Climatic and environmental records from the Far East Rongbuk ice core, Mt. Qomolanglema (Mt. Everest)

Introduction

The Qinghai-Tibetan (Q-T) Plateau is one of the most imposing topographic features on the surface of the Earth. The sensible and latent heat flux released over the Q-T Plateau drives Asian monsoon circulation and strongly influences global circulation patterns. The sensible and latent heat flux released over the Q-T Plateau drives Asian monsoon circulation and strongly influences global circulation patterns. The Qinghai-Tibetan (Q-T) Plateau is one of the most imposing topographic features on the surface of the Earth. The sensible and latent heat flux released over the Q-T Plateau drives Asian monsoon circulation and strongly influences global circulation patterns. The sensible and latent heat flux released over the Q-T Plateau drives Asian monsoon circulation and strongly influences global circulation patterns.

Methodology

During the Sino-American Expedition to Mt. Qomolangma (Mt. Everest) in May 1997, a 41 m ice core was recovered from an elevation of 6500 m from the northern branch firn basin of the Far East Rongbuk (FER) Glacier near Mt. Qomolangma (Mt. Everest). The ice core was dated down to AD 1814 by counting δ18O values recorded in the FER ice core, to investigate the climatic and environmental change during recent 200 years in Mt. Qomolangma (Mt. Everest). The location of the Far East Rongbuk (FER) Glacier near Mt. Qomolangma (Mt. Everest) (27°59’N, 86°55’E) on the boundary of the two climatic regions combined with the high elevation of the site, well above the influence of the boundary layer, provides a unique opportunity to describe and understand paleoclimatic and environmental change in remote regions (Figure 1). This paper focuses on the major ion and oxalate concentrations and δ18O values recorded in the FER ice core, to investigate the climatic and environmental change during recent 200 years in Mt. Qomolangma (Mt. Everest).

In May 1997, a 41-meter core was collected from a site at 6500 m a.s.l. on the Far East Rongbuk Glacier, approximately 13 km north of the peak of Mt. Qomolangma (Mt. Everest) (Figure 1). FER is a valley glacier that stretches approximately 9 km and has an ~150 m thick accumulation zone as measured at several locations with a portable radio echo sounding instrument. The ice core was recovered
from a relatively flat portion of the accumulation zone, where glacier flow is assumed to be fairly laminar, using a new light weight drill (Eclipse). Reoccupation of previous survey sites reveals that FER glacier retreat has averaged 7.4 m/yr since 1966, similar to other larger valley glaciers on the north slope of Mt. Qomolangma (Mt. Everest) (Zheng et al., 1975; Ren et al., 1998).

The ice core was processed in the field in a dedicated clean trench excavated in the snowpack. All core handling and processing was performed by personnel wearing non-particulating clean suits, face masks, and polyethylene gloves. After the outer 10–20 mm of the core was scraped, the clean core was then cut into 0.04 m sections and placed in 60 ml polyethylene cups. The ice core samples were melted just prior to analyses for major ion concentrations via ion chromatography on a Dionex 4000 i at the University of New Hampshire using methods described elsewhere (Buck et al., 1992). Oxygen isotope samples (δ¹⁸O) were measured on a Finnigan MAT-252 mass spectrometer at the Cold and Arid Regions Environmental and Engineering Research Institute, Chinese Academy of Sciences.

Current mean annual temperature (based on standard 10 m depth temperature) is -7°C at the core site. As commonly observed throughout almost all elevations on the Q-T Plateau and Himalayas (Mayewski et al., 1981; Thompson et al., 1988), melting does occur during at least a portion of the year. Preservation of annual signals in the FER δ¹⁸O plus several chemical species (e.g., Na⁺, Ca²⁺, SO₄²⁻, Mg²⁺) series can be assumed based on calibration with known bomb horizons (+/- one year counting error at this level) identified in the well-preserved total β-activity profile from this core (Qin et al., 2000). Melting is, therefore, assumed to be confined within annual layers, thus, preserving the annual signal used to date the record. We counted 184 annual layers back through the entire 41m ice core, indicating that our record spans the time period from 1814 to 1997 (a resolution of 5.5 samples per year). Details of the dating methods are described by others (Hou et al., 1999; Qin et al., 2000). The average annual accumulation is 224 mm (ice equivalent). Variations of δ¹⁸O and major ion concentrations with time are shown in Figure 2.

Results and discussion

Climatic records in FER ice core

In central Himalayas, δ¹⁸O values increase from winter to spring reflecting change in temperature of condensation, and the lowest δ¹⁸O values occur in late summer indicating that δ¹⁸O is controlled by an “amount effect” during the monsoon season (Wushiki, 1977; Wake et al., 1995; Kang et al., 2000). Although the seasonal variations of δ¹⁸O are influenced by both temperature and precipitation, the long-term δ¹⁸O records (e.g. since the Last Glacial Age) in ice cores from tropical regions are still used as a proxy for temperature (Thompson et al., 1995b, 1997). Recent studies indicate that the change of long-term δ¹⁸O in the Dasuopu Ice Core from central Himalayas is consistent with temperature change over the Northern Hemisphere (NH) (Thompson et al., 2000). Apparently, not only the fluctuations, but increasing tendency of FER δ¹⁸O also agrees with those of NH temperature (Jones et al., 1986; Qin et al., 2000) (Figure 3), and with those of Guliya ice core (35°17'N, 81°29'E) from the northwestern Q-T Plateau (Yao et al., 1996b) (Figure 4). It may be deduced that the FER δ¹⁸O changes are mainly dominated by “temperature effect”. Based on this, the climatic change will be discussed.

Figure 1 Sketch map of the Qinghai-Tibetan Plateau, the Far East Rongbuk glacier and the drilling site in 1997.

Figure 2 The concentration variations of chemical species with time in the Far East Rongbuk ice core.
The fluctuations of FER δ18O indicate that there are 5 cold periods and 5 warm periods from the beginning of the 19th century to the 1980s (Table 1; Figure 3 and 4). Since the beginning of 19th century, climate change recorded by FER δ18O is generally consistent with temperature change in the NH. The exception is that the third warm period and the fourth cold period in the earlier 20th century are contrary to that of the NH. This may be caused by the difference between regional and hemispheric climate change. The FER δ18O changes have a good agreement with that from Guliya ice core (Figure 4), suggesting that climate changes are consistent in regional scale over the Q-T Plateau from the beginning of 19th century. Especially, the 20th-century warming trend recorded in the FER ice core agrees with that of plateau-wide warming recovered from other ice core records over the Q-T plateau (Yao et al., 1995; Qin et al., 2000; Thompson et al., 2000). Compared to the monsoon precipitation from northwest India adjacent to Mt. Qomolangma (Mt. Everest) (Sontakke et al., 1996) (Figure 3), high δ18O values from FER ice core correspond to low precipitation in 1870s and the earlier 20th century, and low δ18O values to high precipitation during 1880s~1890s. This inverse association between FER δ18O and precipitation may be expected given the influence of precipitation amount on δ18O (Wushiki, 1977; Rozanski et al., 1992; Wake et al., 1995; Kang et al., 2000).

Since 1930s, summer precipitation in India fluctuated slightly and its tendency is not evident, however, the δ18O fluctuation is larger (Figure 3). Over the last 40 years, there are also some reverse tendencies between FER δ18O and annual precipitation from Xigeza Meteorological Station (e.g. in 1960s, 1970s, and 1980s) (Figure 5), as well as some similar trends between FER δ18O and annual temperature (e.g. in the earlier 1970s and in the middle of 1980s). These may indicate that FER δ18O records are influenced by both precipitation and temperature, or even more than these two factors in the last decades. A recent study (Qin et al., 2000) points out that controls on the isotopic fractionation of moisture deposited on the southern margin of the Plateau have changed over the last few decades. The increase in temperature over the last few decades (Kang et al., 1998) may have led to changes in atmospheric circulation that have resulted in a decrease in moisture flux to the Plateau. Especially since 1980s, influence for FER δ18O may be not simply limited to precipitation and temperature but more complicated, and should be the combination of many factors and processes.

### Environmental records in FER ice core

#### Variations of atmospheric dust

The high concentration of major ions recorded in the FER ice core occur simultaneously and their variation tendency is similar (Figures 2, 6). Ca2+, Mg2+ and SO42− in snow and ice come from crustal dust in the central Himalayas (Wake et al., 1993; Kang et al., 2000), thus, the variation of atmosphere dust in Mt. Qomolangma (Mt. Everest) since the beginning of 19th century can be discussed via Ca2+, Mg2+ and SO42−.

<table>
<thead>
<tr>
<th>Events</th>
<th>Cold period</th>
<th>Warm period</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>(AD)</td>
<td>(years)</td>
</tr>
<tr>
<td>1</td>
<td>~1826</td>
<td>1826–1838</td>
</tr>
<tr>
<td>2</td>
<td>1858–1874</td>
<td>1875–1886</td>
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<tr>
<td>3</td>
<td>1886–1898</td>
<td>1898–1921</td>
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<td>4</td>
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### Table 1 Warm and cold periods recovered by δ18O in the Far East Rongbuk ice core

<table>
<thead>
<tr>
<th></th>
<th>Cold period</th>
<th>Warm period</th>
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<tbody>
<tr>
<td>Period (AD)</td>
<td>Duration (years)</td>
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</tr>
<tr>
<td>1</td>
<td>~1826</td>
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</tbody>
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The variations of atmosphere dust since 19th century recorded in the FER ice core are summarized in Table 2. It shows that the periods of intense dust are during the 1830s to 1840s, and the 1880s and 1960s. However, a period of less dust, from the end of 19th century to late 1920s, was the longest one, in which there was a secondary stage of intense dust around 1903. Since 1960s, the dust storm events, which happened in Northwestern China most strongly and extensively in 1961, 1983 and 1984 (Qian et al., 1997), are synchronously recorded in the FER ice core (dust maxima years in Table 2), suggesting that the dust storms occurred in mid-Asia may be transported to the Northern slope of the Mt. Qomolangma (Mt. Everest), though this process depends on dust transportation from north to south, while dust is generally transported from west to east or from northwest to southeast in mid-Asia (Xu et al., 1997). Dust in FER ice core may also be influenced by spring dust storms from southwestern Asia. Because of the extensive dust storms in southwestern Q-T Plateau (e.g. 10~20 days of dust storms per year) (Lian, 1997), dust
storms from the Plateau are perhaps the one of the most likely sources for ice core crustal species. Additionally, local rock around the FER Glacier may also be dust source. In summary, the change of dust species reflects combination of sources from mid-Asia, southwestern Asia, and southwestern Q-T Plateau, as well as local rock. Among these sources, the dust storms from the vast mid-Asia may be dominant.

Oxalate records from FER ice core

Oxalate is one of the carboxylic acids. Despite their potentially important role in tropospheric chemistry (Galloway, 1982; Keene et al., 1986), few investigations of snow carboxylic acids have been conducted in remote areas, particularly in Asia. From the FER ice core record, the most outstanding feature is the $C_2O_4^{2-}$ increase since the beginning of the 20th century (Figures 2, 6, 7). Fluctuations of $C_2O_4^{2-}$ are small in the 19th century (Figure 7), probably the result of seasonal fluctuations. However, in the 20th century, especially during the 1950s~1980s, many peaks of $C_2O_4^{2-}$ concentrations occur and the fluctuations are very dramatic, indicating that short episodes with high $C_2O_4^{2-}$ concentrations account for elevated mean values in 20th century.

Carboxylic acids come from vegetation emissions and biomass burning (Talbot et al., 1988; Andreae et al., 1988; Legrand et al., 1992), the oxidation of various alkenes (Jacob, 1986) and anthropogenic emissions (Talbot et al., 1988; Legrand et al., 1995). The increasing tendency of $C_2O_4^{2-}$ concentrations in the 20th century is not consistent with that of $Ca^{2+}$, which is representative of atmospheric dust (Kang et al., 2000) in Himalayas (Figure 2), suggesting that the source of $C_2O_4^{2-}$ has very little relation to atmospheric dust. $NH_4^+$ come partly from biogenic sources in Himalayas (Mayewski et al., 1983; Davidson et al., 1986) but the general trend of variations is not obvious (Figure 7). However, the perturbations of $C_2O_4^{2-}$ are larger than those of $NH_4^+$ in the 20th century, suggesting that the 20th century increase of $C_2O_4^{2-}$ concentrations is not mainly related to biogenic sources and thus caused by other sources. Ice core $\delta^{18}O$ has increased since the beginning of 20th century and the general

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**Table 2 Intense and lesser dust periods recovered by crustal ions in the Far East Rongbuk ice core.**

<table>
<thead>
<tr>
<th>Period of intense dust</th>
<th>Period of lesser dust</th>
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<tbody>
<tr>
<td>1827-1843</td>
<td>1827</td>
</tr>
<tr>
<td>1852-1865</td>
<td>1860-1864</td>
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<tr>
<td>1878-1893</td>
<td>1885-1893</td>
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<td>1928-1934</td>
<td>1930-1943</td>
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<tr>
<td>1943-1949</td>
<td>1941-1943</td>
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Figure 5 The comparison of $\delta^{18}O$ in the Far East Rongbuk ice core and temperature, precipitation at Xigaze (the fine line is annual value, the coarse line is 11 points smoothing value).

Figure 6 The 11 points smoothing curve of chemical species in Far East Rongbuk ice core (black parts are higher than average value).

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higher C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentrations, especially those peaks due to short emissions and may represent natural background values, while concentrations in the 19th century were not influenced by industrial pollution could not be controlled. After that, industrial pollution has been severely controlled and lots of factories for producing oxalate had closed all over the world (Hong, 1997). This may be the middle of the 20th century, industrial production of oxalate was high (e.g. in Germany) (Hong, 1997). Thus we assume that C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentration (e.g. the strength of the emissions by vegetation or the oxidation of various alkenes) but is not necessarily the dominant factor for higher C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentrations in the 20th century, notably in the 1950s~1980s. Oxalate has been mass-produced by industry since the 1940s (e.g. in Germany) (Hong, 1997). Thus we assume that C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentrations in the 19th century were not influenced by industrial emissions and may represent natural background values, while higher C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentrations, especially those peaks due to short episodes in 1950s~1980s, may be caused by industrial emissions. Though the transportation and deposition for these industrial C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} pollutants from source regions to mountain glaciers is not well known, the ice core C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} record from Mt. Qomolangma (Mt. Everest) provides a unique opportunity to assess the contribution of anthropogenic emissions to background C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentration. In the middle of the 20th century, industrial production of oxalate was high and the pollution could not be controlled. After that, industrial pollution has been severely controlled and lots of factories for producing oxalate had closed all over the world (Hong, 1997). This may be the cause of the lower C\textsubscript{2}O\textsubscript{4}\textsuperscript{2-} concentrations in the 1990s.

Acknowledgments

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References


(Ren Chinese)


(In Chinese)


