

Evaluating the 11-year solar cycle and short-term ^{10}Be deposition events with novel excess water samples from the EGRIP project

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Abstract. ^{10}Be is produced by the interaction between galactic cosmic rays (GCR) and solar energetic particles (SEP) with the Earth's atmospheric constituents. The flux of GCR is modulated by the varying strength of the magnetic fields of the Earth and the Sun. Measurement of ^{10}Be concentrations from polar ice cores is thus a
20 valuable tool to reconstruct the variations of the geomagnetic field and solar activity levels. The interpretation of ^{10}Be records is, however, complicated by non-production related effects on the ^{10}Be deposition rate caused by climate/weather induced variability. Furthermore, volcanic eruptions have been proposed to lead to short-term ^{10}Be deposition enhancements. In this study, we test the use of excess meltwater from continuous flow analysis (CFA) to measure ^{10}Be , allowing less time-consuming and more cost-effective sample preparation. We compare
25 two records obtained from CFA and discrete samples from the EGRIP S6 firn core, reaching back to 1900 CE. We find that the two records agree well and that the ^{10}Be record from CFA samples agrees as well as the discrete samples with other records from Greenland. Furthermore, by subtracting the theoretically expected GCR-induced signal, we investigate the high-frequency variability of the ^{10}Be records from Greenland and Antarctica after 1951
30 CE, with focus on SEP events and volcanic eruptions. Finally, we use the ^{10}Be records from Greenland and Antarctica to study the 11-year solar cycles, allowing us to assess the suitability of the CFA samples for the reconstruction of solar activity. This result opens new opportunities for the collection of continuous ^{10}Be records with less time-consuming sample preparation while saving an important portion of the ice cores for other measurements.

35 1 Introduction

^{10}Be is produced in the atmosphere by the interaction between high-energy cosmic ray particles (galactic cosmic rays – GCR) and atmospheric atoms. About 65% of atmospheric ^{10}Be nuclides are produced in the stratosphere (Heikkilä et al., 2013; Heikkilä et al., 2009; Masarik & Beer, 1999), where they can be assumed to be well mixed
40 due to an average residence time of 1-2 years (Heikkilä et al., 2008). Upon binding to sulfate aerosols (e.g. Igarashi et al., 1998), ^{10}Be is removed from the atmosphere by wet and dry deposition, and can be measured in environmental archives, such as ice cores from Greenland and Antarctica. Using the ECHAM5 general circulation

model coupled with the aerosol module HAM, Heikkilä et al. (2009) showed that the dominant component of ^{10}Be in polar ice cores is stratospheric ^{10}Be , making up to about two thirds of the signal preserved in ice cores from Greenland and Antarctica. In consequence, they argued that deposition fluxes of ^{10}Be reflect well the changes in the global average ^{10}Be atmospheric production rate.

The flux of GCRs reaching the Earth is modulated by the magnetic fields of Sun and Earth. The production of ^{10}Be thus anticorrelates with the strength of the helio- and geomagnetic fields allowing us to use ^{10}Be from ice cores to reconstruct solar activity (e.g. Beer et al., 1990; McCracken et al., 2004; Muscheler et al., 2007; Steinhilber et al., 2012; Vonmoos et al., 2006), and the variations in the geomagnetic field dipole moment (e.g. Muscheler et al., 2005; Raisbeck et al., 1985; Raisbeck et al., 2006).

In the last decade it has also been shown that solar storms can leave a significant imprint in ^{10}Be from Greenland and Antarctica. These events can, in fact, lead to short-lived peaks in the radionuclide concentrations (Mekhaldi et al., 2015; Miyake et al., 2019; Miyake et al., 2015; O'Hare et al., 2019; Paleari et al., 2022). Mekhaldi et al. (2021) modeled the theoretically expected globally averaged annual production rate of ^{10}Be since the 1950s using neutron monitor data, and the theoretical production induced by ground level enhancements (GLEs), i.e. solar energetic particle (SEPs) events that cause sudden increases in ground-based neutron monitor count-rates. The strongest directly observed GLE - GLE no.05 from 1956 – caused, for instance, an increase of only about 5% in the annual ^{10}Be production rate. Considering that the 11-year solar cycle is estimated to cause a variability of $\pm 15\%$ to $\pm 35\%$ (e.g. Baroni et al., 2011; Paleari et al., 2022; Pedro et al., 2012, Mekhaldi et al., 2021), the signal of modern events cannot be unequivocally distinguished from the variability caused by the 11-year cycle, and climate and local weather influences (Mekhaldi et al., 2021). For the instrumental period no unambiguous SEP signal could be detected in ice cores so far, not even in seasonal ^{10}Be data (e.g. Pedro et al., 2011; Zheng et al., 2020).

Moreover, as proposed by Baroni et al. (2011, 2019), peaks in ^{10}Be concentrations in ice cores may sometimes be linked to stratospheric volcanic eruptions. The Agung and Pinatubo eruptions, for instance, are estimated to have caused an increase in ^{10}Be deposition of 66% and 35% relative to the baseline radionuclide concentration in the Vostok ice core from Antarctica (Baroni et al., 2011).

Especially for detecting the short-lived spikes caused by SEPs in the past, high-resolution measurements are required. These are often labor-intensive and limited by the availability of sufficient ice-core sample material. In this study, we aim at testing the use of excess meltwater samples usually discarded from continuous flow analysis (CFA) for the measurement of cosmogenic ^{10}Be . This method could provide the potential for high-resolution and cost-efficient ^{10}Be sampling without the need for competing for the valuable ice. We collected the meltwater from the CFA system at the Institute for Climate and Environmental Physics at the University of Bern (Switzerland). The system is used to measure impurities in ice cores (e.g. Erhardt et al., 2022), which requires the use of pristine ice. The system is equipped with a melthead designed to only capture the inner part of the ice core for the measurement of impurities, preventing the ice from being possibly contaminated by the surrounding atmosphere (Erhardt et al., 2019). The outer part can in theory be used for cosmogenic radionuclide measurements as they are less prone to contamination. This technique has already been used to collect a short record of about 40 years from the EGRIP ice core to study an extreme SEP event that hit Earth 9,125 years BP (Paleari et al., 2022), estimated to be two orders of magnitude larger than modern GLEs. While suitable for the assessment of the existence of the

peak in ^{10}Be concentrations, it was not possible to analyze in depth and ultimately quantify the uncertainties related to this method. That is because the record was only compared to ice core records from other locations from Greenland and Antarctica, therefore characterized by different climate and weather conditions. Here, we present a new ^{10}Be record from the S6 firn core from EGRIP (East Greenland Ice core Project, see Fig. 1 for the location), covering the depth from about 4 to 26m, corresponding approximately to the period between 1900 and 2008 CE. To assess the suitability of the CFA samples to measure cosmogenic radionuclides and retrieve a solar signal, ^{10}Be records obtained from CFA and discrete firn samples from the EGRIP S6 core (Zheng et al., 2023) are compared. The use of two ^{10}Be records from the same core allows us to directly assess the quality and possible differences between the sampling methods.

In addition, we investigate the causes of short-term and non-GCR related ^{10}Be variability in the ^{10}Be records from EGRIP S6 and other available records from Greenland and Antarctica. More specifically, we aim to investigate whether we can detect the signal of large SEP events (or GLEs) and volcanic eruptions. Furthermore, we investigate the preservation of the 11-year solar cycle in the ^{10}Be records.

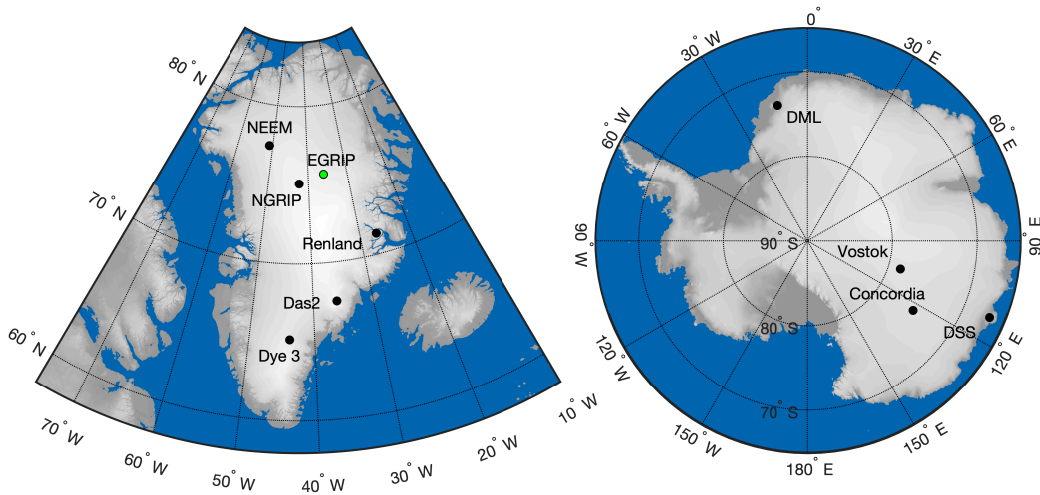


Figure 1. Location of the EGRIP site (marked with a green circle) and other ice core locations from Greenland (left) and Antarctica (right) discussed in this work.

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2 Methods

2.1 Collection and preparation of the CFA samples

The CFA samples were collected during the melting campaign carried out at the Institute for Climate and Environmental Physics at the University of Bern (Switzerland) in October 2019. During the campaign, 36x36 mm vertical sticks of ice of 1m in length were continuously melted on a melthead. The melthead is designed to only inject meltwater from the inner part of the core for measurement (26x26 mm, e.g. Erhardt et al., 2022), while the meltwater from the outer part of the core, otherwise discarded, is pumped towards centrifuge tubes for continuous ^{10}Be sampling through a 3 m long tube in Teflon. To collect the water, we used 50ml centrifuge tubes where 0.1 mg ^9Be carrier (from Scharlau (1000 mg/l; $\text{Be}_4\text{O}(\text{C}_2\text{H}_3\text{O}_2)_6$ in HCl 2%)) was previously injected.

The preparation of the samples for Accelerator Mass Spectrometry (AMS) measurements was carried out at the Department of Geology at Lund university (Sweden). Each ~50ml sample was directly precipitated with NH₄OH. This allows us to avoid the use of ion exchange chromatography (IEC), thereby saving about 3 hours of preparation per 12 samples and a significant amount of materials (e.g., ion exchange columns). A schematic summarizing the preparation of ¹⁰Be samples with and without IEC at Lund University can be found in Nguyen et al. (2021). After centrifugation, the precipitate was transferred to a quartz crucible. During this step, consecutive samples were paired to obtain samples of ~100g with 0.2mg of ⁹Be carrier. The resulting samples have an average length of 25 cm, corresponding to an average time resolution of 1.2 years, with a minimum of 0.7 years and maximum of 2 years, depending on annual layer thickness and firm density. The crucibles were then placed into an oven where the temperature is slowly raised to 850° C, allowing the oxidization of Be(OH)₂ to BeO. Finally, 1mg of Niobium was added and the mixture was pressed into target holders for measurements. The measurements were carried out at the Laboratory of Ion Beam Physics at ETH, Zurich (Switzerland). The record was blank corrected (¹⁰Be/⁹Be blank ratios: ~6% of the ¹⁰Be/⁹Be samples ratio; average ¹⁰Be/⁹Be of the blank samples: 0.007 x 10⁻¹²). The measured ¹⁰Be/⁹Be ratios were normalized to the ETH Zurich in-house standards S2007N and S2010N, which were both calibrated relative to the ICN 01-5-1 standard (¹⁰Be/⁹Be = 2.709 × 10⁻¹¹ nominal) (Christl et al., 2013).

2.2 Depth attribution of the samples

The depth-sample relationship has to be reconstructed since the continuous melting does not allow a precise measurement of the depth range covered by each CFA ¹⁰Be sample. During sample collection it is possible to know in which vial there is a transition between two consecutive 1m sections, and the samples in between are assumed to contain equidistant sections of firm in terms of ice-equivalent depth ignoring any short term fluctuations in melt speed. This is probably not accurate as the layer thickness is not constant. Taking also into account the time needed for the meltwater to flow from the melthead to the centrifuge tubes (usually 20-30 seconds), and that was calculated at the beginning of each melting session, we assume the uncertainty related to the depth attribution to be <6 cm (3-4 months on average). We thus consider this to not affect significantly our results (see Section 3).

2.3 Timescale of the EGRIP S6 core

The timescale for the EGRIP S6 core is constructed by counting the annual layers visible in the CFA dataset of the core. To constrain the annual layer count and align the EGRIP S6 age model with the main core, tie points of the EGRIP-GICC05 age scale (Mojtabavi et al., 2020) were identified in the electrolytic meltwater conductivity records at 21.53, 23.94 and 30.63 m depth and 73, 87 and 125 years b2k (before 2000 CE) respectively. In addition, the drilling year was used as an age constraint at the surface. The counting was performed using the StratiCounter algorithm (Winstrup et al., 2012). Between the first tie-point and the surface age, the time scale is adjusted within the annual layer counting uncertainty based on work by Zheng et al. (2023) who identified the timescale offset (i) between the EGRIP discrete firm ¹⁰Be data and the theoretical ¹⁰Be production rate referred from the sunspot numbers and (ii) between accumulation rates (layer thickness) and precipitation. For consistency, in this study we adopt the updated chronology by Zheng et al. (2023).

3 ^{10}Be record from CFA excess water samples

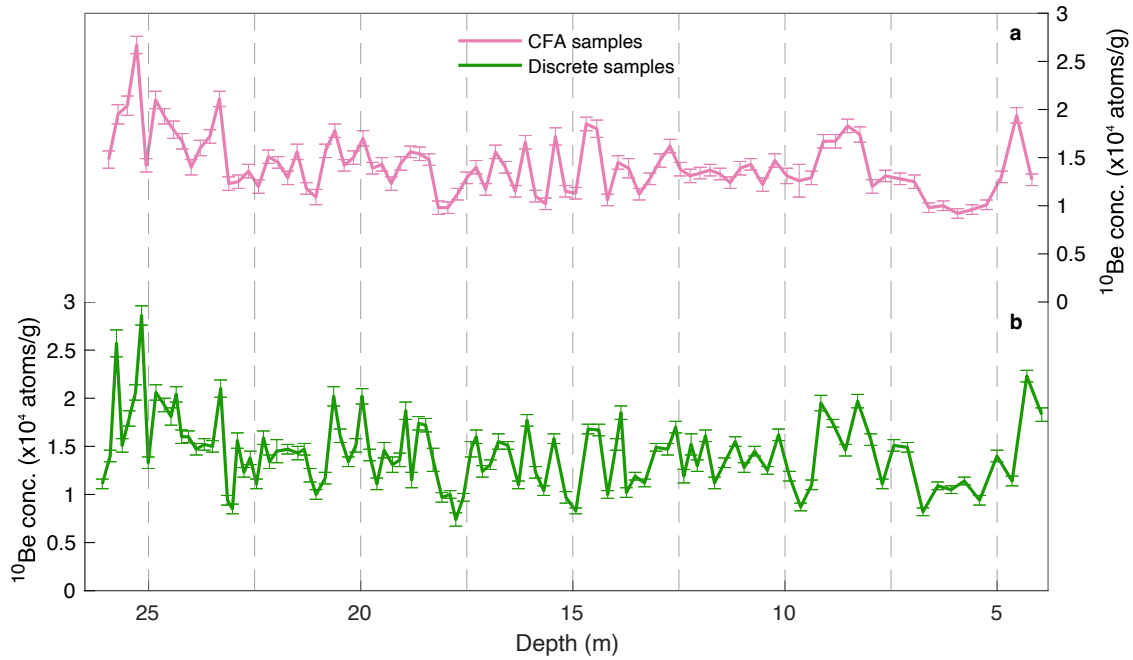


Figure 2. ^{10}Be concentration record from CFA excess water samples (panel a) and ^{10}Be concentrations from
 155 discrete firn samples (panel b - Zheng et al., 2023) from the EGRIP S6 core. The records are plotted versus the
 depth of the core. The error bars represent the 1σ measurement uncertainty.

The ^{10}Be record from EGRIP S6 CFA excess water samples is shown in pink in Figure 2a. The ^{10}Be concentrations
 were measured in firn from about 4 to 26 m of depth, encompassing the period 1900-2008, according to the
 160 chronology by Zheng et al. (2023). The ^{10}Be concentrations have an average of $1.43 (\pm 0.31) \times 10^4$ atoms/g.
 Figure 2b shows the ^{10}Be concentrations from discrete firn samples from the same core. The average ^{10}Be
 concentration of the discrete record is $1.42 (\pm 0.32) \times 10^4$ atoms/g (Zheng et al., 2023). The meltwater samples
 have, on average, a measurement uncertainty of 5%, similar to the ^{10}Be measurements carried out on the discrete
 firn samples (Zheng et al., 2023), despite the smaller size (100g for CFA samples versus ca. 140g for discrete
 165 samples).

As the sampling rates of the two S6 records are different, the ^{10}Be records need to be virtually resampled to be
 directly compared. The ^{10}Be concentrations from CFA samples and from discrete samples were interpolated
 linearly at steps of 0.1 year and the yearly ^{10}Be concentrations were calculated by averaging over 10 consecutive
 datapoints. The two yearly resampled ^{10}Be records agree well ($r=0.77$, $p<0.01$ - computed using a t-test). If we
 170 resample the discrete record to match the resolution of the CFA ^{10}Be record, we obtain a correlation coefficient
 of 0.71 ($p<0.01$). It can, however, be pointed out that the first ~ 8 m of the CFA excess water record (between ~ 4
 and ~ 12 m of depth) is seemingly more autocorrelated compared to the deeper part of the record and relative to
 the discrete ^{10}Be record, the latter also smoother in the first ~ 8 m of the record compared to the deeper part. The
 sampling method and rate can likely affect the degree to which the two records agree. ^{10}Be deposition is known
 175 to be highly seasonal, with maxima closely related to the enhanced stratosphere-troposphere exchanges (STE)
 causing increased descent of ^{10}Be from the stratosphere, where most of the production takes place, to the

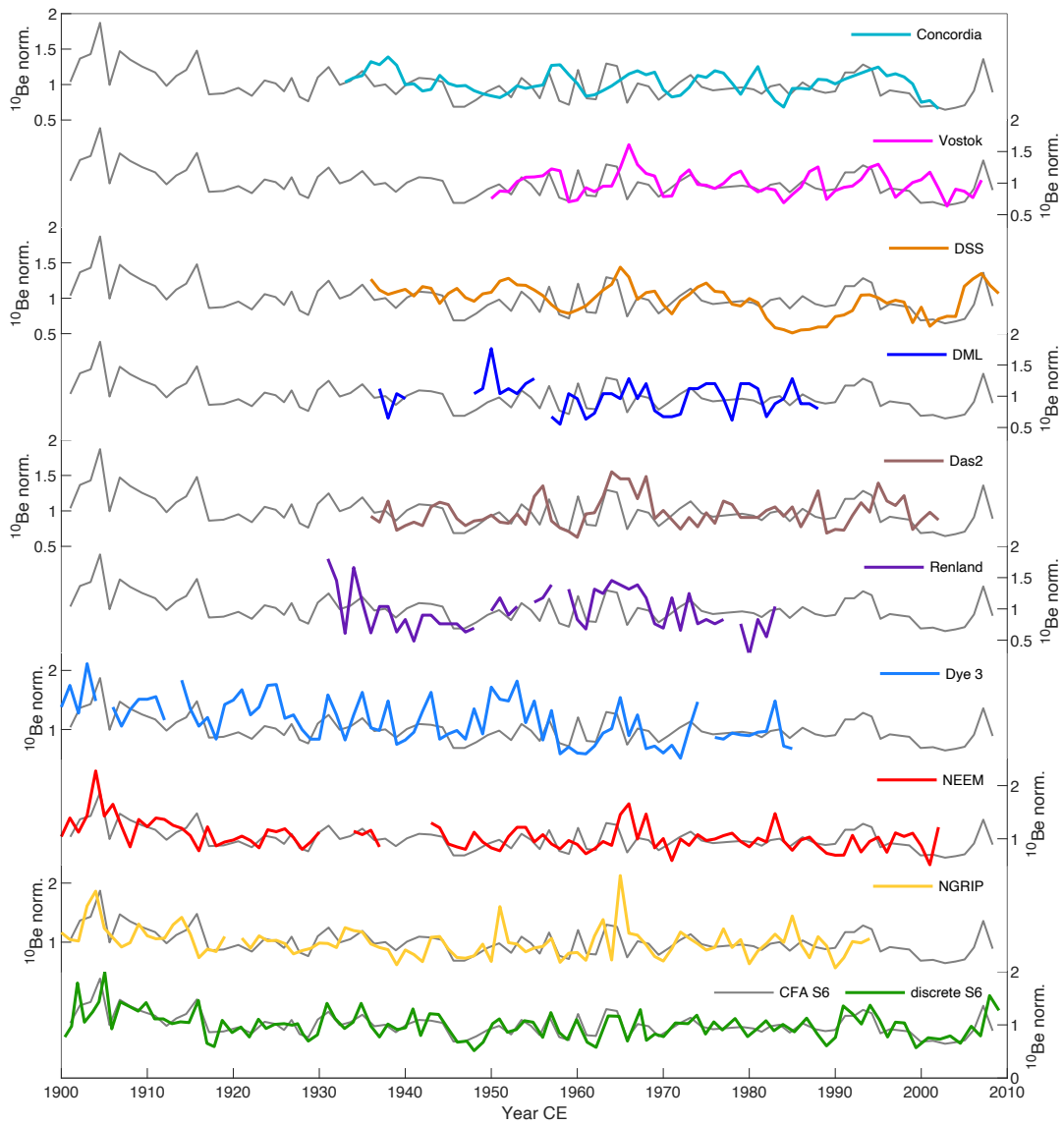
troposphere (Stohl et al., 2003; Zheng et al., 2020; Spiegl et al., 2022; Heikkilä and Smith, 2013; Pedro et al., 2011a). Moreover, one of the main complications of dealing with CFA systems is the possible smoothing of the signal locked in the ice (Erhardt et al., 2022; Kaufmann et al., 2008; Mekhaldi et al., 2017; Sigg et al., 1994). ^{10}Be ions may attach to the walls of the plastic tube connecting the melthead to the centrifuge tubes, as ^{10}Be precipitates in a basic environment. Therefore, it is possible that some atoms may precipitate on the plastic walls and get remobilized at a later stage. Loss of ^{10}Be to the centrifuge tube walls is considered unlikely since the ^9Be carrier (slightly acidic) was added to the centrifuge tubes before collecting the samples, and the $^{10}\text{Be}/^9\text{Be}$ ratio is in theory preserved. It is however most likely that the smoothing in the shallower part of the CFA ^{10}Be record is caused by analytical issues related to the upward wicking of meltwater in the very porous snow during melting, which affects the shallow most parts of the core.

4 Comparison with other records from Greenland and Antarctica

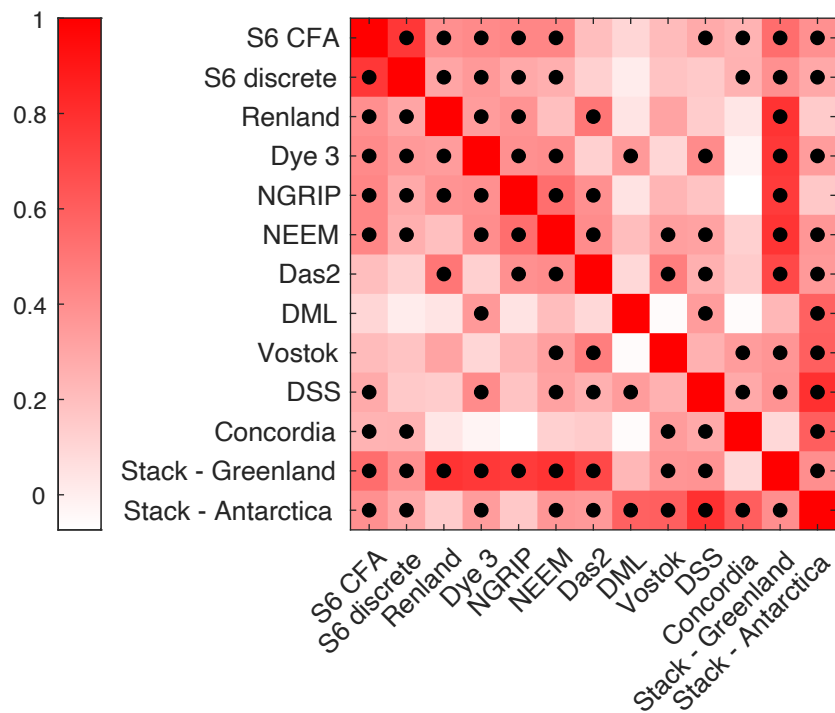
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To investigate whether the CFA excess water samples are suitable to preserve the ^{10}Be production rate signal as robustly as discrete firm samples, the ^{10}Be records from EGRIP S6 are compared to other available ^{10}Be records from Greenland and Antarctica. We compare the ^{10}Be record from CFA samples to ^{10}Be concentration records from the NGRIP (Berggren et al., 2009), NEEM (Zheng et al., 2021), Dye 3 (Beer et al., 1990), Renland (Aldahan et al., 1998), Das2 (Pedro et al., 2012), DML (Aldahan et al., 1998), DSS (Pedro et al., 2012), Vostok (Baroni et al., 2011) and Concordia (Baroni et al., 2011) cores. The NEEM record is available at sub-annual resolution, and thus, we calculated mass-weighted means of each year. Figure 3 shows the comparison between the CFA ^{10}Be record (grey line) and the other records from Greenland and Antarctica starting in 1900 CE. The degree to which the CFA ^{10}Be record agrees with the other ^{10}Be records can be seen in the correlation matrix in Figure 4. For this analysis, we used the CFA and discrete ^{10}Be records resampled at 1 year resolution, to match the timescale of the other records. To calculate the correlation coefficients, the records were normalized to their mean for the period 1949-1985, representing the time span shared by all records.

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205 *Figure 3. Normalized ^{10}Be concentration records from the EGRIP S6 CFA samples in comparison to the normalized ^{10}Be records from discrete samples (Zheng et al., 2023), NGRIP (Berggren et al., 2009), NEEM (Zheng et al., 2021), Dye 3 (Beer et al., 1990), Renland (Aldahan et al., 1998), Das2 (Pedro et al., 2012), DML (Aldahan et al., 1998), DSS (Pedro et al., 2012), Vostok (Baroni et al., 2011) and Concordia (Baroni et al., 2011) starting in 1900 CE.*



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Figure 4. Correlation matrix (Pearson correlation coefficient) between the ^{10}Be records from EGRIP S6 (CFA and discrete), Renland, Dye 3, NGRIP, NEEM, Das2, DML, Vostok, DSS, Concordia, and the Antarctic and Greenland stacks (excluding the EGRIP S6 records) since 1900 CE. Dots represent significant correlations ($p < 0.05$).

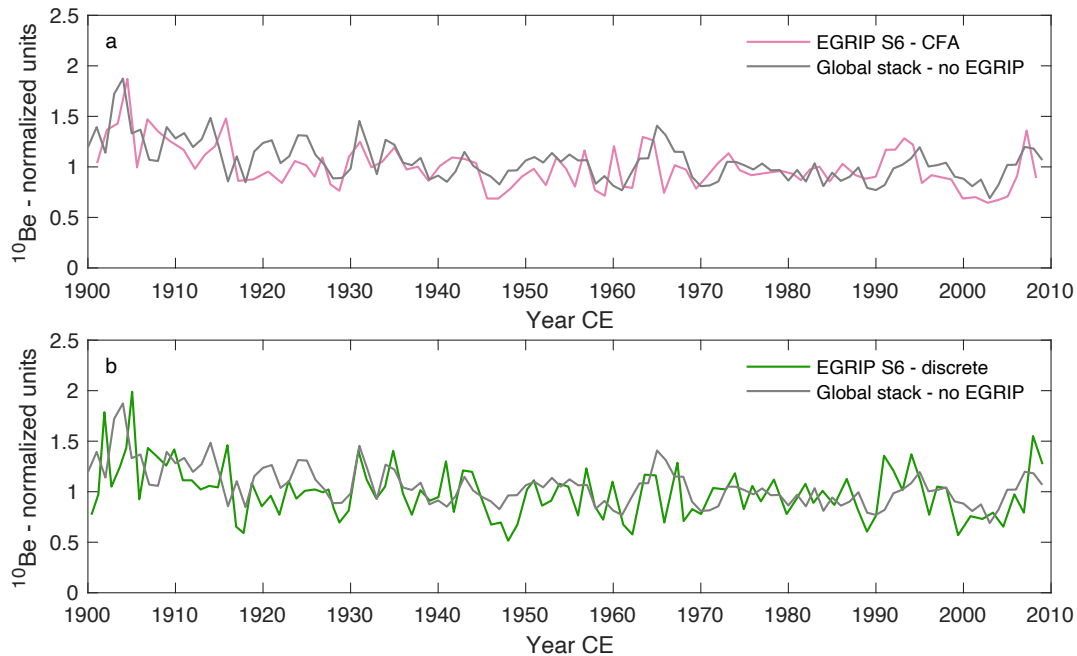
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In general, the EGRIP CFA ^{10}Be record shows similar but somewhat higher correlation coefficients with the other records from Greenland and Antarctica, compared with the discrete EGRIP ^{10}Be record. While the higher correlation coefficients can likely be attributed to the higher autocorrelation inherent to the smoothing effect of CFA sampling in the shallower part of the record, our results indicate that the signal measured in the CFA samples is reproducing the common radionuclide signal in Greenland and Antarctica as well as the discrete firm samples. The EGRIP S6 records from CFA and discrete firm samples agree well with the ^{10}Be records from Greenland, except for Das2 (See Figure 4). This could be explained by the very high ^{10}Be concentrations in the Das2 record during the 1960s, corresponding to a pronounced minimum in accumulation rate (Pedro et al., 2012). Moreover, the differences between the EGRIP S6 records and the Das2 record in the early 1990s likely affect the correlation coefficient. We obtain correlation coefficients of 0.54 ($p < 0.01$) and 0.39 ($p < 0.01$) between the stack of ^{10}Be records from Greenland (excluding the EGRIP S6 records) and the S6 records from CFA and discrete samples, respectively. We obtain correlation coefficients of 0.39 ($p < 0.01$) and 0.29 ($p < 0.01$) between the stack of records from Antarctica and the S6 records from CFA and discrete samples, respectively.

Differences between the records could arise from non-production related biases such as measurement and sampling uncertainty, meteorological influences on the transport and deposition of ^{10}Be en-route and on site, as well as local re-distribution of snow by surface winds. For instance, Heikkilä and Smith (2013) found a strong correlation between ^{10}Be deposition and the North Atlantic Oscillation (NAO) in Greenland, with a positive correlation on the east coast and negative on the west coast.

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235 CFA and discrete ^{10}Be records are then compared with the stack of the normalized ^{10}Be records from Greenland (excluding EGRIP S6 records) and Antarctica (Figure 5). The calculation of a stack with several records from different sites allows us to reduce the noise by isolating their common signal which is expected to more closely reflect the atmospheric production signal.



240 *Figure 5. Comparison between the global stack calculated using the normalized ^{10}Be records from Greenland and Antarctica (NGRIP, NEEM, Renland, Dye 3, Das2, DML, DSS, Vostok and Concordia) and the normalized ^{10}Be records from EGRIP S6 CFA (panel a) and discrete (panel b) samples.*

The CFA ^{10}Be record has a correlation coefficient of 0.60 ($p < 0.01$) with the global ^{10}Be stack, while the ^{10}Be record from discrete firm samples has a correlation coefficient of 0.42 ($p < 0.01$).

5 Potential SEP and volcanic signals in ^{10}Be

5.1 EGRIP S6 records

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^{10}Be records from ice cores are valuable tools to reconstruct solar activity prior to the advent of spaceborne measurements, ground-based instrumental records and sunspot observations. Nevertheless, the interpretation of ^{10}Be records is complicated by the influence of climate, local weather noise (e.g. Pedro et al., 2011) and the stochastic occurrence of volcanic eruptions (Baroni et al., 2011, 2019).

255 Here, we investigate outliers in the two ^{10}Be records from the EGRIP S6 firm core to assess whether it is possible to infer their cause. To do so, the normalized globally averaged ^{10}Be production rate caused by GCR (Mekhaldi et al., 2021) was virtually resampled at the resolution of the CFA and discrete ^{10}Be records. This resampled production rate was subtracted from the two normalized ^{10}Be records from EGRIP S6 for the period from 1951 (constrained by the advent of neutron monitors) to 2008 assuming that this removes the variability induced by the

260 11-year solar cycle. We then standardized the residuals to z-scores, i.e. the number of standard deviations by which the residuals are above or below their mean value. We repeated the experiment using the modeled normalized ^{10}Be deposition flux over latitudes 60-90N caused by GCR and including the transport modeled using the parametrization from Heikkilä et al. (2009) (Mekhaldi et al., 2021). From this point on, we will refer to $\text{Res}_{\text{recordname_prod}}$ and $\text{Res}_{\text{recordname_transport}}$. The standard scores of the resulting residuals are shown in Figure 6 (CFA samples in pink, and discrete firn samples in green) and a summary of the results is shown in Table 1.

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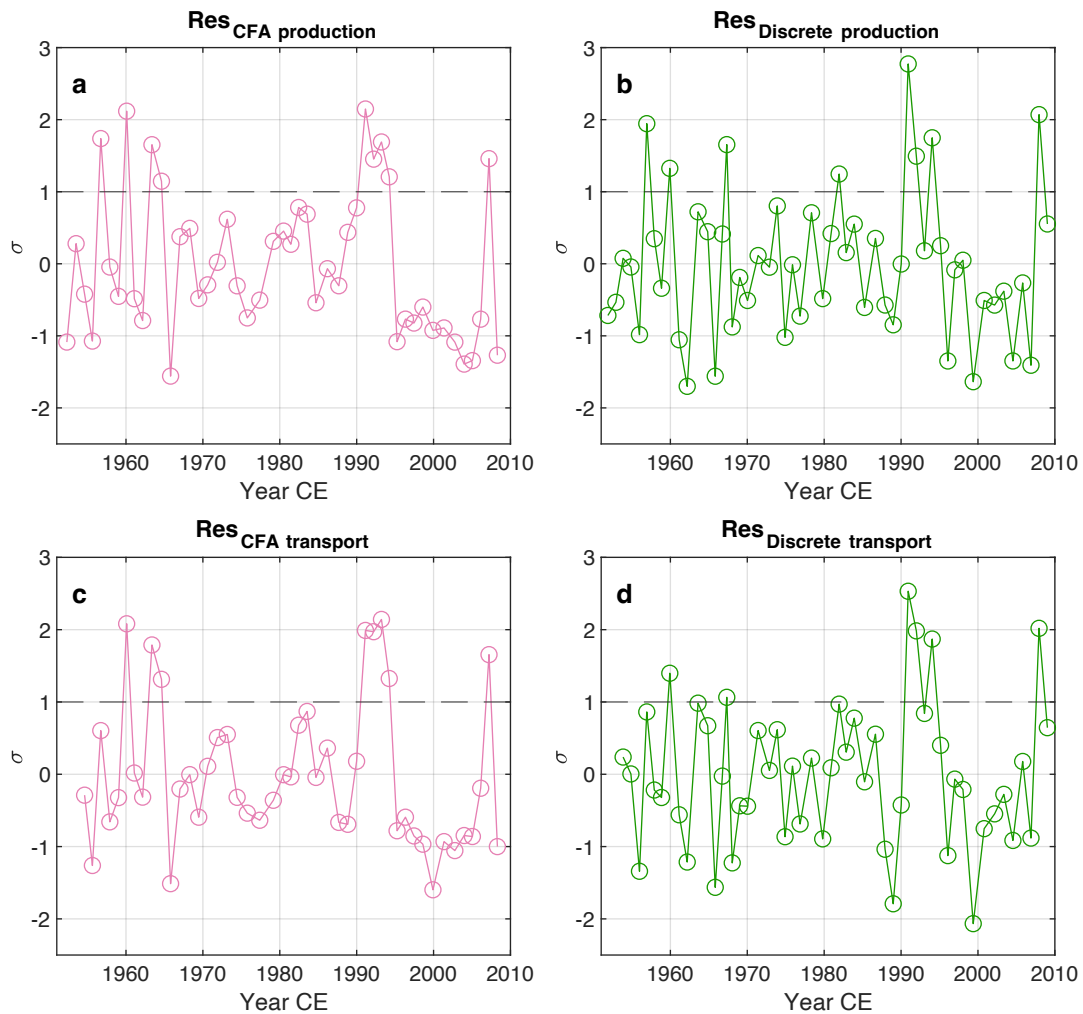


Figure 6. Standard scores for the EGRIP S6 ^{10}Be records. Panel a and b show the standardized residuals obtained by subtracting the normalized theoretical ^{10}Be global production rate from the normalized EGRIP S6 ^{10}Be records since 1951 (CFA ^{10}Be sample residuals in pink, discrete ^{10}Be sample residuals in green). Panel c and d show the same as panel a and b, but for the ^{10}Be deposition flux including the transport over latitudes 60-90N (Mekhaldi et al., 2021).

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Although we expect some of these results to be exclusively caused by noise inherent to the data, values exceeding 1σ in the years following a stratospheric volcanic eruption and/or a solar storm may be more likely. Several of the major stratospheric volcanic eruptions from the last 70 years have been identified in sulfate records from

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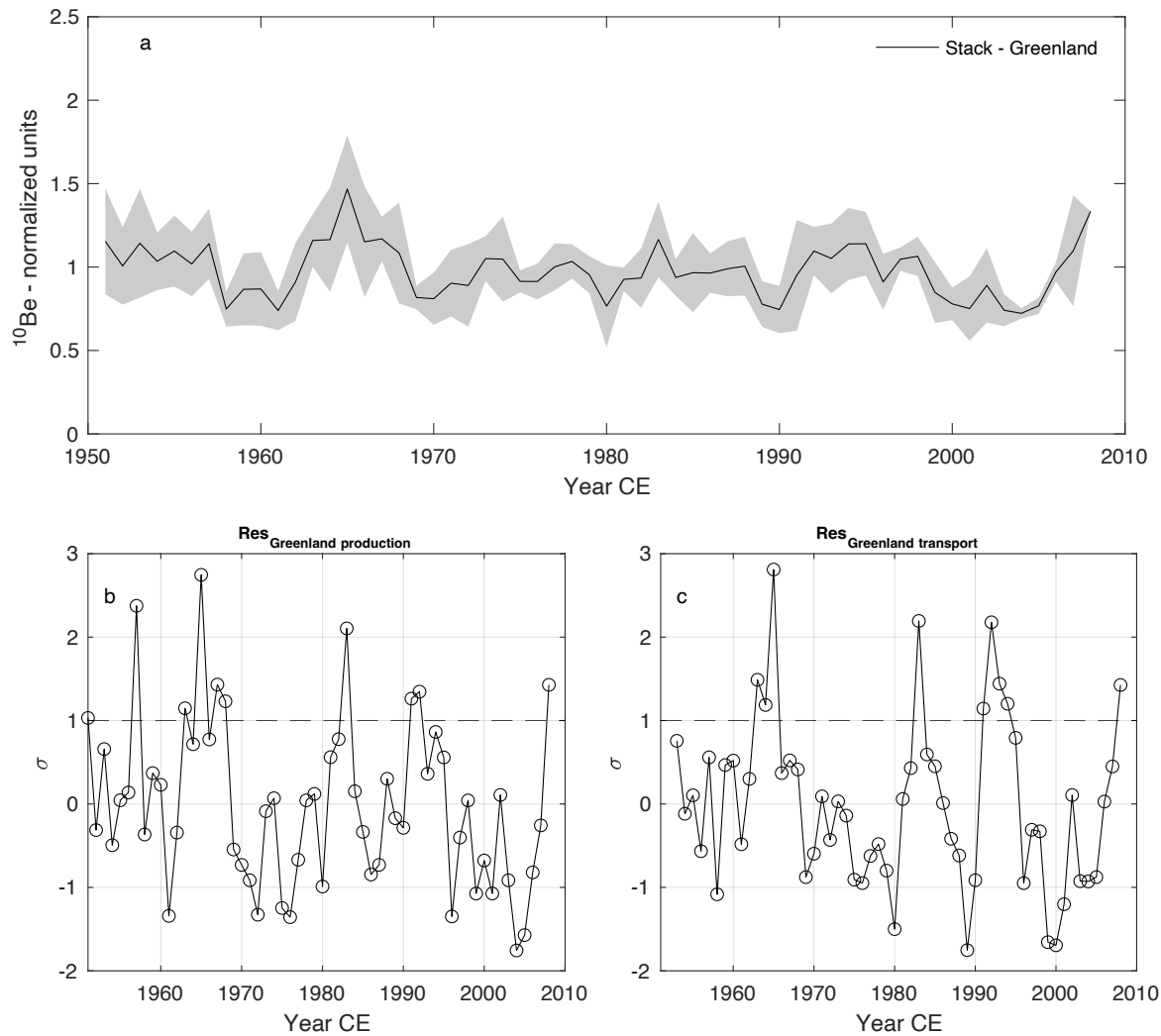
Greenland in 1957, 1964, 1982-84 and 1991 (Sigl et al., 2015; Zielinski et al., 1997). Some of the eruptions were traced to, for instance, Agung (in 1963), El Chichón (in 1982) and Pinatubo (in 1991). Taking into account timescale uncertainties and the lag due to the transport of ^{10}Be from the stratosphere, we would expect to find values exceeding 1σ in the years of the eruptions and/or 1 to 2 years after the eruptions. The results of the analysis are summarized in Table 1. Some of the results are coeval with the timing of the Agung and Pinatubo eruptions, albeit the Pinatubo eruption is generally not particularly prominent in the other ^{10}Be records from Greenland (see Figure 3) and the uncertainty of the S6 chronology may affect our interpretation.

Although it is reasonable to expect values exceeding 1σ to be related to stochastic noise, we want to investigate whether GLE no.05 from 1956 left any traces in the ^{10}Be records from ice cores. GLE no.05 is the event that yielded the most annual production of ^{10}Be over the last 70 years of instrumental data, with an estimated increase of +5.1% in the modeled annual ^{10}Be global production rate (Mekhaldi et al., 2021). The analysis of the standard scores shows that values exceeding 1σ can be identified in 1957 in $\text{ReS}_{\text{CFA_prod}}$ and $\text{ReS}_{\text{discrete_prod}}$. We find an increase in ^{10}Be concentrations of 21% in 1957 in the CFA record, and an increase of 42% in the discrete record. The increases were calculated relative to the average ^{10}Be concentration over solar cycles 19 (excluding the targeted years). However, we cannot exclude that the increase in ^{10}Be concentration is caused mostly by the natural variability of ^{10}Be with a more modest contribution of the SEP events and/or volcanic eruptions.

5.2 ^{10}Be stacks from Greenland and Antarctica

In the following section we analyzed the standard scores of the residuals obtained by subtracting the normalized theoretical ^{10}Be production rate (by GCR) from the Greenlandic and Antarctic stacks. As volcanic eruptions may impact ^{10}Be records from Greenland and Antarctica differently, the analysis of the standard scores has been performed separately on the stacks from the two regions.

The stack from Greenland (see Figure 7a) has been calculated using all the records from Greenland from Fig. 3 normalized to their respective mean during the period 1951-1985, representing the time span shared by all the records. Figure 7b shows the standard scores calculated on the residuals obtained by subtracting the modeled normalized ^{10}Be production rate (Mekhaldi et al., 2021) from the Greenland ^{10}Be stack ($\text{ReS}_{\text{Greenland_prod}}$). Figure 7c shows the standard scores calculated on the residuals obtained by subtracting the normalized ^{10}Be modeled deposition flux including the transport in the northern hemisphere (Mekhaldi et al., 2021) ($\text{ReS}_{\text{Greenland_transport}}$).



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Figure 7. Standard scores for the difference between the Greenland ^{10}Be stack and the theoretical GCR-induced production rate. The stack (panel a) was calculated using the normalized ^{10}Be records from Dye 3, NGRIP, NEEM, Das2, Renland and EGRIP S6 (CFA and discrete) since 1951. Panel b shows the standardized residuals obtained by subtracting the modeled ^{10}Be global production rate from the stack. Panel c shows the same as panel
 310 b but using the normalized ^{10}Be deposition flux modeled (taking transport into account) over latitudes 60-90N (Mekhaldi et al., 2021).

The standard scores are above -2σ , in agreement with the results from the EGRIP S6 records. The years in which the standard scores exceed 1σ are summarized in Table 1. We find an increase of 16% in ^{10}Be concentrations in
 315 year 1957 relative to the baseline (average ^{10}Be concentration over solar cycle 19). However, the increase in ^{10}Be concentrations is coeval with a peak detected in the NEEM sulfate record (Sigl et al., 2013). Therefore, the presence of a ^{10}Be increase in ice cores from Antarctica would lend support to the possible contribution of GLE no.05 to the ^{10}Be deposited in Greenland and Antarctica in 1957. We carried out the same analysis on the stack calculated from the Antarctic records. We subtracted the normalized global ^{10}Be production rate from the stack

320 of DSS, DML, Concordia and Vostok normalized ^{10}Be records and calculated the standard scores. The stack and standard scores of the residuals are shown in Figure 8.

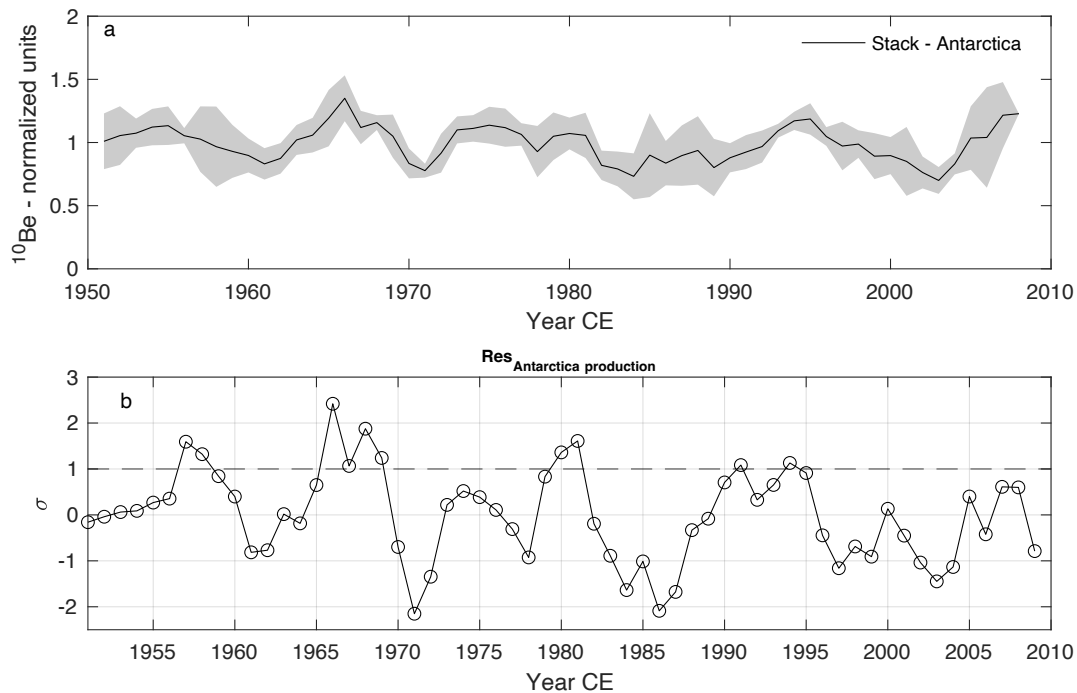


Figure 8. Standard scores for the Antarctic stack. The stack (panel a) was calculated using the normalized ^{10}Be records from Vostok, DML, Concordia and DSS from 1951. The standard scores were calculated on the residuals
 325 obtained by subtracting the normalized modeled ^{10}Be global production rate from the stack (panel b).

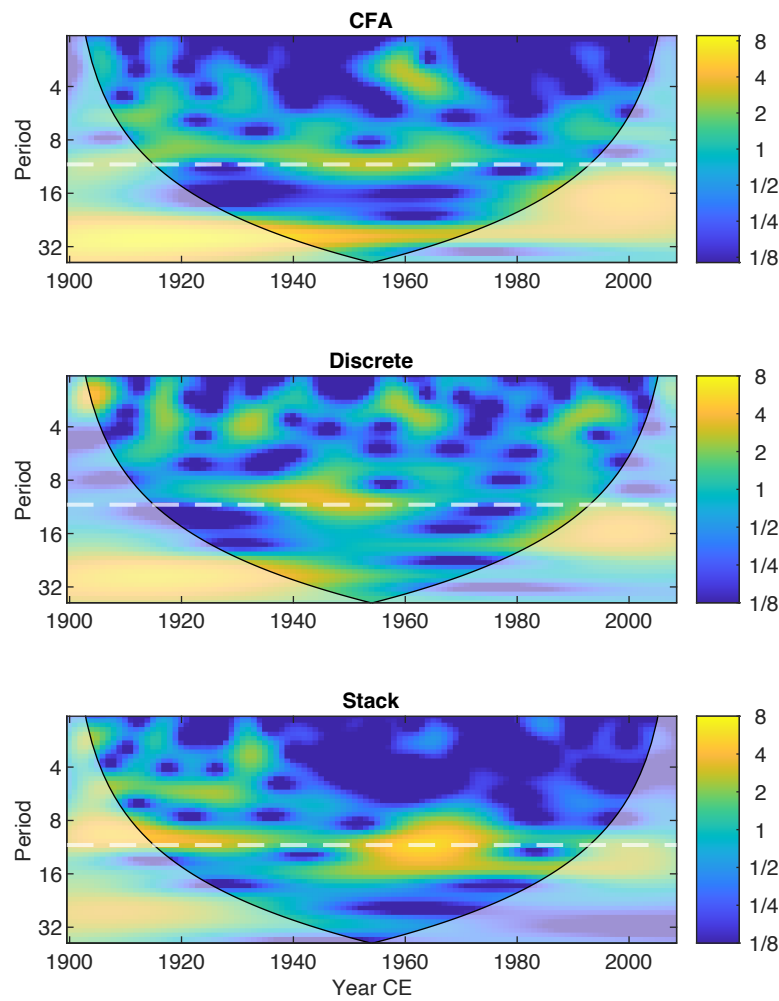
The results of the analysis are summarized in Table 1. The increase in 1994 can be linked to a possible volcanic influence on the ^{10}Be transport and/or deposition (Baroni et al. 2011). We do not find any evidence of the influence of the El Chichón eruption in the Antarctic stack, although the eruption left an imprint in some Antarctic sulfate
 330 records (Sigl et al., 2015). We find an increase relative to the baseline of 4.3% in 1957, lower than the increase modeled by Mekhaldi et al. (2021) and the increase recorded in the stack of records from Greenland.

Records	1-2 σ	2-3 σ
^{10}Be EGRIP S6 - CFA	1957, 1963, 1965, 1992, 1993, 1994, 2007	1960, 1991
Including transport:	1963, 1965, 1991, 1992, 1994, 2007	1960, 1993
^{10}Be EGRIP S6 - discrete	1957, 1960, 1967, 1982, 1992, 1994	1991, 2008
Including transport:	1960, 1967, 1992, 1994	1991, 2008
^{10}Be stack – Greenland	1963, 1967, 1968, 1991, 1992, 2008	1957, 1965, 1983
Including transport:	1963, 1964, 1991, 1993, 1994, 2008	1965, 1983, 1992
^{10}Be stack - Antarctica	1957, 1958, 1967, 1968, 1969, 1980, 1981, 1991, 1994	1966

Table 1. Summary of the years during which we find standard scores exceeding 1σ from the GCR-production
 335 corrected EGRIP S6 records, Greenland and Antarctic stacks.

6 The 11-year solar cycle

To assess the suitability of CFA meltwater samples to reconstruct solar activity, we investigate the presence of the 11-year solar cycle in the data. The wiggles of the 11-year solar cycles are estimated to cause a variability of about $\pm 15\%$ to $\pm 35\%$ in the ^{10}Be records from Greenland and Antarctica (e.g. Baroni et al., 2011; Heikkilä et al., 2009; Mekhaldi et al., 2021; Paleari et al., 2022). To investigate the presence of the 11-year cycle in the data, we performed continuous wavelet transform (Grinsted et al., 2004) on the ^{10}Be records from CFA and discrete samples, and on the global ^{10}Be stack calculated using the records from NGRIP, NEEM, Dye 3, Renland, Das2, DML, DSS, Vostok and Concordia. The continuous wavelet power spectrums are shown in Figure 9.



345

Figure 9. Continuous wavelet power spectrum (Grinsted et al., 2004) of the ^{10}Be concentration records from CFA samples (top panel) and discrete samples (middle panel) from EGRIP S6, and of the global stack (bottom panel) calculated including the records from NGRIP, NEEM, Dye 3, Renland, Das2, DML, DSS, Vostok and Concordia. The white dashed line denotes the frequency of the 11-year cycle.

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Figure 9 shows that the 11-year solar cycle is present in the EGRIP S6 records. In particular, the figure highlights that the 11-year solar cycle is preserved in the CFA and discrete records spanning the period 1900-1960, especially so in the CFA record. Conversely, it is not observed in the more recent part of the record, likely due to higher background noise. As expected from the increased signal-to-noise ratio, Figure 9 also confirms that the 11-year cycle is well preserved throughout the global stack. Moreover, Figure 9 highlights that the discrete ^{10}Be record is characterized by more prominence of higher-frequency noise (period < 4 years), as opposed to the CFA ^{10}Be record, likely due to the smoothing caused by analytical issues inherent to the CFA system (Erhardt et al., 2022; Kaufmann et al., 2008; Mekhaldi et al., 2017; Sigg et al., 1994).

There also exists a 22-year cycle whereby the Sun reverses polarity every 11-year cycle. This is evidenced in neutron monitor measurements that show that the GCR flux at Earth has a polarity-dependent shape. More specifically, 11-year cycles take the form of a peak under negative polarity and the form of a plateau under positive polarity (e.g., Poopakun et al., 2022; Lockwood et al., 2001). Bearing this difference in mind, we separate our analysis between periods characterized by negative and positive polarity (Fig. 10a and 10b, respectively). Because we are now interested in the shape of the cycles as seen in neutron monitor data that are anticorrelated with solar activity, we now consider solar maxima as the onset of solar cycles to investigate the peak shape in the radionuclide data. The ^{10}Be records from Greenland (except EGRIP S6) and Antarctica were split into time windows for each period and normalized to their respective mean, and therefore used to create a global stack (blue line). The blue envelopes denote the standard deviation of the records. The stacks obtained from the modeled ^{10}Be production rate from Mekhaldi et al. (2021) are shown as a dashed line for comparison. The global stack is compared to the CFA stack (magenta) and discrete firn samples stack (green) from EGRIP S6.

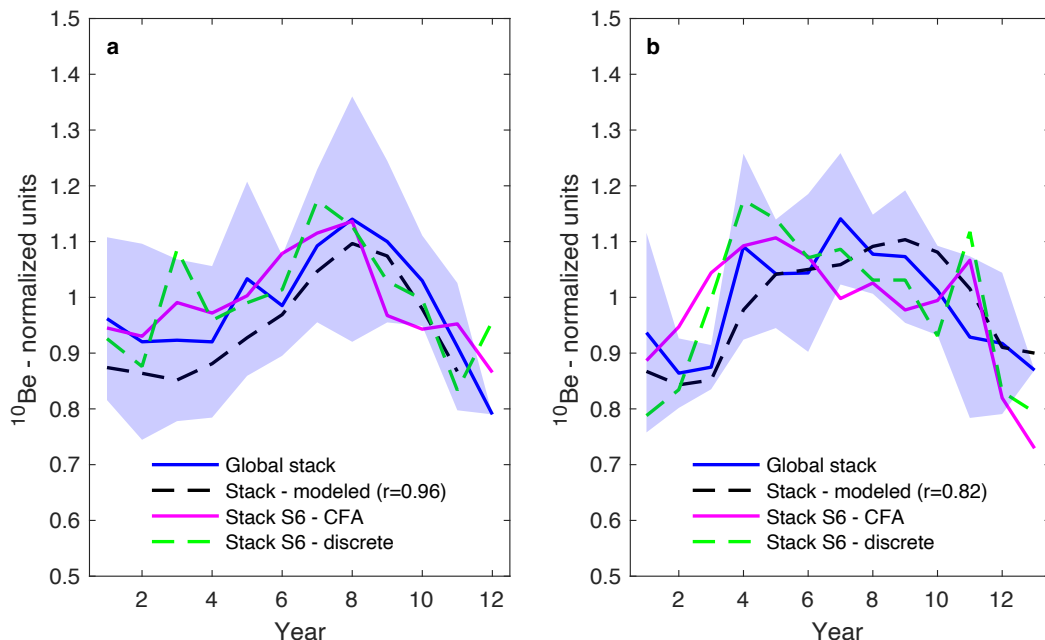


Figure 10. Panel a shows the global stack of the periods of negative polarity of the Sun's magnetic field, as indicated by neutron monitors (in blue, the blue envelope represents the standard deviation). The stacks from CFA and discrete samples from EGRIP S6 are shown as a solid magenta line and a dashed green line, respectively. The global stack includes the ^{10}Be records from NEEM, NGRIP, Dye 3, Das2, Renland, DSS, Vostok, DML and Concordia cores. The stacks are compared to the modeled global annual production rate (Mekhaldi et

al., 2021; dashed black line, the correlation coefficient with the global stack ($p < 0.01$) is denoted in the legend). Panel b shows the same for periods of positive polarity.

380 The global and modeled stacks agree very well in both shape and amplitude for both periods of negative (Fig. 10a) and positive (Fig. 10b) polarity. Similarly, the stacks from discrete and CFA samples agree very well ($r=0.74$ for negative polarity (Fig. 10a) and $r=0.87$ for positive polarity (Fig. 10b), $p < 0.01$), although the S6 CFA stacks are smoother than the discrete stacks. However, CFA and discrete stacks agree well, showing that the two records show a very similar signal. This result is promising, as it shows the suitability of CFA samples for solar activity
385 reconstructions.

7 Conclusions

In this study, we present the ^{10}Be record obtained from excess water from CFA measurements of the EGRIP S6
390 firn core. The record, spanning from 1900 to 2008, has been compared to another ^{10}Be record obtained from discrete firn samples from the same core. Although some smoothing due to analytical issues inherent to the CFA system may be present in the shallower part of the core, the CFA ^{10}Be record agrees well with the ^{10}Be record from discrete firn samples and with other ^{10}Be records from Greenland.

By subtracting the global ^{10}Be production rate variability caused by the 11-year solar cycle, we investigate the
395 possible causes of excursions in the EGRIP S6 ^{10}Be records and in the stacks of Greenlandic and Antarctic records. We find that values exceeding 1σ in the standardized residuals are coeval with the timing of some of the major volcanic eruptions that are also detected in sulfate records (Zielinski et al., 1997; Sigl et al., 2015), such as the Agung (1963) and Pinatubo (1991) eruptions. Moreover, values exceeding 1σ in the residuals in 1957 may be consistent with the occurrence of GLE no.05. However, an increase in sulfate concentrations was detected in
400 records from Greenland according to Sigl et al. (2013), illustrating the challenges to separate the causes for small radionuclide excursions. We find an increase of 16% in the stack from Greenland and in 4.3% the stack from Antarctica in 1957. This study shows that presently volcanic eruptions and GLEs influences cannot be unambiguously distinguished from the internal variability in annually-resolved ^{10}Be records, in agreement with previous studies (e.g. Pedro et al., 2012; Mekhaldi et al, 2021).

405 Finally, we analyzed how the 11-year solar cycles are preserved in the discussed records. Our results show that the 11-year solar cycles are well preserved in CFA and discrete ^{10}Be records from EGRIP S6. By stacking periods of different polarity of the Sun's magnetic field separately it is possible to see their different shape in ^{10}Be data, as expected according to the flux of the GCR at Earth. The stacks from CFA ^{10}Be samples agree well with the stacks from discrete ^{10}Be samples from EGRIP S6, thus showing that CFA excess water samples are suitable for
410 the reconstruction of solar activity. These results open the possibility of collecting continuous and high-resolution ^{10}Be records with a more time-efficient sampling and preparation, while saving an important portion of valuable ice for other measurements.

Data availability

415 The ^{10}Be data generated in this study is provided in the supplementary material.

Author contribution

R.M and C.P. designed the project. C.P. sampled and prepared the EGRIP CFA ^{10}Be samples, performed the analysis, and wrote the manuscript. F.M. contributed to the interpretation of the data. M.C. contributed with the measurement of ^{10}Be samples. T.E. helped with the collection of the EGRIP CFA samples and the interpretation of the data. M.H. helped with the sampling of the EGRIP S6 core. All authors contributed to the discussion.

Competing interests

The authors declare that they have no conflict of interest.

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