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Assessment of the record of the 1982 El Chichón eruption as preserved in Greenland snow

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Abstract. Variability in the SO$_4^{2-}$ and Cl$^-$ time series for the 1980s from 12 shallow snow pits across the Greenland ice sheet is used to evaluate the record of the 1982 El Chichón eruption and the potential for recording a moderate northern equatorial eruption in a single Greenland ice core. Composition of volcanic glass found in spring 1983 snow in one of the pits in the Summit region matches that from El Chichón glass, thereby verifying the deposition of material from the eruption. High Na$^+$ and Cl$^-$ concentrations in this same layer probably represent deposition of the reaction products of halite and H$_2$SO$_4$ as observed in the stratosphere following the eruption. These findings and the presence of a Cl$^-$ signal in five of the other pits indicate that the Cl$^-$ aerosol component of some eruptions has the potential to remain aloft for at least 1 year after the eruption. Some of these aerosols may be adsorbed onto tephra particles. Distinct SO$_4^{2-}$ peaks that can be confidently linked to El Chichón were found in only 50% of the records developed through subseasonal sampling. However, in several other pits an elevated baseline, thought to represent more lengthy periods of El Chichón aerosol deposition as opposed to deposition from a single snowfall event, were observed. Smoothing of the original data by the calculation of annual SO$_4^{2-}$ flux resulted in the presence of high flux values between 1982 and 1984 (years thought to be affected by El Chichón aerosol deposition) in 9 of the 12 pits. These results suggest that a single ice core from anywhere in Greenland may record a signal from a northern equatorial eruption of magnitude similar to that of El Chichón about 75% of the time; this is despite the overall high levels of SO$_4^{2-}$ deposition from anthropogenic sources that now make identification and quantification of the volcanic SO$_4^{2-}$ portion of the record more difficult than obtaining the same data for preindustrial volcanism. Nevertheless, composite records from all the pits sampled yielded stratospheric loading (~20 Mt) and optical depth ($\tau = 0.13$) estimates similar to stratospheric and satellite-based measurements following the eruption. Equally high SO$_4^{2-}$ concentration and flux values in snow from 1980 to 1982 reflect deposition from the series of middle to high northern latitude volcanic eruptions 1979-1981.

1. Introduction

Between March 28 and April 4, 1982, the plinian eruptions of El Chichón, Mexico (17ºN, 121ºW), injected large quantities of debris into the northern equatorial stratosphere with the eventual dispersion over the entire globe. Estimates of the amount of SO$_2$ released vary from 8 [Bluth et al., 1993] to 13 Mt (i.e., 10$^{12}$ g or Tg) [Evans and Kerr, 1983], leading to the initial production of about 12 [McCormick et al., 1995] to 20 Mt of H$_2$SO$_4$ [Evans and Kerr, 1983; Hofmann and Rosen, 1983; Rampino and Self, 1984]. Thus El Chichón may have injected more sulfur-rich aerosols into middle levels of the atmosphere than any eruption in this century except for the 1991 eruption of Mt. Pinatubo. Subsequently, it was expected that evidence of the atmospheric loading of the El Chichón aerosols would be preserved in polar snow through the presence of high peaks in SO$_4^{2-}$ [e.g., Legrand and Delmas, 1987; Mayewski et al., 1993; Zielinski et al., 1994], and particularly in Arctic snow, given the 17ºN location of the volcano. Sur-
 surprisingly, there is no discernible signal in the biennial \( \text{SO}_4^{2-} \) record from the Greenland Ice Sheet Project 2 (GISP2) ice core that can be confidently linked to the El Chichón eruption [Zielinski et al., 1994; Zielinski, 1995].

To determine whether or not there is evidence of the El Chichón eruption in Greenland snow and ultimately to assess the consistency (both temporally and spatially) and magnitude of the signal, we evaluated the variability in snow chemistry at a subseasonal resolution in a series of shallow snow pits across Greenland. The majority of the pits are located in the Summit region (i.e., within 75 km of the GISP2 drilling site), although we include the evaluation of several pits in northern and southern Greenland (Figure 1). The results of this study are used to answer three interrelated questions in the use of polar snow and ice cores to evaluate past volcanism.

The first critical question we address is how likely it is for the chemical signal of a moderate-sized equatorial eruption to be present in a single ice core collected from Greenland, assuming that the chemical record we see in the snow pits would be identical to that found in an ice core collected at the same site. In order for a volcanic signal to exist in an ice core, several criteria must be met. Circulation patterns must bring the volcanic cloud over the coring site followed by deposition of the aerosols via a snow event(s). Finally, there must be minimal postdepositional modification of the volcanic acids that would result in a missing signal for a particular eruption (e.g., wind scouring without redeposition at the site in question). Clausen and Hammer [1988] used multiple Greenland ice cores to evaluate the variability in the Icelandic Laki (A.D. 1783; \( \sim 200 \text{ Mt H}_2\text{SO}_4 \)) and Indonesian Tambora (A.D. 1815; \( \sim 200 \text{ Mt H}_2\text{SO}_4 \); see also Rampino et al. [1988]) eruptions, but these are the largest sulfur-producing eruptions during historical time (i.e., an order of magnitude greater than El Chichón). As a result, signals for these two eruptions have been observed in all northern hemisphere ice cores that we are aware of. Robock and Free [1995] essentially stacked volcanic records from eight northern hemisphere ice cores that span the last 150 years and found that only the Alaskan Katmai eruption (A.D. 1912; 30-40 Mt H\(_2\)SO\(_4\) [e.g., McCormick et al., 1995]) is consistently preserved in these records. Katmai is directly upwind from Greenland and other ice caps in the Arctic, and this situation would enable aerosols to be readily transported to those deposition sites, with the probability of some tropospheric transport. Although the El Chichón eruption is moderate in the overall scheme of climate-forcing volcanism, it still may have cooled global climate by \( 0.35^{\circ}\text{C} \) once the warming effects of the coincident El Niño were accounted for [Angell, 1988]. Identifying the likelihood of missing similar magnitude, equatorial eruptions in ice cores is important in the evaluation of the interannual to decadal variability of past climate. Our results can best be applied to evaluating the climatic impact of past volcanism from the Central America/Caribbean region, areas whose volcanoes are known to have been much more active and explosive in prehistoric time in comparison with the present [e.g., Siebe et al., 1996].

The second question of importance is how much the signal:noise ratio of a volcanic \( \text{SO}_4^{2-} \) peak decreases with detailed, subseasonal sampling. To address this question, we will compare the annual signals from these same pits to test whether the years following the El Chichón eruption are characterized by higher overall \( \text{SO}_4^{2-} \) deposition in comparison with the years before the eruption or beyond the suspected residence time of the aerosols in the atmosphere. In essence, the com-
comparison of annual signals versus subseasonal signals is a form of smoothing of the original data set, which should increase the signal-to-noise ratio. We will be able to determine whether such a smoothing results in a more consistent signal across the Greenland ice sheet from the El Chichón eruption, even though there was not a distinct El Chichón signal in the highly smoothed, biannual sampling of the GISP2 core.

The third question we are able to provide insight into is how well we can use recent eruptions to calibrate the volcanic record in ice cores. For instance, using the snow and ice records of the El Chichón and Pinatubo eruptions to quantify the stratospheric loading of each eruption [e.g., Hammer et al., 1980; Zielinski, 1995] would enable direct comparisons to be made with the atmospheric loading estimates derived from direct stratospheric [e.g., Hofmann and Rosen, 1983] and satellite-based measurements [e.g., Bluth et al., 1993]. The potential problem with this scenario is that for recent eruptions, anthropogenic sulfur emissions may be of such a magnitude that the $SO_4^{2-}$ signal in recent snow from moderate eruptions like El Chichón may not be sufficient to yield a distinct spike in the record. By evaluating the series of snow pits across Greenland and developing composite concentration and flux records we will be better able to identify the range in values of volcanic deposition on the Greenland ice sheet for comparison with the values obtained for eruptions prior to the increase in $SO_4^{2-}$ deposition from anthropogenic sources [Zielinski, 1995]. Adding support to the idea that high levels of anthropogenic $SO_4^{2-}$ deposition on the ice sheet [e.g., Mayewski et al., 1986, 1990] may inhibit the detection of a volcanic signal is the absence of a clear signal from the deposition of Pinatubo aerosols (~2-3 times greater mass than El Chichón [McCormick et al., 1995]) in snow deposited on Greenland and on other Arctic ice caps during the years after the eruption. However, there is a Pinatubo signal in snow at the south pole [Dibb and Whitlow, 1996], snow that is not as affected by anthropogenic sources as is Greenland [Whitlow et al., 1992]. Should we be able to extract a composite volcanic signal for El Chichón, we will estimate the stratospheric loading of $H_2SO_4$ aerosols and the resulting optical depth [e.g., Zielinski, 1995] for comparison with previously published estimates, including satellite-derived estimates.

2. Aerosol Distribution and Other Products of the El Chichón Eruption

The expected timing of the deposition of any chemical signal in Greenland snow from El Chichón may be inferred by the global distribution of the aerosol cloud. (Note that a special issue of Geophysical Research Letters, volume 10, 1983, presented many of the initial results of the atmospheric impact of the eruption.) The main aerosol cloud circumnavigated the globe around the latitude of the eruption within about 3 weeks [Robock and Matson, 1983], eventually extending throughout the northern tropics by the end of May 1982. At that time the cloud consisted of two separate layers: one from the tropopause (12-16 km) to about a height of 20 km and the other centered around 25 km [Hofmann and Rosen, 1983]. Most of the cloud above 20 km remained in the tropics until late August and September 1982, when some intermittent northward transport occurred [Hofmann and Rosen, 1983]. Eventually, atmospheric transport mechanisms dispersed the cloud to the point where it covered the entire northern hemisphere by early 1983 [Hofmann, 1987]. Consequently, we did not expect to see aerosol deposition in Greenland until possibly fall 1982. However, the lower aerosol layer reached the midlatitudes (41°N) by June 1982 [Hofmann and Rosen, 1983]; thus some aerosols could have reached Greenland during summer/fall 1982, given ideal circulation patterns.

In addition to the sulfur component, several other products of importance to our study were released during the El Chichón eruption. Mankin and Coffey [1984] reported increased HCl levels in the stratosphere over northern midlatitudes for several months following the eruption. Using this information, they suggested a total stratospheric loading of $0.04 \times 10^8$ Mt HCl from the eruption itself, a 40% increase over preeruption levels. Woods et al. [1985] used scanning electron microscopy (SEM) and X-ray energy spectroscopy on particles collected in the stratosphere over the western United States during April and May 1982 to identify the presence of halite in the eruption cloud. Halite was no longer observed after July 1982, as most of the halite particles probably converted to $Na_2SO_4$ and HCl through reactions with $H_2SO_4$ within the first month following the eruption [Woods et al., 1985]. Complete conversion probably occurred within 8 months after the eruption. As a result, evidence of aerosol deposition on the Greenland ice sheet from El Chichón may be through peaks in both $SO_4^{2-}$ and Cl$^-$. The mineralogical component of the eruption (i.e., tephra) remained a significant part of the stratospheric aerosol cloud over the Americas until October 1982 [Gooding et al., 1983]. The northward spread of tephra over almost all of the northern hemisphere may have occurred within 6 months following the eruption. Tephra was observed consistently at middle to high latitudes (45-75°N) into late October, although parts of the ash component spread to 60°N by July 1982. About 85% of the ash sampled at latitudes of >45°N in October of 1982 was in the 2-5 μm (intermediate diameter) size range, with another 11% in the 5-10 μm size range [Gooding, 1983]. The timing of the presence of tephra in the Arctic atmosphere and deposition on the Greenland ice sheet was documented in two separate studies. De Angelis et al. [1985] identified the presence of volcanic glass from El Chichón in snow collected in June 1983 from the Dye 3 site in southern Greenland. How-
ever, they suggest that there may have been deposition as early as June 1982. Shapiro et al. [1984] collected mineral matter believed to be from El Chichón during a tropopause fold event over west central Greenland on March 23, 1983. Small droplets (diameter of < 0.5 μm) thought to be H₂SO₄ from El Chichón were found on the same filter. On the basis of these data it is possible that volcanic glass traveling in the stratosphere may have been deposited on the Greenland Ice Sheet anytime between midsummer 1982 and the spring or summer 1983. Any tephra transported tropospherically could possibly have reached Greenland within several weeks of the eruption, given ideal synoptic conditions. Identifying tephra from El Chichón in the same layer as a SO₂⁻ or Cl⁻ peak verifies the presence of aerosols from the eruption in Greenland snow.

3. Methods of Analysis

All snow pit samples were collected with personnel in clean suits to prevent possible contamination following the procedures outlined by Mayewski et al. [1987]. Samples remained frozen until the time of analysis. Anions and cations were analyzed using a Dionex model 2010 ion chromatograph. Duplicate samples and blanks were frequently analyzed to detect any possible contamination, as described by Buck et al. [1992]. All SO₄²⁻ values in this study have been corrected for the sea-salt component, which is usually <5% of total SO₄²⁻. Insoluble microparticle concentrations and size distributions between 0.65 and 12.88 μm were obtained with an Elzone 280PC particle counter in a class 100 clean room. Meltwater samples were filtered through a Nucleopore polycarbonate membrane filter (0.4 μm pore size) in an attempt to locate volcanic glass. Individual particle analyses were undertaken with an automated Jeol JXA-8600 electron microprobe. The chronology developed for all pits in this study was based on the seasonal variability (i.e., summer peaks and winter troughs) of specific parameters such as δ¹⁸O and H₂O₂. A total of nine pits were analyzed in the Summit region, one pit was analyzed from northwestern Greenland, and two pits were analyzed from southern Greenland (Figure 1). Snow density measurements were taken on the same layer as chemistry and microparticle samples in pit 90-1 to determine the annual flux of specific species. Density gradually increased from mean values of 315 kg/m³ in the upper 250 cm to 385 kg/m³ between 250 and 500 cm to 450 kg/m³ between 500 and 600 cm. This general trend is similar to density trends observed in surface snow throughout Greenland (R. Alley, personal communication, 1993); thus these mean values were used to determine flux values in all other pits analyzed. Resulting errors for flux values are thought to be between 5% and 10%.

4. Glaciochemical Time Series

4.1. Pit 90-1, Summit Region

One of the most detailed analyses in this study was on the 6 m pit, 90-1, located at the Atmospheric Sampling Camp (ATM), southern Summit region (Figure 1). Sampling interval in this pit was 5 cm, yielding about 15 samples/yr for chemistry and insoluble microparticle analysis. The record extends from summer 1990 to spring/early summer 1981 based on counting of peaks in δ¹⁸O and H₂O₂ (Figure 2). Spring 1982, the time of the El Chichón eruption, is represented by snow at about the 550 cm depth.

Several interesting signals are prevalent in the time series of the various ions and insoluble microparticles (Figures 2b and 2c). Initially, our interest was drawn to the very high microparticle peak around 500 cm (corresponding to deposition late in 1982 or early 1983, Figure 2b) because the high microparticle loading could be indicative of a large amount of volcanic glass deposition [e.g., Fiacco et al., 1993]. Although the SEM/microprobe analysis of meltwater from this section of the pit indicated that some of these particles were contamination from the stainless steel sampling tool, two populations of glass shards were found (Figure 3 and Table 1).
Glass A has a rhyolitic to trachydacitic composition with a high alkalic component (Na$_2$O and K$_2$O; Table 1) characteristic of the El Chichón eruption. Glass B is much different, having a highly rhyolitic composition (Table 1). Glass A shards were much smaller on average, with long axes often in the 2-4 μm size range, whereas glass B shards had long axes that were often 10-20 μm (Figure 3).

Verification of the source of glass A comes from direct comparison with the composition of El Chichón glass that we analyzed, as well as previous analyses of El Chichón glass (Figure 4 and Table 1). The composition of individual glass grains in pit 90-1 consistently overlaps that of El Chichón glass, and the mean values are almost identical (Figure 4). Further, the mean composition of the glass we analyzed closely matches the mean value of glass analyzed from El Chichón pumice [Luhr et al., 1984], thus verifying El Chichón as the source. A late fall 1982 to early spring 1983 time of glass deposition also fits well with the timing of the northward distribution of tephra [Gooding et al., 1983] and the possible presence of El Chichón mineral matter in a tropopause fold event over Greenland in spring 1983 [Shapiro et al., 1984]. The presence of El Chichón glass in the same layer as large spikes in Na$^+$ and Cl$^-$ (Figure 2c) has some interesting ramifications that we elaborate on below.

Identification of the source of glass B in this same layer is problematic. Many of the other large eruptions in 1981 (e.g., Alaid, Pagan, and Nyamuragira; see Table 2) are characterized by a basaltic composition [i.e., McClelland et al., 1989], thereby eliminating them as a possible source. The 1982 eruption of Galunggung is another possible source, but this volcanic system also is generally less silicic than our glass B [i.e., McClelland et al., 1989]. Consequently, we feel that the most plausible source of glass B is the rapid tropospheric transport of

### Table 1. Average Major Elemental Composition of Volcanic Glass Found at the 495-500 cm Depth of the 90-1 Snow Pit Compared With Glass From El Chichón

<table>
<thead>
<tr>
<th></th>
<th>Pit 90-1</th>
<th>Pumice</th>
<th>Pumice</th>
<th>Stratosphere$^b$</th>
<th>Snow</th>
<th>Pit 90-1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glass A</td>
<td>Glass A</td>
<td>Glass A</td>
<td>Glass A</td>
<td>(n=15)</td>
<td>(n=6)</td>
<td>(n=5)</td>
</tr>
<tr>
<td>(n=8)</td>
<td>(n=9)</td>
<td>(n=16)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SiO$_2$</td>
<td>71.2 (1.0)$^c$</td>
<td>71.1 (1.0)</td>
<td>68.0 (1.7)</td>
<td>69.4 (1.3)</td>
<td>69.2 (0.5)</td>
<td>78.1 (1.2)</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>0.3 (0.3)</td>
<td>0.4 (0.1)</td>
<td>0.3 (0.1)</td>
<td>0.3 (0.1)</td>
<td>0.4 (0.2)</td>
<td>0.2 (0.2)</td>
</tr>
<tr>
<td>Al$_2$O</td>
<td>14.8 (0.3)</td>
<td>15.5 (0.6)</td>
<td>15.9 (0.4)</td>
<td>17.9 (0.6)</td>
<td>17.1 (0.2)</td>
<td>11.9 (0.4)</td>
</tr>
<tr>
<td>Fe$_2$O$_3$</td>
<td>1.7 (0.6)</td>
<td>1.8 (0.2)</td>
<td>1.6 (0.2)</td>
<td>1.4 (0.4)</td>
<td>1.9 (0.5)</td>
<td>1.9 (0.4)</td>
</tr>
<tr>
<td>MgO</td>
<td>0.4 (0.3)</td>
<td>0.3 (0.2)</td>
<td>0.3 (0.1)</td>
<td>0.4 (0.1)</td>
<td>0.4 (0.1)</td>
<td>0.1 (0.2)</td>
</tr>
<tr>
<td>CaO</td>
<td>1.8 (0.5)</td>
<td>2.0 (0.2)</td>
<td>2.1 (0.2)</td>
<td>1.9 (0.3)</td>
<td>1.8 (0.2)</td>
<td>1.5 (0.3)</td>
</tr>
<tr>
<td>Na$_2$O</td>
<td>4.4 (0.6)</td>
<td>2.9 (0.9)</td>
<td>4.6 (0.5)</td>
<td>3.9 (0.8)</td>
<td>4.5 (0.7)</td>
<td>3.4 (1.4)</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>5.2 (0.7)</td>
<td>6.0 (0.4)</td>
<td>5.1 (0.2)</td>
<td>4.7 (0.4)</td>
<td>4.8 (0.3)</td>
<td>3.0 (1.0)</td>
</tr>
</tbody>
</table>

$^a$Reprinted from *Journal of Volcanology and Geothermal Research*, 23, Luhr et al., The 1982 eruptions of El Chichón volcano, Chiapas, Mexico: Mineralogy and petrology of the anhydrite-bearing pumices, Copyright 1984 with kind permission from Elsevier Science-NL, Sara Burgerhartstraat 25, 1055 KV Amsterdam, The Netherlands. Total Fe reported as FeO.

$^b$Reprinted with permission from *Nature* (De Angelis et al., 317, 52-54), Copyright 1985, Macmillan Magazines Limited and from M. De Angelis. Total Fe reported as FeO.

$^c$Value in parentheses is 1σ.
Figure 4. Ternary diagram comparing the composition of the volcanic glass found in pit 90-1 with that from the El Chichón eruption.

tephra to Greenland from the small May 22, 1983, eruption of Bezymianny, Kamchatka. Eruptions of Bezymianny over the last few decades appear to be becoming more silicic than prior eruptions, and in fact, the 1956 event appears to have tapped a rhyolitic magma source [Palais and Sigurdsson, 1989]. Furthermore, the overall larger size of glass B shards compared with those of glass A supports a more proximal or directly upwind source, like Bezymianny, for glass B. Because evaluation of the products of the 1956 Bezymianny eruption indicate that this volcanic system contains a source of Cl⁻ [Palais and Sigurdsson, 1989], we cannot rule out completely the possibility that some of the chemical signal found in this section of pit 90-1 is a result of the 1983 Bezymianny eruption. However, the small size of that eruption (10⁷ m³ ash erupted to only 5-6 km altitude) and the probable ESE transport of material [McClelland et al., 1989] lead us to believe that most of the chemical signal in 1982/1983 snow in pit 90-1 is related to El Chichón aerosol deposition.

Identifying the presence of El Chichón debris in the Summit region of Greenland through the existence of volcanic glass enables us to more readily characterize the SO₄²⁻ and Cl⁻ signals from the eruption. In the case of the SO₄²⁻ signal the first impression is that there is no anomalous spike in pit 90-1 (Figure 2b). It is only the large spike around spring 1987 that is very prominent. That enhancement could be related to aerosols from the 1986 eruptions of St. Augustine and Chikurachki in the northern Pacific (Table 1). However, it should be noted that there is a broad, but subdued, peak in SO₄²⁻ that corresponds to fall/winter 1982 and possibly very early spring 1983 (i.e., just prior to the time when El Chichón tephra was deposited). This peak is not larger

Table 2. Explosive Eruptions That May be Responsible for Peaks in SO₄²⁻ and Cl⁻ Recorded in Greenland Snow Pits During the 1980s

<table>
<thead>
<tr>
<th>Year</th>
<th>Date</th>
<th>Volcano</th>
<th>Latitude</th>
<th>Longitude</th>
<th>VEI</th>
</tr>
</thead>
<tbody>
<tr>
<td>1979</td>
<td>Feb. 8</td>
<td>Westdahl, Alaska</td>
<td>54.5°N</td>
<td>164.6°W</td>
<td>3?</td>
</tr>
<tr>
<td>1979</td>
<td>Nov. 13</td>
<td>Sierra Negra, Galapagos</td>
<td>0.8°S</td>
<td>91.2°W</td>
<td>3</td>
</tr>
<tr>
<td>1980</td>
<td>May 18</td>
<td>St. Helens, Washington</td>
<td>46.2°N</td>
<td>122.2°W</td>
<td>5</td>
</tr>
<tr>
<td>1980</td>
<td>Aug. 7</td>
<td>Gareloi, Alaska</td>
<td>51.8°N</td>
<td>178.8°W</td>
<td>3?</td>
</tr>
<tr>
<td>1981</td>
<td>March 24</td>
<td>Okmok, Alaska</td>
<td>53.4°N</td>
<td>168.1°W</td>
<td>3?</td>
</tr>
<tr>
<td>1981</td>
<td>April 30</td>
<td>Alaid, Kuril Islands</td>
<td>50.9°N</td>
<td>155.6°E</td>
<td>4</td>
</tr>
<tr>
<td>1981</td>
<td>May 15</td>
<td>Pagan, Mariana Islands</td>
<td>18.1°N</td>
<td>145.8°E</td>
<td>4</td>
</tr>
<tr>
<td>1981</td>
<td>Dec. 25</td>
<td>Nyamuragira, Zaire</td>
<td>1.4°S</td>
<td>29.2°E</td>
<td>3</td>
</tr>
<tr>
<td>1982</td>
<td>March 28</td>
<td>El Chichón, Mexico</td>
<td>17.4°N</td>
<td>93.2°W</td>
<td>4+</td>
</tr>
<tr>
<td>1982</td>
<td>April 3</td>
<td>El Chichón, Mexico</td>
<td>17.4°N</td>
<td>93.2°W</td>
<td>5</td>
</tr>
<tr>
<td>1982</td>
<td>May 17</td>
<td>Galunggung, Java</td>
<td>7.3°S</td>
<td>108.1°E</td>
<td>4</td>
</tr>
<tr>
<td>1983</td>
<td>July 23</td>
<td>Colu (Uma Uma), Indonesia</td>
<td>0.2°S</td>
<td>121.6°E</td>
<td>4</td>
</tr>
<tr>
<td>1985</td>
<td>Nov. 13</td>
<td>Ruiz, Columbia</td>
<td>4.9°N</td>
<td>75.3°W</td>
<td>3</td>
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<tr>
<td>1986</td>
<td>March 27</td>
<td>St. Augustine, Alaska</td>
<td>59.4°N</td>
<td>155.4°W</td>
<td>4?</td>
</tr>
<tr>
<td>1986</td>
<td>Nov. 20</td>
<td>Chikurachki, Kuril Islands</td>
<td>50.3°N</td>
<td>155.4°E</td>
<td>4?</td>
</tr>
</tbody>
</table>

From Simkin and Siebert [1994]. VEI, volcanic explosivity index.
than the annual spring peaks in SO$_4^{2-}$ [Whitlow et al., 1992] that are clearly observed along the entire record in this pit. In fact, this bump has the appearance of an additional spring peak, but it is clear from the $\delta^{18}$O and H$_2$O$_2$ records (Figure 2a) that it is not such a peak. We suggest that much of the SO$_4^{2-}$ in this broad peak is associated with deposition of the earliest El Chichón aerosols that reached the polar regions. This suggestion is supported in our later discussion on the annual flux of SO$_4^{2-}$.

Perhaps the most distinct glaciochemical signals in this pit are the coincident peaks in both Na$^+$ and Cl$^-$ in the same sample that contained the volcanic glass (Figures 2b and 2c). Interestingly, the Na:Cl $\mu$eq ratio for this sample is 1:1, not 0.85, as would be the case if the Na$^+$ and Cl$^-$ were solely from sea-salt aerosols. The most likely cause for these high Na and Cl concentrations is the deposition of halite or, more likely, deposition of the by-products of halite dissolution following reactions with H$_2$SO$_4$ in the stratosphere (i.e., Na$_2$SO$_4$ and HCl [Woods et al., 1985]). The presence of stratospheric halite following the eruption could be the source of the halite products, and once these products were able to reach the northern polar regions, the very low relative humidity may have allowed the HCl component to stay gaseous for longer periods of time (R. Chuan, personal communication, 1992). Such a scenario would have prevented complete removal before deposition on the ice sheet. Alternatively, Varekamp et al. [1984] indicated that there was abundant Na (mean of 335 ppm) and SO$_4$ (mean 780 ppm) within materials adsorbed onto tephra grains from the El Chichón eruption. Thus deposition of the Na$_2$SO$_4$ and HCl species recorded in the 90-1 pit could be a result of adsorption onto the tephra grains we located. Rose [1977] also observed the scavenging of HCl by tephra grains following the 1974 Fuego eruption, although his samples were close to the vent. Moreover, he estimated that only 17% of the Cl$^-$ emitted during the eruption was adsorbed onto tephra grains and thus quickly scavenged from the volcanic plume. Regardless of the actual process, it is apparent that there is a well-preserved Cl$^-$ signal in this particular pit from the El Chichón eruption.

4.2. Other Snow Pits in the Summit Region

Six pits dug during summer 1987 have records that extend at least back to summer or early spring 1982. The record for pits 87-2, 87-4, and 87-5 (southern Summit region, Figure 1) end in early 1982, while the record for pit 87-1 extends back to summer 1977, and the records for pits 87-6 and 87-7 (northern Summit region, Figure 1) extend back to summer 1979 (Figure 5). The two pits in the northern Summit region dug in 1990 (90-2 and 90-3; see Figure 1) have records that extend back to summer 1982 and to early 1983, respectively (Figure 5).

There is evidence of the deposition of both SO$_4^{2-}$ and Cl$^-$ aerosols from El Chichón in the many pits excavated in the Summit region, although the characteristics of these signals are not identical. This variability ranges from a lack of an anomalous signal, to more of a broad signal, as in pit 90-1, to a distinct spike above that of the seasonal peaks between late 1982 and early to late 1984 (Figure 5). A 2-year lag in deposition of aerosols in Greenland from an equatorial eruption is not unrealistic [Zielinski, 1995], especially considering that stratospheric SO$_4^{2-}$ levels over the western United States were still 5 times background levels in January 1984 [Snetsinger et al., 1987]. Consequently, we infer that any peaks found in snow deposited in 1984 are primarily from the El Chichón eruption. The fact that some pits may have a signal in 1984, whereas others do not, may be a function of the heterogeneity of the volcanic plume over Greenland during the 2 years it was over the region. In addition, there is evidence of volcanic aerosol deposition from the abundant northern hemisphere eruptions beginning in 1979 and lasting to spring 1981 (Table 2). Several key aspects of our data set, as shown in Figure 5, are now briefly presented.

Evidence of the deposition of El Chichón SO$_4^{2-}$ aerosols in the form of distinct peaks is found in four of the eight pits from the Summit region, with the possibility of a signal in two of the other pits. Peaks in SO$_4^{2-}$ that probably are representative of El Chichón aerosols are found in pit 87-2 (fall 1984, winter 1983/spring 1984), in pit 87-4 (summer/fall 1984), in pit 87-5 (fall 1982), and in pit 90-3 (winter to spring 1984) (Figure 5). Slight enhancements in the SO$_4^{2-}$ spring peak of 1984 in pits 87-1 and 87-6 may be from El Chichón aerosols, but the magnitude of the enhancement is small and not very convincing of a volcanic input. There is no clear volcanic signal in pit 87-7 related to El Chichón, but the elevated SO$_4^{2-}$ baseline may be from a lengthy period of aerosol deposition as opposed to rapid deposition from perhaps a single snowfall. It is possible that postdepositional reworking of the snow by wind may have spread out the original signal, but we are not aware of any studies that would verify such a scenario. This may be the same situation that we see in the record from pit 90-1. Similarly, there is not a distinct peak in pit 90-2 within the likely time period for deposition of El Chichón debris.

Evidence for the deposition of the Cl$^-$ component of El Chichón is not found in as many pits as is that for the SO$_4^{2-}$ component, but the signal is much more distinct where it is found. Very distinct Cl$^-$ spikes or at least enhanced seasonal spikes are observed in pit 87-1 during the winter 1982/spring 1983 snow and in pits 87-4 and 87-6 during the winter 1983/spring 1984 snow (Figure 5). Less preservation of the Cl$^-$ signal than of the SO$_4^{2-}$ signal is not unexpected, because the greater solubility of the Cl$^-$ species in volcanic plumes results in quicker condensation and scavenging of the Cl$^-$ aerosols by supercooled water in relation to that for sulfur-bearing aerosols [e.g., Tabazadeh and Turco, 1993].
Deposition of volcanic aerosols from other eruptions during the 1980s is observed through the presence of several spikes in either $\text{SO}_4^{2-}$ or $\text{Cl}^-$. There are spikes in both $\text{SO}_4^{2-}$ and $\text{Cl}^-$ in pits 87-2 and 87-4 within winter 1981/spring 1982 snow as well as a very broad spring peak in each species. These signals could be a result of the 1981 spring eruptions in the northern Pacific region (i.e., Alaid and Okmok; see Table 1). A large spike in $\text{Cl}^-$ in pit 87-7 within summer 1980 snow and a very broad $\text{SO}_4^{2-}$ peak within fall 1980 snow may be a result of deposition from both tropospherically and stratospherically transported material from the 1980 Mount St. Helens eruption [e.g., Sedlacek et al., 1983; Kent and McCormick, 1984]. High concentrations during late 1986 and early 1987 in either $\text{Cl}^-$ (pit 90-2) or $\text{SO}_4^{2-}$ (pit 87-7) could be a result of the St. Augustine and Chikurachki eruptions from the northern Pacific (Table 1).

4.3. Northern and Southern Greenland

There is a discernible $\text{SO}_4^{2-}$ signal from El Chichón in both the northern Greenland pit 88-1 and the southern Greenland pit 88-6 (Figure 5). Both of these signals occur in the winter 1982/spring 1983 seasons, and they are the two largest spikes observed in each pit. In addition, there is a coincident $\text{Cl}^-$ spike in each pit, although the $\text{Cl}^-$ signal in pit 88-6 is more obvious in comparison with the $\text{Cl}^-$ time series for the rest of the pit than is the same signal in pit 88-1. Pit 88-6 also has broad $\text{SO}_4^{2-}$ and $\text{Cl}^-$ spikes within late winter/spring 1987 snow. Again, the St. Augustine and Chikurachki eruptions would be the most likely candidates for those more recent signals. A broad but low peak at the bottom of pit 88-1 coincides with spring 1981 snow and thus could reflect deposition from the northern Pacific eruptions of 1981 (Table 1).

In addition to Pit 88-6 in southern Greenland, a shallower pit was dug near the 20D ice coring site [e.g., Lyons et al., 1990]. This second pit (20D) [Mayewski et al., 1987] only spans the 2-year period from summer 1984 to summer 1982 (Figure 6). The $\text{SO}_4^{2-}$ record for spring 1983 is characterized by a very broad, enhanced baseline that may be a result of El Chichón aerosol deposition prior to deposition of the spring peak. The spring peak of 1984 also appears to be very broad with an initial smaller peak. The $\text{Cl}^-$ record displays several spikes during winter 1982/1983. We cannot rule out that this increase in $\text{Cl}^-$ is from El Chichón, considering the timing of the northern hemisphere dispersion of...
the stratospheric cloud. However, the shortness of the record from this pit makes it difficult to place the SO$_4^{2-}$ and Cl$^-$ records in perspective in comparison with the rest of the 1980s. Nevertheless, Mayewski et al. [1987] postulated that the high amount of SO$_4^{2-}$ and Cl$^-$ found in this pit compared with mean values in the 20D ice core was from deposition of El Chichón aerosols. They also used the data from this pit and the technique of Hammer et al. [1980] to suggest a stratospheric loading of 14 Mt H$_2$SO$_4$. We will expand on this estimate in a later section by using all the pits sampled in this study.

4.4. Annual Flux

To further evaluate the El Chichón signal, we calculated the annual flux of SO$_4^{2-}$, thereby smoothing the original data sets. This method eliminates any potential dependency of the SO$_4^{2-}$ record on accumulation, although Yang et al. [1996] showed that accumulation does not have an impact on SO$_4^{2-}$ concentrations. More important, flux values are necessary to estimate the amount of stratospheric loading and climate-forcing potential of a volcanic eruption from the snow and ice record [e.g., Zielinski, 1995]. We only evaluated the SO$_4^{2-}$ flux record because of the greater inconsistency in the Cl$^{-}$ signal. The annual flux of SO$_4^{2-}$ for each snow pit is shown in Figure 7.

The most prominent characteristic of the flux data set is that maximum flux years are frequently found between 1982 and 1985, the years most likely to be influenced by the deposition of El Chichón aerosols (noting that our year is from summer to summer, and thus 1985 includes summer/fall 1984). In fact, one of these 3 years contains the highest annual flux in 9 of the 12 pits (i.e., 1982-1983, 3 pits; 1983-1984, 5 pits; and 1984-1985, 1 pit) or 75% of the pits. In almost every pit the year with the highest flux value had a high SO$_4^{2-}$ concentration peak or contained an elevated SO$_4^{2-}$ baseline (as represented by the presence of broad, subdued peaks) in that same year. Aerosols from the May 1982 Galunggung eruption may contribute to the high flux 1982-1985 but would probably be very minor in comparison with the El Chichón component. Reported maximum height of the Galunggung cloud was only 16 km, and most of the cloud appeared to move into the southern hemisphere [McClelland et al., 1989]. Maximum flux values also are found in 1981-1982 (two pits) and 1980-1981 (one pit). Maxima in these years probably reflect enhanced deposition of volcanic aerosols from the volcanic activity along the Northern Pacific Rim in 1979-1981 (Table 2). However, we must be cautious in using these results, as seven of the pits sampled failed to extend back to summer 1981, thus preventing a complete evaluation of the 1980s in all pits, and despite the prevalence of maximum values during 1982-1985, the magnitude of the flux in years with the greatest SO$_4^{2-}$ flux commonly is not much higher than that for other years (e.g., pit 88-1; see Figure 7).

Spatial variability in the magnitude of the annual flux signals does not have a consistent trend. Highest peak flux values are found in the southern Summit region (pits 87-5 and 90-1; see Figure 7), but on the other hand, the lowest flux values are found in the southern Summit region as well (pits 87-1 and 87-2). There is not a large difference in the flux values among the one pit in northern Greenland (88-1, Figure 7) and the two pits from southern Greenland (88-6 and 20D, Figure 7). Furthermore, there is no consistent east-west trend in flux values across the divide, as maximum flux values in pits 87-7 and 90-2 (east of the divide) are intermediate in comparison with other pits, whereas flux values in pit 87-5 (east of the divide) are generally among the highest. Perhaps the only consistent spatial pattern that can be identified is that maximum flux values in pits from the southern Summit region and southern Greenland occur in 1982-1983 or 1983-1984, whereas those from the northern Summit region and northern Greenland occur anywhere between 1980-1981 and 1984-1985. However, we again caution that the record in only two of the pits from the southern Summit region and southern Greenland extend back beyond summer 1982. As a result there does not appear to be a preferred site in Greenland that would be more likely to preserve a northern equatorial eruption relative to another site.

4.5. Composite Records

As a final evaluation of the El Chichón signal we combined the annual concentration and flux records for each pit for each year to develop a composite record of SO$_4^{2-}$ deposition for the period 1979-1989 as presented in the box plots in Figure 8. Pit 20D is not included in this analysis. Annual concentrations for this time period clearly show the influence of the abundant volcanism in the early 1980s. Median values for 1979-1980 to 1984-1985 are the highest of the period, with 1980-1981 being the highest (Figure 8a). The 1982-1983 median values are the second highest, but 11 pits are incorporated into that analysis, as compared with just four pits for 1980-1981. Composite SO$_4^{2-}$ flux values further show the influence of volcanism in the early 1980s, with median

Of critical importance to the interpretation of the El Chichón signal is the magnitude of the SO₄²⁻ flux values during the 1980s, values that include the volcanic component, the biogenic component, and especially the anthropogenic component. Median flux values range from 12 kg/km²/yr for 1988-1989 to a high of 29 kg/km²/yr for 1980-1981, with most median values in the 22-25 kg/km²/yr range (Figure 8b). These values are a continuation of the recent increase in SO₄²⁻ deposition in the Summit region from anthropogenic sources [Mayewski et al., 1986, 1990]. SO₄²⁻ deposition is now up to twice the background levels of several centuries ago and up to 25% greater than background SO₄²⁻ levels at the turn of the century [Mayewski et al., 1986, 1990]. If we assume that the years 1987-1990 are relatively free of volcanic input, then it appears that background SO₄²⁻ deposition is of the order of 18-20 kg/km²/yr for the 1980s (given that 1988-1989 appears to be anomalously low). This result would mean that volcanic SO₄²⁻ flux from El Chichón may be estimated by subtracting these background levels from the total of 26-28 kg/km²/yr in the years 1982-1983 and 1983-1984 (Figure 8b). This calculation yields a volcanic SO₄²⁻ flux of about 6-8 kg/km²/yr or a total volcanic flux from El Chichón of about 12-16 kg/km² for these 2 years.

If these assumptions are correct, then a total volcanic SO₄²⁻ flux from El Chichón of 12-16 kg/km² over a 2-year period is almost identical to that estimated in the biyearly sampling of the GISP2 core for the three eruptions of 1902, of which Santa Maria was the largest (i.e., a volcanic SO₄²⁻ flux of 14 kg/km² over a 2-year period [Zielinski, 1995]). Zielinski's [1995] maximum estimate of the stratospheric loading of the 1902 eruptions was 41
Mt H₂SO₄ aerosols, based on the hemispheric distribution of bomb fallout [Clausen and Hammer, 1988], resulting in a maximum optical depth (τ) of about 0.28 by using the relationship between stratospheric mass and optical depth of Stothers [1984]. Calibration of the estimated stratospheric loading of volcanic eruptions in the GISP2 core with the estimated optical depths from Sato et al. [1993] suggested that the ice core record may overestimate the stratospheric loading for some eruptions [Zielinski, 1995]. Thus the use of a correction factor resulted in intermediate and minimum optical depths of 0.13 and 0.06, respectively, for the 1902 eruptions. If we use these same values for the El Chichón eruption, we obtain an estimated maximum τ of 0.28 for El Chichón, a value that agrees well with the 0.3 optical depth estimate presented by Hofmann [1987]. Our intermediate τ estimate of 0.13 (i.e., 20 Mt) agrees better with other estimations of τ, such as the 0.15 value (~20 Mt) given by Rampino and Self [1984] and Rampino et al. [1988], and with other H₂SO₄ loading estimates (i.e., ~16 Mt, once SO₂ is converted to H₂SO₄ [Bluth et al., 1993] and ~12 Mt [McCormick et al., 1995]). Similarly, our τ estimate of 0.13 is appropriate, given the maximum range of optical depths found at >60°N in late 1982 (0.15-0.20) and in early 1983 (0.10-0.15) [Sato et al., 1993]. Our minimum τ estimate of 0.06 matches the global mean τ of ~0.06 given by Sato et al. [1993].

5. Discussion

Our findings provide valuable information that addresses the questions we presented on (1) the variability of the El Chichón signal, (2) the signal:noise ratio as a function of sampling resolution, and (3) the possible influence that high levels of anthropogenic SO₄²⁻ may have on hindering the detection of an El Chichón signal and quantification of that signal. However, we initially point out that identifying volcanic glass from the El Chichón eruption in pit 90-1 verifies the deposition of material from the eruption in the Summit region of Greenland. Tephra was previously found around the Dye 3 drilling site in southern Greenland; thus debris from the eruption is now definitely known to have been deposited on at least central and southern Greenland. In the case of the record in pit 90-1 the tephra is found in the same layer of snow that contains a Na:Cl ratio, in microequivalents of 1:1, unlike that for sea salt. This signal probably reflects deposition of the reactive components of halite with H₂SO₄ (i.e., Na₂SO₄ and HCl [Woods et al., 1995]), because halite was observed in the stratosphere following the eruption. Transport to and the eventual deposition on the ice sheet of the halite components may be via adsorption onto tephra grains. The appearance of Cl⁻ in this snow pit as well as in several other pits indicates that Cl⁻ aerosols are able to remain in the stratosphere for possibly up to one year after the eruption and are able to travel in the stratosphere well beyond the region immediately around the vent despite the fact that they are easily scavenged [e.g., Tabazadeh and Turco, 1993]. The appearance of distinct Cl⁻ spikes in another ice core from Greenland that can be easily matched to known volcanic eruptions [Lyons et al., 1990] further supports this idea. Whether the magnitude of the stratospheric Cl⁻ that reaches the polar region is sufficient to have a major impact on heterogeneous chemical reactions (as in ozone depletion) in the upper atmosphere is another question. The small number of Cl⁻ signals found in this study certainly does not support the notion of a large mass of El Chichón-produced Cl⁻ aerosols remaining in the polar stratosphere 1 year after the eruption. These results provide information on the potential longevity of Cl⁻ aerosols in the stratosphere and serve as additional evidence for the presence of El Chichón aerosols in Greenland snow.

We now focus our attention on the SO₄²⁻ record and the variability of the El Chichón signal. Peaks in SO₄²⁻
concentration that can be confidently linked to the El Chichón eruption are identified in only 50% of the pits sampled. Enhancements in the spring $SO_4^{2-}$ signal in two other pits are not great enough to confidently link them to a volcanic input. These results could easily explain why there is not a distinct signal from El Chichón in the GISP2 core. However, there are several pits in which $SO_4^{2-}$ concentrations within snow deposited between 1982 and 1984 are noticeably higher than background levels. Concentrations in these sections of the pits are equivalent to spring peaks in $SO_4^{2-}$ (e.g., pit 90-1), but they are found in sections of the pit that do not correspond to spring snow. We believe that these sections with broad peaks in $SO_4^{2-}$ are the result of a lengthy period of El Chichón aerosol deposition as opposed to deposition from a single snowfall event. This finding allows us to say that when evaluating a Greenland ice core at a subseasonal resolution, one can expect to find a signal from a northern equatorial eruption of a magnitude similar to that of El Chichón about two thirds of the time, at least under atmospheric conditions like the present. We will elaborate on this aspect of the record later.

The probable deposition of El Chichón aerosols over a period of months as opposed to scavenging by a single snowfall is evidenced further by our results from the annual $SO_4^{2-}$ flux calculations. We found that by smoothing the snow pit record via the calculation of an annual flux we were able to increase the detectability of an El Chichón signal. Maximum annual flux during the years 1982-1983, 1983-1984, and 1984-1985 occur in 9 of the 12 pits sampled, suggesting that one has a 75% chance of observing a peak in annual $SO_4^{2-}$ flux in a single ice core from Greenland, given a northern equatorial eruption of the magnitude of El Chichón. However, we use caution in putting forth this statement, as not all the pits sampled (Figure 7) extended back to the early 1980s and flux values in the years 1980-1982 were also quite high. In fact, volcanic activity during the 5-year period from 1980 to 1985 is abundant, especially in comparison with the later part of the 1980s. The high number of middle to high northern hemisphere eruptions in the early 1980s has increased the concentration of $SO_4^{2-}$ in Greenland snow, and thus higher levels of $SO_4^{2-}$ flux (Figure 8) to the ice sheet in the early 1980s compared to those in the late 1980s makes isolation of the El Chichón signal much harder than it would be if those eruptions had not occurred. The combined stratospheric loading of all of these northern hemisphere eruptions in the early 1980s is only about 1 Mt [Kent and McCormick, 1984; McCormick et al., 1993], but there is a high probability that some portion, if not most, of the $SO_4^{2-}$ deposited from these eruptions reached Greenland via tropospheric transport. In fact, McCormick and Trepte [1987] showed that optical depths in the Arctic began to rise prior to the time that El Chichón aerosols reached the region, but they suggested that this rise may be from the northward transport of aerosols from the equatorial Nyamuragira eruption in late 1981 (Table 1). Sedlacek et al. [1983] also measured high amounts of stratospheric $SO_4^{2-}$ in the high latitudes of the northern hemisphere from these Northern Pacific eruptions. Thus this 5-year period of high volcanic $SO_4^{2-}$ deposition could be another reason that the biannual sampling of the single GISP2 core could not isolate an El Chichón signal.

We used the concentration and annual flux time series of $SO_4^{2-}$ to suggest that there is a 66-75% chance of observing the volcanic signal from a northern equatorial eruption of a magnitude similar to that of El Chichón in a single Greenland ice core, depending on sampling resolution. However, these numbers are a function of modern-day atmospheric conditions, when background $SO_4^{2-}$ levels are much higher than those prior to the past 2 centuries. We can now evaluate how much of an impact this has on deciphering the El Chichón signal by first determining what percentage of the total $SO_4^{2-}$ flux in the composite record is from El Chichón. We can then compare that percentage to the percentage of the total $SO_4^{2-}$ flux that comes from volcanic eruptions that occurred prior to the major increase in atmospheric $SO_4^{2-}$ from the industrial revolution. Provided that our assumptions are correct, we estimated that volcanic $SO_4^{2-}$ flux from El Chichón is 6-8 kg/km²/yr over the 2-year period 1982-1984. The 6-8 kg/km²/yr is only about one third the total $SO_4^{2-}$ deposition or about 40% of background levels. Most explosive eruptions analyzed in the GISP2 core over the last 2100 years have volcanic fluxes that are at least 50% of the total $SO_4^{2-}$ flux recorded at the time of the individual event and often over 66% of background $SO_4^{2-}$ flux levels. Because background $SO_4^{2-}$ flux levels over the last 2100 years (and especially prior to the industrial period) are generally 6.5-7.5 kg/km²/yr from the continuous biyearly sampling in the GISP2 core, sulfur-producing eruptions smaller than El Chichón, like the 1730 Lanzarote event (Canary Islands), may have deposited about 5 kg/km²/yr of $SO_4^{2-}$ on the Greenland Ice Sheet. This value is 75% of background levels at that time, resulting in a very clear peak in $SO_4^{2-}$ concentrations even with the biyearly smoothing of the continuous analyses [Zielinski et al., 1994; Zielinski, 1995]. Similarly, the Coseguina (Nicaragua) eruption, an eruption that may have been poor in sulfur despite the large volume erupted [Self et al., 1989], also deposited about 5 kg/km²/yr of $SO_4^{2-}$ or about 60% of background levels in the mid-1800s. There is a distinct spike for the Coseguina eruption in the GISP2 $SO_4^{2-}$ record. Large eruptions like Tambora and Kuwae (A.D. 1450s) may have volcanic $SO_4^{2-}$ fluxes that are 2-3 times background levels for the time of those eruptions. If pre-A.D. 1900 background levels existed now, then the El Chichón signal would be about equal to background levels as opposed to being only about 40% of background levels. The most viable explanation for these numbers is not that the El Chichón eruption was that much smaller than many
of the explosive eruptions over the last 2100 years, but that identifying (via a distinct spike in the SO$_4^{2-}$ record) and quantifying the volcanic component of the signal in Greenland snow is complicated by the high amount of anthropogenic SO$_4^{2-}$ that is now being deposited on the ice sheet. Mayewski et al. [1987] alluded to the possible mixing of anthropogenic and volcanic SO$_4^{2-}$ in the 20D pit.

These findings may be part of the reason why a signal from the large 1991 Pinatubo eruption has not been detected in Greenland snow as of yet (at least to the best of our knowledge) despite greater sulfur output from Pinatubo (~30 Mt SO$_2$ for Pinatubo versus ~12 Mt SO$_2$ for El Chichón [McCormick et al., 1995]). One could hypothesize that the more equal global distribution of material from the eruption in the Summit region, volcanic glass in snow from spring 1983 verifies the deposition of Pinatubo aerosol cloud [McCormick et al., 1995] in comparison with the greater northern hemisphere distribution of the El Chichón cloud resulted in similar aerosol loadings in the northern polar region that would be available for deposition on the ice sheet. If this situation was true, then the chance of preserving an identifiable Pinatubo signal would be very similar to that for El Chichón given present anthropogenic SO$_4^{2-}$ levels. Thus it is possible that there is a Pinatubo signal in recent Greenland snow, but it may exist more as a period with higher SO$_4^{2-}$ baseline levels as opposed to a distinct SO$_4^{2-}$ peak.

Our suggestion that anthropogenic SO$_4^{2-}$ can complicate the identification of a volcanic signal would imply that the 66-75% chance of observing a volcanic signal from a northern equatorial eruption in a single ice core from Greenland possibly is low. How much too low is uncertain, but given the noise that occurs in the record, as evidenced in the high-resolution sampling, the actual probability of recording such an eruption may still be around 75%. However, this still should be considered excellent, as there is no other medium that would preserve the far-reaching impact of hemispheric to global volcanism at such a level. The fact that our estimates of the stratospheric loading and the resulting optical depths for the El Chichón eruption (i.e., our intermediate estimates of ~20 Mt, $\tau = 0.13$) are close to estimates put forth by others further exemplifies the reliability of snow and ice core records in paleoclimatic research. However, our findings also show that evaluation of the volcanic record at different sampling resolutions, whenever possible, provides the information needed to better acquire reliable results.

6. Conclusions

We used a series of snow pits across the Summit region of Greenland and several from northern and southern Greenland to evaluate the variability in the record of the 1982 El Chichón eruption. The identification of volcanic glass in snow from spring 1983 verifies the deposition of material from the eruption in the Summit region. Evidence of the presence of components derived from the dissociation of halite through reactions with H$_2$SO$_4$ [Woods et al., 1985] in the same layer as the volcanic glass indicates that the Cl$^-$ component of an equatorial volcanic eruption is able to reach the polar stratosphere, possibly adsorbed onto silicate grains. Distinct spikes in SO$_4^{2-}$ that we feel are related to El Chichón aerosol deposition were observed in only half the pits sampled, although periods of enhanced SO$_4^{2-}$ concentrations in the form of broad subdued peaks (i.e., elevated baselines) were found in other pits. These data suggest that a northern equatorial eruption of the magnitude of El Chichón could be recorded about two thirds of the time in a single ice core from Greenland if sampled continuously at a subseasonal resolution. Annual SO$_4^{2-}$ flux calculations essentially smooth the record, thus reducing the noise and increasing the chance of observing an El Chichón signal to about 75% of the time. However, the increase in anthropogenic SO$_4^{2-}$ complicates identification and quantification of the El Chichón signal, as does the abundant middle to high northern latitude volcanism of the early 1980s.

This study provides information on the possible variability in detecting a volcanic signal from moderate northern equatorial eruptions in Greenland ice and snow. Although we estimate that such a signal would be recorded in a single ice core about 75% of the time, the use of polar snow and ice to develop the chronology of past climate-forcing volcanism and quantify the magnitude of the forcing remains the best method to undertake an evaluation of volcanism-climate linkages for time periods prior to recent technological advances (i.e., satellite coverage). The limitations presented in this study should be beneficial to paleoclimatologists and modelers who use ice core records of volcanism in the evaluation of past climate change.

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