Deep ice as a geochemical reactor: insights from iron speciation and mineralogy of dust in the Talos Dome ice core (East Antarctica)

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Abstract. Thanks to its insolubility, mineral dust is considered a stable proxy in polar ice cores. With this study we show that below an ice-depth of 1000 m, the Talos Dome ice core (TALDICE, Ross Sea sector of East Antarctica) presents displays evident and progressive signs of post-depositional processes affecting the mineral dust record below 1000 m deeps. We applied a suite of established and cutting-edge techniques to investigate the properties of dust present in the Talos Dome ice core TALDICE, ranging from concentration and grain-size to elemental-composition and Fe-mineralogy. Results show that through acidic/oxidative weathering, the conditions of deep ice at Talos Dome promote the dissolution of specific minerals and the englacial formation of others, deeply-ffecting dust primitive features. The expulsion of acidic atmospheric species from ice-grains and their concentration in localized environments is likely the main process responsible for englacial reactions and is related with ice re-crystallization. Deep ice can be seen as a "geochemical reactor" capable of fostering complex reactions which involve both soluble and insoluble impurities. Fe-bearing minerals can efficiently help exploring be used to explore such transformations.

1. Introduction

Antarctic ice cores are a valuable archive which allowed to reconstruct the climatic history of the Earth during the last 800,000 years through ice cores (Wolff et al., 2010) and even further back in time thanks to ancient ice outcropping at blue-ice fields (Yan et al., 2019). Mineral dust is one of the most extensively studied many proxies that is possible to investigate in ice cores. Its importance stems from its role in the Earth’s climate system: production, transport and deposition of dust are controlled by climate-related processes, but at the same time dust affects the climate (Maher et al., 2010). Ice cores have
been fundamental for understanding the role of atmospheric dust in past climatic periods. Studying the properties of dust trapped in ice cores, it has been possible to obtain information about on how the influence of climate influences on the dust cycle (Delmonte et al., 2004) and about the effects of dust under different climatic regimes (Mahowald et al., 1999; Wolff et al., 2006; Potenza et al., 2016).

It is known that the physical and compositional properties of dust trapped in ice cores are influenced by climatic, environmental and atmospheric processes (Sugden et al., 2009; Delmonte et al., 2017; Markle et al., 2018). The dust concentration of dust in ice is strictly controlled by climate (Delmonte et al., 2002), while dust grain size is related to its atmospheric transport (Delmonte et al., 2017; Albani et al., 2012a) and geochemistry to dust sources (Delmonte et al., 2004).

In the last years, a growing number of studies focused on the fact that the proxies used to infer paleoclimatic information from ice cores are not imperishable (Barnes et al., 2003; De Angelis et al., 2013; Ohno et al., 2016; Eichler et al., 2019; Baccolo et al., 2021). After the incorporation in snow, the proxies are affected by alterations occurring produced by after their incorporation into snow, which are defined as post-depositional processes. Generally, As a general rule, the older the ice and the stronger the influence of such changes progressively increases with ice age and depth (De Angelis et al., 2013; Tison et al., 2015), despite some occur in the surficial part of ice sheets, such as loss of volatile species, fractionation in firm and wind-scouring (Blunier et al., 2005; Town et al., 2008; Buizert et al., 2013).

Post-depositional processes recognized to act in deep ice are related to three main causes: (1) the interaction between ice flow and bedrock (Goossens et al., 2016); (2) the metamorphism of ice, intended as the set of physical transformations to which ice is subject at increasing pressure and age (Faria et al., 2010); 3- the diffusion of impurities (Barnes et al., 2003). An irregular bedrock can produce stratigraphic disturbances affecting the original ice succession for several hundred meters. On the contrary, ice metamorphism and englacial diffusion act at ice-crystal a smaller scale, comparable to the size of ice-crystals (mm-cm scale) (Rempel et al., 2002; Faria et al., 2010). The effects of such phenomena on the proxies investigated in ice cores are not yet fully understood, but improving their comprehension will be essential be topical considering the "Oldest Ice" challenge (Fischer et al., 2013). One aspect that still needs to be addressed is the role of depth and pressure in post depositional processes. From the top to the bottom of Antarctic ice cores, temperature-pressure conditions change significantly, promoting different and diverse physical and/or chemical reactions (Tison et al., 2015).

Dust is considered relatively immobile and stable in ice and its concentration has been used to synchronize deep ice cores when other proxies deteriorated (Ruth et al., 2007; Kawamura et al., 2017). However, mineral particles may be altered not untouched by post-depositional changes. Impurities in ice are affected by the small-scale relocation resulting from ice metamorphism,
i.e., which in deep ice consists in the growth and re-crystallization and orientation of ice grains (Faria et al., 2010; Marath and Wettlaufer, 2020). Being most of the impurities incompatible with respect to the ice lattice (Wolff, 1996), during the re-crystallization they the impurities are expelled from the ice crystalline structure and accumulated at ice grain junctions or within intra-grain micro-inclusions (Mulvaney et al., 1988; De Angelis et al., 2005; Sakurai et al., 2017; Stoll et al., 2021). The accumulation of soluble and insoluble impurities forms eutectic mixtures whose pressure melting point is below the ice temperature, promoting the localized formation of allowing for the local formation of liquid brines which lead favor to in situ chemical reactions (Fukazawa et al., 1998; De Angelis et al., 2005, 2013; Sakurai et al., 2017). Such small scale environments are dominated by sulphur-rich acidic species, strongly affected by re-mobilization and concentration because of their high incompatibility with the ice lattice (Mulvaney et al., 1988; Wolff, 1996; Fukazawa et al., 1998). The interaction between acidic brines and concentrated impurities, including dust, leads to acid-base reactions (Traversi et al., 2009; Ohno et al., 2016). Considering dust With respect to dust, the most common reactions which-happening in ice have been recognized to take place into ice are the dissolution of carbonates, the precipitation of gypsum (Ohno et al., 2006; Iizuka et al., 2008; Eichler et al., 2019) and of other uncommon sulphates (Ohno et al., 2014), or and the englacial formation of secondary iron minerals (De Angelis et al., 2013; Baccolo et al., 2021). Below 3000 m deep in the EPICA Dome C ice core, in very deep englacial environments also the re-precipitation of carbonates has been reported, suggesting that once acidity is consumed, additional reactions take place (Tison et al., 2015).

The iron fraction of dust is particularly sensitive to englacial transformations. De Angelis et al. (2013) have identified secondary Fe-bearing minerals in the deep part of the EPICA Dome C ice core, while Eichler et al. (2019) have identified-detected in the EPICA Dronning Maud Land ice core a few particles whose-with a Raman signature was compatible with jarosite, a Fe-K sulphate forming through acidic from mineral weathering (Papike et al., 2006). This finding has been confirmed by a thorough investigation of the Talos Dome ice core (TALDICE, East Antarctica), where jarosite has been found below the depth of 1000 m deep and interpreted as the result of acidic, water-limited weathering of dust (Baccolo et al., 2021). Since Fe biogeochemistry is strongly coupled with the global carbon cycle, this element receives considerable attention by the ice core community (Wolff et al., 2006; Conway et al., 2015; Hooper et al., 2019) and methods have been developed to measure its concentration and speciation in ice cores (Spolaor et al., 2013; Burgay et al., 2019; Conway et al., 2015). Nevertheless, But until now, the effects of post-depositional processes in ice cores on its geochemistry have still been only poorly investigated.
This aim of the present work is to describe and discuss a set of geochemical evidences showing that below 1000 m deep the dust record of TALDICE is affected by post-depositional transformations which alter its particle-size distribution, elemental composition, and mineralogy. A detailed analysis of speciation and mineralogy of the Fe-fraction of dust reveals that deep ice at Talos Dome acts as a geochemical reactor, favoring reactions and transformations which involve both soluble and insoluble impurities how climate and post-depositional processes affect dust in the Talos Dome ice core (TALDICE, East Antarctica), with a focus on Fe speciation and mineralogy.

The TALDICE ice core

TALDICE has been drilled at Talos Dome (72°49'S, 159°11'E; 2315 m a.s.l.), a peripheral accumulation dome in the Ross Sea sector of East Antarctica (Frezzotti et al., 2004). It is 1620 m long and its ice age at the depth of 1438 m (depth refers to the distance between the ice surface and the considered section of the core) is ~150 ky BP according to the AICC2012 chronology (Veres et al., 2013; Bazin et al., 2013), but there is evidence that the water stable isotope record is preserved up to ~1550 m deep, where the estimated ice age is ~343 ky BP, corresponding to Marine Isotopic Stage 10.1 (Crotti et al., 2021). Extensive research has been conducted on the dust record of TALDICE. Its concentration trend during the last climatic cycle reflects what observed at other East Antarctic sites, i.e. high atmospheric loads in glacial periods and lower ones during interglacials. A peculiarity of TALDICE is given by the influence of local Antarctic dust sources, corresponding to the ice-free sites of Victoria Land region, close to Talos Dome (Delmonte et al., 2010; Baccolo et al., 2018b).

2. Materials and methods

2.1. The TALDICE ice core

TALDICE has been drilled at Talos Dome (72°49'S, 159°11'E; 2315 m a.s.l., Fig.1), a peripheral ice dome in the Ross Sea sector of East Antarctica (Frezzotti et al., 2004). It is 1620 m long and its ice age at 1438 m deep (depth refers to the distance between the ice surface and the considered section of the core) is ~150 ky BP according to the AICC2012 chronology (Veres et al., 2013; Bazin et al., 2013), but there is evidence that the water stable isotope record is preserved up to ~1550 m deep, where the estimated ice age is ~343 ky BP, corresponding to Marine Isotopic Stage 10.1 (Crotti et al., 2021). Dust concentration in TALDICE during the last climatic cycle reflects high atmospheric loads in glacial periods and lower ones during interglacials, as observed at other East Antarctic sites (Delmonte et al., 2010). A peculiarity of TALDICE is given by the influence of local Antarctic dust sources, corresponding to the ice-free sites of the Victoria Land region, close to Talos Dome (Albani et al., 2012a; Baccolo et al., 2018b).

1.1.2. Sample preparation

Fifty-four samples were prepared using 191 TALDICE ice sections of TALDICE (~25x3x2 cm each one 25 cm long). They consist of insoluble mineral particles extracted from meltwater and deposited on filtration membranes. The
preparation took place in a clean room (ISO6) installed at the University of Milano-Bicocca. Ice sections were decontaminated with three baths in ultra-pure MilliQ water (Merck Millipore). They were stored in clean tubes under a ISO5 laminar flow bench until complete melting. Meltwater was split in two aliquots, one (~10 mL) for Coulter counter, the remnant for synchrotron radiation analysis.

The aliquots for Coulter counter (CC), corresponding to single ice sections, were stored in plastic clean cuvettes (rinsed with MilliQ water) and added with a NaCl solution (final Na$^+$ concentration of samples ~1% m/m). The solution was prepared using MilliQ water and high purity solid NaCl and before use it was filtered with 0.22 μm pore-size filters. This procedure makes the liquid sample electrically conductive, a requisite for Coulter counter analyses. Final Na$^+$ concentration of samples was ~1% m/m. The aliquots for synchrotron radiation analysis were merged considering multiple ice sections (191 in total), so as to prepare 54 samples with a total dust consisting each in a mass of suspended dust of at least 1 μg (estimated through Coulter counter). Merged samples were filtrated using hydrophilic PTFE membranes (ø 13 mm, pore-size 0.4 μm). Before filtration, membranes were rinsed for two weeks in a high purity HNO$_3$ solution (concentration m/m 5%, weekly renewed). Filtration is done with a micro-pipette to concentrate the particles on the membrane in the smallest possible area. Filtration was operated with a micro-pipette, so as to concentrate the particles on the membrane in the smallest possible area (Macis et al., 2018). After filtration the membranes were placed in dedicated clean PTFE holders and sealed in plastic bags.

Samples were prepared considering the entire length of the core: 7 Seven samples correspond to the Holocene (0-673 m, 0-11.7 ky BP), 7 to the last deglaciation (674-827 m, 11.7-18 ky BP), 5 to MIS2 (828-951 m, 18-30 ky BP), 15 to MIS3 (952-1259 m, 30-60 ky BP), 23 to MIS4 (1260-1292 m, 60-80 ky BP), 6 to MIS5 (1293-1418 m, 80-146 ky BP), 1 from to MIS6 (1419-1438 m, 146 154 147 ky BP) and 11 to the deep part of TALDICE not dated by AICC2012 chronology (1439-1620 m), but partially dated by the new TALDICE-deep chronology (Crotti et al., 2021).

**1.2.2.3. Coulter counter**

To determine the concentration and grain size of insoluble dust particles, the Coulter counter method was used. The method is well-known for the analysis of insoluble particles in ice cores. It relates the changes in the electrical conductance of meltwater with the size and number of insoluble particles suspended into the suspension (Delmonte et al., 2002). Samples were measured with a Beckman Multisizer 4, equipped with a 30 μm orifice to measure the concentration of particles between 0.6 and 18 μm divided into 256-400 channels. Details are found in Delmonte et al. (2002).

**1.3.2.4. Synchrotron radiation spectroscopic measurements**

The application of synchrotron light to determine the elemental and mineralogical composition of TALDICE dust, performed at beamtime B18 of the Diamond Light source (Cibin et al., 2019). A glovebox was reconstructed and connected
to the experimental chamber of the beamline to handle the samples in clean conditions. Many additional precautions are adopted to limit contamination and increase the signal to noise ratio: the application of plastic sheets inside the experimental chamber to limit radiation backscattering; the defocus of the incidental beam to illuminate the largest part of the samples; the preservation of high-vacuum during the acquisition. Further details are available in Baccolo et al., (2018a).

1.3.1.2.4.1. **X-ray fluorescence spectroscopy**

Major elements (Na, Mg, Al, Si, K, Ca, Ti, Mn, Fe) in dust were investigated through synchrotron radiation X-ray fluorescence spectroscopy, using synchrotron radiation as the excitation source (Iida, 2013). Samples were irradiated with a 10 keV incident beam (cross section ~1x1 mm) for 600 s and the fluorescence signal was acquired with a silicon drift detector, allowing the quantification of the following elements: Na, Mg, Al, Si, K, Ca, Ti, Mn, Fe. Analytical accuracy is evaluated analyzing NIST standard reference materials (SRM 2709a); it decreases from light to heavy elements (standard deviation of the replicates for Na is 25%, 10% for Fe). Recovery factors are evaluated comparing certified concentrations of SRMs with calculated values; they range from 85% to 115% except for Ca and Na (133% and 129%). Full details, including analytical performance, are given in Baccolo et al. (2018a, b). Elemental concentrations were converted into oxides concentrations and closed to 100%, following an established practice (Rudnick and Gao, 2003).

1.3.2.2.4.2. **X-ray absorption spectroscopy**

Speciation and mineralogy of the Fe fraction of TALDICE dust were investigated through X-ray absorption near edge structure spectroscopy (XANES), performed at the Fe K-edge transition, that is the excitation energy of the innermost electrons of Fe. XANES relates the spectral features of X-ray absorption spectra to chemical and molecular characteristics of specific elements. The sample is irradiated with a monochromatic beam of photons whose energy finely changes with time. The response of the sample depends on features such as oxidation, coordination and mineralogy (Calvin, 2013). For each sample three measurements were carried out, acquiring the fluorescence signal of the samples at steps of 0.15 eV and considering the interval between 7000 and 7400 eV. Spectra were calibrated, normalized and averaged using the Athena software (Ravel and Newville, 2005). Three spectral features were gathered: (1) the energy of the Fe K-edge transition; (2) the energy of the pre-edge peak centroid; 3- the intensity of the pre-edge peak; see (Figure S1) for details (Baccolo et al., 2018b). The energy of both pre-edge peak and Fe K-edge transition are directly related to the oxidation state of Fe, while the intensity of the pre-edge peak depends on its coordination and symmetry considerations (Berry et al., 2003).

1.3.3.2.4.3. **Relative abundance of Fe-bearing minerals**

Comparing dust samples with the ones corresponding to 14 Fe-mineral references (biotite, chlorite, glaucophane, goethite, hematite, hornblende (ferro-hornblende), jarosite, magnetite, muscovite, fayalite, pargasite, pyrite, schoerlite, siderite), it was...
possible to estimate the contribution of single Fe-bearing minerals into the samples (Shoenfelt et al., 2018). XANES spectra of reference minerals were collected following the same procedure protocol adopted for TALDICE samples (Figure S2). XANES spectra of ice core dust were reproduced through ordinary least square regression (OLS), using linear combinations constructed with mineral spectra (Figure S3). For each sample all the combinations defined by 4, 3, 2 and 1 mineral references (1456 combinations per sample) were calculated through ordinary OLS least square regression and the best one (in terms of R-squared, it always exceeded 0.9) was selected to represent the sample. In some cases, the second best-fit combination had an R-squared close to the best-fit one, but the difference between the two combinations always regarded the less abundant of the 4 selected references, with negligible effects on the data analysis and interpretation. The combinatoric package of the Athena software was used (Ravel and Newville, 2005). Following the procedure adopted by Shoenfelt et al. (2018) and Liu et al. (2018), the relative abundance of Fe-bearing minerals was estimated considering the % linear coefficients of the selected combinations obtained from OLS.

2.3. Results and discussion

2.3.1. The TALDICE dust record

Figure 1 shows the dust record of TALDICE. Considering the last climatic cycle (i.e., the Holocene and MIS 2-3-4), some well-known features characterizing the relationships between the atmospheric dust cycle and Antarctic climate are recognized (Fig. 1). The most evident is the negative correlation between dust concentration and δ¹⁸O (1m resolution Stenni et al., 2011). In accordance with the suppression of dust production and transport from remote sources during interglacials (Albani et al., 2012b), the mean dust concentration in TALDICE Holocene ice is ~25 ng g⁻¹, while during MIS2, corresponding to the last glacial maximum, it exceeds 300 ng g⁻¹, as a consequence of the massive activation of South American sources (Sugden et al., 2009) and to the enhanced atmospheric transport toward Antarctica (Markle et al., 2018).

The shift from interglacial to glacial conditions not only affects dust concentration, but also its grain size, as revealed by the FPP (fine particle percentage) and CLPP (coarse local particle percentage) indexes. The first one is the relative concentration of particles between 0.6 and 2 μm with respect to the 0.6 - 5 μm interval; CLPP is the ratio between the concentration of particles between 5 and 10 μm and the total concentration of particles between 0.6 and 10 μm. During the Holocene, FPP has a mean value of 50%, while during MIS2 it increases to 63%, revealing that under glacial conditions dust particles deposited at Talos Dome are relatively smaller than in interglacial periods. Similar evidence has already been reported from other East Antarctic sites (Dome C, Vostok) and interpreted considering that dust transported to East Antarctica during glacials is subject to long-range and high-altitude atmospheric pathways, allowing for the efficient removal of coarse particles (Delmonte et al., 2002). CLPP has a mean value of 19% during the Holocene and of 6.5% during MIS-2. This index is indicative of the relative abundance of particles larger than 5 μm, which are related to local Antarctic sources (Albani et al., 2012a; Baccolo et al., 2018b). The decrease of the index in glacial
periods must be interpreted in relative terms. CLPP is lower in glacials not because of a reduction of coarse particles, but due to an increase of the fine ones from South America (Baccolo et al., 2018b), indicative of the MIS4 displays features similar to MIS2, while MIS5.5 is similar to the Holocene, confirming increased deposition of fine dust during glacial culminations and reduced fluxes of relatively coarse dust during peak interglacial warmth. Going in the deeper layers of the core, the relationship relative abundance of particles larger than 5 μm, which, given the coarse diameter, are related to local Antarctic sources (Albani et al., 2012a; Baccolo et al., 2018b). The decrease of the index during glacial periods has to be interpreted in relative terms. It has been observed that the absolute activity of local Antarctic sources is only partially influenced by climate (Baccolo et al., 2018b). CLPP is lower in glacials not because of a reduction of coarse particles, but because of an increase of the fine ones from South America. MIS4 presents features similar to MIS2. During MIS5.5, the previous peak interglacial warmth, dust concentration reaches a low associated with high values of δ¹⁸O and CLPP, compatible with a regionalization of the atmospheric dust cycle in the Talos Dome area, similarly to the Holocene. Between dust and climate is less evident. Fig. 1 and Tab. 1 show—it can be appreciated—that below 1430 m deep, dust concentration exceeds 100 ng g⁻¹ with dampened oscillations. CLPP exceeds 20-% and FPP drops to below values between 45- and 35-%. Looking at highlighting that in deep TALDICE dust particles are coarser. This can be interpreted as an increase of dust grain size is interpreted as a consequence of dust aggregation in deep ice, a process already observed in Antarctic ice below 2500 m deep—a post depositional process occurring in deep ice (Lambert et al., 2008; De Angelis et al., 2013).

### Table 1. Dust concentration values in TALDICE. For each climatic period mean concentrations are reported along with standard deviations.

| Period (kyr BP) | Holocene | Degl. | MIS2 | MIS3 | MIS4 | MIS5 | MIS6 | MIS7 | MIS8-9 | deep part |
|----------------|----------|-------|------|------|------|------|------|------|--------|-----------|
| Conc (ng g⁻¹)  | 26±18    | 104±25| 317±174| 80±31| 116±59| 61±55| 158±190| 182±302| 121±73| 110±123 |
| FPP (%)        | 50±4     | 57±8  | 63±8 | 55±7 | 53±5 | 52±5 | 54±7 | 45±9 | 35±10 | 36±9      |
| CLPP (%)       | 19±7     | 14±7  | 6.5±3.4 | 10±5 | 9.6±3.9| 13±8 | 9.1±5.2| 21±14 | 21±6  | 24±11     |
Figure 1. The mineral dust record of TALDICE the Talos Dome ice core. From the upper to the lower curve: (a) total dust concentration of insoluble particles presenting a diameter between 0.6 and 10 μm (Baccolo et al., 2021); (b) FPP (fine particle percentage); (c) CLPP (coarse local particle percentage); (d) δ¹⁸O (Stenni et al., 2011). Light blue bands highlight glacial culminations (marine isotopic stages MIS 2-4-6-8), the red one bands the Holocene and MIS marine isotopic stage 5.5, 7.5 and 9.5. In the upper-left corner the position of Talos Dome in Antarctica.

Size distributions of insoluble particles also point to dust aggregation. Dust from the upper part of the core (Figure 2e-h), presents a mode between 1 and 2.5 μm and a tail of particles larger than 5 μm, suggesting a mix from remote and local sources (Albani et al., 2012a). Dust from deep TALDICE (Figure 2j-l) is characterized by a higher abundance of coarse particles, a lack of fine ones and modal values exceeding 4 μm. Such features are not encountered in shallow sections of Antarctic ice cores, where the effects of post-depositional alterations are limited (Royer et al., 1983; Delmonte et al., 2002; Wegner et al., 2015); they result from in situ aggregation of particles in deep ice (Lambert et al., 2008; De Angelis et al., 2013; Baccolo et al., 2021).

Post-depositional processes in TALDICE ice also influence the climatic significance of its dust record. This is shown in Fig. 2a-d, where the correlation between dust and ice stable isotopes is analyzed. The correlation is high during the last climatic cycle, but decreases in older periods. This is revealed by R-squared decreases from 0.66 during the last cycle, to less than 0.3 in the deep core. The same conclusion is drawn looking at the trajectory describing the evolution of the δ¹⁸O – dust pair. During the first cycle it well reproduces the transition from glacial to interglacial conditions, including and other events, such as the Antarctic Cold Reversal and the partition of the last glacial period into MIS 2-3-4. This is
partially true for the previous climatic cycle, but the correlation decreases (R-squared from 0.66 goes to 0.49). The degradation continues in the deepest part of TALDICE, where the coefficient does not exceed 0.25 and the trajectories follow irregular paths, highlighting a substantial decoupling of dust concentration and isotopic signals. This is confirmed by Crotti and colleagues (2021) who showed that at Talos Dome below 1548 m deep, the climatic signals are not preserved, doesn’t exceed 0.25 and the trajectories follow irregular paths, highlighting a substantial decoupling of dust concentration and isotopic signals. This is confirmed by a study (Crotti et al., 2021) showing that at Talos Dome below 1578 m deep, climatic signals are not preserved.

**Figure 2.** Panels a-d: linear correlation between $\delta^{18}$O and dust concentration (log(logarithmic scale)) in the TALDICE Talos Dome ice core. Blue bands correspond to the 95% confidence level interval of the linear regression calculated considering the last climatic cycle; dashed lines highlight the 95% confidence level interval of the linear regression calculated considering the last climatic cycle (panel a). The orange curve is the trajectory showing the evolution of the $\delta^{18}$O - dust pair (arrows from shallow to deep) with time (red arrows). It was obtained applying a first order (40-% sample window) Savitzky-Golay filter to both variables. Panel e-I: dust grain size distributions from the Talos Dome ice core TALDICE.

### 2.2.3.2. Major element composition of mineral dust

Comparing TALDICE dust composition during the last climatic cycle resembles the signature of Post Archean Australian Shale (PAAS, Taylor and McLennan, 1985; Fig. 3) with geochemical references; it is noticed that the best agreement is found with the Post Archean Australian Shale reference (PAAS) (Taylor and McLennan, 1985), not with the average Upper Continental Crust reference (UCC) (Rudnick and Gao, 2003). With respect to the Upper Continental Crust (UCC, Rudnick and Gao, 2003), PAAS is depleted in mobile major oxides (such as CaO and Na$_2$O) and enriched in Fe-Ti and Al-oxides as a result of chemical weathering. UCC refers to the whole upper continental crust, while PAAS is in fact representative of surficial sedimentary rocks subject to chemical weathering (Taylor and McLennan, 1985), while UCC of the whole upper...
The continental crust is richer in residual oxides while lacks labile soluble oxides. The similarity between TALDICE dust and PAAS is not unexpected, since atmospheric dust is produced at the Earth surface, where sedimentary and weathered rocks dominate.

Another feature emerging from Fig. 3, is represented by the change of dust composition along with depth in the core. Some oxides show significant variations with depth (see also Fig. S4). The ones showing the most evident trends are SiO$_2$ (increasing) and CaO (decreasing). Their, which pass from an average concentration varies between 64% and 1.7% in the Holocene to 74% and 0.4% in the deep part of the core, respectively. Other oxides showing minor variations are: Na$_2$O (decreasing), MgO (decreasing), Al$_2$O$_3$ (decreasing) and K$_2$O (increasing). TiO$_2$, MnO and Fe$_2$O$_3$ are stable. These variations are related to depth and not affected by climatic cycles, only by depth. This is an indication that they are likely related to post-depositional processes and not to primary changes of dust sources. The effect of disturbance from the bedrock must also be discarded since the ice stratigraphy at Talos Dome is uninterrupted until 15478 m deep (Crotti et al., 2021). Ca, Mg and Na, the elements showing the strongest decrease, are mobile and typically affected by chemical weathering (Nesbitt & Young, 1982). Their reduction suggests a progressive alteration of dust with depth. In particular, the deepest samples lack MgO and CaO, indicating carbonate dissolution. The reaction between acidic species and carbonates and the consequent precipitation of gypsum, are well-known post-depositional processes in deep ice (Ohno et al., 2006; Iizuka et al., 2008; Traversi et al., 2009; Eichler et al., 2019). Our evidences suggest that this reaction also occurs at Talos Dome. The increase of SiO$_2$ is likely relative and reflects the progressive loss of labile species.

### 3.3. Iron oxidation and coordination symmetry

XANES reveals a progressive oxidation of Fe with depth (Fig. 4). Figure 4a shows that in the first 1000 m of the core, Fe in dust consists of a mixture of Fe$^{2+}$ and Fe$^{3+}$, reflecting the typical composition of mineral aerosol (Schroth et al., 2009). Deep samples show a pure Fe$^{3+}$ signature. About Fe-coordination, samples display an octahedral symmetry (coordination number 6), with secondary inputs from other geometries. This is also in accordance with observations concerning aerosols (Wilke et al., 2001; Formenti et al., 2014). Figure 4b and c focus on Fe oxidation. Panel c shows the variation of the K-edge energy transition along the core. It increases by ~2 eV, pointing to increase confirms the oxidation of Fe in mineral dust in deep ice. The process is rather continuous, regardless of the climatic oscillations, suggesting that it is related to post-depositional changes. This is confirmed by the similarity of the oxidation trend and the growth of ice grains with depth (Fig. 4c). There are two exceptions: (1) in the deepest part of the core the Fe K-edge reaches a stable energy, pointing to a complete oxidation; (2) in correspondence with MIS2 and 4 the trend shows two slowdowns, as if oxidation was inhibited.
The partial increase of Fe$^{2+}$ during glacial culminations MIS 2 and 4 is related to the transport of fresh glacial dust from South America (Spolaor et al., 2013), from South America only partially oxidized due to limited atmospheric exposure (Spolaor et al., 2013; Shoenfelt et al., 2017). Another reason for this increase is the chemical weathering of dust in TALDICE. In particular, the deepest samples lack of MgO and CaO, pointing to carbonate dissolution. The reaction between acidic species and carbonates, with the consequent precipitation of gypsum and other soluble sulphates, is one of the most investigated post-depositional processes in deep ice (Ohno et al., 2006; Iizuka et al., 2008; Traversi et al., 2009; Eichler et al., 2019). Our evidence confirms that this reaction also occurs at Talos Dome and for the first time they are detected in relation to the compositional changes of insoluble dust. The increase of SiO$_2$ is likely relative and reflects the progressive loss of labile species.

Table 2. Average major element composition of TALDICE dust. Data are expressed as % mass fractions of oxides. For each climatic period mean values are reported with standard deviations.

| Period (kyr BP) | Holocene | Degl. | MIS2 | MIS3 | MIS4 | MIS5 | MIS6 | deep part |
|-----------------|----------|-------|------|------|------|------|------|-----------|
| Na$_2$O (m/m %) | 3.1±1.1  | 2.1±1.0 | 1.9±0.5 | 2.0±0.5 | 4.3±3.3 | 2.3±1.0 | 0.2 | 1.6±0.9   |
| MgO (m/m %)    | 1.6±0.4  | 2.0±1.0 | 3.0±1.6 | 0.8±0.3 | 0.8±1.1 | 0.4±0.3 | 0.5 | 0.6±0.8   |
| Al$_2$O$_3$ (m/m %) | 20.7±5.3   | 19.7±4.8 | 23±2.5 | 16.3±3.9 | 23.5±8.3 | 14.7±3.9 | 17.2 | 13.5±5.3   |
| SiO$_2$ (m/m %) | 63.9±5.0 | 65.9±6.3 | 62.9±3.4 | 72±3.6 | 63.9±4.5 | 73.4±3.9 | 72.9 | 74.2±5.6   |
| K$_2$O (m/m %) | 1.6±0.3  | 1.8±0.5 | 1.7±0.3 | 2.0±0.4 | 2.6±1.7 | 2.3±0.5 | 2.5 | 2.4±0.5   |
| CaO (m/m %)    | 1.7±0.4  | 1.3±0.5 | 1.0±0.4 | 0.9±0.4 | 0.4±0.2 | 0.4±0.2 | 0.3 | 0.4±0.3   |
| TiO$_2$ (m/m %) | 1.5±0.4  | 1.6±1.3 | 0.9±0.2 | 0.9±0.2 | 0.4±0.2 | 0.9±0.3 | 0.8 | 1.0±0.2   |
| MnO (m/m %)    | 0.06±0.02 | 0.05±0.01 | 0.06±0.02 | 0.04±0.02 | 0.05±0.04 | 0.04±0.01 | 0.04 | 0.04±0.02 |
| Fe$_2$O$_3$ (m/m %) | 5.8±0.6 | 5.6±1.3 | 5.4±1.1 | 5.1±0.8 | 4.0±1.0 | 5.5±0.6 | 5.6 | 6.3±1.5   |
2.3. Iron oxidation and coordination

The most evident result from XANES concerns the progressive oxidation of Fe with depth. This is appreciated through different approaches in Fig 4. Figure 4a shows that the Fe fraction of dust in the first 1000 m of the core consists in a mixture of Fe$^{2+}$ and Fe$^{3+}$, reflecting the typical composition of mineral aerosol (Schroth et al., 2009). Deep samples show a pure Fe$^{3+}$ signature. With respect to coordination, samples display an octahedral symmetry (coordination number 6), with secondary inputs from other geometries. This is also in accordance with observations concerning aerosols (Wilke et al., 2001; Formenti

![Figure 4a](image1.png)

**Figure 3.** Major element in dust from the TALDICE Talos Dome ice core. Data are shown considering all major element oxides (panel a), excluding SiO$_2$ (panel b) and SiO$_2$ and Al$_2$O$_3$ (panel c). UCC (upper continental crust, Rudnick and Gao (2003)) and PAAS references (post-archean Australian shale, REF) are shown for comparison (Rudnick and Gao, 2003; Taylor and McLennan, 1985).

for the slowdown of Fe oxidation during glacials is that in these periods the amount of dust deposited at Talos Dome acts as a buffer and partially neutralizes the acidity of ice, and consuming reactive species, as proposed for the Dome Fuji (Ohno et al., 2005, 2006 and EDML ice cores (Ohno et al., 2005, 2006; Eichler et al., 2019), and inhibits dust oxidation. Figure 4c shows that the growth of ice grains is also temporarily inhibited during MIS2 probably because of grain boundary pinning by insoluble particles as
Figure 4. X-ray absorption near edge structure spectroscopy (XANES) results from the analysis of dust in the TALDICE Talos Dome ice core. Panel a: analysis of the Fe K- pre-edge spectral region. The intensity and energy position of the pre-edge peak are shown following the scheme proposed by Wilke et al. (2001). Ellipses and numbers in brackets refer to the coordination of Fe, vertical lines to its two oxidation states; the orange curve represents the trajectory of the samples along the core, it was calculated as in Fig. 2 (arrows from shallow to deep). Panel b: a comparison between the energy position of the pre-edge peak and the main K-edge transition (see Fig. S1 for details). Panel c: the energy position of the Fe K-edge transition (blue line) vs. isotope composition of ice (red line), from (Stenni et al., 2011) and ice grain size (yellow line), from Montagnat et al. (2012); the arrow indicates the observed but not quantified ice crystals larger than 40 cm found below the depth of 1481 m (deep); blue bands highlight MIS 2, 4 and 6. For each panel one point presents mean error bars (not visible in panel c because of scale).

During glacial culminations (MIS 2, 4 and 6) the dominant source of dust transported to Antarctica was Southern South America (Delmonte et al., 2004, 2010). One of the processes responsible for the increased dust emission during glacials, is glacial activity producing deflatable sediments (Sugden et al., 2009). Glacial sediments are geochemically fresh and thanks to the limited atmospheric exposure they are only partially oxidized (Shoenfelt et al., 2017), suggested by Durand et al. (2006) because of pinning on ice re-crystallization by the high concentration of insoluble particles (Durand et al., 2006).

Something similar pattern is visible in MIS4, while ice corresponding to MIS6 does not present neither an Fe$^{2+}$ recovery, nor a decrease of ice grain size (Figure 4c), probably because in deep ancient ice in situ oxidation of Fe-minerals and ice metamorphism re-crystallization are too advanced.

On the contrary, in interglacial ice atmospheric acidity is more available for post-depositional reactions favoring oxidation and weathering, also thanks to a more efficient ice re-crystallization (Iizuka et al., 2008; Eichler et al., 2019).

2.4.3.4. Iron mineralogy

Fe-mineralogy results are shown in Fig. 5, Fig. 6 and Tab. 3. Only minerals whose average relative abundance (rel. ab.) exceeds 2% have been considered to the aims of in the discussion.
3.1.1.3.4.1. Hornblende and jarosite

Minerals showing the most evident trends are hornblende and jarosite. Hornblende dominates samples in the first 1000 m of TALDICE with a decreasing trend (Fig. 6a), on the contrary, jarosite is present only below 1000 m deep and its concentration increases with depth (Fig. 6h). Trends related to these minerals involve the large parts of the entire core regardless of climatic conditions; we interpret them as a consequence of post-depositional processes. Hornblende, Ca$_2$(Fe$_{2+4}$Al)(Si$_7$Al)O$_{22}$(OH)$_2$, after goethite, is the second most abundant Fe-mineral in Holocene ice (rel. ab. 19.2%), but it rapidly decreases with depth and in the deepest part of TALDICE it is almost absent (rel. ab. 0.7%). Jarosite shows an opposite behavior. It is not present in the shallow part of TALDICE, it appears below 1000 m deep, becoming the dominant Fe-bearing mineral below 1400 m (rel. ab. 50.3%). The two trends are linked. Hornblende is a common ferrous mineral present in South American dust (Shoenfelt et al., 2018) and in general is one of the dominant Fe$_{2+}$-bearing minerals in global aerosol (Schroth et al., 2009; He et al., 2020). On the contrary, jarosite, KFe$_3$(SO$_4$_2)(OH)$_6$, a ferric sulphate, is not common in atmospheric dust. Jarosite is well-known for being a weathering product and its widespread identification in the deep layers of TALDICE has been regarded as an evidence of weathering affecting dust in deep ice (Baccolo et al., 2021). The concurrent decrease of hornblende and increase of jarosite confirm Fe oxidation. Hornblende seems to be the principal mineral whose dissolution leads to the consumption of Fe$_{2+}$, while jarosite build-up drives the accumulation of Fe$_{3+}$. The other minerals presenting a ferric-ferrous component, with the exception of magnetite, also show a decreasing trend (muscovite, siderite, pyrite) and are almost absent in the deepest part of the core.

**Table 3.** Average relative abundance of Fe-bearing minerals in TALDICE dust. Data are expressed as % abundances.

| Period (kyr BP) | Holocene | Degl. 11.7-19 | MIS2 19-31 | MIS3 31-58 | MIS4 58-68 | MIS5 68-132 | MIS6 132-154 | deep part |
|----------------|----------|---------------|-----------|-----------|-----------|-------------|-------------|----------|
| Hornblende (%) | 19.2     | 12.3          | 19.1      | 2.6       | 0         | 4.1         | 8.4         | 0.7      |
| Muscovite (%)  | 11.5     | 12.9          | 18.0      | 13.0      | 7.6       | 2.0         | 0           | 0        |
| Siderite (%)   | 2.9      | 3.9           | 0.4       | 4.2       | 1.9       | 2.1         | 0           | 1.1      |
| Magnetite (%)  | 4.8      | 3.8           | 0         | 0.8       | 35.2      | 5.6         | 0           | 0.9      |
| Goethite (%)   | 30.8     | 56.0          | 37.4      | 60.3      | 30.8      | 45.5        | 31.7        | 42.9     |
| Hematite (%)   | 15.7     | 8.3           | 20.6      | 2.6       | 0         | 0           | 0           | 0        |
| Pyrite (%)     | 5.9      | 1.8           | 0.5       | 1.8       | 3.3       | 1.9         | 1.7         | 2.1      |
| Jarosite (%)   | 0        | 0             | 0         | 10.5      | 14.3      | 38.8        | 58.1        | 50.3     |
| Others (%)     | 9.1      | 1.0           | 4.0       | 4.1       | 7.0       | 0           | 0           | 2.0      |
Beside the trends involving the entire ice core, some minerals show an additional pattern in correspondence of MIS2. In some cases, it is a relative maximum (muscovite, hematite), in others a minimum (siderite, pyrite). Given the correspondence with MIS2, such features are interpreted as climate-related signals. Considering the Holocene siderite and pyrite, FeCO$_3$ and FeS$_2$ respectively, constitute about 3% each of Fe-minerals in TALDICE ice, but during MIS2 their abundance drop to 0.5%. A shift in mineralogy between the Holocene and MIS2 is expected, as during MIS2 dust deposited on East Antarctica mostly comes from Patagonia (Delmonte et al., 2004, 2010), while during the Holocene it is provided from the ice-free outcrops of Northern Victoria Land (Delmonte et al., 2010; Baccolo et al., 2018b). The shift of mineralogy between MIS2 and Holocene is related to this. In MIS2 dust is supplied to Talos Dome mostly by Patagonian sources (Delmonte et al., 2004, 2010), while in the Holocene its origin is local from Northern Victoria Land (Delmonte et al., 2010; Baccolo et al., 2018b). Minerals which distinguish local Holocene dust from glacial Patagonian one, are siderite and pyrite. In the Holocene they constitute about 3% each of the Fe-minerals in TALDICE, but during MIS2 they drop to 0.5%. The presence of siderite and pyrite in Holocene dust at Talos Dome agrees with the geology of Victoria Land, where they are relatively common accessory minerals, owing to the basaltic/doleritic nature of local rocks (Sturm and Carryer, 1970; Dow and Neall, 1974). In addition to the geologic context, also atmospheric transport could partially explain their absence during MIS2.

Figure 5. Pie-charts showing Fe-mineralogy of TALDICE dust from the Talos Dome ice core. Each chart refers to a climatic period. Marine isotopic stages 4 and 6 have been excluded because of the low number of samples corresponding to these periods.
are easily oxidized when exposed to the atmosphere. Their lack in MIS2 can be related to their oxidation during the long-range transport from South America (Shi et al., 2012). On the contrary, in the Holocene siderite and pyrite did not undergo chemical reactions because of proximal dust sources and short-range atmospheric transport.

3.1.3.4.3. Muscovite

Muscovite reaches its highest rel. ab. during MIS2, presenting a mean value of 18%. Dust deposited during MIS2 is rich in muscovite and hematite. Thanks to the aerodynamic shape of its crystals, muscovite, an Al-phyllosilicate, is common in atmospheric dust subject to long-range transport and is one of the most abundant minerals deposited on East Antarctica during MIS2 (Delmonte et al., 2017; Paleari et al., 2019). The abundance of this mineral in glacial ice is likely the main responsible reason for the slowdowns observed along the oxidation trend of Fe (Fig. 4c). Fe in muscovite is present as both ferric and ferrous iron. During MIS2, muscovite is the only mineral, with hornblende, presenting a ferrous component, since siderite, magnetite and pyrite are almost completely absent (Fig. 6).

Something similar is observed in MIS4, when the second slowdown in Fe oxidation is observed, again corresponding with an increase of muscovite. In this second case both the slowing of oxidation and the increase of muscovite are less evident, probably because of the stronger, more advanced weathering influence of oxidation at greater depth. Muscovite is completely absent below 1300 m deep, indicating that it is affected by weathering in deep ice, similarly to hornblende. Muscovite dissolution probably supplies a fraction of the K required. Below 1300 m deep, muscovite is completely absent, regardless of the climatic period, suggesting that also this mineral is affected by weathering in deep ice, such as hornblende. Being muscovite an Al-K silicate, its dissolution probably supplies a fraction of the K required for jarosite precipitation.
Figure 6. The variation abundance of the major Fe-bearing minerals identified in TALDICE dust with respect to depth. Dashed lines correspond to a 5-point moving average, solid grey lines to linear trends. In panel d, one sample-point is out of scale (black arrow, magnetite relative abundance 70%). Panel i shows the weathering index, defined as the % ratio between jarosite concentration and the sum of jarosite, hornblende, muscovite and hematite.

3.1.4. Hematite

Hematite, $\text{Fe}_2\text{O}_3$ (a Fe-oxide), is a weathering product in soils under dry and warm conditions, it is typical in tropical regions while it is rarely encountered in cold and wet climates (Schwertmann, 1988). Along In TALDICE it is mostly found during in MIS2, when the dust signature is fully South American (Delmonte et al., 2010). During glacial culminations it has been proposed that an additional source other than Patagonia, the Puna-Altiplano dry region in the tropical Andes, activates in South America, supplying dust to Antarctica. This is the Puna-Altiplano dry region, located in the tropical Andes (Delmonte et al., 2010). In the Puna-Altiplano, where hematite is widely present (Aubry et al., 1996). Accordingly, our results show an excess of hematite during in MIS2 (mean rel. ab. 21%), supporting this hypothesis. A previous study focusing on inner East Antarctica observes a higher abundance of hematite during the Holocene than in MIS2 (Paleari et al., 2019). The difference could be related to the geographic position of the two sites and to the relative influence of secondary sub-tropical sources during the Holocene. Below 1300 m deep, hematite is not observed, suggesting that this mineral is not stable in deep ice at Talos Dome. It is known that under acidic conditions (pH~4) hematite is not stable and...
dissolve\(^2\) (Schwertmann and Murad, 1983; Zolotov and Mironenko, 2007), leading to the precipitation of jarosite and goethite (Papike et al., 2006).

3.4.5. **Goethite**

Considering the entire core, goethite, FeO(OH), is the dominant Fe-mineral in dust, with a mean rel. ab. of 46% (the second is jarosite, 18-% rel. ab.). The distribution of goethite with depth is rather uniform (Fig. 6e). This is indicative of the fact that two things: (1) goethite is a common Fe-bearing phase in mineral aerosol (Formenti et al., 2014) and that (2) goethite is stable in the englacial environment, regardless of depth. The second point is corroborated by previous studies showing that at low temperature and in acidic wet environments, goethite is the most stable Fe oxide-hydroxide (Schwertmann and Murad, 1983; Zolotov and Mironenko, 2007).

3.4.6. **Magnetite**

Magnetite, Fe\(_3\)O\(_4\), has a The mean rel. ab. of magnetite, a ferric-ferrous oxide, isof about 3-% and does not show evident trends; only a single sample presents a high concentration (70.4-%). Without considering this anomalous value, possibly related to contamination or to the presence of a micrometeorite (Rochette et al., 2008), the mean value drops to 2-%. Despite its low rel. ab., magnetite is relatively stable along TALDICE and it is not depleted in the deepest part, suggesting that it is not affected by oxidation and/or dissolution, as the other ferrous minerals considered in this study. This confirms its higher chemical resistance to acidic oxidation when compared to other Fe-minerals (Moncur et al., 2009).

3.4.7. **The weathering index**

An index was developed to summarize the information from the different Fe-minerals. It is defined the weathering index and corresponds to the % ratio between the rel. ab. of jarosite and the sum between the rel. ab. of jarosite, hornblende, muscovite and hematite. The trend of the index varies between 0% in the upper part of TALDICE and 100% below 1300 m deep (Fig. 6i), reflecting the progressive weathering of dust, which consist in the consumption of some ferrous minerals and the precipitation of jarosite.

3.2. **Englacial weathering of dust**

When considering the last climatic cycle—in particular the pair Holocene-MIS2, which corresponds to the first 950 m of TALDICE, the ice core— the chemical and physical properties of TALDICE dust are fully interpretable in the light of the well-known effects of major climatic swings on atmospheric dust. This is true for concentration, grain size and geochemistry, suggesting that the dust record is not significantly affected by post-depositional processes within this depth interval. Things change moving down: below 1000 m deep, trends not related to climatic oscillations appear. One of the most evident is the presence—increase of jarosite. The mineral is first observed at 1000 m deep, and its concentration progressively increases toward the core bottom (Fig. 5 and Fig. 6h).
Jarosite is a Fe-K hydrated sulphate and results from chemical weathering. It has never been reported in global mineral aerosol. These features, together with its increasing abundance, support the hypothesis that jarosite is a product of englacial diagenesis related to dust chemical weathering (Baccolo et al., 2021). Thanks to the conditions under which jarosite precipitates, its formation returns supplies information about the deep englacial environment and in particular about the occurrence of limited amounts of liquid water (ice pre-melting). Liquid water in deep ice has been extensively predicted by theories (Rempel et al., 2002; Ng, 2021) and in a few cases confirmed by observations (Fukazawa et al., 1998), but is still a disputed argument (Eichler et al., 2019). The precipitation of jarosite is an strong indirect evidence for the presence of highly concentrated acidic liquid water solutions in deep ice (pre-melting), since this ferric mineralsulphate forms when limited amount of acidic (pH < 4) aqueous solutions rich in solutes (also known as brines) interact with Fe-bearing minerals (Zolotov and Shock, 2005; Papike et al., 2006). At Talos Dome ice temperature at 1000 m deep, where jarosite appears, is -25.5°C (Rix & Martin personal communication). Combining this information with the phase diagram of the sulfuric acid-water pair solution (Beyer et al., 2003), it is possible to estimate that acidic brines forming in deep ice at Talos Dome ice grain junctions have a sulfuric acid concentration between 15 and 25% m/m. Liquid water in deep ice has been predicted by theories (Rempel et al., 2002; Marath and Wettlaufer, 2020; Ng, 2021) and confirmed by observations (Fukazawa et al., 1998). The englacial formation of jarosite is a further confirmation of deep ice pre-melting. Moreover, it is also indicative that liquid water found in deep ice is strongly acidic and oxidative.

The relationships between ice re-crystallization and grain growth of deep ice are responsible for the concentration of impurities in deep ice is not yet completely understood (Eichler et al., 2019; Stoll et al., 2021). Our study suggests that such correlation actually exists and that it has effects on dust geochemistry since Fe oxidation and ice grain growth share a similar trend in TALDICE (Fig. 4c). The formation of the acidic brines required for jarosite precipitation is in fact probably related to ice re-crystallization, since the latter has been invoked as the responsible for the concentration of impurities in localized environments and for that can promote the occurrence of such brines in deep ice through the local lowering of pressure melting point (De Angelis et al., 2013; Tison et al., 2015). This is supported by the similar trends of Fe oxidation and ice grain growth (Fig. 4c). The acidity of deep brines is explained in the light of the strong incompatibility of acidic atmospheric species with respect to the ice molecular lattice (Wolff, 1996) and of their concentration at grain boundaries (Mulvaney et al., 1988), leading to the formation of acidic fluids.

The formation of jarosite not only alters the original mineralogical assemblage of dust in TALDICE, but it can also be related to the anomalies in dust grain-size observed in deep (Figure 2e-l). Jarosite is well-known for creating a cementing matrix during weathering (Long et al., 1992). Its precipitation promotes the aggregation of mineral particles, altering the original granulometric signals. This has also been confirmed by microscopic observations,
as shown in (Baccolo et al., 2021). Considering all these elements, it is likely that chemical weathering also affects dust concentration, explaining the decrease of correlation with ice isotopic composition in the deep part of the ice core (Fig. 2).

Jarosite formation is not the only dust-related geochemical anomaly process occurring in deep TALDICE. Another one anomaly regards is the oxidation of Fe present in dust (Fig. 4). The oxidation of Fe-minerals under acidic conditions is a well-known weathering pathway (Jones et al., 2014) which, among the others, leads to jarosite precipitation (Papike et al., 2006). Fe oxidation is also confirmed by the decline of many ferrous minerals in the deep part of the core (This is the case for hornblende, muscovite, siderite and pyrite; see Fig. 5 and 6 and Tab. 3). It is worth highlighting that the oxidation of pyrite, a process already known to occur in ice (Raiswell et al., 2018), supplies additional acidity to the englacial environment, further favoring jarosite precipitation. The disappearance of hematite in deep TALDICE, which is not stable under acidic and wet conditions (Schwertmann and Murad, 1983), further supports oxidative/acidic weathering in deep ice at Talos Dome. The weathering index (Fig. 6i) summarizes these mineralogical changes and highlights the progressive weathering of dust with depth.

The trends of An index was developed to summarize evidence from Fe-mineralogy: it is defined as the % ratio between the rel. ab. Of jarosite and the sum between rel. ab. of jarosite, hornblende, muscovite and hematite; it is shown in Figure 6i. The index varies between 0 % in the upper part of TALDICE (pristine mineral assemblage) and rapidly increases to 100 % between 1000 and 1300 m, reflecting dust weathering.

The trend of some elements major element oxides in TALDICE dust in the deep part of TALDICE (Tab. 2, Fig. 3) agrees with the scenario described above. The decrease of Ca, Na and Mg with depth is interpreted as an effect of acidic weathering. In presence of acidic aqueous solutions, mobile and soluble elements are easily mobilized (Nesbitt and Young, 1982). It is worth mentioning that Al oxide is also affected-depleted in deep by depletion in deep TALDICE dust. This suggests that not only the labile and soluble fraction of dust undergoes weathering but also the most stable fractions of dust such as aluminosilicates undergo weathering. This is confirmed by the identification in the deep part of the Dome Fuji ice core-Antarctic ice of secondary aluminum-sulphate (Ohno et al., 2014). To further investigate such aspects, a comparison between soluble and insoluble impurities in deep TALDICE would be desirable. Currently data about ionic records from TALDICE are available, but they concern the upper part of the core, not the deep one (Schüpbach et al., 2013; Mezgec et al., 2017).

TALDICE presents a number of peculiarities if compared to other East Antarctic ice cores: deposition of local dust presenting a basaltic signature (Baccolo et al., 2018a), relatively warm temperatures, and a strong oceanic influence. They Such features can partially explain why post-depositional processes affecting dust are so notable at this site. Basaltic-doleritic Domesust deposited at Talos Dome has a local basaltic doleritic signature (Baccolo et al., 2018a) and basalts are easily weatherable rocks, also even at low temperature (Li et al., 2016; Niles et al., 2017). In addition, Talos Dome is located
near the Southern Ocean and receives a considerable amount of marine aerosol rich in reactive acidic species (Iizuka et al., 2013; Mezgec et al., 2017). A further factor to consider is ice temperature. Talos Dome is located near the coast and its climate is tempered by the ocean. When considering the same depth, ice temperature at Talos Dome is about 15-20 °C warmer than at inner sites (Talalay et al., 2020) and temperature is an essential parameter for the relocation of impurities in deep ice (Marath and Wettlaufer, 2020).

All these local features are likely related to the ice-metamorphism observed in deep TALDICE (Montagnat et al., 2012), where ice crystals up to 40-50 cm have been observed below 1480 m deep. High ice-temperature and metamorphism could partially explain why dust alteration is so relevant in TALDICE, while at inner sites the original properties of dust seem preserved at greater depth and further back in time (Delmonte et al., 2004; Kawamura et al., 2017). Replicating this study at Inner Antarctic sites will be essential to distinguish the processes depending on the local characteristics of single sites from the ones more deeply related to ice depth and age.

**3.3.6. Conclusions and perspectives**

This study provides a first description of dust chemical weathering in deep polar ice. Dust grain size, concentration, mineralogy and composition of dust are all affected by post-depositional processes in TALDICE. Fe speciation and mineralogy investigated through synchrotron radiation are efficient probes to explore such transformations in Antarctic ice. The englacial precipitation of jarosite, the oxidation of Fe in dust, the concurrent decline of ferrous minerals (hornblende, pyrite, siderite, muscovite) and of hematite, and the depletion of some major elements (Ca, Mg, Na) suggest that below 1000 m deep, dust in TALDICE is affected by acidic-oxidative weathering. The latter results from the formation of acidic brines which the interaction of dust with dust in highly saline and acidic brines deep ice, leading to geochemical reactions. The production of such brines is likely related to ice re-crystallization and to the accumulation of impurities in highly localized environments where the interaction between soluble and insoluble species is favored. From this perspective deep Antarctic ice can be seen as a "geochemical reactor", capable of promoting the precipitation of secondary minerals and the concurrent dissolution of others.

These findings pose issues to the interpretation of future deep Antarctic ice cores. Because of diffusive processes, the isotopic signal of ice, as well as other proxies (Barnes et al., 2003), is progressively deteriorated with depth (Jones et al., 2017), making difficult to isolate the climatic content of deep ice cores. Being considered relatively immobile and stable, dust has been regarded as a candidate to overcome these difficulties. This study demonstrates shows that also dust-related signals in ice cores, traditionally considered stable and resistant to post-depositional processes, are significantly touched altered in deep ice, at least in TALDICE by post-depositional transformations which need to be addressed.
It would be desirable to replicate the present study considering deep ice cores from inner East Antarctica where it would be possible to unveil additional processes related to the conditions found at 2000-3000 m ice deep in the ice. Another future possible implementation will is be the concurrent analysis of soluble and insoluble impurities, including elements other than Fe. This will help to investigate the dissolution and precipitation of primary and secondary minerals and identify the geochemical reactions behind responsible for these transformations, paving the way for the development of englacial geochemistry.

Data availability. Data used to the aims of the present study are available in the Supplementary Material. Full XANES spectra are found in the PANGAEA open repository, DOI: 10.1594/PANGAEA.924114

Author contributions. GB conceived the idea of this work. GB, BD, EDS prepared the samples and performed Coulter counter analyses. GB, GC, DH, AM carried out X-ray absorption and fluorescence analyses. GB interpreted the data and wrote the manuscript with contributions from all the coauthors.

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