

An ice-core-based record of biomass burning in the Arctic and Subarctic, 1750–1980

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ABSTRACT

Ammonium records from 3 ice cores, 20D and GISP2 (Greenland) and Mt. Logan (Yukon), covering the period from 1750 to the 1980s are analyzed. For each data set, samples with NH_4^+ concentrations greater than one standard deviation above the mean value also tend to be enriched in NO_3^- and K^+ , similar to the chemical composition of aerosols from aged biomass burning plumes. We believe the NH_4^+ spikes originate from biomass burning events. There is not a one to one correspondence between documented large fires and NH_4^+ spikes, nor are specific annual layers with elevated NH_4^+ concentrations often found in more than one core. However, frequency of NH_4^+ spikes increase during periods of more extensive and intensive biomass burning in the NH_4^+ source areas for the ice core sites. The 20D and GISP2 records are characterized by increased spike frequency from 1790 to 1810 and from 1830 to 1910. This latter time coincides with a period of increased biomass burning documented in the historical fire records for northern North America. In contrast to both Greenland ice core records, the Mt. Logan NH_4^+ record shows periods of increased spike frequency from 1770–1790, 1810–1830, 1850–1870 and 1930–1980. The poor agreement between the Mt. Logan record and the records from Greenland suggests that another source area, perhaps Siberia, may be the dominant summertime source area for NH_4^+ spikes in Mt. Logan snow.

1. Introduction

Ammonium concentration records from three high latitude ice cores were examined. Two of the core sites are located on the Greenland icecap, 20D (65.01°N, 44.87°W, 2625 m asl), and GISP2 (72.2°N, 37.80°W, 3200 m asl). The 3rd site, Mt. Logan, in Yukon, Canada (60.58°N, 140.58°W, 5346 m asl) is a very high altitude, subarctic site (Fig. 1). The ice core records studied cover the period from 1750 to the 1980s.

The ammonia component of the nitrogen cycle has been studied intensely in recent years (e.g., Langford et al., 1992; Hegg et al., 1988; Stedman and Shetter, 1983). Ammonia is the dominant

atmospheric gaseous base available for neutralizing H_2SO_4 and HNO_3 . Since both H_2SO_4 and HNO_3 are increasing due to anthropogenic activity, the role of NH_3 is becoming more important. A recent review of NH_3 sources and sinks concluded that continents are the dominant source of NH_3 (Stedman and Shetter, 1983; Langford et al., 1992). Ammonia emissions are positively correlated to temperature, hence are highest in the summertime (LeBel et al., 1985; Langford et al., 1992).

A major impediment to NH_3 and NH_4^+ atmospheric studies has been the difficulty of accurately measuring ambient concentrations of either component (Langford et al., 1992; Griffith et al., 1991; Silvente and Legrand, 1993). The atmospheric lifetime of NH_3 is relatively short in the boundary layer (a few days) due to rapid incor-

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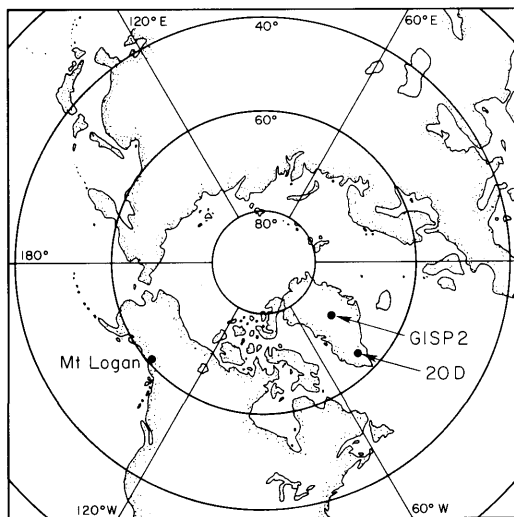


Fig. 1. Location map showing the ice core sites: Mt. Logan (ML) (Yukon), 20D and GISP2 (Greenland).

poration and removal in acidic aerosols, mineral dust aerosols or cloud water. The terrestrial surface source and the reactivity of NH_3 , combine to yield concentration profiles in the atmosphere which rapidly decrease with height. Most NH_3 is returned to the surface as an NH_4^+ salt relatively near the initial source (LeBel et al., 1985; Langford et al., 1992). However, the fraction that does reach the free troposphere can be transported long distances (Langford et al., 1992).

Biomass burning is probably the dominant process which transports high concentrations of NH_3 and NH_4^+ above the boundary layer. Turbulent mixing and convective columns associated with large fires can lift smoke to altitudes greater than 3 Km (Radke et al., 1991; Andreae et al., 1988; Shipham et al., 1992). Recent studies suggest that biomass burning is the dominant source of NH_4^+ for the free troposphere in the arctic and subarctic region of the Northern Hemisphere in the summertime (LeBel et al., 1991; Talbot et al., 1992). If this is true, the NH_4^+ spike records recovered from high latitude ice cores could provide records of biomass burning in the primary summertime aerosol source areas for the different coring sites. Ammonium records from 20D, Mt. Logan and GISP2 were compared to northern hemisphere

biomass burning records to determine if the ice core records can provide a fire activity history.

2. Methods

Cores from 20D, Mt. Logan and GISP2 were collected and processed using stringent precautions and ultraclean procedures to minimize contamination (Mayewski et al., 1986; Holdsworth et al., 1988; Buck et al., 1992). The 115 m 20D and 103 m Mt. Logan cores were sampled at high resolution, averaging 6.3 and 6.4 samples a^{-1} , respectively, for the length of each core. GISP2 samples are approximately biannual, although some sections of the core were sampled at higher resolution (10 samples per year). The discontinuous detailed sections from the GISP2 core (25% of the top 80 m) allowed the seasonality of the chemical species to be determined. Eight ions (Na^+ , NH_4^+ , K^+ , Mg^{2+} , Ca^{2+} , Cl^- , NO_3^- , SO_4^{2-}) were determined for each sample by suppressed ion chromatography using a 0.5 ml sample loop in the Glacier Research Group laboratory (Buck et al., 1992).

Samples from Mt. Logan, GISP2 and the lower 37 mm of 20D (older than 1850) were run within 2 h of initial melting. Samples from the upper portion of the 20D core had been refrozen prior to NH_4^+ analysis. Ammonium values for these samples have been corrected by subtracting a procedural blank. Comparison of data from duplicate aliquots from the GISP2 core, one analyzed immediately and one refrozen prior to analysis, indicates higher NH_4^+ values for refrozen samples. The increase in NH_4^+ concentration in refrozen samples seems to be independent of the initial concentration and has never exceeded $5 \mu\text{g l}^{-1}$ (Unpub. data, GRG). Based on this, refreezing could not cause the large NH_4^+ spikes in the 20D data which are the focus of this paper.

All three cores were dated by multiple techniques. Annual oscillations in $\delta^{18}\text{O}$, soluble ionic chemistry, and correlation of SO_4^{2-} spikes to known volcanic eruptions were used to date the 20D and Mt. Logan cores (Holdsworth et al., 1984, 1991; Mayewski et al., 1986, 1993a). The dating produced annually resolved chronologies back to 1688 at Mt. Logan and 1769 at 20D. The GISP2 core was dated utilizing a combination of

physical stratigraphy, $\delta^{18}\text{O}$, ECM (electrical conductivity measurement), dust, and chemistry as described in Meese et al. (in press).

3. Results

The detailed records from 20D and Mt. Logan are both characterized by NH_4^+ spikes superim-

posed on a relatively uniform, low background (Fig. 2). Median values for the three sites are very similar (Table 1). Inspection of the detailed profiles from all three cores shows that the maximum NH_4^+ concentrations occur in "summertime" snow and minimum concentrations in "wintertime" snow. (Summer and winter are defined in reference to the $\delta^{18}\text{O}$ profiles). In addition, some years are characterized by unusual large NH_4^+

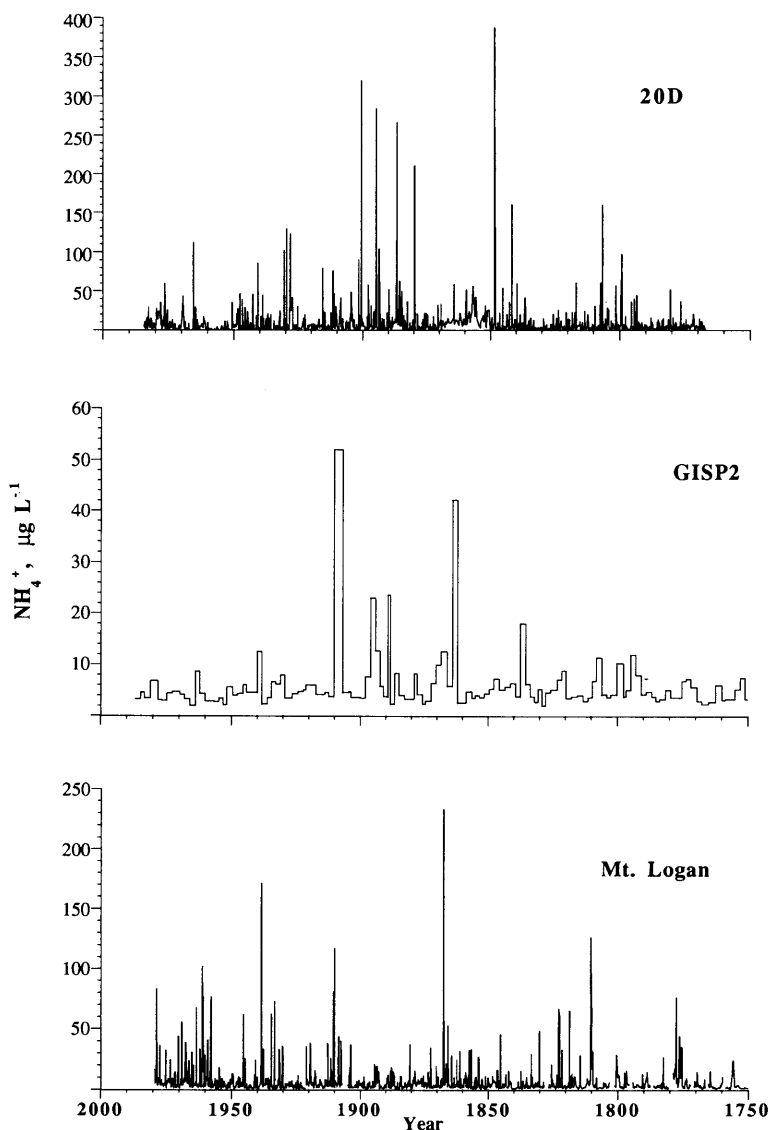


Fig. 2. Detailed records from 20D, GISP2 and Mt. Logan; concentrations ($\mu\text{g l}^{-1}$).

spikes which consist of one or two "summertime" samples with concentrations that are at least a factor of 5 greater than the maximum summertime NH_4^+ concentration of neighboring years (Fig. 3). These unusual spikes are the events of interest.

Annual averages were calculated from the 20D and Mt. Logan detailed data to generate a data set with no seasonal NH_4^+ signal. These derived data sets and the bi-yearly GISP2 data are also characterized by NH_4^+ spikes, defined as NH_4^+ values greater than the mean plus one standard deviation

(Table 1, Fig. 4). All the unusual NH_4^+ events in the 20D and Mt. Logan detailed records survived averaging over annual intervals and still appear as spikes in the smoothed record.

75% of the 20D NH_4^+ spikes and 70% of the GISP2 NH_4^+ spikes are concentrated in the interval from 1830 to 1910. (The GISP2 record shows lower spike concentrations due to the 2-year sampling resolution.) The period from 1790 to 1810 at both sites is also characterized by 2 spikes (Fig. 4). The similarity in the records at 20D and

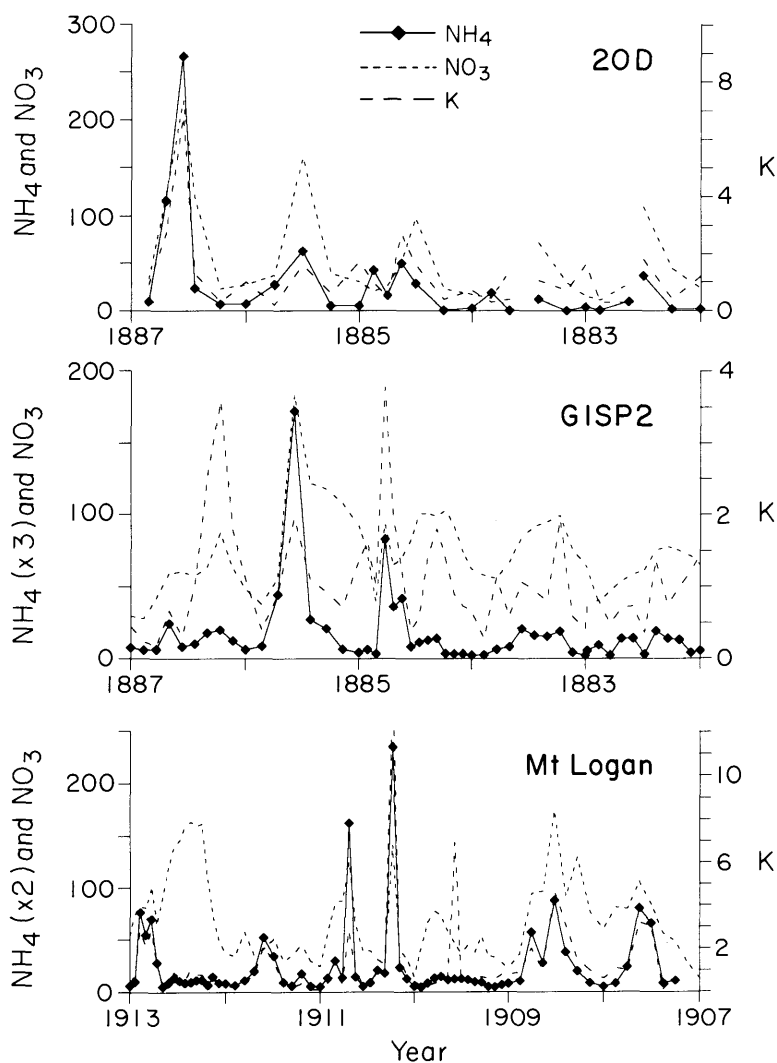


Fig. 3. Plots of detailed NH_4^+ , K^+ , and NO_3^- data over 5-year intervals from 20D, GISP2 and Mt. Logan; concentrations ($\mu\text{g l}^{-1}$).

GISP2 is encouraging because the 2 sites are near the crest of the Greenland ice sheet, separated by 8° of latitude. This similarity suggests that they have common summertime aerosol source areas.

Ammonia spikes in the Mt. Logan record are present in the intervals 1770–1790, 1810–1830, 1850–1870, and 1930–1980 (Fig. 4). 68 % of the NH_4^+ spikes in the Mt. Logan record occur in the

Table 1. Summary statistics for NH_4^+ concentrations, $\mu\text{g l}^{-1}$, based on yearly averages

Site	Mean	σ	Median
Mt. Logan	6.9	6.74	3.5
20-D	9.31	11.83	3.52
GISP	5.89	5.2	4.24

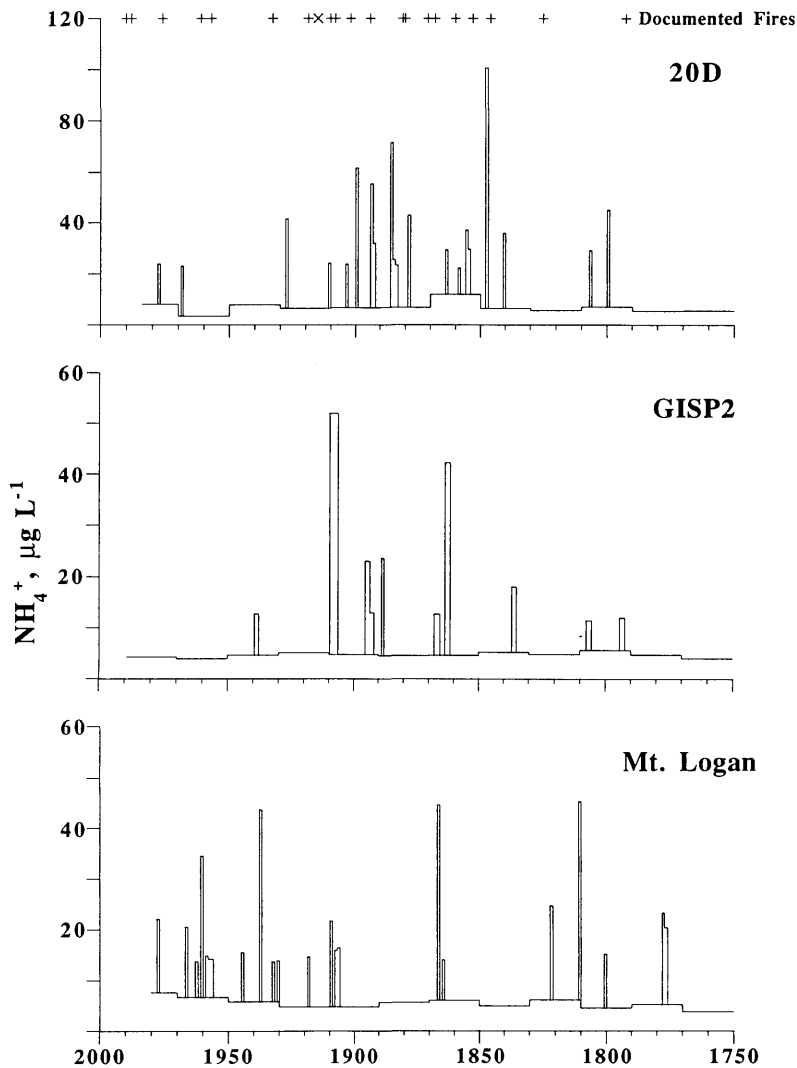


Fig. 4. Plot of NH_4^+ spikes ($>\text{mean} + 1\sigma$) for 20D, GISP2, and Mt. Logan. The baseline NH_4^+ values plotted with the spikes are 20-year averages calculated without the years with spikes. Years with large ($>500,000$ acres) fires in northern North America (+) and Siberia (x) are indicated above the 20D plot.

interval 1900 to 1980, when few are seen in Greenland. The dissimilarity in the timing of the NH_4^+ spikes between Mt. Logan and the 2 Greenland sites strongly suggests that the ice core records from the 2 areas are recording events from different NH_4^+ source areas.

4. Discussion

The large NH_4^+ spikes observed in the ice cores from the three sites are similar to the large, short duration NH_4^+ spikes that have been documented in summer layers in the GRIP ice core from Summit (Legrand et al., 1992). The GRIP samples with elevated NH_4^+ concentrations also had enhanced concentrations of organic acids, especially formate (Legrand et al., 1992). Elevated concentrations of NH_4^+ and organic acids characterize aged biomass burning plumes in the free troposphere (Andreae et al., 1988; Talbot et al., 1992), leading Legrand et al. (1992) to hypothesize that the layers in the GRIP core represent biomass burning events. Organic acids were not analyzed in samples from the three sites discussed here. However, other characteristics of the soluble ion chemistry in the samples with enhanced NH_4^+ concentrations also suggest a biomass burning source.

In addition to higher NH_4^+ and organic acid concentrations, aerosols in aged biomass burning plumes contain elevated concentrations of K^+ , NO_3^- , and SO_4^{2-} (Andreae et al., 1988; Talbot et al., 1992). When the large NH_4^+ spikes from the 3 ice core records are examined at subseasonal resolution; K^+ and NO_3^- concentrations tend to increase in the same samples with higher NH_4^+ concentrations (Fig. 3). Chemical similarity between the composition of the ice core NH_4^+ spikes and aerosol enrichments observed in aged biomass burning plumes suggests that the ice core NH_4^+ spikes record biomass burning events.

General west to east circulation in the upper troposphere at the latitudes of the three ice core sites suggests that northern North America is probably the dominant source area for Greenland and Siberia is a likely source for Mt. Logan (Davidson et al., 1993; Holdsworth et al., 1992). These two areas, Canada and Siberia, are the dominant biomass burning source areas of all the circumpolar countries, with estimated annual burn areas of over 3 million acres in Siberia and over

2 million acres in Canada (Stocks, 1991). Minor inputs to Greenland from Europe and Siberia are also possible due to disruptions in the zonal flow (Davidson et al., 1993). Additional possible source regions impacting Mt. Logan include the neighboring areas of Alaska and British Columbia. Satellite imagery and aircraft sampling have shown that forest fire plumes travel quickly and maintain their visual and chemical identity for at least several days (Robock, 1991; Westphal and Toon, 1991; Ferrare et al., 1990; Talbot et al., 1992; Wofsy et al., 1992); therefore, NH_4^+ spikes source areas do not have to be in close proximity to the core sites.

The unusual high NH_4^+ spikes preserved in the ice core record are the result of a progression of special events: (1) a large, very intense fire (minimum size is not known) must occur in the source area and generate a $\text{NH}_3/\text{NH}_4^+$ rich plume that is lofted rapidly to the free troposphere; (2) wind trajectories must transport the plume quickly to the core site; (3) dilution/dispersion or scavenging of the plume enroute must be minimal; (4) the aerosols in the plume must be deposited and preserved at the ice core site. These special events mean that an ice core record will preserve only a small fraction of large biomass burning events that occur in its summertime source area. However, the number of spikes preserved per unit time should reflect the relative frequency of fires in the source area.

The NH_4^+ spike records from 20D and GISP2 were compared to the fire record for northern North America. Unfortunately, very poor to no historical fire records exist for northern Canada or Alaska prior to the early 1900s (since historical records are biased toward populated areas). Fig. 4 shows 20 documented intense fires (> 500,000 acres burned) which occurred in North America since 1750 and a large Siberian fire in 1915 (Plummer, 1912; Pyne, 1982; Fuller, 1991). As there are no historical or recent fire records for Siberia, probably the dominant source region for Mt. Logan (see below), a similar comparison is not possible.

The sparsity of spikes since 1910 in both Greenland records suggest low biomass burning activity in the source area (Table 2, Fig. 4). This coincides with the beginning of active fire suppression in 1910 in northern North America which resulted in a decrease in fire activity (Van Wagner,

Table 2. Number of individual years with spikes for each 20-year interval for 20D, Mt. Logan and GISP2; ccd-core drill date 1989-GISP2; 1984-20D; and 1980-Mt. Logan

Time	Number of Spikes		
	20D	GISP2	Mt. Logan
1750–1769	0	0	0
1770–1789	0	0	2
1790–1809	2	2	1
1810–1829	0	0	2
1830–1849	2	1	0
1850–1869	4	2	2
1870–1889	5	1	0
1890–1909	4	3	2
1910–1929	2	0	2
1930–1949	0	1	4
1950–1969	1	0	6
1970-cdd	1	0	1

1988; Stocks, 1991). Anthropogenic acidification of the northern hemisphere atmosphere since 1900, which is recorded in Greenland snow by increasing SO_4^{2-} and NO_3^- concentrations (Mayewski et al., 1993a) may be accentuating the decrease through production of ammonium aerosols which are largely scavenged before reaching Greenland (Stedman and Shetter, 1983; Legrand et al., 1992). The most recent NH_4^+ spikes at 20D may reflect the increase in total number of acres burned in North America since the seventies (Van Wagner, 1988; Fuller, 1991; Stocks, 1991).

Although quantitative data about forest fires prior to 1920 does not exist, historical accounts and analysis of tree fire scars indicate that there were numerous, large, extensive forest fires during the 1800s and early 1900s in northern North America (Plummer, 1912; Williams, 1989; Pyne, 1982; Leavitt, 1991; Suffling, 1991; Alexander, 1980; Johnson, 1980; Tande, 1980; Payette, 1980). This period of greater biomass burning is reflected in the 20D and GISP2 NH_4^+ records by the high number of NH_4^+ spikes during the 1830–1900 period (Fig. 4).

The Mt. Logan NH_4^+ spikes do not coincide with spikes in the two Greenland records. This agrees with the earlier observation that North America is not a dominant source region for air over Mt. Logan (Holdsworth et al., 1991; Mayewski et al., 1993a). Central and eastern

Siberia are possible source areas; plumes from the 1987 fire in central Siberia were observed in satellite images over Alaska (Robock, 1991). In addition, aged biomass burning plumes from Siberia were sampled over Alaska during ABLE 3A (Shipham et al., 1992; Wofsy et al., 1992). The Mt. Logan NH_4^+ spike record suggests there have been 4 periods of higher biomass burning in the source area, 1770–1790, 1810–1830, 1850–1870 and 1930–1980. The decreasing frequency of intense forest fires in Alaska and Canada since 1920 and the increasing frequency of NH_4^+ spikes in the Mt. Logan data, suggests that local biomass burning is of minor importance to this record.

If as suggested above, the NH_4^+ spike frequency record preserves a generalized biomass burning record of the summertime source area, this record may also be used as a general indicator of the existence of climatic conditions conducive to biomass burning. The North American record since 1920 has been dominated by fire suppression and fire control which probably has modified forest fire response to climatic factors. The recent increase in North American fires is thought to be climate driven, corresponding to dry summer weather (Suffling, 1991; Fuller, 1991; Stocks, 1991; Schindler et al., 1990). Current biomass burning response may be enhanced due to the tinder accumulated during the preceding decades of fire suppression. It has also been suggested that the North American fires in the 1800s were largely climate driven, the number of fires increasing during droughts (Suffling, 1991; Tande, 1980; Alexander, 1980; Schindler et al., 1990; Payette, 1980). The mid to late 1800s is considered to be the end of the Little Ice Age (Lamb, 1982; Mayewski et al., 1993b). Fire intensity and frequency in the 1800s was amplified by extensive burning to clear land for agriculture as the North American populus rapidly expanded westward (Pyne, 1982). In addition, logging practices of that era left much waste behind which served as fire fuel, and sparks from the railroad increased the possibility of ignition (Plummer, 1912; Leavitt, 1991; Williams, 1989; Tande, 1980).

If as suggested by Stocks (1991), most of the fires in Siberia are caused by lightning (due to the remoteness and low population density of the area), the record from Mt. Logan may be a better example of the climate-forest fire relationship that could be used with longer ice core records.

5. Conclusions

Correspondence between the biomass burning history of northern North America since the 1800s and NH_4^+ spike records from 20D and GISP2 suggests that these ice cores at least qualitatively record the frequency of intense, large biomass burning in northern North America. It may be possible to extend the relationship further based on the general coincidence between dry summer conditions and increased fire intensity and activity (Van Wagner, 1988). This suggests that the intervals characterized by a high frequency of NH_4^+ spikes may also correspond to periods of dry summer weather in the source area.

The difference between the NH_4^+ spike record from Mt. Logan and the two records from Greenland suggests that Siberia and northern North America have very different biomass burning histories. The NH_4^+ record from Mt. Logan suggests that Siberia has experienced an increase in intense, extensive biomass burning since the late

1800s. The average NH_4^+ values from 1930 to 1970 are the highest in the 230 year record, and also represent the longest time period of high average NH_4^+ values.

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