Short-Term Meteorological and Environmental Signals Recorded in a Firn Core from a High-Accumulation Site on Plateau Laclavere, Antarctic Peninsula

Kirstin Hoffmann-Abdi 1,2,* , Francisco Fernandoy 3, Hanno Meyer 1, Johannes Freitag 4, Thomas Opel 1, Joseph R. McConnell 5 and Christoph Schneider 2

1 Alfred Wegener Institute, Helmholtz Centre for Polar and Marine Research, Research Unit Potsdam, Telegrafenberg A45, 14473 Potsdam, Germany; Hanno.Meyer@awi.de (H.M.); Thomas.Opel@awi.de (T.O.)
2 Geographisches Institut, Humboldt-Universität zu Berlin, Unter den Linden 6, 10099 Berlin, Germany; christoph.schneider@geo.hu-berlin.de
3 Facultad de Ingeniería, Universidad Andrés Bello, Viña del Mar 2531015, Chile; francisco.fernandoy@unab.cl
4 Alfred Wegener Institute, Helmholtz Centre for Polar and Marine Research, Am Alten Hafen 26, 27568 Bremerhaven, Germany; Johannes.Freitag@awi.de
5 Division of Hydrologic Sciences, Desert Research Institute, 2215 Raggio Parkway, Reno, NV 89512, USA; Joe.McConnell@dri.edu
* Correspondence: Kirstin.Hoffmann@awi.de

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Abstract: High-accumulation sites are crucial for understanding the patterns and mechanisms of climate and environmental change in Antarctica since they allow gaining high-resolution proxy records from firn and ice. Here, we present new glacio- and isotope-geochemical data at sub-annual resolution from a firn core retrieved from an ice cap on Plateau Laclavere (LCL), northern Antarctic Peninsula, covering the period 2012–2015. The signals of two volcanic eruptions and two forest fire events in South America could be identified in the non-sea-salt sulphur and black carbon records, respectively. Mean annual snow accumulation on LCL amounts to 2500 kg m−2 a−1 and exhibits low inter-annual variability. Time series of δ18O, δD and δ13C excess show no seasonal cyclicity, which may result from (1) a reduced annual temperature amplitude due to the maritime climate and (2) post-depositional processes. The firn core stratigraphy indicates strong surface melt on LCL during austral summers 2013 and 2015, likely related to large-scale warm-air advection from lower latitudes and temporal variations in sea ice extent in the Bellingshausen-Amundsen Sea. The LCL ice cap is a highly valuable natural archive since it captures regional meteorological and environmental signals as well as their connection to the South American continent.

Keywords: Antarctic Peninsula; firn cores; stable water isotopes; glacio-chemistry; high accumulation; surface melt

1. Introduction

The Antarctic Peninsula (AP) has long been considered as one of the “hot spots” of global climate warming since near-surface air temperatures on the AP have increased by more than 3 °C since the 1950s [1–3]. The rapid atmospheric warming of the AP during the 20th century has been accompanied by a significant increase in precipitation and snow accumulation [4–6]. In addition, oceanic warming has caused the destabilization and disintegration of ice shelves in the AP region and consequently, the acceleration of glacier discharge towards the ocean contributing to global sea level rise [7–11]. The most recent 100-year warming trend on the AP is unusual, but not unprecedented in the context of natural climate variability over the past two millennia [12,13]. The observed atmospheric and oceanic changes in the AP region have mainly been attributed to phase-changes of...
the Southern Annular Mode (SAM), which is the dominant mode of atmospheric variability in Southern Hemisphere high latitudes [14–17]. Since the mid-1970s, the SAM has shifted towards its positive phase, especially in austral summer and autumn [15,18]. This has been associated with a strengthening of the circumpolar vortex and, hence, an intensification (15–20%) and poleward migration of the westerlies over the Southern Ocean [14,15,19,20]. For the northern AP region, this implies a reduced cold-air advection from the south, but an enhanced transport of warm and moist air from the Southern Ocean towards its western coast [3,18,20].

Furthermore, sea ice extent in the Bellingshausen-Amundsen Sea at the western side of the AP has decreased throughout all seasons during recent decades, contrasting the generally positive Antarctic sea ice trends [21]. Sea ice variations in the Bellingshausen-Amundsen Sea are primarily driven by changes in the position and strength of the Amundsen Sea Low (ASL), which governs the direction and strength of the meridional flow over this sector [22]. Sea ice in the Weddell Sea on the eastern side of the AP shows seasonally different trends and its temporal variability is related to changes in the strength of the Weddell Gyre [21,23]. Both the depth of the ASL and the strength of the Weddell Gyre, and hence sea ice variability in the AP region, are influenced by phase-changes of large-scale climate modes, i.e., SAM and the El Niño Southern Oscillation [22–25].

Recent studies have drawn a new picture of climate change in the AP region, suggesting little or no air temperature change since the late 1990s [26,27]. The reversed air temperature trend has been linked to the increased advection of cold air by easterly to south-easterly winds because of greater cyclonic activity in the northern Weddell Sea [26,28].

In order to understand the recent climate variability of the AP region, long-term high-resolution data such as air temperature, precipitation and accumulation rates are essential. However, their continuous observation only started in the second half of the 20th century, making the assessment of regional changes on different timescales difficult [2,29]. Proxy data, such as glacio-chemical data from firm and ice cores, may partly compensate for the lack of direct observations. The northern AP is a suitable region for such studies due to its very high snow accumulation rates of up to >3000 kg m⁻² a⁻¹, which allow the preservation of climate and environmental signals on sub-annual timescales [30,31]. Previous studies of Fernandoy et al. [31,32] have demonstrated that firm cores from the ice cap on Plateau Laclavere (LCL) can provide reliable climate proxy data. LCL was explored for the first time in austral summer 2010. Since then, more than ten firm cores of up to 20 m depth have been obtained from different altitudes and locations (Figure 1). In addition, more than 300 precipitation samples have been collected at Bernardo O’Higgins station (OH) since 2008 in order to study the relation between near-surface air temperatures at OH and LCL, stable water isotope signals in LCL firm cores and precipitation reaching the northern AP. Fernandoy et al. [31] found a seasonally varying relationship between δ¹⁸O of precipitation and near-surface air temperatures at OH station for the period 2008–2014 and linked this observation to the formation of an inversion layer in the lower troposphere on the western side of the northern AP during months with sea ice coverage. Variations in the duration and extent of the sea ice cover also leads to seasonally varying lapse rates, which should be considered when comparing stable water isotope records from LCL with near-surface air temperatures [31]. A seasonal cyclicity was most clearly pronounced in the firm core δ excess records during 2008–2009, but no seasonality was found in δ¹⁸O or δD, which both show no correlation with air temperatures at the nearby Bellingshausen (BH) and OH stations. Based on the calculation of backward trajectories Fernandoy et al. [31,32] identified the Bellingshausen Sea (south of 60° S) and the South Pacific Ocean (50°–60° S) as the main moisture sources for precipitating air masses reaching LCL.
Here, we present high-resolution stratigraphic, glacio-chemical and stable water isotope data from a firn core drilled on Plateau Laclavere at the northern tip of the AP. In addition, precipitation samples collected at Bernardo O’Higgins station in 2015–2017 are
analysed for their stable water isotope composition extending the previously existing precipitation dataset for the site derived by Fernandoy et al. [30,31]. Since LCL is a high-accumulation site, our aim is to analyse whether short-term changes in meteorological conditions (e.g., warm-air events) are recorded by the firn core. We also investigate its potential to preserve southern hemispheric environmental signals (e.g., volcanic eruptions, forest fires) with a focus on the connection to the South American continent. The strong influence of the adjacent oceans (mainly the Bellingshausen Sea; to a lesser extent, the Weddell Sea) on the climate of the northern AP and the likely impact of post-depositional processes (wind drift, surface melt) on the proxy signals preserved in the firn core are assessed. The study provides an important basis for further paleoclimatic studies in the high-accumulation region of the northern AP based on the analysis of glacio-chemical firn core records.

2. Materials and Methods

2.1. Fieldwork and Laboratory Analysis

The firn core OH-12 was retrieved within the frame of a Chilean-German field campaign conducted on LCL in January 2016. LCL is located on the northernmost end of the AP at an altitude of about 1100 m above sea level (a.s.l.) and about 19 km south-east of OH station (12 m a.s.l.; Figure 1). It is about 3 km wide and 6 km long and has a flat surface. It is located at the divide between the warmer and moister western side and the colder and drier eastern side of the AP [32]. The ice cap reaches a maximum ice thickness of about 350 m [33]. The mean annual air temperature at the site is about −7.5 °C (2009–2014) [31].

The firn core OH-12 was drilled at about 1090 m altitude down to 19.93 m depth using a portable solar-powered and electrically operated ice-core drill (Backpack Drill; icedrill.ch AG; Figure 1). It was first stored at the Chilean Antarctic Station “Profesor Julio Escudero” on King George Island, South Shetland Islands, and then shipped to the Alfred Wegener Institute, Helmholtz Centre for Polar and Marine Research (AWI) in Bremerhaven, Germany. At the ice-core processing facilities of AWI Bremerhaven, the core was visually inspected for ice layers and lenses and their thicknesses were documented. Furthermore, high-resolution (<1 mm) density measurements were performed using X-ray microfocus computer tomography (ICE-CT; [34]). The core was then sampled at 0.05 m resolution (n = 414) and analysed for stable water isotopes at the ISOLAB Stable Isotope Facility of AWI Potsdam, Germany. Stable water isotopes were measured with cavity ring-down spectrometers L2130-i and L2140-i (Picarro Inc.) coupled to an auto-sampler (L2130-i: PAL HTC-xt, CTC Analytics AG; L2140-i: Picarro Autosampler, Picarro Inc.) using six repeated injections, from which the first three were discarded. The stable water isotope raw data were corrected for linear drift and memory effects following the procedures suggested by van Geldern and Barth [35]. Linear regression analysis was applied to calibrate the drift- and memory-corrected stable water isotope data using four different in-house standards that have been calibrated to the international VSMOW2 (Vienna Standard Mean Ocean Water)/SLAP2 (Standard Light Antarctic Precipitation) scale. Stable water isotope ratios are reported in per mil (‰) versus VSMOW2. Precision of the measurements is ±0.08‰ for δ18O and ±0.5‰ for δD (number of measured standards: 540).

In addition, analysis of stable water isotopes applying the described procedure was performed for precipitation samples collected at OH station between December 2015 and December 2017 (n = 133). Precipitation sampling took place according to the method described by Fernandoy et al. [31]. However, a significant number of the collected samples (n = 50) apparently have been isotopically modified, most likely during storage and transport, as they exhibited exceptional and unrealistic d excess values (≤ −10‰ down to −149‰). Hence, these samples were excluded from further data analysis and interpretation [31]. The considered data (n = 83) were then used to further constrain the Local Meteoric Water Line (LMWL) which has previously been derived by Fernandoy et al. [31,32] for the study site using stable water isotope data of recent precipitation collected at OH
station from January 2008 to March 2009 ($n = 139$) and from March to November 2014 ($n = 72$).

Furthermore, OH-12 was analysed in very high resolution (better than 10 mm) for hydrogen peroxide (H$_2$O$_2$), black carbon (BC) and various chemical elements at the Trace Chemistry Laboratory of the Desert Research Institute (DRI) in Reno, Nevada, USA. Measurements were conducted according to methods described by Röthlisberger et al. [36] and McConnell et al. [37]. For the analyses, longitudinal subsections of the core (32 mm x 32 mm in size) were melted continuously on a chemically inert, ultra-clean melt head (Continuous Flow Analysis [CFA]; [37]). Chemical elements were measured using two Thermo Finnigan Element2 High Resolution-Inductively Coupled Plasma-Mass Spectrometry (HR-ICP-MS) instruments. BC was analysed with a Single Particle Soot Photometer intercavity laser system (SP2, Droplet Measurement Technologies). In addition, values of sea-salt sodium (ssNa) and non-sea-salt sulphur (nssS) and the nssS/ssNa ratio were calculated from Na and S concentrations following the approaches of Röthlisberger et al. [38] and Sigl et al. [39], respectively, and using relative abundances from Bowen [40] (Equations (S1) and (S2) in the Supplementary Materials). While BC is used as a proxy for biomass burning and industrial pollution [41,42], the combined analysis of the nssS and nssS/ssNa records can provide information about the occurrence of volcanic eruptions [43,44].

Corrected and calibrated δ$^{18}$O and δD values were used to calculate the d excess (d excess = δD−8×δ$^{18}$O; [45]) and to derive the δ$^{18}$O–δD relationship (δD = m × δ$^{18}$O + n) for OH-12. The d excess is a proxy for the conditions in the moisture source region, where lower sea surface temperatures and higher relative humidity would induce lower d excess values and vice versa [46–48]. The δ$^{18}$O–δD relationship of OH-12 was compared to the Global Meteoric Water Line (GMWL) defined by Craig [49], and the site-specific LMWL derived from event-based precipitation samples as described above. Finally, high-resolution profiles of density, δ$^{18}$O, δD and d excess as well as of all glacio-chemical parameters were generated.

2.2. Age Model Construction and Time Series Analysis

OH-12 has been dated using annual layer counting (ALC) of high-resolution H$_2$O$_2$ measurements. H$_2$O$_2$ is photochemically produced in the atmosphere under the influence of solar UV radiation and water vapor, and therefore it reflects the annual insolation and air temperature cycles [50,51]. It is washed out of the atmosphere by precipitation and hence reaches the Earth’s surface by wet deposition. Thus, H$_2$O$_2$ typically exhibits clear seasonal cycles following cyclic changes in solar radiation, making it a suitable parameter for dating [52–54].

Using the derived age model, time series of ice layer thickness and stable water isotopes of OH-12 were constructed. Based on the high-resolution density record, accumulation rates were calculated for the study site. The records of OH-12 stratigraphy and stable water isotope composition were then inter-compared as well as related to time series of meteorological parameters and sea ice extent (SIE) in the adjacent Bellingshausen-Amundsen and Weddell Seas.

Unfortunately, there are no available in situ measurements of meteorological parameters from LCL. The automatic weather stations (AWS) nearest to LCL are located at OH and BH stations and provide meteorological records at daily resolution from 1973 (BH) and 1985 (OH) onwards. They are available from the Global Surface Summary of the Day (GSOD) of the National Centers for Environmental Information (NCEI; available at: https://www.ncei.noaa.gov; last access: 17 June 2021). Monthly means of AWS daily data are calculated only when at least 90% of the data are present in the respective month. However, both datasets contain considerable gaps. Therefore, we use data from the European Centre for Medium-Range Weather Forecasts (ECMWF) ERA5 Reanalysis (1979 onwards, spatial resolution: 31 km, temporal resolution: 1 h; available at: https://www.ecmwf.int/en/forecasts/datasets/reanalysis-datasets/era5; last access: 16
April 2021) [55]. Monthly ERA5 reanalysis data were extracted from the grid points closest to OH and BH stations as well as closest to the OH-12 drill site on LCL (Figure 1) through the KNMI Climate Explorer (available at: https://climexp.knmi.nl/; last access: 16 April 2021) [56]. Coordinates and altitudes of the selected ERA5 grid points and the corresponding sites are given in Supplementary Table S1. For the analysis of OH-12 time series, we focus on the following meteorological parameters from the closest ERA5 grid point: precipitation, near-surface and 850 mbar air temperatures, zonal and meridional components of the 850 mbar wind. From the two wind components at 850 mbar geopotential height, which roughly corresponds to the altitude of LCL, we calculated wind directions and derived wind roses for the study site (Equation (53)).

Data on SIE in the Bellingshausen-Amundsen and Weddell Seas, defined as the sections of the Southern Ocean extending from 130° W–60° W and 60° W–20° E, respectively, were obtained from the National Aeronautics and Space Administration (NASA; available at: https://earth.gsfc.nasa.gov/cryo/data/arcticantarctic-sea-ice-time-series; last access: 25 April 2021) [57,58].

Cross-correlation analyses between OH-12 and ERA5 time series and regional SIE were performed using monthly means. Pearson correlation coefficients (r) and p-values (p) were calculated at the 95% confidence level (α = 0.05).

3. Results
3.1. Firn Core Age Model

Due to the generally high snow accumulation on LCL, the H2O2 record is well preserved in the OH-12 firn core, showing clear seasonal cycles in the upper 16 m, albeit a pronounced double peak structure (Figure 2). The drill date of OH-12 (14 January 2016) was used as first tie point for deriving the age model. H2O2 is positively related to accumulation rates and inversely related to temperature, i.e., low accumulation rates and/or higher air and thus snow temperatures favor the degassing of H2O2 from the snowpack [50,54,59]. Therefore, the minima in summer might be caused by missing snow precipitation and/or higher temperatures. In addition, surface melt can lead to the redistribution of H2O2 in the snowpack causing a staircase shape of the peaks, which however is not the case in OH-12 above 16 m depth. Since at the time of the drilling of OH-12 (mid-January 2016), the uppermost double peak had already formed, we set 1 January to the first maximum of the H2O2 double peak (from the top) for each year identified between 0 m and 16 m depth. Below 16 m depth, the H2O2 record seems to be altered by melt water percolation leading to a staircase shape of the peak and hence an unclear seasonal cyclicity (Figure 2). The firn core stratigraphy exhibiting several ice layers in this part of the core may provide evidence for surface melt and percolation events (see Section 3.2, Figure 4a). Hence, 1 January was set to the first peak (from the top) that is higher than the mean value of the entire record period (1.96 μM) at about 18.2 m depth (Figure 2). Below that depth, the age scale was derived by linear extrapolation using the depth–age relationship between the previous two clearly identifiable peaks (years 2012 and 2013). In total, OH-12 covers four full years (2012–2015; Figure 2). The estimated dating uncertainty is ±2 months above 16 m depth and increases to ±10 months in the bottom part of the core due to the percolation biased H2O2 record.
To improve the ALC-based age model, we accounted for precipitation intermittency at the study site using ERA5 reanalysis data, since environmental signals are only recorded in the snow when there is a precipitation event [60–62]. First, we tested, how well ERA5 reanalysis data capture the seasonal variability of precipitation in the northern AP region. For this, we compared the ERA5 and AWS precipitation records from BH and OH stations on a monthly scale as well as based on the seasonal cycle calculated using the respective overlapping period between both datasets (AWS/ERA5 OH: 1985–2020; AWS/ERA5 BH: 1979–2020). For both stations, correlations are statistically significant and quite high for monthly means (r = 0.68 for BH, r = 0.71 for OH with p < 0.05) as well as for the seasonal cycle (r = 0.59 for BH, r = 0.54 for OH with p = 0). Hence, we conclude that ERA5 reanalysis data reliably reflect the seasonal variability of synoptic precipitation in the northern AP region. However, the influence of the orography on the annual precipitation distribution which is particularly important on LCL is not fully captured by ERA5. Furthermore, the ERA5 precipitation record comprises both solid and liquid precipitation, with the latter not contributing to accumulation on LCL. Finally, wind-driven removal and relocation of freshly fallen snow can partially alter the annual accumulation distribution at the site. Despite these biases, the age model of OH-12 can be improved by considering the annual precipitation distribution from the ERA5 grid point closest to the firm core drill site to best represent the annual accumulation distribution on LCL. Based on this, we calculated for each single year covered by OH-12 the percentage of each month in annual precipitation. Subsequently, the thickness of each annual layer was calculated and multiplied by the monthly percentage precipitation shares in the respective year. Hence, for each year, we obtained the monthly layer thicknesses and thus could derive a depth–age relationship. The age of firm core samples within each monthly layer could then be computed by linear interpolation. The precipitation-based weighted age scale is slightly shifted compared to the ALC-based age scale depending on the core section (maximum offset: 33 days; Figure 3). In the following, we use the precipitation-based weighted age scale for the construction of OH-12 time series and subsequent statistical analyses.
3.2. Firn Core Stratigraphy and Accumulation Rates

The high-resolution density–depth profile of OH-12 (Figure 4a) shows a clear increase in density with depth reflecting snow compaction processes and the metamorphosis from snow to firn. Linear regression across the entire density–depth profile reveals the snow–firn density transition (550 kg m⁻³) at about 8.4 m depth. Furthermore, firn core OH-12 is characterised by the presence of numerous ice lenses and layers with thicknesses varying between 0.6 mm and 40 mm. This is also visible in the high-resolution density–depth profile that exhibits distinct maxima (≥800 kg m⁻³) with values up to the density of pure ice (917 kg m⁻³) in the respective core sections (Figure 4a). More than 75% of all ice lenses and layers have a width of <10 mm. Most of them are probably not related to surface melt, but instead are formed by wind ablation and wind scouring processes or infiltration, percolation and refreezing of liquid precipitation. In addition, hoar frost formation on the snow surface also plays a role. Hoar frost layers in OH-12 are between 2 mm and 25 mm thick. In order to analyse the firn core stratigraphy for the occurrence of surface melt events, we only consider ice layers most likely indicating melt, i.e., near-horizontal ice layers with a continuous lateral extension and little or no air bubbles [63,64]. They vary between 1 mm and 40 mm in thickness with a mean value of 8.5 mm (Figure 4b). The detailed thickness distribution of the melt layers is shown in Supplementary Figure S1. Most prominent melt layers are found at 3.8 m (39 mm), 4.9 m (35 mm) and 13.5 m depth (40 mm) corresponding to April/May 2015, February 2015 and April 2013, respectively (Figure 4a,b).

Based on the identified melt layers and the weighted age scale, the total melt layer thickness (MLT) was calculated for each month within the period covered by OH-12 (Figure 4b), revealing a clear seasonality with MLT maxima occurring during summer and MLT minima during winter months. The years 2013 and 2015 show the highest MLT values reaching up to 83 mm and 104 mm in austral late summers 2013 and 2015, indicating strong surface melt processes at the OH-12 drill site (Figure 4b). Based on the high-resolution density measurements of OH-12, snow accumulation at the study site was calculated for each full year identified in the core (Table 1). On average, snow accumulation amounts to about 2500 kg m⁻² a⁻¹ with only little variations for the period 2012–2015 (on average ±8%; Table 1). The maximum accumulation rate is observed in 2013 (2890 kg m⁻² a⁻¹), whereas the minimum value is reached in 2015 (2260 kg m⁻² a⁻¹).
Figure 4. (a) Ice layer stratigraphy (blue lines) and high-resolution density–depth profile (grey; 1500-point running mean in red) of firn core OH-12. (b) Total monthly melt layer thickness (MLT; bold blue line) for the period covered by OH-12 calculated from ice layers indicating surface melt (light blue lines; details in the text) using the weighted age scale.

Table 1. Annual accumulation rates calculated for the OH-12 drill site for the period 2012–2015.

<table>
<thead>
<tr>
<th>Year</th>
<th>Accumulation Rate (kg m$^{-2}$ a$^{-1}$)</th>
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<tbody>
<tr>
<td>2012</td>
<td>2260</td>
</tr>
<tr>
<td>2013</td>
<td>2470</td>
</tr>
<tr>
<td>2014</td>
<td>2890</td>
</tr>
<tr>
<td>2015</td>
<td>2390</td>
</tr>
<tr>
<td>2012–2015</td>
<td>2500</td>
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3.3. Firn Core Records of Black Carbon (BC), Non-Sea-Salt Sulphur (nssS) and the Non-Sea-Salt Sulphur—Sea-Salt Sodium Ratio (nssS/ssNa)

The high-resolution depth profiles of BC, nssS and nssS/ssNa exhibit distinct maxima at certain depths (Figure 5). BC reaches maximum values of 11.6 ng g$^{-1}$ and 12.7 ng g$^{-1}$ at 3.3 m (May/June 2015) and 16.5 m depth (June/July 2012), which are more than 20 times higher compared to the mean value of 0.5 ng g$^{-1}$ (Figure 5a). The combined analysis of the nssS and nssS/ssNa records reveals three remarkable maxima compared to the mean values of 18.6 ng g$^{-1}$ and 0.06, respectively (calculated excluding negative values): a single peak at about 3.3 m depth (May/June 2015; nssS: 72.2 ng g$^{-1}$; nssS/ssNa: 1.2) that coincides with the first BC maximum (from the top), a double peak between 9.3–10.1 m depth (November 2013–March 2014; maximum nssS: 71.2 ng g$^{-1}$; maximum nssS/ssNa: 0.95) and a long-stretched peak between 16.9–18.5 m depth (November 2011–May 2012; maximum nssS: 86.9 ng g$^{-1}$; maximum nssS/ssNa: 0.4; Figure 5b). The maxima in the BC, nssS and nssS/ssNa records point towards cross-regional environmental events which left an imprint on the LCL ice cap and can thus be used as independent tie points for confirming the weighted age scale of OH-12 (see Section 4.1).
Figure 5. High-resolution depth profiles of (a) BC (black), (b) non-sea-salt sulphur (nssS; green) and the non-sea-salt sulphur–sea-salt sodium ratio (nssS/ssNa; orange) of firn core OH-12. Southern hemispheric events assigned to the remarkable maxima identified in the BC, nssS and nssS/ssNa records are indicated (for details, see Section 4.1).

3.4. Firn Core and Precipitation Stable Water Isotopes

In general, within each collection period (Table 2), the precipitation samples are well distributed over the recorded years, i.e., there is more than one sample for each month (Supplementary Figure S2). Altogether, the three sampling periods cover 49 months, with each month sampled four times except March, which was sampled five times. However, as described in Section 2.1 and by Fernandoy et al. [31]. Several samples had to be excluded from the statistical analysis of stable water isotope data (Supplementary Figure S2). Accordingly, Figure 6a shows the stable water isotope composition of all considered precipitation samples collected at OH station between 2008 and 2017 in comparison to the stable water isotope composition of OH-12. The stable water isotope composition of the 2015–2017 precipitation samples is very similar to that of the previously collected samples although covering a wider range (from −22.1‰ to −0.9‰ for δ¹⁸O and from −181.6‰ to −11.8‰ for δD; Table 2). The 2015–2017 samples have mean δ¹⁸O and δD values of −9.2‰ and −74.5‰, respectively, and a mean δD of −1.0‰. Mean δ¹⁸O, δD and δD values of all precipitation samples collected between 2008 and 2017 (n = 294) amount to −9.4‰, −74.4‰ and 0.8‰, respectively. Based on all precipitation samples, a better constrained LMWL has been calculated for the study site: δD = 7.76 × δ¹⁸O − 1.52 (R² = 0.97, p = 0; Figure 6a).
Table 2. Basic statistics of the stable water isotope composition of precipitation samples collected at Bernardo O’Higgins station from January 2008 to March 2009, from March to November 2014 and from December 2015 to December 2017. For each sampling period the number of samples (n) considered for the statistical analysis together with the total number of collected samples (in brackets) is given.

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<tbody>
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<td>n (samples)</td>
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<td>72 (94)</td>
<td>83 (133)</td>
<td>294 (366)</td>
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<tr>
<td>Stable water isotope composition</td>
<td>( \delta^{18}O ) (‰)</td>
<td>( \delta D ) (‰)</td>
<td>( d ) excess (‰)</td>
<td>( \delta^{18}O ) (‰)</td>
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<td>26.4</td>
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</table>

Figure 6. (a) \( \delta^{18}O–\delta D \) relationship of all considered precipitation samples collected at Bernardo O’Higgins station (OH) between 2008 and 2017 (n = 294; coloured dots) compared to the \( \delta^{18}O–\delta D \) relationship of firm core OH-12 (n = 414; white dots). The Global Meteoric Water Line (GMWL) is indicated in blue. The Local Meteoric Water Line (LMWL) established for the study site by Fernandoy et al. [31,32] is shown as a dashed red line and the LMWL derived in this study as a solid red line. For each \( \delta^{18}O–\delta D \) relationship, the equation, the coefficient of determination \((R^2)\) and the \( p \)-value \((p) \) are given. (b) Time series of \( \delta^{18}O \), \( \delta D \) and \( d \) excess of OH-12 constructed based on the weighted age scale. High-resolution data are shown as light-coloured lines and monthly means as bold lines.

Stable oxygen and hydrogen isotopes of OH-12 are highly correlated \((R^2 = 0.99, p = 0; \text{Figure 6a})\). The slope of the \( \delta^{18}O–\delta D \) relationship (7.94) is close to that of the Global Meteoric Water Line (GMWL) [49] and is of the same order of magnitude as the slope of the site-specific LMWL \((m = 7.76)\). However, intercepts differ significantly (OH-12: -6.01; LMWL: -1.52; GMWL: +10), which is also reflected by the position of the OH-12 samples in the \( \delta^{18}O–\delta D \) plot (Figure 6a). All firm core samples are located above the LMWL and mostly below the GMWL, implying higher \( d \) excess values compared to the event-based precipitation samples from OH station.

\( \delta^{18}O \) and \( \delta D \) values show a high sub-annual variability ranging between \(-7.0\%o\) and \(-19.4\%o\) and between \(-49.3\%o\) and \(-148.0\%o\), respectively (Table 3). Mean \( \delta^{18}O \) and \( \delta D \) values for the period covered by OH-12 are \(-12.4\%o\) and \(-92.0\%o\), respectively. \( d \) excess values vary between \(1.4\%o\) and \(15.2\%o\) with a mean value of \(6.8\%o\) (Figure 6b and Table 3).
Inter-annual differences in the stable water isotope composition of OH-12 (Table 4) are less pronounced than those on the sub-annual timescale. However, the year 2015, which contains thick ice layers, exhibits slightly higher mean annual δ values than the other years. In contrast, this is not the case for the year 2013, despite the presence of exceptionally thick ice layers. Differences in the mean annual d excess are negligible between the four years.

Table 3. Basic statistics of the stable water isotope composition of firm core OH-12 (n = 414).

<table>
<thead>
<tr>
<th></th>
<th>δ¹⁸O (%)</th>
<th>δD (%)</th>
<th>d Excess (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Min</td>
<td>−19.4</td>
<td>−148.0</td>
<td>1.4</td>
</tr>
<tr>
<td>Mean</td>
<td>−12.4</td>
<td>−92.0</td>
<td>6.8</td>
</tr>
<tr>
<td>Max</td>
<td>−7.0</td>
<td>−49.3</td>
<td>15.2</td>
</tr>
<tr>
<td>Sdev</td>
<td>2.4</td>
<td>19.2</td>
<td>2.1</td>
</tr>
</tbody>
</table>

Table 4. Annual minimum, mean and maximum values and standard deviations of stable water isotopes of firm core OH-12 for the period 2012-2015 (January–December) calculated based on the weighted age scale.

<table>
<thead>
<tr>
<th>Year</th>
<th>2012</th>
<th>2013</th>
<th>2014</th>
<th>2015</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>δ¹⁸O</td>
<td>δD</td>
<td>d excess</td>
<td>δ¹⁸O</td>
</tr>
<tr>
<td></td>
<td>(%)</td>
<td>(%)</td>
<td>(%)</td>
<td>(%)</td>
</tr>
<tr>
<td>Min</td>
<td>−16.0</td>
<td>−123.1</td>
<td>3.2</td>
<td>−18.5</td>
</tr>
<tr>
<td>Mean</td>
<td>−12.5</td>
<td>−92.8</td>
<td>6.9</td>
<td>−12.4</td>
</tr>
<tr>
<td>Max</td>
<td>−9.5</td>
<td>−63.6</td>
<td>13.3</td>
<td>−7.2</td>
</tr>
<tr>
<td>Sdev</td>
<td>1.5</td>
<td>13.0</td>
<td>2.1</td>
<td>2.8</td>
</tr>
</tbody>
</table>

3.5. Cross-Correlation Analysis

For cross-correlation analysis between OH-12 time series and ERA5 near-surface air temperature at LCL, we first compared the ERA5 and AWS temperature records from BH and OH stations. For both stations, the records are highly and statistically significantly correlated on a monthly scale as well as with respect to the seasonal cycle calculated as previously described (see Section 3.1; r ≥ 0.9, p < 0.001 for BH and OH). Hence, we consider ERA5 reanalysis data to reliably reflect the regional temperature variability. We accounted for the elevation difference between the OH-12 drill site and the closest ERA5 grid point (Δh = 812 m; Supplementary Table S1) by calculating a monthly lapse rate using the ERA5 records of near-surface and 850 mbar air temperature. Cross-correlations were calculated based on monthly means considering only the period with full-year records for OH-12 (2012–2015; Table 5). Monthly means of OH-12 were computed using the weighted age scale.

δ¹⁸O and δD values of OH-12 are not related to ERA5 near-surface air temperatures at LCL (Table 5). Instead, statistically significant correlations with ERA5 temperatures exist for both d excess and MLT (d excess: r = −0.4, p < 0.01; MLT: r = 0.5, p < 0.0001). Furthermore, δ¹⁸O and δD values of OH-12 are highly inter-correlated (r > 0.99, p = 0), but both show no negative relation to the d excess (Table 5). Monthly MLT is neither correlated with δ¹⁸O or δD, nor with the d excess. Moreover, there is no relationship of δ¹⁸O, δD and d excess with SIE in the close-by Bellinghausen-Amundsen and Weddell Seas for the period 2012–2015 (Table 5). Instead, we found a statistically significant negative correlation of MLT with SIE in both sectors (r = −0.6, p = 0).
Table 5. Results of cross-correlation analysis for monthly records of $\delta^{18}O$, $\delta D$, $d$ excess and total melt layer thickness (MLT) of firn core OH-12 and monthly time series of ERA5 near-surface air temperature at Plateau Laclavere, sea ice extent (SIE) in the Bellingshausen-Amundsen (BA) and Weddell Seas (Weddell). Prominent correlations with a low $p$-value ($p < 0.01$, $\alpha = 0.05$) are marked bold.

<table>
<thead>
<tr>
<th></th>
<th>$\delta^{18}O$</th>
<th>$\delta D$</th>
<th>$d$ excess</th>
<th>$T_{mean}$ ERA5</th>
<th>SIE BA</th>
<th>SIE Weddell</th>
</tr>
</thead>
<tbody>
<tr>
<td>MLT</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$r$</td>
<td>0.07</td>
<td>0.07</td>
<td>0.04</td>
<td>0.53</td>
<td>-0.59</td>
<td>-0.56</td>
</tr>
<tr>
<td>$p$</td>
<td>0.66</td>
<td>0.64</td>
<td>0.77</td>
<td>0.0001</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>$\delta^{18}O$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$r$</td>
<td>1</td>
<td>0.99</td>
<td>-0.03</td>
<td>0.06</td>
<td>0.03</td>
<td>0.02</td>
</tr>
<tr>
<td>$p$</td>
<td>0</td>
<td>0</td>
<td>0.86</td>
<td>0.67</td>
<td>0.85</td>
<td>0.91</td>
</tr>
<tr>
<td>$\delta D$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$r$</td>
<td>1</td>
<td>0.08</td>
<td>0.02</td>
<td>0.05</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>$p$</td>
<td>0</td>
<td>0.59</td>
<td>0.88</td>
<td>0.75</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td>$d$ excess</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$r$</td>
<td>1</td>
<td>-0.38</td>
<td>0.18</td>
<td>0.10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$p$</td>
<td>0</td>
<td>0.01</td>
<td>0.21</td>
<td>0.52</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4. Discussion

4.1. Environmental Signals Recorded in Firn Core OH-12

OH-12 is the first firn core from LCL with available high-resolution glacio-chemical data. Hence, for the first time, it is possible to assess whether and how signals of atmospheric emissions such as from volcanic eruptions, biomass burning and anthropogenic pollution are recorded and preserved in an ice cap on the northern AP. We consider the distinct maxima identified in the records of BC, nssS and nssS/ssNa to be real, since we did not find clear coincidences between enhanced concentrations of BC, nssS and nssS/ssNa and remarkable maxima in the high-resolution depth records of density and ice (melt) layer thickness (Figure 7). However, in case of vertical movement of these chemical species due to meltwater percolation, the impact on the timing of the peaks found in their depth profiles would likely be diminished by the very high annual snow accumulation on LCL.

Figure 7. Comparison of OH-12 high-resolution depth records of (a) black carbon (BC; black), (b) non-sea-salt sulphur (nssS; green) and (c) the non-sea-salt sulphur–sea-salt sodium ratio (nssS/ssNa; orange) with the high-resolution density–
depth profile (grey; 1500-point running mean in bold) and the ice layer stratigraphy (blue). Ice layers indicating surface melt are highlighted in red.

In the nssS and nssS/ssNa records, three distinct maxima were identified, two of which could be assigned to well-known volcanic eruptions in the Southern Hemisphere. The long-stretched peak of medium-high amplitude at about 16.9–18.5 m depth likely reflects the Volcanic Explosivity Index (VEI) 5 eruption of the Puyehue-Cordón Caulle volcano in Southern Chile (June 2011–April 2012) [65]. Koffman et al. [66] found sulphate and ash from this event in surface snow and shallow firm core samples from the West Antarctic Ice Sheet (WAIS) Divide region. According to Koffman et al. [66], the ash cloud of the Puyehue-Cordón Caulle eruption circled the globe between 40° S and 60° S four times from 4 June to 20 July 2011, transporting ash and sulphate primarily with the westerlies rapidly to Antarctica. Hence, it is very likely that sulphate deposition also took place on the LCL ice cap, although the sulphate peak in OH-12 does not occur directly in austral winter 2011, as observed on the WAIS Divide [66], but rather in austral spring and summer 2011/2012. This may be due to the relatively high dating uncertainty in the lower part of the firm core. Different to explosive stratospheric eruptions, the Puyehue-Cordón Caulle eruption did not cause a sustained deposition of aerosols (i.e., over 1–3 years), but transport and deposition of ash and sulphate primarily occurred through the troposphere and were constrained to the -2–3 weeks following the eruption [66]. This also suggests that the nssS and nssS/ssNa maxima at 9.3–10.1 m depth (austral summer 2013/2014) in OH-12 are not related to the Puyehue-Cordón Caulle eruption.

Long-range transport of volcanic aerosols in the Southern Hemisphere also was observed after the Calbuco VEI 4 eruption in Southern Chile in April 2015 [67–69] that resulted in a record size of the ozone hole over Antarctica in October 2015 [70]. The volcanic aerosol plume of Calbuco was mainly advected eastward in the Southern Hemisphere (across northeastern South America and South Africa) latitudinally bounded by the subtropical barrier and the polar vortex, but potentially reached polar latitudes about 4 months after the eruption (60° S) [67,68]. Fernandoy et al. [31] showed that LCL clearly has been influenced by air masses originating over the South American continent during the preceding years 2008–2014. Hence, we infer that the nssS and nssS/ssNa maxima in OH-12 at about 3.3 m depth are connected to the Calbuco eruption. However, further confirmation of the eruption sources will require microphysical investigations of the respective core sections.

Attributing the origin of the exceptionally high BC peaks at about 3.3 m and 16.5 m depth in OH-12 is challenging (Figure 5a). The explosion at the Brazilian Antarctic station Comandante Ferraz (CF), located in Admiralty Bay on King George Island (Figure 1a), on 25 February 2012 could have caused or contributed to the distinct BC maximum at 16.5 m depth. According to Evangelista et al. [71], the 450 m high plume of smoke rapidly dispersed in NE-E direction travelling over King George Island before reaching the Drake Channel and passing the northern tip of the AP. Hence, the Comandante Ferraz fire seems a probable source for the deposition of BC on the LCL ice cap. Furthermore, vast South American forest fires could be a major ash source for the two BC peaks in OH-12. Generally, (tropical) biomass burning is the main source of atmospheric BC in Antarctica [72]. Several studies have shown that emissions from biomass burning in (tropical) South America reached as far as to the northern AP [73] and Dronning Maud Land [74]. Most recently, Jumelet et al. [75] provided evidence that biomass burning events are capable of injecting significant amounts of BC particles up to the lower stratosphere from where they can be transported towards higher latitudes. Hence, the vast forest fires in Torres del Paine in southern Chile, which lasted from the end of December 2011 until March 2012, could have contributed to the BC peak at 16.5 m depth. The BC maximum at 3.3 m depth coincides almost exactly with the nssS and nssS/ssNa peaks of the Calbuco eruption. It has likely been caused by BC emitted from forest fires, which took place in the Argentinian province Chubut (Patagonia) between mid-February and the beginning of April 2015, i.e.,
at the time of the Calbuco eruption. Both forest fires, in Torres del Paine in 2011/12 and in Argentinian Patagonia in 2015, are among the most devastating forest fires documented in these regions [76–78] and hence, are likely to have left an imprint on the LCL ice cap. However, other BC sources than biomass burning such as industrial emissions cannot be ruled out to have contributed to the deposition of BC on the LCL ice cap. Recent model simulations show that emissions from fossil fuels and biofuels account for more than 70% of the BC deposition over James Ross Island in the present day [79].

In summary, the environmental signals found in the glacio-chemical records of OH-12 provide clear evidence for a connection between the northern AP and the South American continent.

4.2. Accumulation Rates

Annual snow accumulation rates (Table 1) derived from OH-12 are in line with previous accumulation estimates made by Fernandoy et al. [32] for the same site for 2008 and 2009. However, for the period 2010–2014, Fernandoy et al. [31] reported remarkably (i.e., up to 1800 kg m\(^{-2}\) a\(^{-1}\)) lower accumulation rates than obtained from OH-12 for the period 2012–2015. Furthermore, the mean snow accumulation rate calculated from OH-12 for 2012–2015 (2500 kg m\(^{-2}\) a\(^{-1}\)) is about 700 kg m\(^{-2}\) a\(^{-1}\) higher than that estimated for 2008–2014 (1770 kg m\(^{-2}\) a\(^{-1}\)) [31]. We propose that the differences in annual snow accumulation between our study and Fernandoy et al.’s [31] originate from the different dating approaches used.

Fernandoy et al. [32] applied ALC to firn core d excess records as they show a clear seasonal cyclicity in 2008–2009. However, from 2010 onwards, a clear seasonal cyclicity has been neither found in the b values nor in the d excess records of LCL firn cores. Therefore, Fernandoy et al. [31] matched measured firn core d excess data to d excess data synthetically derived from meteorological observations by comparing both datasets for similarities in their seasonal variability. Since we consider the H\(_2\)O\(_2\)-based age model of OH-12 to be better constrained, we assume the firn core age model of Fernandoy et al. [31] to have a dating error of +1 year, leading to a substantial underestimation of the annual snow accumulation between 2010 and 2014. We did not find any considerable changes in precipitation amounts during the period 2008–2015 in the AWS and ERA5 precipitation records of BH and OH stations and the OH-12 drill position that could explain the remarkably lower snow accumulation on LCL in the years 2010–2014. In the context of generally high accumulation rates, also no evidence for increased wind drift and thus enhanced removal and relocation of snow at the study site during 2008–2015 has been found in the ERA5 record of 850 mbar wind speed for the OH-12 drill position. In summary, we conclude a constantly high annual snow accumulation of about 2500 kg m\(^{-2}\) a\(^{-1}\) on LCL as derived from OH-12 to be realistic.

4.3. Comparison of Precipitation and Firn Core Stable Water Isotopes

Generally, precipitation reaching LCL exhibits a lighter stable water isotope composition and higher d excess values than precipitation at sea level from OH station (Tables 2 and 3). On average, for 2014—the only year with continuous records for both OH precipitation (12 m a.s.l.) and OH-12 (1090 m a.s.l.)—\(\delta^{18}O\) values of OH-12 are about 2.6% (\(\delta D\) about 13%) lower than those of event-based precipitation (Tables 2 and 3). This is in line with Fernandoy et al. [31] who found a seasonally varying \(\delta^{18}O\)-altitude gradient between OH station and LCL of up to \(-2.4\%\) km\(^{-1}\). D excess values of OH-12 are about 8% higher than those of OH precipitation. Furthermore, stable water isotope records of OH-12 show a lower temporal variability compared to OH precipitation (Supplementary Figure S3). However, in general, there is only little correspondence between both records. The observed differences are likely caused by several processes. First, the orographic effect, i.e., the uplift of air masses from sea level to about 1100 m altitude, facilitates the fallout of the heavier stable water isotopes near the coast and during transport to the plateau (“altitude effect”) [45,80] and hence can partly explain the lower \(\delta^{18}O\) and \(\delta D\) values in the firn core.
Second, post-depositional processes, such as diffusion, wind-pumping, sublimation, melting and refreezing, might alter the stable water isotope composition of the snowpack. Due to the very high accumulation rates at the study site, the exposition time of freshly fallen snow to the atmosphere is limited. Hence, diffusion is considered to play only a minor role, also because of the inhibiting effect of the ice layers (high-density layers) [31,62,81]. The same is principally true for sublimation and condensation at the snow surface [82,83]. However, LCL is often enclosed in clouds and hence, isotopic exchange between the snowpack and the surrounding moist air masses is probable. Condensation of water vapor at the snow surface and in the uppermost firm layers or sublimation of snow/firm into the atmosphere allow for secondary fractionation, i.e., it potentially leads to an enrichment in heavier isotopes and a decrease in the d excess of the snowpack [82,84–86]. Consequently, this may reduce the annual amplitude of the stable water isotope records of OH-12 compared to OH precipitation. Isotopic modifications due to wind-forced firm ventilation (“wind-pumping”) and/or deep-air convection can most likely be excluded due to the high snow accumulation rates. They cause the rapid advection of the snow away from the surface and through the ventilated zone [85,86], and preclude intense firm metamorphism and the formation of large vertical cracks in the firm that both foster deep-air convection [87]. LCL is certainly affected by surface melt reflected by the presence of melt layers in OH-12 in austral summers 2012–2015. However, we did not find a correlation between MLT and stable water isotope records of OH-12 (Table 5) and the melt layers are mostly confined to the summer seasons. Hence, we assume that the stable water isotope composition of OH-12 is not substantially altered by infiltration, percolation and refreezing of surface meltwater throughout the core, but that possible smoothing effects are limited to the melt-affected seasons only.

Due to the exposed position of the study site, we suggest wind drift, i.e., removal and redistribution of snow, to have the largest impact on the stable water isotope composition of snow and firm, at least on sub-seasonal timescales [88,89]. Wind drift certainly leads to spatial mixing of snow with potentially different stable water isotope compositions. First, this may smooth the isotopic signal of the snowpack compared to the isotopic signal of precipitation and lead to the generally low temporal variability of the OH-12 stable water isotope composition [90,91]. Secondly, the temporal correspondence between stable water isotopes in the snowpack on LCL and OH precipitation may be disrupted. Fernando et al. [31] describe the formation of an inversion layer in the lower troposphere (up to 350 m a.s.l.) on the western side of the northern AP during months with sea ice coverage. This may lead to a temporary de-coupling between LCL and OH station, as the stable atmospheric conditions associated with the inversion inhibit the uplift of moist air masses from sea level to LCL at 1100 m a.s.l. Finally, the dating uncertainty of ±2 months in the upper and ±10 months in the lower part of the firm core may also reduce the temporal correspondence between the stable water isotope records of OH-12 and OH precipitation.

4.4. Relation between Firn Core and Meteorological Records

Post-depositional processes are likely one of the major causes for the absence of a correlation between δ18O and δD of OH-12 and ERA5 near-surface air temperatures at the site. Wind drift and the presence of a cloud cover during most of the year may certainly reduce seasonal differences in the stable water isotope composition of the snowpack as previously described (see Section 4.3). Moreover, the strongly maritime character of the northern AP climate and the existence of an inversion layer during months with sea ice coverage leading to seasonally varying lapse rates may reduce the annual amplitude in near-surface air temperatures at the site [31,92]. The annual amplitude of ERA5 near-surface air temperature at BH and OH stations is about 8.2 °C and 11.4 °C (1979–2020), respectively. For the OH-12 drill site, the altitude-corrected ERA5 record of near-surface air temperature reveals an annual amplitude of about 10.3 °C (1979–2020). Such low annual oscillations, typical for coastal sites, are hardly captured in the stable water isotope com-
position of the snowpack on LCL. A weak or absent correlation between stable water isotopes in firm and near-surface air temperatures has been also observed for other coastal Antarctic regions [93,94]. Instead, the finding of a statistically significant negative correlation between firm core δ excess and ERA5 near-surface air temperatures at LCL has been similarly obtained by Fernandoy et al. [31,32] for the study site. Depending on air humidity and wind speed, enhanced sublimation rates at higher air temperatures potentially leading to a decrease in the δ excess as previously described (see Section 4.3) may account for the observed negative correlation. In addition, changes in moisture origin and moisture transport paths, kinetic fractionation during ice-crystal formation at supersaturation as well as changes in equilibrium fractionation rates at low temperatures are suggested as possible explanations [90,95].

Differently to the period 2008–2009 [32], we did not find any seasonal cyclicity in the δ excess record of OH-12 (Figures 6b and 8b). As for δ18O and δD, this may reflect the strong impact of post-depositional processes. Furthermore, the δ excess of evaporated moisture and hence of precipitating air masses transported to a certain site is related to sea surface temperature and relative humidity in the source region [46]. Hence, the absence of a seasonal cycle in the δ excess record of OH-12 may suggest a low inter- and intra-annual variability of the contribution of different local and distal moisture sources to precipitation reaching LCL. In general, the westerly air flow exhibits the strongest influence on the study site as corroborated by wind roses derived for the individual years 2012–2015 as well as for the entire period covered by OH-12 (Supplementary Figure S4). The stable flow of the westerlies (43 out of 48 months in 2012–2015) is rarely interrupted by northerly (2 months) and southerly winds (3 months). Hence, we hypothesise that only if northerly and southerly wind directions predominate, sufficient moisture from distal source regions can reach the study site. In order to further constrain the origin of precipitating air masses reaching LCL, the calculation of backward trajectories would be necessary; that, however, is beyond the scope of this study. Nevertheless, Fernandoy et al. [31,32] identified the Bellingshausen Sea as the most important moisture source for precipitation at LCL based on backward trajectory analysis. This is in line with the dominance of the westerlies at the study site, which preferably foster moisture transport from the adjacent Bellingshausen Sea towards LCL. The dominant influence of local moisture sources on LCL may additionally explain the absence of a seasonal cyclicity in both the δ18O and δD as well as δ excess records of OH-12, since the stable water isotope composition of precipitation reaching a certain location is a function of the temperature difference between the moisture source and the precipitation site [48,96]. Hence, for LCL, the meteorological conditions at both the moisture source and sink may change in parallel due to their proximity, having no effect on the stable water isotope composition of precipitation arriving at the study site.

However, moisture from other sources, e.g., the South Pacific Ocean, the South American and Antarctic continents, can certainly influence the site, reflected by exceptional stable water isotope values [32]. For example, in austral autumn 2015, the high-resolution time series of δ18O and δ excess exhibit their absolute extrema with respect to the period 2012–2015 (absolute maximum in δ18O: ~7.0‰; absolute minimum in δ excess: 1.4‰; Figures 6b and 8a,b). From 18 to 27 March 2015, an exceptionally strong heat event followed by an extreme precipitation event occurred in northern and central Chile, extending south to continental Antarctica. Maximum air temperatures were measured at Esperanza station (EP) at the northern tip of the AP (17.5 °C on 24 March 2015; Figures 1 and 8d) as a result of a widespread foehn event occurring on the leeward side of the AP [92,97]. The foehn event was triggered by an intense northwest–southeast oriented atmospheric river originating over the subtropical Pacific Ocean and reaching the northern AP on 23 March 2015. The synoptic conditions favouring the development of the atmospheric river (a deep low-pressure centre over the Amundsen-Bellingshausen Sea and a northwest–southeast oriented blocking ridge over the southeast Pacific Ocean) prevailed between 21 and 25 March 2015 [92,97]. During this period, exceptionally warm and moist air
from the subtropical Pacific was largely advected towards the northern AP. Maximum near-surface air temperatures were also recorded at windward stations (e.g., Arturo Prat, San Martin and Carlini stations) [92], although not prominently visible at BH and OH stations. Considering the dating uncertainty of ±2 months in the upper part of the core, we conclude that the absolute extrema in the stable water isotope records of OH-12 (Figures 6b and 8a,b) are likely associated with the large-scale advection of warm air towards the northern AP during this event. This is supported by findings of Fernandoy et al. [32] who demonstrated that moisture originating from lower latitudes of the South Pacific Ocean exhibits higher δ18O, but lower d excess values compared to proximal moisture sources in the Bellingshausen-Amundsen Sea (south of 60° S).

The atmospheric river also caused strong surface melt throughout the northern AP [98]. There are two thick melt layers (21 mm and 39 mm; Figures 4b and 8c) present in the OH-12 firn core in austral autumn 2015 which coincide with the δ18O and d excess extrema (Figures 6b and 8a,b). Hence, we hypothesise that these ice layers correspond to surface melt occurring on LCL during the described atmospheric river event.

In general, the stratigraphic record of OH-12 indicates that LCL is regularly affected by surface melt events (Figures 4 and 8c). The advection of warm air masses from lower latitudes leading to an increased sensible heat flux probably plays the most important role in surface melt on LCL [32,99]. This is supported by the statistically significant positive correlation of MLT of OH-12 with ERA5 near-surface air temperatures at the site (r = 0.5, p < 0.0001; Table 5). In addition, increased insolation on cloud-free days and hence the absorption of solar radiation by the snowpack can further trigger surface melt [100]. However, due to the exposed position of the study site, high wind speeds may occur year-round (Supplementary Figure S4). Hence, depending on air humidity, the absorbed solar radiation may not necessarily lead to strong surface melt but rather to an increased latent heat flux, i.e., enhanced evaporation/sublimation.
4.5. Relation between Firn Core Records and Sea Ice Extent

SIE variability in the adjacent oceans especially influences the availability of proximal and distal moisture for precipitation reaching LCL [2]. Reduced SIE will favour the transport of proximal (higher-latitude) oceanic moisture towards the northern AP with generally higher $\delta^{18}O$, but lower $d$ excess values due to the short distance between moisture source and sink. Increased SIE fosters both the influence of more (local) continental moisture with lower $\delta^{18}O$ but higher $d$ excess values, and also the contribution of distal (lower-latitude) oceanic moisture with variable characteristics depending on the conditions in the respective moisture source regions [32,96,101]. However, a correlation between SIE in the adjacent oceans and stable water isotopes of OH-12 is missing or diminished due to the previously described processes (see Sections 4.3 and 4.4.). Instead, it appears that inter-annual changes in SIE in the Bellingshausen-Amundsen and Weddell Seas (Figure 8e) play a role in the occurrence of strong surface melt events on LCL during the summer season. Figures 8c and 8e visualise the anti-correlation between MLT and SIE in both the Bellingshausen-Amundsen Sea and the Weddell Sea ($r > -0.6$, $p = 0$; Table 5). The observed anti-correlation may partly result from the fact that changes in MLT and SIE are both governed by variations in the magnitude of atmospheric and oceanic warming during the summer season. However, surface melt and SIE may be also directly related. In general, a decrease in SIE leads to more open water and lower albedo favouring stronger...
warming of the atmosphere and, thus, enhancing the sensible heat flux. This in turn favours surface melt. Accordingly, high MLT values in austral summers 2013 and 2015 coincide with lower SIE, especially in the Bellingshausen-Amundsen Sea, compared to the other two years (Figure 8c,e). In 2013, SIE in the Bellingshausen-Amundsen Sea is up to 21% and 34% lower during DJF and MAM, respectively, compared to 2012 and 2014. In 2015, it is reduced by up to 14% during DJF and up to 16% during MAM (Figure 8e). The respective differences in SIE in the Weddell Sea are less pronounced. In 2013, SIE is reduced by up to 11% during both DJF and MAM, whereas in 2015, it is even slightly increased compared to 2012 and 2014 (up to 4% during DJF and 10% during MAM; Figure 8e). Hence, we conclude that reduced SIE in the close-by Bellingshausen-Amundsen Sea likely favours the occurrence of strong surface melt events on LCL. However, warm-air advection from lower latitudes remains the main driver, as evidenced by the exceptional heat event in March 2015 likely causing strong surface melt on LCL.

5. Conclusions

In this study, we examined a 20 m long firn core retrieved from Plateau Laclavere located at the northern tip of the Antarctic Peninsula in austral summer 2016. Annual layer counting applied to the well-preserved high-resolution H2O: record of OH-12 revealed that the firn core covers a period of four full years (2012–2015).

We show that distinct maxima in the BC, nssS and nssS/ssNa records of OH-12 are probably linked to natural atmospheric emissions originating on the South American continent. Sources for BC are likely vast forest fire events in Chile and Argentina in 2011/2012 and 2015, and for nssS and nssS/ssNa, the eruptions of the southern Chilean volcanoes Puyehue-Cordón Caulle and Calbuco in 2011–2012 and 2015, respectively.

Snow accumulation in 2012–2015 was calculated to about 2500 kg m⁻² a⁻¹ on average with low inter-annual variability (on average ±8%), confirming LCL as a high-accumulation site.

Comparison between stable water isotopes of OH-12 and precipitation samples from Bernardo O’Higgins station reveals limited similarities. This implies that the isotopic signal of precipitation reaching the study site is modified during and/or after transfer into the snowpack, mainly due to orographic effects and post-depositional processes. Stable water isotope records of firn core OH-12 show no seasonal cyclicity either in the δ values or in the δ excess, most likely because of the maritime character of the northern AP climate and the impact of local post-depositional processes.

The study site is strongly influenced by the westerlies. Hence, we conclude that moisture reaching LCL primarily originates from proximal oceanic sources. The influence of moisture from more distal sources (e.g., South Pacific Ocean) is reflected in an exceptional stable water isotope composition of the snowpack. We demonstrate that stable water isotopes of OH-12 likely record an extreme warm-air event occurring in the northern AP region in austral autumn 2015. An intense atmospheric river prevailing over the Pacific Ocean between 21 and 25 March 2015 caused the large-scale advection of exceptionally warm and moist air from the subtropical Pacific towards the northern AP and caused record high air temperatures throughout the region. Concurrently with this event, δ¹⁸O and δ excess values of OH-12 reach their respective absolute maximum and minimum in the covered period.

Furthermore, the stratigraphy of OH-12 indicates strong surface melt events on LCL, especially in austral summers 2013 and 2015. Warm-air advection from lower latitudes towards the northern AP is probably the most important trigger for surface melt on LCL. Thick melt layers found in OH-12 in austral summer 2015 provide evidence that surface melt on LCL was likely related to the atmospheric-river induced warm-air event prevailing in the region in March 2015. In addition, we show that seasonal and inter-annual variations in SIE in the nearby Bellingshausen-Amundsen Sea likely play a role in the occurrence of surface melt on LCL.
In summary, we demonstrate that the ice cap on LCL as a high-accumulation site records meteorological and environmental signals in its glacio- and isotope-geochemical composition on sub-annual timescales and documents a clear connection to the South American continent. However, longer records from a deeper ice core will be necessary to reliably assess whether stable water isotopes from the ice cap on LCL can provide insights into past and present climate changes in the northern AP region on multi-decadal to multi-centennial timescales.

**Supplementary Materials:** The following are available online at www.mdpi.com/article/10.3390/geosciences11100428/s1, Equation (S1): Calculation of ssNa values, Equation (S2): Calculation of nssS values, Equation (S3): Calculation of wind directions at 850 mbar geopotential height, Supplementary Table S1: Coordinates and altitude of the geographic location and the nearest ERA5 grid point for the Antarctic stations Bernardo O’Higgins (OH), Bellingshausen (BH), Esperanza (EP) and Comandante Ferraz (CF), as well as for the OH-12 drill site on Plateau Laclavere (LCL), Supplementary Figure S1: Thickness distribution of OH-12 ice layers that are indicating surface melt, Supplementary Figure S2: Monthly distribution of precipitation samples collected at Bernardo O’Higgins station from (a) January 2008 to March 2009, from (b) March to November 2014, and from (c) December 2015 to December 2017, Supplementary Figure S3: Comparison of the (a) stable oxygen and hydrogen isotope and (b) d excess records of firn core OH-12 and precipitation samples collected at Bernardo O’Higgins station (OH) in the year 2014, Supplementary Figure S4: Wind roses for the study site on Plateau Laclavere derived for the single years 2012–2015 as well as for the entire period based on ERA5 monthly records of the zonal and meridional components of the 850 mbar geopotential height wind extracted for the grid point closest to the OH-12 drill site.

**Author Contributions:** F.F. designed the study and carried out the fieldwork, supported by K.H.-A. and H.M. K.H.-A. and H.M. performed the stable water analyses of firn core OH-12. K.H.-A. and J.R.M. performed the glacio-chemical analyses of OH-12. High-resolution density profiles of OH-12 were measured by K.H.-A. and J.F. K.H.-A. was responsible for data analysis, interpretation and writing of the manuscript, supported by H.M., C.S., F.F. and T.O. All authors contributed to the data interpretation and the preparation of the final manuscript. All authors have read and agreed to the published version of the manuscript.

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**Data Availability Statement:** The data will be made publicly available via https://www.pangaea.de.

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