Anthropogenic sulfate and Asian dust signals in snow from Tien Shan, northwest China

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ABSTRACT. Snow samples were collected from a 0.5 m snowpack at Glacier No. 1 and near Bogda Feng, eastern Tien Shan, northwest China. Samples that were melted in the field were analyzed for chloride, nitrate, sulfate, sodium, potassium, magnesium, calcium, and microparticles. Eight samples were returned frozen and were analyzed for the above ions plus ammonium, acetate, formate, methylsulfonate, and hydrogen peroxide. There was no significant difference in measured major ion concentrations between the melted and frozen samples. Measured cations in both sets of samples were two to three times greater than measured anions. Calcium and sodium are the dominant cations while sulfate is the dominant measured anion. High ion burdens are associated with dusty layers in the snowpack, indicating that dust from the vast arid regions of central Asia is the dominant source for major ions in Tien Shan snow. The significant increase in sulfate and decrease in the cation:anion ratio in Bogda Feng snow relative to Glacier No. 1 snow suggest that anthropogenic emissions from Ürümqi are an important source of sulfate to precipitation downwind from the city.

INTRODUCTION

The chemical composition of precipitation in eastern China is strongly influenced by fossil fuel combustion as well as agricultural and cultural practices. The primary source for the high levels of sulfate measured in precipitation is the oxidation of sulfur dioxide released during combustion of coal in small- and medium-sized furnaces and coal-burning stoves (Zhao and Sun, 1986; Galloway and others, 1987). However, few measurements of the chemical composition of precipitation in western China have been undertaken. The vast extent of glacierized area and thick seasonal snowpacks in the high mountains of western China provide several locations from which to recover continuous, high-resolution precipitation chemistry records without the expense and effort required to operate continuous monitoring stations.

The collection and chemical analysis of snow samples recovered over a wide spatial area have already proven extremely useful for determining, for example, the vertical separation of monsoonal and westerly derived air masses in northwest India (Mayewski and Lyons, 1983), the influence of anthropogenic pollution on precipitation chemistry in Canada (Schmeisser and others, 1985; McBean and Nikleva, 1986), Greenland (Mayewski and others, 1987), the Soviet Union (Belikova and others, 1984), and Spitsbergen (Semb and others, 1984), and the spatial variation and sources of major ions in precipitation in central Asia (Wake and others, 1990), central Greenland (Mayewski and others, 1990) and the world (Lyons and others, 1990). Further, understanding the spatial variation in snow chemistry and determining the relative importance of local versus regional chemical inputs are fundamental steps in confidently identifying glacier basins from which representative long-term glaciochemical records can be recovered.

As part of an ongoing glaciochemical research program in the mountains of central Asia, snow samples were collected from two different watersheds in the Tien Shan mountain range of northwest China. The chemical composition of these samples shows the influence of Asian dust and anthropogenic emissions on levels of selected carbon, nitrogen, and sulfur species in snow from the Tien Shan.

SITE DESCRIPTION

Snow samples were collected from Glacier No. 1 at the headwaters of the Ürümqi River (hereafter referred to as...
Glacier No. 1) and near Bogda Feng. Both locations are in the eastern Tien Shan in northwest China; the sample sites are separated by 120 km (Fig. 1). The Tien Shan is bounded on the north by the Junggar, Pesk Sary Ishikotrau and Muyun Kum Deserts, on the south by the Taklamakan Desert and on the east by the Gobi Desert. Very little is known concerning the chemical composition of surface desert materials in this region (Dregne, 1968). The gravelly, medium- and fine-textured soils are probably calcareous. Soil salinity becomes greater toward and within deserts; many depressions within the deserts are very saline as a result of evaporation of water that collects in the depressions in spring (Dregne, 1968).

**Glacier No. 1**

Glacier No. 1 lies approximately 50 km northeast of the Rear Gorge (Houxia), which is the site of the Yaojin steel factory and a cement plant. Coal and iron are mined in the surrounding hills. Meteorological data from the Tien Shan Glaciological Station (5 km up valley from Houxia) indicates that daytime up-valley winds transport air from the Rear Gorge towards Glacier No. 1. Open valleys several hundred meters above the tree line characterize the headwaters of the Ürümqi River. Glacial coverage is limited to small cirque and hanging glaciers; the valley also holds extensive moraine deposits. Glacier No. 1 faces northwest and consists of two branches (east and west arms); its area is 1.84 km².

**Bogda Feng**

Bogda Feng (5570 m a.s.l.) lies approximately 60 km west, and commonly downwind, from Ürümqi, which is a city of one million people and the economical and industrial center of Xinjiang Uygur Autonomous Region. Glaciers are restricted predominantly to the upper slopes of Bogda Feng. The open, glacier-free valleys to the northwest of Bogda Feng, where samples were collected, were covered with an ≈0.5 m deep snowpack in April 1989.

**METHODS**

**Sample collection**

Extreme care was taken at all times during sample collection, handling, transport, and analysis to assure sample integrity. Non-particulating clean suits and hoods, plastic gloves and particle masks were worn during all sampling procedures. Prior to use, all sample tools and containers were rinsed, soaked, and rinsed again with Millipore Milli-Q™ deionized water. Analyses of duplicate samples as well as transport and laboratory blanks demonstrate that sample contamination during sample transfer, transport, and subsequent analytical procedures was negligible.

In mid-April 1989, samples of the entire seasonal snowpack (≈0.5 m of snow) were recovered from snow pits excavated on the east and west arms of Glacier No. 1,
Table 1. Snow samples collected from the Tien Shan for chemical analysis

<table>
<thead>
<tr>
<th>Date</th>
<th>Elevation m a.s.l.</th>
<th>Number of samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glacier No. 1.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>snow pit 1</td>
<td>11 April</td>
<td>3960</td>
</tr>
<tr>
<td>snow pit 2</td>
<td>12 April</td>
<td>4140</td>
</tr>
<tr>
<td>Bogda Feng</td>
<td></td>
<td></td>
</tr>
<tr>
<td>snow pit 3</td>
<td>16 April</td>
<td>3450</td>
</tr>
<tr>
<td>snow pit 4</td>
<td>17 April</td>
<td>3550</td>
</tr>
</tbody>
</table>

and from two snow pits approximately 7 km northwest of Bogda Feng (Table 1). Samples were collected over glacier ice at Glacier No. 1 and over the ground at Bogda Feng. Just before sample collection, the outer 0.30 m of the snow-pit wall was scraped away using a precleaned acrylic scraper. Snow-pit samples were collected over a continuous vertical section at 40 mm intervals using a custom-made sample tool. The samples were placed in 125 ml sample containers, melted in the field and transferred into leakproof 20 ml vials for transport to the University of New Hampshire (termed “melted” samples). Eight sets of duplicate near-surface snow samples (0 to 0.20 m from the upper snow surface) were collected (three from Glacier No. 1 and five from Bogda Feng) and placed in insulated shipping containers with pre-frozen eutectic mixtures. These samples were dispatched to UNH in a frozen state and are referred to as “frozen” samples.

Temperature of the snowpack at 0.10 m depth during the day ranged from -6 to -10°C at Glacier No. 1 and was constant at -4°C at Bogda Feng. Air temperature measured during snow-pit sampling periods was at or below 0°C.

Laboratory analysis

Snow samples were analyzed for anions (chloride, sulfate, and nitrate) and cations (sodium, ammonium, potassium, calcium, and magnesium) using a Dionex™ Ion Chromatograph Model 2010 with computer-driven autosampler in the laboratories of the Glacier Research Group at the University of New Hampshire. Snow samples were not filtered prior to analysis. Ammonium is only quoted for analyses conducted on frozen samples. The anion system employed a Dionex™ AS4A anion exchange column in conjunction with an anion guard column and a 1.7 mM NaHCO₃/2.08 mM Na₂CO₃ eluent. The cation system consisted of cation-I and cation-II fast columns with an 18 mM HCl eluent with 0.33 mM diaminopropionic acid monohydrochloride (DAP). A sample loop of 0.5 ml was used in both systems. Ten samples were analyzed in duplicate using two separately drawn samples. This procedure resulted in a standard deviation from the mean of less than 5% for anions and less than 7% for cations. Microparticle concentrations over a size range of 0.65 to 12.88 μm were measured using an Elzone 280PC particle counter.

Acetate (CH₃COO⁻), formate (CHOO⁻), methane-sulfonate (CH₃SO₃⁻) and hydrogen peroxide (H₂O₂) analyses were undertaken only on frozen samples. Samples were analyzed for acetate, formate and methanesulfonate using a Dionex QIC ion chromatograph with a Dionex™ AS4 anion exchange column and a 0.4 mM Na₂CO₃ eluent. Peroxide was analyzed using the flurometric technique described in Lazrus and others (1985).

RESULTS

Physical stratigraphy

The physical stratigraphy of the snowpack was similar at both sample locations even though the snowpack overlay glacier ice at Glacier No. 1 and ground at Bogda Shan. 0.15 to 0.20 m of snow from two separate snowstorms in early April (termed near-surface snow) lay on top of 0.30 to 0.50 m of depth hoar (Fig. 2). Two to three dust layers 10 to 20 mm thick were apparent in all four snow pits. At the elevations where snow samples were collected (Table 1) the entire snowpack melts during the summer ablation period. The snowpack in April, therefore, contained only snow deposited during winter 1988-89 and spring 1989, and thus represented approximately six months’ accumulation. Total water equivalent of the snowpack varied from 0.10 to 0.17 m.

Chemistry of frozen samples

The near-surface snow samples which remained frozen until just before analysis show a wide variation in concentration of chemical species measured (Table 2). However, much of the cation content is probably solubilized when the samples are melted. Total measured cations are two to three times greater than measured anions.

The ion balance of the frozen samples was calculated using the microequivalent per kilogram (μEq kg⁻¹) concentrations of all measured ions:

\[ AC = [Na⁺] + [NH₄⁺] + [K⁺] + [Mg²⁺] + [Ca²⁺] \]
\[ - [Cl⁻] - [NO₃⁻] - [SO₄²⁻] \] (I)

Ammonium concentration in the frozen samples accounts for 75% of nitrogen measured and 12% of the total cation burden. Acetate and formate account for 8% of the total anion burden. Acetate displays a relatively constant concentration in the frozen samples while formate shows a wide range of concentrations which correlate well with variations in sulfate (r² = 0.93; n = 8). In snow samples with relatively high total ionic burden (> 300 μEq kg⁻¹)
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Fig. 2. Major ion concentration, physical stratigraphy, and microparticle concentration profiles in samples collected from snow pit no. 1 at 3960 m a.s.l. on the west arm of Glacier No. 1. Glacier ice underlay the 0.5 m snowpack. Note the strong correlation between high concentrations of Na\(^+\), Ca\(^{2+}\) and SO\(_4^{2-}\) with high microparticle concentrations and visible dust layers in the snowpack.

Table 2. Ion concentrations in frozen samples of near-surface snow

<table>
<thead>
<tr>
<th>Type of snow</th>
<th>Na</th>
<th>NH(_4)</th>
<th>K</th>
<th>Mg</th>
<th>Ca</th>
<th>CH(_3)COO</th>
<th>HCOO</th>
<th>Cl</th>
<th>NO(_3)</th>
<th>SO(_4)</th>
<th>H(_2)O(_2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dirty</td>
<td>164</td>
<td>20.0</td>
<td>2.8</td>
<td>18.1</td>
<td>285</td>
<td>2.1</td>
<td>21.7</td>
<td>58.7</td>
<td>9.3</td>
<td>158</td>
<td>0.4</td>
</tr>
<tr>
<td>dirty</td>
<td>226</td>
<td>24.4</td>
<td>3.6</td>
<td>20.6</td>
<td>243</td>
<td>1.9</td>
<td>22.2</td>
<td>69.1</td>
<td>11.3</td>
<td>206</td>
<td>0.4</td>
</tr>
<tr>
<td>dirty</td>
<td>218</td>
<td>23.3</td>
<td>5.6</td>
<td>27.1</td>
<td>350</td>
<td>1.8</td>
<td>21.4</td>
<td>68.3</td>
<td>11.0</td>
<td>198</td>
<td>0.4</td>
</tr>
<tr>
<td>clean</td>
<td>6.1</td>
<td>15.0</td>
<td>0.5</td>
<td>2.5</td>
<td>45.9</td>
<td>2.7</td>
<td>4.4</td>
<td>2.1</td>
<td>4.2</td>
<td>13.1</td>
<td>9.5</td>
</tr>
<tr>
<td>clean</td>
<td>28.3</td>
<td>11.6</td>
<td>1.0</td>
<td>9.9</td>
<td>127</td>
<td>2.6</td>
<td>8.0</td>
<td>11.2</td>
<td>3.7</td>
<td>35.3</td>
<td>0.5</td>
</tr>
<tr>
<td>clean</td>
<td>52.2</td>
<td>18.3</td>
<td>1.5</td>
<td>10.7</td>
<td>171</td>
<td>2.4</td>
<td>13.0</td>
<td>25.2</td>
<td>5.5</td>
<td>58.6</td>
<td>0.0</td>
</tr>
<tr>
<td>clean</td>
<td>5.7</td>
<td>13.3</td>
<td>0.5</td>
<td>2.5</td>
<td>27.4</td>
<td>2.5</td>
<td>4.9</td>
<td>1.9</td>
<td>4.5</td>
<td>12.4</td>
<td>7.1</td>
</tr>
<tr>
<td>clean</td>
<td>15.2</td>
<td>8.3</td>
<td>0.5</td>
<td>4.9</td>
<td>70.4</td>
<td>4.5</td>
<td>3.4</td>
<td>4.8</td>
<td>2.1</td>
<td>17.2</td>
<td>2.8</td>
</tr>
</tbody>
</table>
formate concentrations are five to 10 times greater than acetate concentrations. In samples with lower ionic burdens, formate accounts for 40 to 60% of total acetate and formate load. Methanesulfonate was below the detection limit of 1 part per billion (ppb).

Hydrogen peroxide concentrations ranged from 0.0 to 9.5 ppb, and show a rapid decrease with increasing sulfate. Samples with relatively high sulfate (> 30 μEq kg⁻¹) contain less than 0.5 ppb hydrogen peroxide while samples with sulfate less than 20 μEq kg⁻¹ show hydrogen peroxide levels greater than 7.1 ppb.

Chemistry of melted samples

The results of chemical analysis for all snow-pit samples melted in the field \( (n = 50) \) appear in Table 3. An ionic balance equation for the melted samples of the form

\[ \Delta C' = [Na^+] + [K^+] + [Mg^{2+}] + [Ca^{2+}] - [Cl^-] - [NO_3^-] - [SO_4^{2-}] \tag{2} \]

was used to determine the balance between measured cations and anions in each sample. This equation ignores the potential contribution from ammonium and organic acids; however, in the frozen samples, ammonium concentrations and combined acetate and formate concentrations were similar and therefore tend to cancel each other out.

In the melted samples, as in the frozen samples, the sum of major cations is two to three times greater than the sum of major anions. Among cations, calcium shows the highest concentration followed by sodium. The predominant anions are represented by ΔC' (Equation (2)) and sulfate.

Linear regression on all melted samples showed that calcium and magnesium correlate well \( (r^2 = 0.97) \), as do calcium and ΔC' \( (r^2 = 0.89) \) and magnesium and ΔC' \( (r^2 = 0.91) \). Sodium correlates well with sulfate \( (r^2 = 0.87) \). All other major anion–cation combinations show a positive correlation with \( r^2 \) ranging from 0.55 to 0.70. All correlation coefficients are significant at the \( P = 0.005 \) level. In addition, high levels of sodium, calcium, chloride, and sulfate are associated with snow that was collected from a layer which contained visible dust and which had high microparticle concentrations relative to layers without visible dust (Fig. 2).

DISCUSSION AND CONCLUSIONS

Frozen samples

The high levels of ammonium are probably a result of widespread use of fecal remains as a fertilizer (Galloway and others, 1987) in the agricultural regions surrounding Ürümqi. Norton (1983) found that high levels of organic acids, especially formate (≈ 10 to 100 μEq kg⁻¹), in precipitation downwind from large urban centers are related to anthropogenic emissions. The high formate levels measured in the frozen snow from Glacier No. 1 and Bogda Feng, and the good correlation between formate and sulfate, suggest that anthropogenic emissions are a source of both species in the eastern Tien Shan. The low levels of methanesulfonate in the snow suggest that biogenically produced dimethyl sulfide (DMS) is not a source of sulfur for this region (e.g. Legrand and others, 1988). Oxidation of sulfur dioxide can occur by reaction with hydrogen peroxide to produce sulfate (e.g. Penkett and others, 1979; Laj and others, 1990). A sharp drop in hydrogen peroxide levels in samples with high sulfate

### Table 3. Measured major ion concentrations in snow-pit samples melted in the field

<table>
<thead>
<tr>
<th></th>
<th>Na⁺</th>
<th>K⁺</th>
<th>Mg²⁺</th>
<th>Ca²⁺</th>
<th>Cl⁻</th>
<th>NO₃⁻</th>
<th>SO₄²⁻</th>
<th>ΔC'⁺⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(ion concentration in μEq kg⁻¹)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glacier No. 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>snow pits (n = 27)</td>
<td>mean</td>
<td>45.7</td>
<td>1.2</td>
<td>9.3</td>
<td>86.9</td>
<td>13.9</td>
<td>4.0</td>
<td>31.9</td>
</tr>
<tr>
<td></td>
<td>standard deviation</td>
<td>85.9</td>
<td>1.3</td>
<td>11.1</td>
<td>98.4</td>
<td>21.2</td>
<td>3.5</td>
<td>53.0</td>
</tr>
<tr>
<td>Bogda Shan</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>snow pits (n = 23)</td>
<td>mean</td>
<td>68.2</td>
<td>2.9</td>
<td>10.0</td>
<td>86.3</td>
<td>12.5</td>
<td>5.3</td>
<td>69.9</td>
</tr>
<tr>
<td></td>
<td>standard deviation</td>
<td>152.5</td>
<td>3.0</td>
<td>11.2</td>
<td>102.7</td>
<td>19.4</td>
<td>3.8</td>
<td>130.4</td>
</tr>
</tbody>
</table>

\[ \Delta C'⁺⁺ = [Na^+] + [K^+] + [Mg^{2+}] + [Ca^{2+}] - [Cl^-] - [NO_3^-] - [SO_4^{2-}] \]
Table 4. Relative contribution of major ions to total ion burden in snow-pit samples melted in the field

<table>
<thead>
<tr>
<th></th>
<th>Σ'</th>
<th>% of total cation charge</th>
<th>% of total anion charge</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>μEq kg⁻¹</td>
<td>Na⁺</td>
<td>K⁺</td>
</tr>
<tr>
<td>Glacier No. 1</td>
<td>286</td>
<td>16</td>
<td>1</td>
</tr>
<tr>
<td>Bodga Shan</td>
<td>334</td>
<td>21</td>
<td>6</td>
</tr>
</tbody>
</table>

Percentages were computed on a sample-by-sample basis and then averaged. Mean values may not add to 100% as a result of rounding.

\[ Σ' = 2([Na⁺] + [K⁺] + [Mg²⁺] + [Ca²⁺]) \]

suggests that oxidation of sulfur dioxide in northwest China occurs via reaction with hydrogen peroxide.

**Melted versus frozen samples**

Analysis of duplicate melted and frozen snow samples indicates there is little difference in the concentration of dominant ions (i.e. sodium, magnesium, calcium, chloride, nitrate, and sulfate) between samples that were melted in the field and those that were kept frozen until just prior to analysis (Fig. 3). Magnesium shows a slight enrichment in the melted samples in six out of eight cases (i.e. ratio greater than 1, Fig. 3); calcium displays a slight depletion in the melted samples in six out of eight cases. However, there exists no consistent trend of enrichment or depletion of any individual ion in the samples that were melted. In fact, the mean ratio for ion concentration in melted and frozen samples falls within one standard deviation of 1 for all ions. This improves our confidence not only in the major ion-concentration data from melted samples presented in this paper, but also in the chemical data from snow collected in the past at several different sites in the mountains of central Asia (Wake and others, 1990).

**Glacier No. 1 snow**

The total ionic charge (Σ') for the melted samples from Glacier No. 1 was calculated using:

\[ Σ' = 2([Na⁺] + [K⁺] + [Mg²⁺] + [Ca²⁺]) \] (3)

This calculation ignores the contribution from ammonium which accounts for ≈10% of the total cation burden in the frozen samples. Σ' is therefore considered a minimum estimate of the total ionic burden in the melted samples.

The percentage contribution of major cations and anions to the total cation and anion burden was determined for each individual sample; the mean values derived from this calculation appear in Table 4. For melted snow-pit samples from Glacier No. 1, calcium and sodium are the dominant cations, accounting for 74% and 16% of the total cation burden, respectively; ΔC' (Equation (2)) accounts for 59%, and sulfate 23%, of the total anion content.

Williams and others (1992) performed major ion and acid-neutralizing capacity (ANC) measurements on snow samples collected from Glacier No. 1 and surrounding...
Williams and others (1992) suggested that the source of correlation in our samples between calcium and ANC was dissolution of CaCO$_3$ rich dust. The good total anion charge. This figure compares well with burden in our snow samples (Table 4). Furthermore, the similarity between our represents primarily carbonate/bicarbonate. The position of the Tien Shan relative to large desert basins with predominantly calcareous material (Dregne, 1968) originates from an ice core recovered by a joint Sino-Japanese program (Han and others, 1989); 12 m of core B12 was sectioned and analyzed in GRG laboratories. The Dunde Ice Cap data come from Thompson and others (1989).

The mean value for sulfate concentration in snow from Glacier No. 1 (Table 2) is among the highest sulfate levels measured in snow from the mountains of central Asia (Fig. 4 and Wake and others, 1990). The high sulfate content in Glacier No. 1 snow combined with an anthropogenic source of sulfate originating from the nearby steel factory suggests that anthropogenic emissions account for at least a portion of sulfate deposited in this region; however, with the available data, this is difficult to quantify.

**Bogda Feng snow**

While the ratio of measured cations to anions in Glacier No. 1 snow is 3.3, the ratio in Bogda Feng snow is only 1.8. This large variation in cation: anion ratio is not due to a decrease in cation concentrations in Bogda Feng snow; rather, it is a result of a doubling of sulfate levels in Bogda Feng snow (Table 2). As with snow from Glacier No. 1, calcium and sodium are the dominant cations, accounting for 64% and 21% of $\Sigma^+$, respectively. However, sulfate accounts for a greater percentage of $\Sigma^+$ in Bogda Feng snow compared to snow from Glacier No. 1 (Table 4). ($\Delta C'$ in Table 4 for Bogda Feng snow was calculated using Equation (2). However, we expect that, for ions listed in Equation (2), $\Sigma^+$ is actually greater than $\Sigma^+$ as a result of the influx of sulfate. Unfortunately our data are not detailed enough to determine the excess anions with an acceptable degree of certainty.)

Among all the major ions, only sulfate displays a significant difference between snow-pit samples collected from Glacier No. 1 and Bogda Feng (Student's t test; confidence interval = 0.90). All other major ions, as well as $\Sigma^+$, show no significant difference between the Glacier No. 1 and Bogda Feng samples, even at a confidence interval as low as 0.70. Only half of the increase of sulfate in Bogda Feng snow can be accounted for by increased dust input (i.e. increase in Na concentration; Table 2). Snow from Bogda Feng contains two to 10 times more sulfate than has been measured in snow from several other mountain regions in central Asia (Fig. 4). The high values of sulfate in Bogda Feng snow compare well with sulfate levels in polluted snow surrounding major industrial regions in the USSR (Belikova and others, 1984). The position of Bogda Feng, 60 km downwind from Ürümqi, combined with the very high levels of sulfate in snow from this region are indicative of an anthropogenic source for a portion of the sulfate in Bogda Feng snow.

Snow from the western Kunlun and the Dunde Ice Cap show mean sulfate concentrations of 20.3 and 18.9, respectively (Fig. 4). These levels of sulfate, in addition to those in snow from Glacier No. 1 and Bogda Feng, are much higher than in any other snow samples recovered from the mountains of central Asia (Fig. 4 and Wake and others, 1990). While the Glacier No. 1 and Bogda Feng sites lie adjacent to centers of anthropogenic emissions, which in part contribute to the observed high levels of sulfate, the western Kunlun and Dunde Ice Cap sites are distant from any major industrial center. The position of the western Kunlun and the Dunde Ice Cap adjacent to large desert basins, (the Tarim basin to north of western Kunlun and the Qaidam basin to west of Dunde Ice Cap), suggests that the high sulfate levels are a result of an
influx of desert-derived material. This, in turn, suggests that glaciochemical records recovered from these regions are difficult to interpret in terms of regional atmospheric chemistry due to swamping of regional chemical signals by local desert-derived materials.

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The accuracy of references in the text and in this list is the responsibility of the author(s), to whom queries should be addressed.