methodological limitations. This particularly concerns detection limits hampering the collection of statistically significant large datasets that allow us to draw generalized conclusions. However, with the introduction of state-of-the-art impurity mapping techniques, laser ablation inductively-plasma mass spectrometry (LA-ICP-MS) has now become a new tool in investigating the localization of impurities in the ice matrix at tens of micron-resolution (Bohleber et al., 2020), extending previous 2D work with this technique (Della-Lunga et al., 2014). For mostly soluble elements such as Na, a high degree of localization at grain boundaries has been observed, while insoluble, dust-related elements can also be found in the grain interiors (Bohleber et al., 2023; Stoll et al., 2023). Initial studies in LA-ICP-MS ice core analysis and in particular impurity mapping have focused on analytically more easily accessible elements such as the metals Na, Mg, Al, Ca, Fe, Sr (Reinhardt et al., 2001; Della Lunga et al., 2014; Sneed et al., 2015; Bohleber et al., 2020; Hoffmann et al., 2024), The range of elements accessible to this technique has been steadily extended, in particular when using a "time-of-flight" mass spectrometer (LA-ICP-TOFMS) that in principle records the entire spectrum of elements (Wilhelms-Dick, 2008; Bohleber et al., 2023). Although the measurement of S and Cl is technically possible with this technique, they both come with their specific analytical challenges: For S, the mass of its primary isotope ³²S is heavily interfered by oxygen signals (¹⁶O¹⁶O+) and Cl suffers from low ionization efficiency in the

hat nach oben verschoben [1]: The localization of impurities within the ice plays a key role in conceptualizing potential diffusion mechanisms. Due to the low eutectic point of sulfuric acid, it was hypothesized that it is mostly located within a network formed by connected triple junctions (the intersection of grain boundaries of three grains) where it can diffuse along the veins subsequently (Mulvaney et al., 1988).

hat gelöscht: For the upper 350 m in the EDC ice core drilled in central Antarctica, Barnes et al. (2003) investigated the effective diffusivity of sodium, chloride and sulfate and found evidence for the diffusion only for chloride and sulfate. Based on their findings,

hat gelöscht: ¶

hat gelöscht:

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

hat gelöscht:

hat gelöscht: in particular

hat gelöscht: high

hat gelöscht:

hat formatiert: Englisch (USA)

hat gelöscht: with

hat formatiert: Englisch (USA)

plasma. At the same time, sulfate, chloride and sodium belong to the relatively more abundant ionic impurity species in ice cores, although their concentrations in Antarctic ice typically remain in the low "parts per billion" (ppb) range when measured as bulk meltwater samples (e.g. Legrand & Mayewski, 1997). Recent evidence suggests that, due to the high degree of spatial localization at grain boundaries, concentrations of Na are substantially increased at the scale of tens of microns, which generally aids the detection and may bring S and Cl within the detectable concentration range of this technique (Bohleber et al., 2024).

(hat formatiert: Englisch (USA)

Here, we report on recent work specifically targeting the mapping of S and Cl signals using LA-ICP-TOFMS within EDC ice core samples from the Holocene and last glacial period. Na serves as a reference element to compare our maps to previous LA-ICP-MS maps. The hypotheses put forward in recent studies served as the main motivation for this work: If the above-mentioned hypothesis by Rhodes et al. (2024) explaining the difference in effective diffusivity between the Holocene and the last glacial holds true, it would mean that impurity localization for S should be distinctly different in ice from these periods. If no such changes can be found, it would remain to be tested if evidence for changes in the vein connectivity with depth appears (Barnes et al., 2003). Similarly, based on mapping the localization of S it should become possible to distinguish between the two scenarios put forward by Ng (2021) – "ice vein only" versus "grain boundary and ice vein contributions". Note that localization is key to determining the plausibility of all these scenarios, and a fully quantitative analysis is not required. We thus focus on delivering information on the localization in the ice matrix rather than obtaining quantitative information on concentrations, which adds considerable complexity and uncertainty (Bohleber et al., 2024).

hat formatiert: Englisch (USA)

hat gelöscht: Note that localization is key to decide on all these scenarios and fully quantitative analysis is not required.

hat gelöscht: over

hat formatiert: Englisch (USA)

2 Methods

180

185

195

200

2.1 LA-ICP-TOFMS ice analysis,

For the purpose of this study, two different systems were used for impurity mapping with LA-ICP-MS. In so doing we make use of an opportunity to discuss maps showing Na, Cl and S signals on ice from two initially unrelated measurement campaigns at the University of Graz and later at the Alfred Wegener Institute Helmholtz Centre for Polar and Marine Research, Bremerhaven (AWI). The newly established cryo-LA-ICP-TOFMS system at AWI consists of a 193 nm excimer laser ablation system capable of repetition rates up to 1 kHz (Iridia by Teledyne Photon Machines, Bozeman, MT, USA) coupled to a ICP-TOFMS (Vitesse ICP-TOFMS by Nu Instruments, Wrexham, UK). The latter includes a segmented reaction chamber, in which He and H2 are used at mL/min flow rates to remove spectral interferences. The laser ablation system is equipped with a cryogenic sample cell (CryoCOBALT Cell) using water-cooled Peltier elements to cool a sample holder. Two temperature sensors and a humidity sensor are used to control the cooling via an integrated software, ensuring that the sample surface temperature stays below -20°C. Temperature stability was better than +/- 1°C over more than 8 hours. An ARIS (Aerosol Rapid Introduction System, Teledyne Photon Machines, USA) is used for fast aerosol transfer to the ICP-TOFMS. Using a NIST612 glass standard, instrumental parameters are optimized for best sensitivity in the mass range from m/z = 23 to m/z=90

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

hat gelöscht: made

hat gelöscht: combine

hat gelöscht: the Alfred Wegener Institute Helmholtz Centre for Polar and Marine Research, Bremerhaven (AWI)

hat gelöscht: "time-of-flight"

hat gelöscht: -

hat formatiert: Englisch (USA)

hat gelöscht: up

(comprising the mass range of primarily interesting analytes for ice, and minimal oxides). NIST612 is used to tune the system for best single-pulse-response (SPR), including tuning the gas flows in the sample chamber as well as the distance between cell and sample surface. For these purposes, NIST glass standards were preferred over artificial ice standards due to easier use and higher homogeneity (Bohleber et al., 2024). Further details on the AWI setup can be found elsewhere (Bohleber et al.,

225 2025).

As a second setup, we combined the cryogenic sample holder employed at Ca'Foscari University of Venice (Bohleber et al., 2020) with the LA-ICP-TOFMS system at the Institute of Chemistry, University of Graz (Niehaus et al., 2024; Lockwood et al., 2024). The latter comprises an Analyte G2 laser ablation system (Teledyne Photon Machines, USA) equipped with a HelEx II ablation chamber compatible with the Venice cryostage. The cryostage was operated in connection to a chiller circulating glycol-water mix cooled to -27°C, achieving sufficient cooling of the sample with the temperature at the surface measuring below -20°C and with no visual evidence of melting. Again, this ablation chamber is connected with an ARIS rapid aerosol introduction system to an ICP-TOFMS (Vitesse ICP-TOF-MS by Nu Instruments, Wrexham, UK). Since using an identical ICP-TOFMS, the two systems are complementary through the differences in the laser ablation instrument. The Graz system was used primarily with spot sizes of 20 and 40 μm over comparatively large areas, while the AWI system was used for obtaining smaller maps at higher resolution analysis at 10 μm spot size. Table 1 summarizes the settings used by both LA-ICP-TOFMS systems.

Table 1: Overview on instrumental settings used for LA-ICP-TOFMS ice core analysis

AWI			
LA (Iridia 1kHz, ARIS)		ICP-TOFMS (Vitesse)	
Wavelength (nm)	193	RF power (W)	1300
Fluence (J cm ⁻²)	4	Auxiliary gas flow (L min-1)	2
Repetition rate (Hz)	500 / 1000	Coolant flow (L min-1)	13
Dosage	10	Nebulizer flow (L min ⁻¹)	1.25
Spot size (µm)	1,5,10	Reaction cell gas (mL min ⁻¹)	7 (He) / 12 (H ₂)
Graz			
LA (Analyte G2, ARIS)		ICP-TOFMS (Vitesse)	
Wavelength (nm)	193	RF power (W)	1450

Repetition rate (Hz)	300	Coolant flow (L min-1)	13
Dosage	10	Nebulizer flow (L min-1)	1.35
Spot size (um)	20. 40	Reaction cell gas (mL min-1)	14 (He) / 6 (H ₂)

Auxiliary gas flow (L min-1)

hat gelöscht:)

hat gelöscht: minimum

hat gelöscht: The entire setup is operated within a class 4 clean room at AWI. On ice samples only the narrowed mass range from m/z=23 up to m/z=90 was acquired.

hat gelöscht:

hat gelöscht: The strategy mainly employed in the present work focused on achieving maximum sensitivity in the lower mass range, by specifically tuning the ICP-TOFMS for high sensitivity in the range of m/z=23 to 90. In doing so, we also carefully tuned parameters of the segmented reaction chamber, including the cell gas flows of He and H_2 in order to maximize the breaking up of $^{16}O^{16}O^{1}$ while limiting the amount of sensitivity decrease to ^{32}S . An ad $(-17)^{11}$

hat nach unten verschoben [6]: Following our previously

hat gelöscht: in

hat formatiert: Englisch (USA)

hat gelöscht: Beam

hat formatiert: Englisch (USA)

Formatiert: Zentriert

hat formatiert: Englisch (USA)

Formatiert: Zentriert

hat formatiert: Englisch (USA)

Formatiert: Zentriert

hat gelöscht: 0

hat formatiert: Englisch (USA)

Formatiert: Zentriert

hat gelöscht: Beam

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

Formatiert: Zentriert

hat formatiert: Englisch (USA)

Fluence (J cm⁻²)

We focused here on measuring maps of the following isotopes: ³⁵Cl, ³⁷Cl and ³²S, ³³S, ³⁴S, with ²³Na used as a reference element with respect to previous studies. The strategy mainly employed for the detection of Cl and S focused on achieving maximum sensitivity in the lower mass range, by specifically tuning the ICP-TOFMS for high sensitivity in the range of m/z=23 to 90. In doing so, we also carefully tuned parameters of the segmented reaction chamber, including the cell gas flows of He and H₂ in order to maximize the breaking up of ¹⁶O¹⁶O⁺ while limiting the amount of sensitivity decrease to ³²S. An additional approach was to reduce the amount of background O coming from the ablation chamber via sublimation of the ice surface, i.e. through the constant background stream of H₂O to the ICP-TOFMS. For this purpose, at the University of Graz, we successfully explored lowering the gas flows inside the ablation chamber, while respecting the limits imposed on this to not negatively affect SPR. With the AWI setup, the ablation chamber has a built-in humidity sensor, which shows that the humidity can be decreased by running additional gas evacuation and refilling cycles. We typically used 20 – 25 such cycles initially and also applied about 30 minutes of constant flushing with dry He. Thereby, the humidity was found to decrease rapidly and then remain at typically around 5-10%, Further details on the analytical challenges of measuring S and Cl are discussed in Sect. 4.1.

2.2 Sample selection

We chose a set of ice core samples both from the Holocene and the last glacial period in the EDC ice core. In order to test the mapping of S and Cl on a broad range of ice conditions and impurity concentrations we also included one sample of the East Greenland ice-core project (EGRIP) core drilled in East Greenland. The glacial EGRIP sample (EGRIP 2286) contains a dustrich cloudy band from the Younger Dryas and thus represents contrasting conditions to the low impurity samples from EDC. It contains insoluble particle concentrations > 250,000 particles/ml (Stoll et al., 2022). This sample selection deliberately excluded ice containing known traces of volcanic eruptions, in order to provide a realistic subsample of typical polar ice conditions. We name subsamples with a hypen to denote their sequence below the top of the bag number, for example subsamples denoted as EDC 1819-2, 1819-3 etc are taken from the respective bag (1819) and are of 8 cm length, i.e. originating from 8 and 16 cm below the top of the bag, respectively. The same applies to EDC 513-2. Table 2 provides an overview, including information on the respective age of the sample. As a general reference for bulk ionic Na⁺ and (non-sea-salt) SOa²⁻ concentrations in EDC we report in Table 2 data for the depths 280.5, 584.1, 999.9 m (Wolff et al., 2006).

Table 2: Overview on ice samples analyzed in this study. The analysis in Graz produced maps larger in size at comparatively coarser spot sizes while the analysis at AWI focused on finer resolution over smaller maps. Bulk ionic values are reported for EDC from Wolff et al. (2006) for adjacent depths (see text).

Core name	Bag n°	Depth	<u>Age</u>	<u>Period</u>	<u>Resolution</u>	Bulk concentration	
A	_	[m]	[ka BP]	_	beam size [µm]	Na / Cl/ SO4 (ppb)	
EDC	513	281.6	9.0	Holocene	10 (AWI), 40 (Graz)	20 / 30/ 85	
EDC	1065	585.2	27.3	Last glacial	20 (Graz)	74 / 106 / 148	
EDC	1819	1000	64	Last glacial	10 (AWI)	90 / 112 / 212	
		•••••	•••••	·····			

hat gelöscht: as well as
hat gelöscht: A
hat gelöscht: for this purpose
hat gelöscht: typically
hat gelöscht: stay
hat formatiert: Englisch (USA)
hat gelöscht: 250
hat formatiert: Englisch (USA)
hat gelöscht: D
hat gelöscht: subsamples
hat formatiert: Hochgestellt
hat formatiert: Tiefgestellt
hat formatiert: Hochgestellt
subsamples of EDC bag 1819, denoted as 1819-2 and 1819-3. These subsamples are of 9.5 cm length and originate from 9.5 and 19 cm below the top of the bag, respectively. Additional maps were recorded on EDC bag 513, subsample 513-1, a 9.5 cm long sample originating from the top of the bag. Following our previously established approach for impurity mapping (Bohleber et al., 2020), the ice surface was decontaminated by scraping with a major-element free ceramic ZrO ₂ blade (American Cutting Edge, USA) immediately before inserting the samples into the ablation chamber. Prior to obtaining maps on the ice, the respective location was also cleaned by pre-ablation. Mosaics of optical images of the ice surface were recorded using the camera integrated with the laser ablation systems. Maps were acquired as adjacent horizontal lines covering a rectangular area (Bohleber et al., 2020). The maps were generated using HDIP (Teledyne Photon Machines). ¶
hat verschoben (Einfügung) [6]
hat formatiert: Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)
hat formatiert: Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)
hat formatiert: Englisch (USA)
hat formatiert[2]
hat formatiert[3]
hat formatiert: Englisch (USA)
hat formatiert [4]
hat formatiert[5]
hat formatiert: Englisch (USA)
hat formatiert[6]
hat formatiert[7]
hat formatiert: Englisch (USA) hat formatiert [2] hat formatiert [3] hat formatiert: Englisch (USA) hat formatiert [4] hat formatiert [5]

[8]

hat formatiert

EGRIP 2286 1256.9 12.1 Younger Dryas 20 (Graz)

Following our previously established approach for impurity mapping (Bohleber et al., 2020), the ice surface was decontaminated by scraping with a major-element free ceramic ZrO₂ blade (American Cutting Edge, USA) immediately before inserting the samples into the ablation chamber. Prior to obtaining maps on the ice, the measured area was also cleaned by preablation. Mosaics of optical images of the ice surface were recorded using the camera integrated with the laser ablation systems. Maps were acquired as adjacent horizontal lines covering a rectangular area (Bohleber et al., 2020). The maps were generated using HDIP (Teledyne Photon Machines).

3 Results

Experimental constraints in measurement time and resources require making a trade-off between the size of the mapped area and the resolution (spot size). Large size maps at comparatively coarse resolution (40 and 20 µm) typically offer a good overview on the degree of impurity localization at grain boundaries versus grain interiors. Finer resolution (10 µm and finer) may be required to investigate impurity variability within grain boundaries and in particular triple junctions. The result section is structured accordingly, going from coarser resolution large size maps to finer resolution detailed maps taking a closer look at grain boundaries and triple junctions.

3.1 Large size maps

365

Figures 1 and 2 show examples for the large size ³²S intensity maps alongside ³⁷Cl and ²³Na recorded with the Graz LA-ICP-TOFMS system. Additional maps are included in the Supplementary Material. As mentioned in the introduction, Na primarily serves as a reference element showing the high degree of localization at grain boundaries, analogous to previous observations in EDC and EGRIP (Bohleber et al., 2020; Bohleber et al., 2023; Stoll et al., 2023). The maps of S reveal a similar picture as for Na, with high localization at grain boundaries and low intensities within the grain interiors. In EGRIP 2286, a sample containing a dust-rich cloudy band from the Younger Dryas, some isolated pixels within the grain interior also show high S values (Fig. 2). The EGRIP 2286 cloudy band sample likely has significantly higher impurity content compared to the EDC 513 Holocene sample, reflected in the higher signal amplitude in spite of less material ablated with the smaller spot size used (20 μm EGRIP 2286 vs 40 μm EDC 513). Both maps show intensity variability along the grain boundaries, indicating that some spatial variability in grain boundary concentrations exists on the scale of a few mm. Note for example lower intensities highlighted in the upper right area of the EGRIP 2286 map (Fig. 2) and higher relative intensities in the lower left of the EDC 513 map (Fig. 1). At 40 μm and 20 μm spot size, respectively, no clear evidence for enhanced concentrations at triple junctions relative to grain boundaries is detected, e.g. via pixels with brighter intensity at triple junctions. For a more quantitative evaluation of this finding, pixels belonging to grain boundaries (including triple junctions) were segmented from the Na and S maps using a watershed algorithm in HDIP analogously to previous work (Bohleber et al., 2023). Pixels representing triple

hat formatiert: Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)

hat formatiert: Englisch (USA)

hat formatiert: Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)

hat formatiert: Englisch (USA)

hat gelöscht: Using the AWI setup, maps were recorded on two subsamples of EDC bag 1819, denoted as 1819-2 and 1819-3. These subsamples are of 9.5 cm length and originate from 9.5 and 19 cm below the top of the bag, respectively. Additional maps were recorded on EDC bag 513, subsample 513-1, a 9.5 cm long sample originating from the top of the bag.

hat formatiert: Englisch (USA)

hat gelöscht: respective

hat gelöscht:

hat formatiert: Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)

hat formatiert: Schriftart: (Standard) +Textkörper (Times New Roman)

Formatiert: Standard

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

Figure 1: Map of EDC bag 513 (40 µm spot, 6.4 x 10.4 mm). Shown are intensities in counts (cts) for ²³Na (top left), ³²S (bottom left) and ³⁷Cl (bottom right) together with the optical mosaic (top right).

hat formatiert: Englisch (USA)

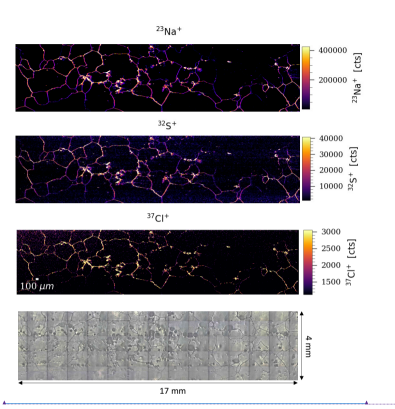


Figure 2: Map of EGRIP bag 2286 (20 μ m spot, 4 x 17 mm). Shown are intensities in counts (cts) for ²³Na (top row), ³²S and ³⁷Cl (middle rows) and the optical mosaic (bottom)

3.2 High-resolution maps obtained with 10 μm spot size

395 Figures 3 – 5 show the results for EDC 1819-2, 1819-3, and 513-1 respectively. An additional map was recorded on EDC1819-3 in one session right after the map on EDC513-1 (shown in the Supplementary Material). The maps show that also at 10 μm spot size, Na, S and Cl are highly localized at grain boundaries. Notably these maps also show signal distributed among different isotopic channels. The map for ³⁵Cl contains only noise in all maps recorded (hence not shown), but strong signal in ³⁷Cl, which indicates the potential formation of Chlor- hydrogen adducts (see Sect. 4.1). As shown in Fig. 3 – 5 for S, ³²S

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)

hat nach oben verschoben [7]: Using the AWI setup, maps were recorded on two subsamples of EDC bag 1819, denoted as 1819-2 and 1819-3. These subsamples are of 9.5 cm length and originate from 9.5 and 19 cm below the top of the bag, respectively. Additional maps were recorded on EDC bag 513, subsample 513-1, a 9.5 cm long sample originating from the top of the bag.

contains most of the signal. Despite the low isotopic abundance, some maps even reveal signal in the isotopes ³³S (on rare occasion) and ³⁴S. Also, at 10 µm spot size, there is no clear evidence for enhanced intensities at or around triple junctions.

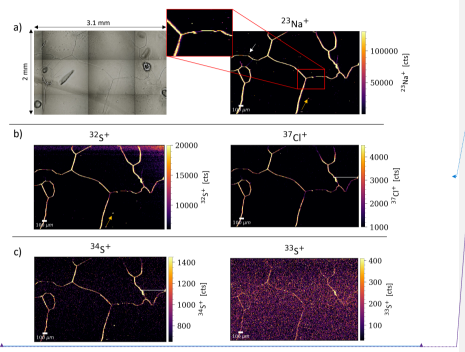


Figure 3: Example map for EDC1819-2, 10 μm spot size. Row a) shows the optical mosaic with dark lines denoting grain boundaries and the map for ²³Na. Note the potential discontinuity in the grain boundary intensity marked by a red square; the same area is shown as a zoom-in section. White and yellow arrows indicate examples of an intra-grain intensity spot for Na only and Na with S, respectively. Row b) contains the maps for the two main channels with signals for S and Cl, ³²S and ³⁷Cl respectively. Under best case mapping performance, other isotopes of sulfur may also show signals, illustrated in row c). The white horizontal line on the right-hand side in the ³⁷Cl and ³⁴S maps is an absence of signal as a result of an autoblanking procedure protecting the detector in the ICP-TOFMS.

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)
Formatiert: Zentriert

hat gelöscht: white circle

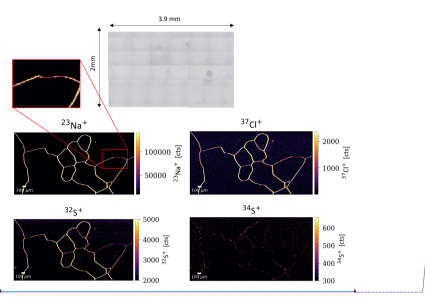


Figure 4: Example map for EDC1819-3, 10µm spot size. The channels for the isotopes ³³S and ³⁵Cl contain noise only and are therefore not plotted. Shown on the top is the optical mosaic. Note the potential discontinuity in the grain boundary intensity marked by a red square in the ²³Na map and the respective zoom-in on the top left.

420

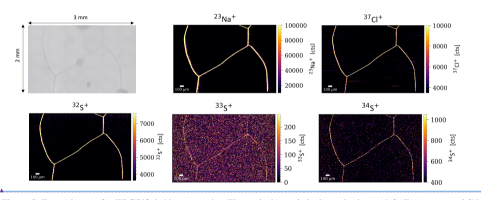


Figure 5: Example map for EDC513-1, $10\mu m$ spot size. The optical mosaic is shown in the top left. Features are visible in each plotted S channel, while Cl remained confined to m/z=37 (see text).

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)

hat gelöscht: white circle

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

Formatiert: Zentriert

3.3 Ultra-high resolution maps obtained with 1 µm spot size

Based on the limited evidence for enhanced impurity localization at triple junctions in maps collected with spot sizes 40, 20 and 10 μ m, an exemplary investigation was performed with even smaller spot sizes. For this purpose, a triple junction was mapped with subsequently finer resolution, with 10 μ m, 5 μ m and 1 μ m spots (Fig. 6). The results for Na show that at 10 and 5 μ m, the triple junction is confined to a single pixel, whereas the maps recorded with 1 μ m show more detail and a region of brighter intensity at the triple junction. Due to the much smaller volume ablated, it was not possible to obtain signals for S, although some signal was detectable for 37 Cl.

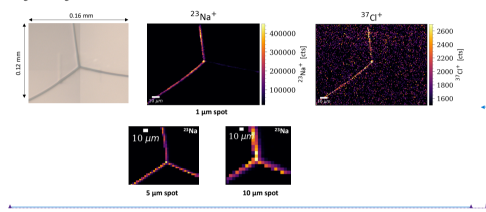


Figure 6: Exemplary investigation into the influence of spot size on detecting enhanced localization at triple junctions versus grain boundaries. The top row shows ultra-high resolution maps of a triple junction recorded at 1_µm spot size, alongside an optical mosaic of the same area. Note that the lower intensity on the right-hand side grain boundary is likely impacted by the scanning-direction, an effect only noticed at this high resolution (see text). The bottom row shows Na maps recorded over the same triple junction with 5 and 10 µm spot size.

4 Discussion

430

4.1 The analytical challenge of detecting sulfur and chlorine in ice with LA-ICP-TOFMS

As demonstrated by the results shown in Fig. 1 – 6, high-resolution mapping of both S and Cl is possible in ice with LA-ICP-TOFMS but come with their own specific challenges. The main hurdle for detecting S is the mass interference by $^{16}O^{16}O^{+}$ on m/z=32, which results in high background levels. As opposed to the analysis of other materials using dry plasma conditions, the $^{16}O^{16}O^{+}$ background cannot be avoided for ice since it is caused to a large degree by ablating the ice matrix itself, resulting

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)

Formatiert: Zentriert

hat gelöscht: Although Na, S and Cl belong to the most abundant elements in polar ice (albeit generally at low ppb bulk concentrations) (Legrand & Mayewski, 1997), initial studies in LA-ICP-MS ice core analysis and in particular impurity mapping have focused on analytically more easily accessible elements such as Na, Mg, Al, Ca, Fe, Sr (Reinhardt et al., 2001; Della Lunga et al., 2014; Sneed et al., 2015; Bohleber et al., 2020; Hoffmann et al., 2024).

hat gelöscht: 7

in a constant stream of H₂O to the ICP-MS. Mitigation strategies involve i) high mass resolution in order to resolve the closely-spaced two peaks the mass spectrum caused by ¹⁶O¹⁶O⁺ and ³²S, ii) breaking up ¹⁶O¹⁶O⁺ by collision or reaction in the segmented reaction cell, iii) mass shifting of ³²S through chemical reaction, e.g. with Oxygen. Strategy iii) is S-specific and typically used in applications with a triple-quadrupole instruments (Martinez-Sierra et al., 2015), whereas i) and ii) can be applied in LA-ICP-TOFMS also without an additional quadrupole mass filter. In a previous pilot study, we were also able to resolve signals of ³²S using a Teledyne Photon Machines G2 laser ablation system (as used in Graz) coupled to an icpTOF 2R (TOFWERK, Thun, Switzerland) (Bohleber et al., 2021). Regarding the detection of Cl, an important side-effect of using the segmented reaction chamber is the formation of Chlor-hydrogen adducts, which explains the lack of signal in the mass channel m/z=35 in our maps (Figure S5 in the Supplementary Material). This channel should contain the majority of the Cl signal based on isotope abundance (about 75% for ³⁵Cl and 25% for ³⁷Cl). However, by monitoring ³⁵Cl and ³⁷Cl whilst analyzing Cl standards, we found conclusive evidence that the entirety of ³⁵Cl signal is mass-shifted to ³⁷Cl by formation of H₂Cl⁺. It is likely that also ³⁷Cl is mass-shifted and detected at ³⁹K instead.

Ultimately, the strategy outlined above, requiring specific tuning of the ICP-TOFMS and controlling the background levels of Oxygen add considerable complexity to the already elaborate nature of high-resolution impurity mapping in ice cores. Future efforts will require making a tradeoff between achieving an adequate degree of sample throughput versus such time-consuming fine-tuned analysis. The latter could be potentially further extended in quantitative analyses through dedicated artificial ice standards. Since not required for this study (see introduction) and due to the already complex measurement strategy, we have not added the quantitative dimension considering the substantial complexity and added uncertainty involved in the use of artificial ice standards (Bohleber et al., 2024). Further extensions to this work for the detection of S could include exploring the use of alternative collision-reaction cell gases, such as Xe or CH₄ (Guillong et al., 2008; Singh et al., 2025).

465

480

The maps showing a signal not just in ³²S but also in the much less abundant (but less interfered) isotopes ³³S and ³⁴S also motivate exploring in more detail the limits to isotopic analysis with LA-ICP-TOFMS. This could involve specific sections with known volcanic eruption, for which the added experimental effort would be warranted. If taken at face value, the relative abundance of the sulfur isotopes agrees within their uncertainty (typically less than 3%) with the natural isotopic abundance, 95, 0.75 and 4% for ³²S, ³³S and ³⁴S, respectively (Meija et al, 2016). We calculated for this purpose the mean intensity at grain boundaries (the location of the signal in the maps) in Fig. 1 and 5 for all three isotopic channels (Table 3).

Table 3: Average of intensities (in cts) at grain boundaries calculated for sulfur isotope channels. The maximal uncertainty for the relative abundance values is 3%.

Sample	Isotope	Average	Uncertainty (1s)	Relative Abundance (%)	
EDC1819-2	32S	21300000	330000	94	
	33S	208000	3700	1	

hat nach oben verschoben [9]: The strategy mainly employed in the present work focused on achieving maximum sensitivity in the lower mass range, by specifically tuning the ICP-TOFMS for high sensitivity in the range of m/z=23 to 90. In doing so, we also carefully tuned parameters of the segmented reaction chamber, including the cell gas flows of He and H2 in order to maximize the breaking up of 16O16O+ while limiting the amount of sensitivity decrease to 32S. An additional approach is to reduce the amount of background O coming from the ablation chamber via sublimation of the ice surface, i.e. through the constant background stream of H2O to the ICP-TOFMS. At the University of Graz, we successfully explored for this purpose lowering the gas flows inside the ablation chamber. while respecting the limits imposed on this to not negatively affect SPR. With the AWI setup, the ablation chamber has a built-in humidity sensor, which shows that the humidity can be decreased by running additional gas evacuation and refilling cycles. We used typically 20 - 25 such cycles initially and also applied about 30 minutes of constant flushing with dry He. Thereby, the humidity was found to decrease rapidly and then stay at typically around 5-10%.

hat gelöscht: ¶

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

A	34S	1240000	15000	5	
EDC513-1	32S	12100000	140000	93	
	33S	134000	2600	1	
A	34S	717000	8300	6	

hat formatiert: Englisch (USA) hat formatiert: Englisch (USA)

4.2 Implications for diffusion studies

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)

Due to the sophisticated nature of LA-ICP-MS impurity mapping in general and mapping S and Cl in particular, the amount of data remains limited. The presented maps have to be regarded as snapshots covering comparatively small areas. Although some spatial variability in the levels of S and Cl at grain boundaries is detected by the larger maps (Fig. 1 and 2), the localization of S and Cl at grain boundaries and triple junctions is found consistently in all maps. Thus, this has to be considered a robust result.

While methodological differences have to be kept in mind, our data are consistent with previous studies investigating S e.g.

hat gelöscht: t

510

hat gelöscht: is

by cryo-SEM and X-ray analysis or Raman spectroscopy, finding S signals in a limited number of samples and measurements primarily at triple junctions, potentially indicating liquid-filled veins (e.g. Mulvaney et al., 1988; Fukazawa et al., 1998). Regarding a crucial point in conceptualizing impurity diffusion along ice veins, the question of a potential enrichment of impurities at triple junctions is much harder to answer based on our data. The exploration of a triple junction at ultra-high resolution (1 μ m) points towards spatial resolution being a key issue in this context: only at this fine scale can the intensity difference be clearly recognized in our example (Fig. 6). At 1 μ m, the investigated triple junction does show a local increase in intensity by about a factor of 1.3 relative to the grain boundary at the lower left. This grain boundary was chosen because the intensity of the other two grain boundaries (at the top and to the lower right of the map) was found to depend on the scan direction used to acquire the map (horizontal versus vertical scans, Fig. S3). This effect was only observed to occur at 1 μ m spot size and illustrates another dimension of complexity in impurity mapping at this high resolution. Adding to this are the higher limits of detection due to the much smaller amount of ablated material ablated at 1 μ m (hence no S signal in the maps) and the higher amount of resources consumed (almost 200k shots were needed for the small map in Fig. 6).

hat gelöscht: Supplementary Material

hat gelöscht:

hat gelöscht: Consequently

hat gelöscht: this exemplarily

hat gelöscht: 1

hat gelöscht: 1

hat gelöscht: there is some evidence for

hat gelöscht: visible

hat gelöscht: if changinges

hat gelöscht: at higher

hat gelöscht:

hat gelöscht:

hat gelöscht: While a

hat gelöscht: are

hat gelöscht: mapped

hat gelöscht: ing

hat gelöscht: with

hat gelöscht: high intensity

hat gelöscht: , with the same dosage 10 for all maps.

The primary bottleneck in terms of resources is measurement time, which also translates into consumption of energy and cell gases, however. As a consequence, we only investigated mapping at 1 µm resolution exemplarily in this work. Yet, dedicated mapping of triple junctions at 1 µm resolution appears feasible for future studies mastering some methodological challenges such as the scan direction influence and resource consumption. Computer-vision assisted mapping may provide a better compromise between resolution and resource consumption in this regard (Larkman et al., 2025a). It is also worthwhile mentioning that the apparent spatial extent of the grain boundaries and triple junctions in the maps can change depending on the applied resolution. At 10 µm spot size, grain boundaries typically extend over 2-3 pixels in width, for 5 µm this reduces to 1-2 pixels. An identical dosage of 10 was used for all maps. For 1 µm, the spatial extent is around 5 pixels, roughly consistent

with the 5 µm maps. Notably, this only refers to the apparent size of surface features in the chemical maps, which has been influenced by sublimation. The 1 µm spot size may be more capable of resolving the fine-scale impurity variability around triple junctions, and shows that some enhancement of Na at triple junctions relative to grain boundaries may exist at that spatial scale. Mulvaney et al., 1988 applied a comparable spatial resolution of around 1x1 µm in their SEM analysis supporting our results. Also in this SEM analysis, such a local enhancement at triple junctions could have pushed signals to the detectable range, while signals at grain boundaries remained undetectable.

555

560

570

580

At this point, our results do not support the notion of vein-dissolved ionic impurities dominating the respective bulk ice core records (e.g. referred to as "scenario 1" by Ng (2021)). Similar to Na, also S and Cl appear to be highly localized at grain boundaries, with a minor enhancement possible in veins at triple junctions. The fact that grain boundaries are populated to a similar degree as triple junctions suggests that diffusion of soluble impurities in ice needs to be treated considering grain boundary surfaces in three dimensions. Here, changes in grain size, e.g. via recrystallization and grain growth, becomes a crucial factor (Ng, 2021). The localization of S and Cl at grain boundaries has significant implications for the interpretation of 1D LA-ICP-MS profiles designed to collect climate-related signals. The impurity-boundary imprint manifests strongly in such measurements and may also become relevant for bulk meltwater measurements, such as with continuous flow analysis (CFA), particularly in ice samples with large average grain sizes (Larkman et al., 2025b).

It has to be noted in this context, however, that dust-rich ice such as the EGRIP 2286 cloudy band sample has already shown. higher concentrations of Na also in grain interiors (Fig. 2 and Bohleber et al., 2023), similar to other EGRIP cloudy band. samples (Stoll et al., 2023). Some cases with Na less abundant in grain interiors are also present in the glacial EDC samples of bag 1819 (Fig. 3 and 4). It is thus likely that S in dust-rich samples is also more prominent in the grain interior. On these grounds, we tentatively note that S and Na appear also in isolated clusters in the grain interior (although some co-localization of S with Na also occurs, e.g. Fig. 3), while Cl is often co-localized with Na (likely indicating NaCl). The fact that some clusters in grain interiors are found in the maps is consistent with evidence for sulfate-bearing micro-inclusions such as salts found in grain interiors (e.g. Ohno et al. 2005, Eichler et al., 2019, Stoll et al., 2022). Notably, inter-method comparisons are always a challenge and LA-ICP-MS cannot directly distinguish dissolved and particulate fractions, other than identifying particles via spatial information on their localization (isolated clustered pixels of high intensity in grain interiors) (Bohleber et al., 2023). In a simplified view, however, depending on the degree to which salts can dissociate and be displaced to grain boundaries due to e.g., grain growth, they could contribute to the overall apparent impurity mobility via diffusion. Similarly, dissolution of minerals resulting from chemical reactions at grain boundaries is a possibility (Baccolo et al., 2021). Accordingly, the physical reality likely is a mix between the two scenarios described in Ng (2021). For our "snapshots" we find a high degree of localization at grain boundaries for Na, S and Cl, with only minor contributions from the grain interiors However, in general the relative contributions from ice veins, grain boundaries and grain interiors can depend on conditions such as climate period and insoluble particle content, and uncharacterized microstructural processes.

hat gelöscht: likely

hat gelöscht: In previous studies

hat gelöscht: was used

hat gelöscht: (Mulvaney et al., 1988)

hat gelöscht: Here

hat gelöscht: at triple junction

hat gelöscht: (Mulvaney et al., 1988)

hat formatiert: Englisch (USA)

hat gelöscht: 2024

hat gelöscht: ed

hat gelöscht: s

hat gelöscht: cases

hat gelöscht: also

hat nach unten verschoben [8]: LA-ICP-MS cannot directly distinguish dissolved and particulate fractions, other than identifying particles via spatial information on their localization (isolated clustered pixels of high intensity in grain interiors) (Bohleber et al., 2023). On

hat gelöscht: this

hat gelöscht: always

hat verschoben (Einfügung) [8]

(**hat gelöscht:** by

hat gelöscht: via

hat gelöscht:),

hat gelöscht: with a

hat gelöscht: highly localized nature of impurities in ice

hat gelöscht: but variable

hat gelöscht: depending

hat gelöscht:

Notably, the situation in deep ice sections, with larger grains and more time spent at higher temperatures, may deviate from the snapshots from relatively shallow depths presented here and remains to be explored, in particular regarding the influence of geochemical reactions leading to dissolution at grain boundaries (Baccolo et al., 2021). Sections with high S-peaks from volcanic eruptions, which have not been measured here, may also prove to be a special case. At least for sections within the Holocene and last glacial period in the EDC core, it is difficult to conceive systematic differences between the data presented here, however, other than the bulk S concentration levels.

625

630

In considering a physical process to explain their empirical constraints on the effective diffusivity of sulfate, based on data from EDC Holocene ice, Barnes et al. (2003) invoked either the diffusion through a connected vein network or a system of discontinuous veins with interconnectivity changes due to grain growth. In recent studies, both Fudge et al. (2024) and Rhodes et al. (2024) find an order of magnitude lower values for the effective diffusivity beyond the Holocene. Possible explanations of the lower effective diffusivity involve i) sulfuric acid not readily diffusing in liquidlike veins (Fudge et al., 2024), ii) changes to an initially inter-connected network of veins and iii) changes in the mechanism of diffusion resulting due to sulfate ions located within the ice microstructure (Rhodes et al., 2024). Diffusion both through veins and grain boundaries appears to be supported by the maps obtained in the present work, while a change to sulfate predominantly located in the grain interiors is not. This means the new maps do not indicate changes in the general character of impurity localization between Holocene and glacial samples, which would support the notion of a change in diffusion mechanism according to (iii). In contrast, the new data are consistent with a recently formulated view that the majority of the diffusion is triggered at shallow depths and that only minor differences occur with depth (Ng et al., 2025). It is important to note that, based on the inevitable snapshot character of the data, it is not possible to make generalized statements, in particular regarding the level of 3D interconnectivity of veins and grain boundaries. As indicated in Fig. 3 and 4, we find some occasional evidence of partially interrupted impurity population at grain boundaries in the glacial maps but not in the Holocene. Notably, the discontinuities among grain boundaries in Fig. 1 are due to air bubbles, which is not the case for Fig. 3 and 4. However, the discontinuities in the impurity population at grain boundaries are rare, and do not concern the triple junctions. This does not support the potential explanations i) nor ii) above, and points towards a potential additional mechanism at work. An investigation taking into account 3D geometric effects and grain size changes may provide further insight. A possible approach to investigate the 3D interconnectivity would be a LA-ICP-MS high resolution study ablating layer by layer subsequently resulting in quasi-3D mapping. The large amount of time and resources needed hamper the practicality of this approach so far, Additional exemplary investigations could be in reach soon, however, similar to the 1 µm mapping performed here, Further semi-empirical investigations could employ a combination of LA-ICP-MS maps with 3D modeling of the grain-scale chemistry (Larkman et al., 2025b; Ng 2024; Ng 2021).

Interestingly, Barnes et al. (2003) found evidence for signal damping only for Cl and S, but not for Na, suggesting that a diffusive process transported sulfate and chloride ions but sodium ions remained fixed. More precisely constraining the

hat gelöscht: Technically, the same is true for

hat gelöscht: s

hat gelöscht: , which have not been measured here

hat gelöscht: by

hat gelöscht: due

hat gelöscht: changes in impurity localization

hat gelöscht: is

hat gelöscht: high

hat gelöscht: practicability

hat gelöscht: ,

hat gelöscht: but similar to the 1 μm mapping exemplary investigations could be in reach soon.

hat gelöscht: 2024

hat gelöscht: ind

hat gelöscht: Constraining better

mobility of Cl would also be important for cosmogenic radionuclide dating with ³⁶Cl (e.g. Muscheler et al., 2005). The maps collected here provide no support for the notion that S and Cl are more mobile than Na₄For the Holocene and glacial maps, which were obtained sequentially in one session (Fig. 5 and Fig. S2), and hence not significantly affected by instrumental drift, we compared the average intensity ratios of S/Na and Cl/Na at grain boundaries but found similar values for the Holocene and glacial samples (although notably, the ratio is strongly influenced if spurious low Na intensity pixels from the grain interiors are included). Tentatively, this result argues against a relative difference, e.g. glacial ice being more depleted in potentially more "diffusive" elements such as S and Cl versus Holocene samples. On the other hand, the co-localization of Na and Cl at grain boundaries could entail processes slowing of diffusion, e.g. by anion-cation trapping decreasing the mobility by electrostatic attraction and eventually leading to the formation of NaCl. This effect was observed in laboratory experiments decreasing the mobility of HCl in ice and was suspected also for other ionic species (Livingston & George, 2001). Since LA-ICP-MS cannot directly distinguish particulates (e.g. salt formation) and dissolved impurity fractions, a combined approach e.g. with cryo-Raman spectroscopy (albeit not for NaCl in ice) may help to shed light on potential chemical reactions taking place within grain boundaries. This would provide important new insight into geochemical reactions in ice and ultimately aid an improved signal interpretation, especially in deep ice conditions such as in the "Oldest Ice".

hat gelöscht:

hat gelöscht:

hat formatiert: Englisch (USA)

Conclusions

680

695

The investigation of impurity localization and ice-impurity interactions has gained new momentum with the introduction of state-of-the-art impurity mapping using LA-ICP-MS. So far, the elemental range in focus with this method has not been extended to the analytically more challenging elements S and Cl. Both are relatively abundant in polar ice and are of significance for paleoclimate reconstructions and dating efforts, but are also known to be affected by post-depositional processes, such as diffusion and chemical reactions. As part of the ongoing progress in advancing LA-ICP-MS, we show here that signals of S and Cl can be detected in Greenland and Antarctic ice using LA-ICP-TOFMS mapping, albeit requiring careful fine-tuning of the instruments. In doing so, it becomes possible to obtain multi-elemental maps at high resolution, up to 10 µm, and, exemplarily even at 1 µm. We observe a high level of localization of S and Cl at grain boundaries, which has already been observed for other elements such as Na. Additionally, there is a dispersed occurrence within grain interiors in dust-rich ice. Evidence for enhancement at triple junctions, and hence in ice veins, is limited to the ultra-high resolution map at 1 µm. The new maps support a view on diffusive transport not only through ice veins but also grain boundaries, but do not show any clear differences in this regard between samples from the Holocene and last glacial period in the EDC ice core. To determine the physical processes behind impurity diffusion, our results emphasize the merit of integrating such direct measurements with studies on empirical evidence and modelling. Such a combined approach promises a new level of insight into impurity signal preservation in polar ice cores.

hat gelöscht:

hat gelöscht: still

hat gelöscht: evolution

hat gelöscht: of

hat gelöscht: this new technique

hat gelöscht: so

hat gelöscht: find

hat gelöscht: , but also

hat gelöscht: some

hat gelöscht:

Data availability

The underlying data will be made available via a public data repository (e.g. Pangaea) after completion of the peer-review process.

Author contributions

Experimental design was developed by PB and DC₂ Measurements were conducted by PB, NS, PL and DC. RR contributed expertise on impurity diffusion. All authors contributed to the discussion of the results and the final version of the manuscript₂

Competing interests

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

Acknowledgements

help in the laboratory. Pascal Bohleber gratefully acknowledges funding from the European Union's Horizon 2020 research and innovation program under the Marie Skłodowska-Curie grant agreement no. 101018266. Nicolas Stoll and Pascal Bohleber gratefully acknowledge funding from the Programma di Ricerche in Artico (PRA). Nicolas Stoll further acknowledges funding from the European Union's Horizon 2020 research and innovation program under the Marie Skłodowska-Curie grant agreement no. 101146092. Co-funded by the European Union (ERC, AiCE, 101088125 and NanoArchive, 101165171). Views and opinions expressed are however those of the authors only and do not necessarily reflect those of the European Union or the European Research Council. Neither the European Union nor the granting authority can be held responsible for them. We

We are grateful to Remi Dallmayr, Raquel Gonzalez de Vega, Lukas Schlatt & Ciprian Stremtan for their valuable input and

References

725

Baccolo, G., Delmonte, B., Di Stefano, E., Cibin, G., Crotti, I., Frezzotti, M., Hampai, D., Iizuka, Y., Marcelli, A., and Maggi, V.: Deep ice as a geochemical reactor: Insights from iron speciation and mineralogy of dust in the Talos Dome ice core (East Antarctica), Cryosphere, 15, 4807–4822, 2021.

730 Barnes, P. R., Mulvaney, R., Robinson, K., and Wolff, E. W.: Observations of polar ice from the Holocene and the glacial period using the scanning electron microscope, Ann. Glaciol., 35, 559–566, 2002.

also thank Hanna Brooks, an anonymous referee and editor Kaitlin Keegan for their help in improving the manuscript.

Barnes, P. R. F., Wolff, E. W., Mader, H. M., Udisti, R., Castellano, E., and Röthlisberger, R.: Evolution of chemical peak shapes in the Dome C, Antarctica, ice core, J. Geophys. Res.-Atmos., 108, D3, 2003.

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)
hat formatiert: Englisch (USA)

hat formatiert: Englisch (USA)

hat formatiert: Schriftart: (Standard) Times New Roman, 10 Pt.

hat gelöscht:

Baccolo, Giovanni, Delmonte, B., Di Stefano, E., Cibin, G., Crotti, I., Frezzotti, M., Hampai, D., Iizuka, Y., Marcelli, A., & Maggi, V. (2021). Deep ice as a geochemical reactor: Insights from iron speciation and mineralogy of dust in the Talos Dome ice core (East Antarctica). Cryosphere, 15(10), 4807–4822. https://doi.org/10.5194/tc-15-4807-2021

Barnes, P. R., Mulvaney, R., Robinson, K., & Wolff, E. W. (2002). Observations of polar ice from the Holocene and the glacial period using the scanning electron microscope. Annals of Glaciology, 35, 559-566.

Barnes, P. R. F., Wolff, E. W., Mader, H. M., Udisti, R., Castellano, E., & Röthlisberger, R. (2003). Evolution of chemical peak shapes in the Dome C, Antarctica, ice core. Journal of Geophysical Research: Atmospheres, 108(D3).

Bohleber, P., Larkman, P., Stoll, N., Clases, D., Gonzalez de Vega, R., Šala, M., ... & Barbante, C. (2024). Quantitative insights on impurities in ice cores at the micro-scale from calibrated LA-ICP-MS imaging. Geochemistry, Geophysics, Geosystems, 25(4), e2003GC011425

Bohleber, P., Stoll, N., Rittner, M., Roman, M., Weikusat, I., & Barbante, C. (2023). Geochemical Characterization of Insoluble Particle Clusters in Ice Cores Using Two-dimensional Impurity Imaging. Geochemistry, Geophysics, Geosystems, 24(2), e2022GC010595

Bohleber, P., Roman, M., Stoll, N., Bussweiler, Y., & Rittner, M.

(2021). Imaging the distribution of elements in antarctic ice cores with LA-ICP-TOFMS. TOFWERK Application Note. Bohleber, P., Roman, M., Šala, M., & Barbante, C. (2020). Imaging the impurity distribution in glacier ice cores with LA-ICP-MS. Journal of Analytical Atomic Spectrometry, 35(10), 2204–2212. Brook, E. J., Wolff, E., Dahl-Jensen, D., Fischer, H., Steig, E. J., & others. (2006). The future of ice coring: international partnerships in ice core sciences (IPICS). PAGES News, 14(1), 6–10. Cullen, D., & Baker, I. (2000). The chemistry of grain boundaries in Greenland ice. Journal of Glaciology, 46(155), 703-706. Della Lunga, D., Müller, W., Rasmussen, S. O., & Svensson, A. (2014). Location of cation impurities in NGRIP deep ice revealed by cryo-cell UV-laser-ablation ICPMS. Journal of Glaciology, 60(223), 970–988. https://doi.org/10.3189/2014JoG131199

Eichler, J., Weikusat, C., Wegner, A., Twarloh, B., Behrens, M., Fischer, H., ... & Weikusat, I. (2019). Impurity analysis and microstructure along the climatic transition from MIS 6 into 5e in the EDML ice core using cryo-Raman microscopy. Frontiers in Earth Science, 7, 20.¶

Fischer, H., Blunier, T., & Mulvaney, R. (2021). Ice Cores: Archive of the Climate System. In Glaciers and Ice Sheets in the Clima ... [9]

hat formatiert: Schriftart: Nicht Fett, Englisch (USA)

hat formatiert: Englisch (USA)

Formatiert: Überschrift 1

- Bohleber, P., Mervič, K., Dallmayr, R., Stremtan, C., and Šala, M.: Argon versus helium as carrier gas for LA-ICP-MS impurity mapping on ice cores, Talanta Open, 11, 100437, 2025.
- Bohleber, P., Larkman, P., Stoll, N., Clases, D., Gonzalez de Vega, R., Šala, M., and Barbante, C.: Quantitative insights on impurities in ice cores at the micro-scale from calibrated LA-ICP-MS imaging, Geochem. Geophys. Geosyst., 25, e2023GC011425, 2024.
 - Bohleber, P., Stoll, N., Rittner, M., Roman, M., Weikusat, I., and Barbante, C.: Geochemical characterization of insoluble particle clusters in ice cores using two-dimensional impurity imaging, Geochem. Geophys. Geosyst., 24, e2022GC010595, 2023.
- 860 Bohleber, P., Roman, M., Stoll, N., Bussweiler, Y., and Rittner, M.: Imaging the distribution of elements in Antarctic ice cores with LA-ICP-TOFMS, TOFWERK Application Note, 2021.
 - Bohleber, P., Roman, M., Šala, M., and Barbante, C.: Imaging the impurity distribution in glacier ice cores with LA-ICP-MS, J. Anal. At. Spectrom., 35, 2204–2212, 2020.
 - Brook, E. J., Wolff, E., Dahl-Jensen, D., Fischer, H., Steig, E. J., and others: The future of ice coring: international partnerships in ice core sciences (IPICS), PAGES News, 14, 6–10, 2006.
 - Cullen, D., and Baker, I.: The chemistry of grain boundaries in Greenland ice, J. Glaciol., 46, 703-706, 2000.
 - Della Lunga, D., Müller, W., Rasmussen, S. O., and Svensson, A.: Location of cation impurities in NGRIP deep ice revealed by cryo-cell UV-laser-ablation ICPMS, J. Glaciol., 60, 970–988, 2014.
- Eichler, J., Weikusat, C., Wegner, A., Twarloh, B., Behrens, M., Fischer, H., and Weikusat, I.: Impurity analysis and microstructure along the climatic transition from MIS 6 into 5e in the EDML ice core using cryo-Raman microscopy, Front.
 - Fischer, H., Blunier, T., and Mulvaney, R.: Ice cores: archive of the climate system, in: Glaciers and Ice Sheets in the Climate System, Springer, 279–325, 2021.
 - Fudge, T. J., Sauvage, R., Vu, L., Hills, B. H., Severi, M., and Waddington, E. D.: Effective diffusivity of sulfuric acid in
- 875 Antarctic ice cores, Clim. Past, 20, 297–312, 2024.

Earth Sci., 7, 20, 2019.

- Fukazawa, H., Sugiyama, K., Mae, S., Narita, H., and Hondoh, T.: Acid ions at triple junction of Antarctic ice observed by Raman scattering, Geophys. Res. Lett., 25, 2845–2848, 1998.
- Fujita, S., Parrenin, F., Severi, M., Motoyama, H., and Wolff, E. W.: Volcanic synchronization of Dome Fuji and Dome C Antarctic deep ice cores over the past 216 kyr, Clim. Past, 11, 1395–1416, 2015.
- 880 Guillong, M., Latkoczy, C., Seo, J. H., Günther, D., and Heinrich, C. A.: Determination of sulfur in fluid inclusions by laser ablation ICP-MS, J. Anal. At. Spectrom., 23, 1581–1589, 2008.
 - Hoffmann, H., Day, J., Rhodes, R. H., Grieman, M., Humby, J., Rowell, I., and Wolff, E.: Laser ablation inductively coupled plasma mass spectrometry measurements for high-resolution chemical ice core analyses with a first application to an ice core from Skytrain Ice Rise (Antarctica), Cryosphere, 18, 4993–5013, 2024.

- 885 Larkman, P., Vascon, S., Šala, M., Stoll, N., Barbante, C., and Bohleber, P.: Faster chemical mapping assisted by computer vision: insights from glass and ice core samples, Analyst, in press, 2025a.
 - Larkman, P., Rhodes, R. H., Stoll, N., Barbante, C., and Bohleber, P.: What does the impurity variability at the microscale represent in ice cores? Insights from a conceptual approach, Cryosphere, 19, 1373–1390, 2025b.
 - Legrand, M., and Mayewski, P.: Glaciochemistry of polar ice cores: a review, Rev. Geophys., 35, 219-243, 1997.
- 890 Livingston, F. E., and George, S. M.: Effect of sodium on HCl hydrate diffusion in ice: evidence for anion–cation trapping, J. Phys. Chem. A, 106, 5114–5119, 2002.
 - Lockwood, T. E., de Vega, R. G., Du, Z., Schlatt, L., Xu, X., and Clases, D.: Strategies to enhance figures of merit in ICP-ToF-MS, J. Anal. At. Spectrom., 39, 227–234, 2024.
 - Martínez-Sierra, J. G., San Blas, O. G., Gayón, J. M., and Alonso, J. G.: Sulfur analysis by inductively coupled plasma-mass spectrometry: a review, Spectrochim. Acta B, 108, 35–52, 2015.
 - Meija, J., Coplen, T. B., Berglund, M., Brand, W. A., De Bièvre, P., Gröning, M., and Prohaska, T.: Isotopic compositions of the elements 2013 (IUPAC Technical Report), Pure Appl. Chem., 88, 293–306, 2016.
 - Mulvaney, R., Wolff, E. W., and Oates, K.: Sulphuric acid at grain boundaries in Antarctic ice, Nature, 331, 247-249, 1988.
 - $\underline{\text{Muscheler, R., Beer, J., Kubik, P. W., and Synal, H. A.: Geomagnetic field intensity during the last 60,000 years based on {}^{10}\text{Be} }$
- 900 and ³⁶Cl from the Summit ice cores and ¹⁴C, Quat. Sci. Rev., 24, 1849–1860, 2005.

895

- Ng, F. S. L., Rhodes, R. H., Fudge, T. J., and Wolff, E. W.: Doomed descent? How fast sulphate signals diffuse in the EPICA Dome C ice column, EGUsphere [preprint], https://doi.org/10.5194/egusphere-2025-1566, 2025.
- Ng, F. S.: Pervasive diffusion of climate signals recorded in ice-vein ionic impurities, Cryosphere, 15, 1787–1810, 2021.
- Ng, F. S.: The grain-scale signature of isotopic diffusion in ice, Cryosphere, 18, 4645-4669, 2024.
- 905 Niehaus, P., de Vega, R. G., Haindl, M. T., Birkl, C., Leoni, M., Birkl-Toeglhofer, A. M., and Clases, D.: Multimodal analytical tools for the molecular and elemental characterisation of lesions in brain tissue of multiple sclerosis patients, Talanta, 270, 125518, 2024.
 - Ohno, H., Igarashi, M., and Hondoh, T.: Salt inclusions in polar ice core: location and chemical form of water-soluble impurities, Earth Planet. Sci. Lett., 232, 171–178, 2005.
- 910 Parrenin, F., Barnola, J. M., Beer, J., Blunier, T., Castellano, E., Chappellaz, J., and Wolff, E.: The EDC3 chronology for the EPICA Dome C ice core, Clim. Past, 3, 485–497, 2007.
 - Reinhardt, H., Kriews, M., Miller, H., Schrems, O., Lüdke, C., Hoffmann, E., and Skole, J.: Laser ablation inductively coupled plasma mass spectrometry: a new tool for trace element analysis in ice cores, Fresenius' J. Anal. Chem., 370, 629–636, 2001.
 - Rempel, A. W., Waddington, E. D., Wettlaufer, J. S., and Worster, M. G.: Possible displacement of the climate signal in ancient
- 915 ice by premelting and anomalous diffusion, Nature, 411, 568–571, 2001.
 - Rhodes, R. H., Bollet-Quivogne, Y., Barnes, P., Severi, M., and Wolff, E. W.: New estimates of sulfate diffusion rates in the EPICA Dome C ice core, Clim. Past, 20, 2031–2043, 2024.

- Severi, M., Becagli, S., Castellano, E., Morganti, A., Traversi, R., Udisti, R., and Steffensen, J. P.: Synchronisation of the EDML and EDC ice cores for the last 52 kyr by volcanic signature matching, Clim. Past, 3, 367–374, 2007.
- 920 Sigl, M., Toohey, M., McConnell, J. R., Cole-Dai, J., and Severi, M.: Volcanic stratospheric sulfur injections and aerosol optical depth during the Holocene (past 11 500 years) from a bipolar ice-core array, Earth Syst. Sci. Data, 14, 3167–3196, 2022.
 - Singh, S., Mucalo, M. R., and Grainger, M. N.: Analysis of sulfur in soil and plant digests using methane as a reaction gas for ICP-MS, Talanta, 281, 126797, 2025.
- 925 Sneed, S. B., Mayewski, P. A., Sayre, W. G., Handley, M. J., Kurbatov, A. V., Taylor, K. C., Bohleber, P., Wagenbach, D., Erhardt, T., and Spaulding, N. E.: New LA-ICP-MS cryocell and calibration technique for sub-millimeter analysis of ice cores, J. Glaciol., 61, 233–242, 2015.
 - Stoll, N., Eichler, J., Hörhold, M., Shigeyama, W., and Weikusat, I.: A review of the microstructural location of impurities in polar ice and their impacts on deformation, Front. Earth Sci., 8, 658, 2021.
- 930 Stoll, N., Hörhold, M., Erhardt, T., Eichler, J., Jensen, C., and Weikusat, I.: Microstructure, micro-inclusions, and mineralogy along the EGRIP (East Greenland Ice Core Project) ice core Part 2: Implications for palaeo-mineralogy, Cryosphere, 16, 667–688, 2022.
 - Stoll, N., Westhoff, J., Bohleber, P., Svensson, A., Dahl-Jensen, D., Barbante, C., and Weikusat, I.: Chemical and visual characterisation of EGRIP glacial ice and cloudy bands within, Cryosphere, 17, 2021–2043, 2023.
- 935 Svensson, A., Bigler, M., Blunier, T., Clausen, H. B., Dahl-Jensen, D., Fischer, H., and Winstrup, M.: Direct linking of Greenland and Antarctic ice cores at the Toba eruption (74 ka BP), Clim. Past, 9, 749–766, 2013.
 - Traversi, R., Becagli, S., Castellano, E., Marino, F., Rugi, F., Severi, M., and Udisti, R.: Sulfate spikes in the deep layers of EPICA-Dome C ice core: evidence
 - Wilhelms-Dick, D.: Enhanced analysis of stratified climate archives through upgrade of Laser Ablation Inductively Coupled
- 940 Plasma Quadrupole to Time of Flight Mass Spectrometry?, Doctoral dissertation, Universität Bremen, 2008.
 - Wolff, E. W., Mulvaney, R., and Oates, K.: The location of impurities in Antarctic ice, Ann. Glaciol., 11, 194–197, 1988.
 - Wolff, E. W., Fischer, H., van Ommen, T., and Hodell, D. A.: Stratigraphic templates for ice core records of the past 1.5 Myr, Clim. Past, 18, 1563–1577, 2022.
- Wolff, E. W., Fischer, H., Fundel, F., Ruth, U., Twarloh, B., Littot, G. C., and Gaspari, V.: Southern Ocean sea-ice extent, productivity and iron flux over the past eight glacial cycles, Nature, 440, 491–496, 2006.
 - Yan, Y., Bender, M. L., Brook, E. J., Clifford, H. M., Kemeny, P. C., Kurbatov, A. V., and Higgins, J. A.: Two-million-year-old snapshots of atmospheric gases from Antarctic ice, Nature, 574, 663–666, 2019.

Seite 5: [1] hat gelöscht	Pascal Bohleber	09.07.25 17:06:00
Seite 6: [2] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
Seite 6: [3] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
Seite 6: [4] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
Seite 6: [5] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
Seite 6: [6] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
Seite 6: [7] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
Seite 6: [8] hat formatiert	Microsoft Office User	16.07.25 14:10:00
Schriftart: (Standard) +Textkörper (Times New Roman), Englisch (USA)	
		10.07.25 10:57:00

I

I

l

I

l

I

l