

# Partitioning of reactive nitrogen ( $NO_y$ ) and dependence on meteorological conditions in the lower free troposphere

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Abstract. Results of continuous nitrogen oxide (NO), nitrogen dioxide (NO<sub>2</sub>), peroxyacetyl nitrate (PAN) and total reactive nitrogen (NO<sub>v</sub>) measurements along with seasonal field campaigns of nitric acid (HNO3) and particulate nitrate (NO<sub>3</sub><sup>-</sup>) measurements are presented for a two-year period at the high-alpine research station Jungfraujoch (JFJ), 3580 m asl. The NO<sub>v</sub> mixing ratio and partitioning is shown to strongly depend on meteorological conditions. Knowledge of these meteorological transport processes allows discrimination between undisturbed (i.e. clean) and disturbed (i.e. influenced by regional pollution sources) free tropospheric (FT) conditions at the JFJ. Median NO<sub>v</sub> concentrations during undisturbed FT periods ranged from 350 pptv (winter, December to February) to 581 pptv (spring, March to May). PAN was found to be the dominant NO<sub>v</sub> species during spring and summer, whereas NO2 was most abundant during autumn and winter. Particulate nitrate was found to contribute significantly to total NO<sub>v</sub> during thermally induced vertical transport. Föhn events, synoptical lifting (e.g. fronts) and thermally induced vertical transport resulted in mixing ratios up to 10 times higher at the JFJ compared to undisturbed FT conditions. Furthermore this meteorological variability of the NO<sub>v</sub> concentration and partitioning often dominated the seasonal variability. As a consequence the use of filters at the JFJ (and other mountainous sites) is crucial for the interpretation of data from such measurement sites. This study presents a further development of meteorological filters for the high-alpine site Jungfraujoch, which also could be modified and adapted to other mountainous measurement sites.

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#### 1 Introduction

Reactive nitrogen compounds play a central role in the chemistry of the troposphere. Nitrogen oxides  $(NO_x = NO +$ NO<sub>2</sub>) are the limiting precursors for O<sub>3</sub> production throughout most of the troposphere, and also directly influence the abundance of the hydroxyl radical concentration in the troposphere (e.g. Crutzen, 1979; Roberts, 1995). Emissions of reactive nitrogen occur primarily as NO, followed by oxidation to NO<sub>2</sub>. Reactive nitrogen, abbreviated NO<sub>v</sub>, is defined as the sum of NO<sub>x</sub> and its atmospheric oxidation products, abbreviated NO<sub>z</sub>. NO<sub>2</sub>, peroxyacetyl nitrate (PAN), nitric acid (HNO<sub>3</sub>) and particulate nitrate (NO<sub>3</sub><sup>-</sup>) are considered to be the most abundant NO<sub>v</sub> compounds in the troposphere, while their relative abundance varies significantly. For instance, NO<sub>2</sub> is often dominant close to sources; PAN tends to be most abundant in regionally polluted air masses where there is more active organic photochemistry; and inorganic nitrates are most abundant in the more remote areas of the troposphere. However, there is significant uncertainty in the levels and speciation of NO<sub>v</sub> in the non-urban troposphere due to a lack of measurements (Carroll and Thompson, 1995).

In the present study, we present a two-year time series of total and speciated NO<sub>y</sub> from the high Alpine Research Station Jungfraujoch (JFJ), Switzerland. This is the first longer-term data series on nitrogen oxides levels in the free troposphere over Central Europe. Ongoing measurements at the JFJ of NO, NO<sub>2</sub> (since June 1991) and total NO<sub>y</sub> (since March 1997) were accompanied by PAN measurements (March 1997 to April 1998) and seasonal measurement campaigns of particulate nitrate and HNO<sub>3</sub> (between July 1997 and April 1998). In contrast to previously published work (Zellweger et al., 2000) the focus of this paper is on the seasonality of NO<sub>y</sub> and its constituents in the undisturbed troposphere over Central Europe. Further attention was also given to the identification of meteorological transport processes that occasionally cause pollution episodes at the JFJ.

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The JFJ is one of the 16 measurement sites of the Swiss National Air Pollution Monitoring Network (NABEL). Furthermore, the JFJ high-alpine station has been the site of an aerosol program since 1988 (Baltensperger et al., 1997; Lugauer et al., 1998; Nyeki et al., 1998). Due to the importance of monitoring long-term trends of gaseous and aerosol parameters in the remote troposphere, the JFJ station has been incorporated into the Global Atmosphere Watch (GAW) program of the World Meteorological Organization (WMO). The JFJ is also part of the combined European GAW virtual baseline station, additionally comprising the Zugspitze (2962 m, Germany) and Sonnblick (3106 m, Austria) highalpine stations.

Although air-quality monitoring networks provide observations of the spatial and temporal variation of trace gases, they usually provide no information on chemistry above ground and higher. Aircraft observations have shown that surface and upper air chemistry are frequently decoupled (Berkowitz et al., 1998). As a consequence, surface measurements are in general only representative of measurements near the ground. In contrast, mountainous stations are often decoupled from the planetary boundary layer and measurements are therefore representative of a larger region. Nevertheless, it is also important to develop methods to detect periods of local or regional pollution at mountainous stations. The influence of different meteorological processes on the NO<sub>v</sub> mixing ratio at the JFJ is specifically considered in this paper in order to better quantify disturbed and undisturbed free tropospheric (FT) conditions, and hence aid in the interpretation of long-term measurements.

## 2 Experimental

Total NO<sub>y</sub> measurements began in March 1997 at the JFJ. While the measurements of NO and NO<sub>2</sub> started in June 1991 and are along with total NO<sub>y</sub> measurements ongoing, PAN measurements were made between March 1997 and April 1998. Measurements of particulate nitrate and HNO<sub>3</sub> were performed during four seasonal field campaigns between July 1997 and April 1998. Because the measurement station and instrumental details have already been described elsewhere (Baltensperger et al., 1997; Weingartner et al., 1999; Zellweger et al., 1999, 2000), only a brief summary of experimental aspects is presented here.

#### 2.1 Measurement site and inlet system

The high alpine research station Jungfraujoch (46°33′ N, 7°59′ E, 3580 m asl) is located on the main crest of the Bernese Alps, Switzerland (e.g. Baltensperger et al., 1997). Two inlet systems, both located on top of the Sphinx building, were used for the measurements of NO, NO<sub>2</sub>, NO<sub>y</sub>, HNO<sub>3</sub>, particulate nitrate and PAN. HNO<sub>3</sub>, particulate nitrate and the aerosol surface area were measured using the

GAW aerosol inlet (Weingartner et al., 1999). In addition to the original inlet system, a polyethylene tube of 3 m length (i.d. 4 mm) was used inside the stainless steel tube. The inlet heating (20°C) resulted in low relative humidity (<10%). The sample flow rate was  $41\,\mathrm{min}^{-1}$ . NO, NO<sub>2</sub>, NO<sub>y</sub>, PAN and CO and O<sub>3</sub> were measured using the NABEL stainless steel inlet with an inside diameter (i.d.) of 8 cm and a total length of 3.1 m. The air flow rate was  $50\,\mathrm{m}^3\,\mathrm{h}^{-1}$ , and a heating system provided a constant gas temperature of  $12^\circ\mathrm{C}$ . NO<sub>x</sub>, NO<sub>y</sub> and PAN instruments were connected directly to the inlet with PTFE tubing.

## 2.2 Instrumentation

NO, NO<sub>x</sub> and NO<sub>y</sub> were measured with a commercially available instrument (CraNOx, Ecophysics) using two chemiluminescence detectors (CLD 770 AL pptv) with temperature-controlled reaction chambers. NO<sub>x</sub> was measured as NO after photolytic conversion (PLC 760). NO<sub>v</sub> species were converted on a heated gold catalyst (300° C) with 2% CO (99.997%, Messer-Griesheim GmbH) as a reducing agent (Bollinger et al., 1983; Fahey et al., 1985, 1986). A full description of the instrumentation including interferences, conversion efficiencies, statistics and sensitivities can be found in Zellweger et al. (2000). An automatic calibration of the instrument was performed every 23 h. The chemiluminescence detectors were calibrated with NO standard gas (5  $\pm$  0.1 ppmv, diluted with zero air to 30 ppbv) and zero air (dew point  $-20^{\circ}$  C), and the conversion efficiencies of both converters were measured for NO<sub>2</sub> by gas phase titration of NO with ozone. When the conversion efficiency dropped below 95% ( $\sim$  every 3 months), the Au catalyst was cleaned with ethanol, acetone and ultrapure water. The conversion efficiencies of the PLC ranged from 45 to 82%. The NO, NO<sub>2</sub> and NO<sub>y</sub> impurity of the zero air was checked daily. The NO standard mixture (NO 99.8% in N<sub>2</sub> 99.999%, Messer-Griesheim GmbH) was traced back to the National Institute of Standards and Technology (NIST) Reference Material. The instrumental detection limit for NO, NO<sub>2</sub> and NO<sub>v</sub> was 50 pptv for 2 min and 20 pptv for 30 min average values, respectively. Overall uncertainties in the measurements were estimated to be  $\pm 5\%$  for NO,  $\pm 10\%$  for NO<sub>2</sub> and  $\pm 9\%$  for NO<sub>v</sub> at ambient levels of 500 pptv (1 $\sigma$ ). They include the precision of the CLDs, the NO standard uncertainty, the conversion efficiencies of the PLC (after linear detrending) and the Au catalyst, and artifact uncertainties of the zero air check. Our NO<sub>x</sub> and NO<sub>y</sub> measurements were compared at the JFJ during spring 1998 with measurements of the School of Environmental Sciences, University of East Anglia (UEA), Norwich (UK), to assess the overall measurement uncertainty. The inlet of the UEA custom-built system consisted of a 1/4 inch PFA tube of 0.2 m length. Further details of the UEA instrument are given in Bauguitte (1998). Agreement of both systems was within 10% (Carpenter et al., 2000), with slightly lower values for our system. For the inter-comparison period between 24 March and 5 April 1998, the following linear relationship for a concentration range of 0 to 2 ppbv was found between the two instruments:

EMPA NO<sub>y</sub> (pptv) = 0.94 \* UEA NO<sub>y</sub> (pptv) + 22.9 pptv ( $r^2 = 0.89$ ).

Slightly lower values in the EMPA system might be attributed to potential losses of HNO<sub>3</sub> in the inlet system. Therefore, the overall uncertainty of NO<sub>y</sub> measurements including inlet losses is estimated to be +9/-15%.

PAN was measured with a commercially available gas chromatograph (GC) and a calibration unit (Meteorologie Consult GmbH) coupled to an electron capture detection unit (ECD). The instrument and the calibration procedure is described in Zellweger et al. (2000). The detection limit was 50 pptv, and the overall measurement uncertainty was estimated to be  $\pm 3\%$  ( $1\sigma$ ).

HNO<sub>3</sub> and particulate nitrate measurements were carried out with the wet effluent diffusion denuder/aerosol collector (WEDD/AC) technique described by Blatter et al. (1994) and Simon and Dasgupta (1995). The detection limit was 10 pptv for HNO<sub>3</sub> and particulate nitrate (10 min sampling time). Details of the instrument are described in Zellweger et al. (1999, 2000).

Both CO and O<sub>3</sub> were continuously monitored with commercially available instruments by the NABEL network (APMA-360, Horiba, for CO; Thermo Environmental Instruments, Model 49C, for O<sub>3</sub>). The detection limit was 30 ppbv for CO and 0.5 ppbv for O<sub>3</sub> (30 min average). More detailed information including measurement uncertainties and the calibration procedure can be found in Zellweger et al. (2000).

The aerosol active surface area concentration was measured using an epiphaniometer (Gäggeler et al., 1989). This instrument detects the active surface area concentration, which represents the total particle surface area accessible to a diffusing molecule (Baltensperger et al., 2001). The data is corrected for the decline of the actinium source ( $T_{1/2} = 21.7$  years) and was inverted using the algorithm presented in Rogak et al. (1991). The raw data (units: counts per second, cps) were converted into an active surface area concentration ( $\mu$ m<sup>2</sup> cm<sup>-3</sup>) using the algorithm described in (Baltensperger et al., 2001). The relative statistical error of the epiphaniometer signal was found to be 7% for median winter concentrations and negligible for summer concentrations.

# 2.3 Definition of meteorological filters

Meteorological processes were recognized as playing an important role in the understanding of measurements obtained at elevated continental sites. Therefore, care was taken to identify and separate periods with a potential perturbation due to local or regional pollution sources. The data was divided into two periods, "undisturbed" and "disturbed" free tropospheric (FT) conditions. The term "disturbed FT" is used to account for the various processes involved during dilution of PBL air with FT air during upward transport. For in-

stance, the contribution of PBL to undisturbed FT air masses was estimated at 14–20% during convection events. This corresponds to a dilution factor of  $\sim$ 5 to 7 of PBL air by FT air masses (Zellweger et al., 2000). In this work, disturbed FT conditions are defined as FT air masses that have been mixed with PBL air masses due to any of the following meteorological conditions: (1) föhn (or chinook) events, (2) synoptical lifting (e.g. frontal systems), and (3) thermally induced vertical transport. These three conditions may significantly influence concentrations of gaseous and aerosol parameters and are crucial for the interpretation of time series obtained at Alpine stations. Data were categorized in the priority (1)>(2)>(3) in order to optimize discrimination, and criteria are further described below.

- Föhn events were excluded from the FT data using a procedure described by Forrer et al. (2000). In addition to the work of Forrer et al. (2000), north föhn (NF) and south föhn (SF) events were distinguished. The following criteria were used to detect föhn events, in which all criteria had to be fulfilled:
  - The absolute north-south pressure difference over the Alps using data from the Stabio (Southern Switzerland) and Schaffhausen (Northern Switzerland) meteorological stations was >2.1 hPa/100 km according to Hoinka (1980).
  - The relative humidity on the lee side of the Alps using data from Altdorf for south-föhn and Locarno-Monti for north-föhn events was <50%, and accompanied by precipitation at the other station.</li>
  - The local wind speed at the JFJ was >10 m s<sup>-1</sup>.
    These criteria were verified by comparison with weather charts, the Swiss Meteorological Institute (SMI) bulletin and 3-D back-trajectories.
- 2. Synoptical lifting (SYN) (e.g. frontal systems) was excluded from the undisturbed FT data by running 3-D back-trajectories calculated by the TRAJEK DWDmodel (Fay et al., 1995) using the wind fields of the Swiss Model (SM). The SM is a hydrostatic numerical weather prediction model with a grid size of  $\sim$ 14 km, which is operationally used at the SMI. The arrival location of the calculated trajectories was the JFJ, and the altitude was set to 700 hPa in the model. The trajectories were then analyzed with respect to their minimum height (corresponding to maximum pressure) 24 h previous to their arrival time. If the trajectory height dropped below the 850 hPa level for at least four hours, the data were excluded from the undisturbed FT data set. These criteria were verified by comparison with data from the Alpine Weather Statistics (AWS) scheme (Schüepp, 1979; Wanner et al., 1998; Forrer et al., 2000).

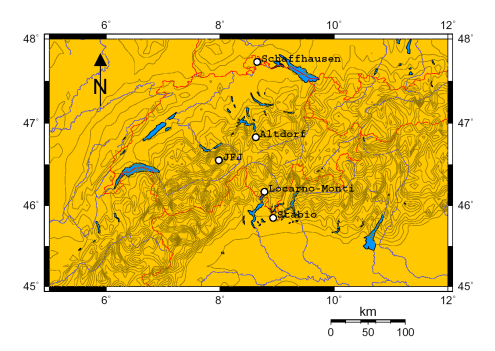


Fig. 1. Map showing the location of the Jungfraujoch and the other stations used for the definitions of the meteorological filters.

- 3. Periods where thermally induced vertical transport (THER) occurred were excluded from undisturbed FT data using the following procedure:
  - Periods with diurnal peak concentrations between 15 February and 15 October were excluded from FT conditions by comparing the mean concentrations of NO<sub>v</sub> for the diurnal periods between 03:00 to 09:00 CET on two consecutive days with the period from 15:00 to 21:00 CET between the morning periods. If NO<sub>v</sub> was ≥50% higher during the afternoon period, only the data from 03:00 to 09:00 CET was considered to represent undisturbed FT conditions. If no NO<sub>v</sub> data was available, the aerosol surface area concentration was used because of its similar diurnal behavior during thermally induced vertical transport at the JFJ (Zellweger et al., 2000). If both NO<sub>v</sub> and aerosol surface area concentration data were missing, the specific humidity (q) was used. For q, the criterion for thermally induced vertical transport was a  $\geq 25\%$  higher concentration during the afternoon in comparison to the morning periods (same time intervals as above).
  - Convective days (anticyclonic or indifferent conditions) from March to September with low wind speed at the 500 hPa level were excluded entirely from the FT data. Such periods were shown to be influenced by convective boundary layer (CBL) air masses throughout the whole day (Zellweger et al., 2000).

A map showing the location of the Jungfraujoch and the other stations used for the evaluation of the above described meteorological filters is presented in Fig. 1. The seasons in this work are defined as spring (March to May), summer (June to August), autumn (September to November), and winter (December to February). The winter half year is defined as from September to February.

#### 3 Results and discussion

Seasonal variation and meteorological transport processes

An overview of monthly median and mean NO<sub>y</sub>, NO<sub>x</sub>, PAN mixing ratios and the NO<sub>x</sub>/NO<sub>y</sub> ratio for the period April 1997 to March 1999 is given in Fig. 2 for the entire data set. Generally higher NO<sub>y</sub> concentrations were observed at the JFJ during spring and summer, due to enhanced vertical transport processes during these seasons, and in the case of PAN higher production rates within the CBL. The monthly mean values of NO<sub>x</sub> and NO<sub>y</sub> were often found to be a factor of two or more higher than the corresponding median values due to the occurrence of relatively short episodes with high concentrations. This can be attributed to periods with transport of polluted air masses from source regions to the JFJ caused by various meteorological processes (Zellweger et al., 2000).

Ground-based measurements of  $NO_y$  at remote sites are relatively sparse in number, and only few data are available. The data presented here is the first longer-term data series

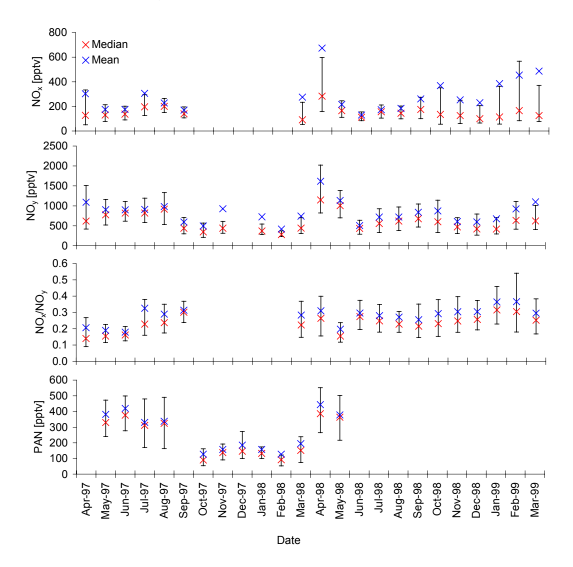
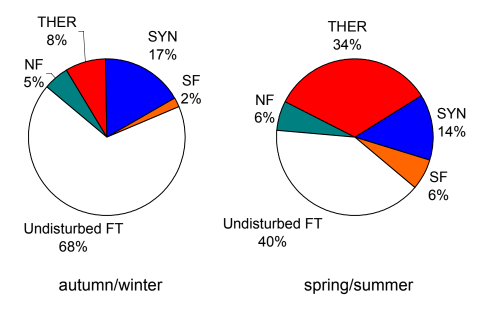


Fig. 2. Monthly mean and median values of  $NO_x$ ,  $NO_y$ , PAN and the  $NO_x/NO_y$  ratio from April 1997 to March 1999. The first and third quartiles are indicated by bars. Only data where >75% of the hourly values were available are shown.

on nitrogen oxides levels in the free troposphere over Central Europe. Such measurements in the undisturbed troposphere can only be performed at high mountainous sites on a continuous basis. The NO<sub>v</sub> concentrations measured at the JFJ are generally higher compared to those at other remote sites. Data from other European sites are not available or not yet published or. However, NO<sub>v</sub> measurements without further speciation except for NO and NO<sub>2</sub> have been performed at the Hohenpeissenberg station (Germany, 985 m a.s.l.) since 1999, and hourly data are available from the World Data Centre for Greenhouse Gases (http://gaw.kishou.go.jp/wdcgg.html). During 1999, NO<sub>v</sub> averaged 3.14 ppbv at the Hohenpeissenberg station, while a yearly average of only 0.88 ppbv was observed at the Jungfraujoch (JFJ). This demonstrates that the two sites are not easily comparable, especially when the difference in altitude of 2595 m between the sites is considered, and suggests that the Hohenpeissenberg station is by far more often influenced by polluted air masses than the JFJ.

Fischer et al. (1998) reported NO<sub>y</sub> measurements at the Izaña Observatory on Tenerife Island. During the Oxidizing Capacity of the Tropospheric Atmosphere (OCTA) campaign, nighttime mean NO<sub>y</sub> concentrations of 370 to 420 pptv were reported. This is slightly below the NO<sub>y</sub> level of mean 653 pptv that is observed during summer for undisturbed FT conditions at the JFJ. The lowest values that have been measured at Izaña are as low as 120 pptv, which is comparable to the lowest values observed at the JFJ during undisturbed FT conditions. At Izaña thermally induced vertical transport was also recognized as playing an important role in the upward transport of polluted air masses. However, the difference between the "polluted" daytime conditions of Izaña with a reported mean NO<sub>y</sub> of 520 to 1300 pptv is often less pronounced compared to the JFJ (mean summer NO<sub>y</sub> of



**Fig. 3.** Frequencies of meteorological situations (NF = north föhn, SF = south föhn, THER = thermally induced vertical transport, SYN = synoptical lifting) for the period between April 1997 and March 1999. Results shown for autumn/winter (September to February) and spring/summer (March to August) half-year periods.

899 pptv during polluted conditions). The period when high daytime NO<sub>y</sub> of 1300 pptv was reported at Izaña was associated with air parcels of continental origin. NO<sub>y</sub> mixing ratios at this or a higher level are also often observed at the JFJ, indicating anthropogenic influences.

In addition, a few aircraft based measurements over Europe are published. However, these measurements often cover only a few hours and were performed at different altitude levels in the troposphere. Hering et al. (1998) described aircraft NO<sub>v</sub> measurements performed over the Swiss plateau during the POLLUMET (POLLUtion and METeorology) study in summer 1993. NO<sub>v</sub> concentrations ranged from 1.12 to 21.52 ppb. The flight altitude was significantly lower (500 to 1300 m a.s.l.) compared to the JFJ. However, this data may also be representative for air masses reaching the JFJ during thermally induced vertical transport (average NO<sub>v</sub> 0.99 ppbv) when dilution with undisturbed FT air during upward transport is considered. Other aircraft measurement campaigns were performed in the upper troposphere. For example, Ziereis et al. (2000) reported an average NO<sub>v</sub> mixing ratio of 0.24 ppbv measured during the POLINAT 2 campaign at the 8000 m level between 49°N and 57°N over the North Atlantic between 19 September and 25 October 1997. During undisturbed FT conditions, NO<sub>v</sub> averaged 0.37 ppbv (median 0.20 ppbv) at the JFJ for the above time period. Thus, the JFJ concentrations are relatively well comparable to the values observed during POLINAT 2.

In the following, the influence of different meteorological situations on the JFJ data set is investigated. Figure 3 shows the frequencies of the meteorological situations described in Sect. 2.3 for the autumn / winter (September to February) and

spring / summer (March to August) half-year of the period from April 1997 to March 1999. As expected, thermally induced vertical transport occurs predominately during spring and summer, whereas frontal systems and föhn events occur all year round.

Table 1 gives an overview of seasonal median and mean NO<sub>x</sub>, NO<sub>y</sub>, PAN, HNO<sub>3</sub>, particulate nitrate, CO, O<sub>3</sub> and the aerosol surface area concentration measured at the JFJ for the period from April 1997 to March 1999. Table 1 includes the whole data set based on hourly values, as well as for undisturbed FT conditions, and disturbed FT conditions. It can be seen from Table 1 that the NO<sub>v</sub> mixing ratio is comparable during the spring and summer months (median NO<sub>v</sub> 766 pptv and 702 pptv, all data) and during the autumn and winter months (median NO<sub>v</sub> 495 pptv and 411 pptv, all data). This is also the case for the NO<sub>x</sub> and PAN mixing ratios, while the aerosol surface area concentration, particulate nitrate and nitric acid showed maximum concentrations during the summer months. The effect of data filtering for NO<sub>x</sub>, NO<sub>y</sub>, PAN and the aerosol surface area concentration is illustrated in Figs. 4a and b, where monthly median mixing ratios are shown for the filtered data set as well as for föhn events, synoptical lifting and thermally induced vertical transport. It can be seen that the meteorological processes described in Sect. 2.3 caused an increase of the mixing ratios as well as of the variability for NO<sub>x</sub>, NO<sub>y</sub> and PAN. The effect was most pronounced for NO<sub>x</sub> as a primary pollutant with a strong vertical concentration gradient. Different behavior was observed for the aerosol surface area concentration, where higher values were observed only for thermally induced vertical transport. The lower values for the aerosol

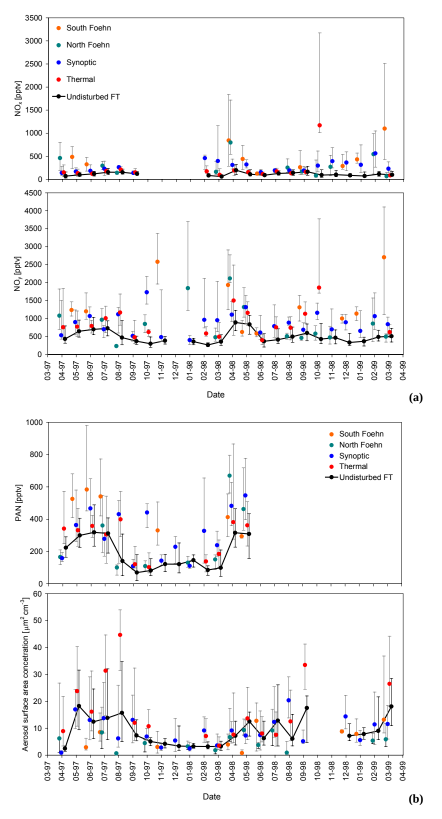
**Table 1.** Seasonal median and mean values of  $NO_x$ ,  $NO_y$ , PAN, HNO<sub>3</sub>, particulate nitrate, CO, O<sub>3</sub>, and the aerosol surface area (AS) concentration with standard deviation (s.d.) and number of measurements (N) based on hourly values at the JFJ for the period from April 1997 to March 1999, for the whole data set, as well as for undisturbed and disturbed FT conditions. (Spring MAM = March to May, etc.)

	All Data				Undisturbed FT				Disturbed FT				
	Median	Mean	s.d.	N	Median	Mean	s.d.	N	Median	Mean	s.d.	N	
	Spring MAM												
NO <sub>x</sub> [pptv]	152	370	769	3860	98	183	289	1631	204	506	958	2229	
NO <sub>v</sub> [pptv]	766	1103	1076	3978	581	748	614	1693	981	1365	1255	2285	
PAN [pptv]	294	346	245	2828	256	287	212	1192	334	389	258	1636	
HNO <sub>3</sub> [ppt]	10	18	19	420	5	17	20	182	14	20	19	238	
NO <sub>3</sub> [pptv]	5	269	754	206	5	24	46	81	5	427	935	125	
CO [ppbv]	168	174	35	4378	160	165	28	1875	173	181	38	2503	
O <sub>3</sub> [ppbv]	58	58	9	4316	57	57	8	1847	59	59	9	2469	
AS [µm <sup>2</sup> cm <sup>-3</sup> ]	8	14	17	3489	8	13	16	1511	8	14	18	1978	
	Summer JJA												
NO <sub>x</sub> [pptv]	161	206	335	3144	133	191	482	1185	177	215	199	1959	
NO <sub>v</sub> [pptv]	702	806	538	3239	529	653	570	1225	822	899	495	2014	
PAN [pptv]	330	355	211	1804	271	286	198	637	369	393	209	1167	
HNO <sub>3</sub> [pptv]	19	70	180	817	17	41	96	235	22	82	203	582	
NO <sub>3</sub> [pptv]	122	161	147	812	68	116	121	275	130	173	155	573	
CO [ppbv]	126	129	28	4288	122	125	28	1592	128	131	27	2696	
O <sub>3</sub> [ppbv]	59	58	9	4261	58	57	10	1587	59	58	8	2674	
AS [µm <sup>2</sup> cm <sup>-3</sup> ]	13	19	17	4151	10	16	16	1565	15	21	18	2586	
		Autumn SON											
NO <sub>x</sub> [pptv]	143	276	494	2234	115	205	355	1289	184	373	624	945	
NO <sub>v</sub> [pptv]	495	738	869	3626	422	553	477	2269	700	1049	1221	1351	
PAN [pptv]	118	145	124	1456	102	119	81	983	142	200	171	473	
HNO <sub>3</sub> [pptv]	5	17	33	765	5	18	36	628	5	11	15	137	
NO <sub>3</sub> [pptv]	5	12	13	761	5	11	14	624	15	15	9	137	
CO [ppbv]	143	149	40	4190	138	146	40	2539	149	154	39	1651	
O <sub>3</sub> [ppbv]	49	49	7	4176	49	49	7	2541	50	50	8	1635	
AS [µm <sup>2</sup> cm <sup>-3</sup> ]	5	10	13	2258	5	7	7	1342	7	15	17	916	
	Winter DJF												
NO <sub>x</sub> [pptv]	115	320	565	2281	87	203	454	1636	409	617	695	643	
NO <sub>y</sub> [pptv]	411	673	797	2997	350	501	659	2233	911	1178	939	764	
PAN [pptv]	115	154	156	1453	105	132	105	1078	149	219	239	375	
HNO <sub>3</sub> [pptv]	11	14	15	762	10	13	14	697	19	26	23	65	
NO <sub>3</sub> [pptv]	11	20	63	763	11	19	64	697	18	31	39	66	
CO [ppbv]	171	179	46	3248	164	169	41	2409	191	207	49	839	
O <sub>3</sub> [ppbv]	46	46	7	4027	47	47	7	3038	43	42	7	989	
AS [µm <sup>2</sup> cm <sup>-3</sup> ]	6	9	13	4267	6	9	13	3157	7	11	14	1110	

surface area concentration during föhn events and synoptical lifting can be explained by precipitation scavenging during these transport events. This mechanism is the major sink for particulate matter as well as for the highly water soluble nitric acid. Thus, the summer maximum of aerosol parameters, particulate nitrate and HNO<sub>3</sub> can be explained by episodes with thermally induced vertical transport processes,

when production of HNO<sub>3</sub> is efficient and wet deposition is of minor importance.

Frequency distributions of  $NO_x$ ,  $NO_y$ , the  $NO_x/NO_y$  ratio, PAN, the aerosol surface area concentration, and CO are shown in Figs. 5a and b. For each species, undisturbed FT conditions, north and south föhn events, synoptical lifting and thermally induced vertical transport data are shown



**Fig. 4.** (a) Monthly median values of  $NO_x$  and  $NO_y$  concentration for the filtered data set as well as for different meteorological situations from April 1997 to March 1999. The first and third quartiles are indicated by bars. (b) Monthly median values of PAN and the aerosol surface area concentration for the filtered data set as well as for different meteorological situations from April 1997 to March 1999. The first and third quartiles are indicated by bars.

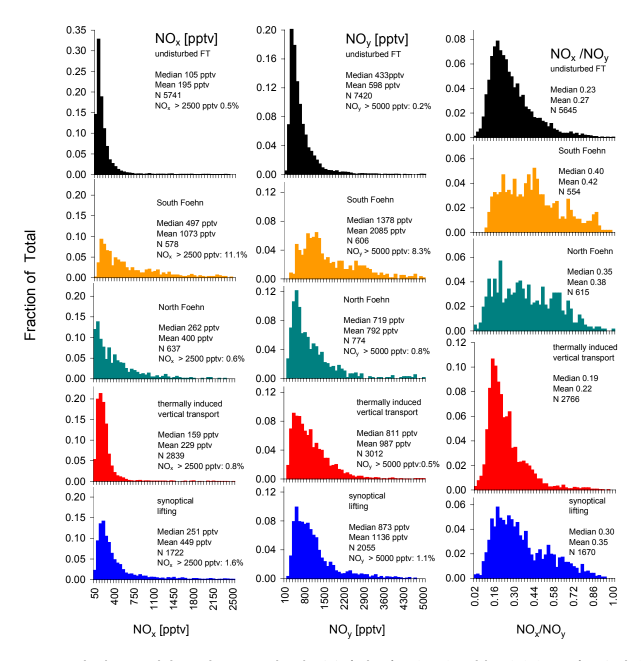
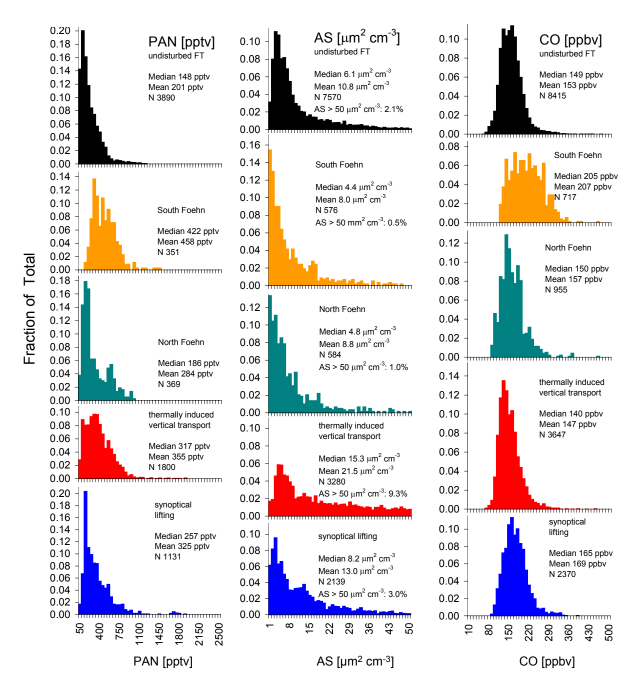


Fig. 5a. Frequency distributions including median, mean and number (N) of values for  $NO_x$ ,  $NO_y$  and the  $NO_x/NO_y$  ratio from April 1997 to March 1999 for the whole data set and different meteorological situations. For each species, the data set was based on hourly values and divided into 50 bins.

separately. The plots in Figs. 5a and b are based on hourly mean values, and for each species the data was divided into 50 bins.

It can be seen from Figs. 5a and b that the mixing ratios of  $NO_x$ ,  $NO_y$ , CO, PAN and the  $NO_x/NO_y$  ratio were elevated during south föhn events, in conjunction with low aerosol concentrations. Similar increases of the CO and  $NO_x$  mixing ratios at the JFJ during south föhn episodes were also observed by Forrer et al. (1999) for the period between April 1996 and November 1997. During the observation period be-

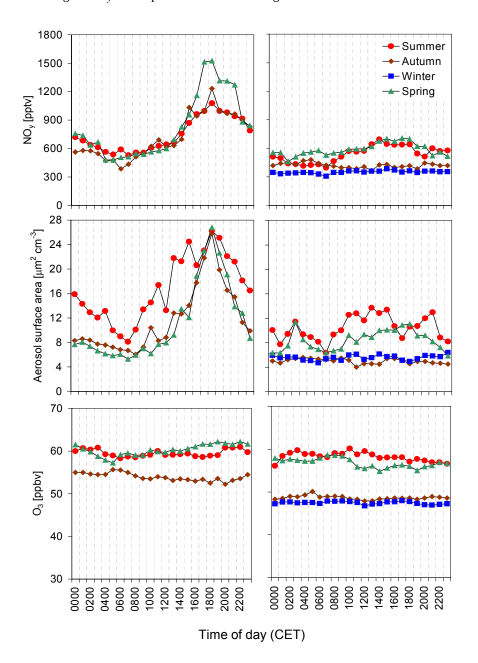
tween April 1997 and March 1999, 4.1% of the data fulfilled the criteria for south föhn. Although these events are less frequent than thermally induced vertical transport or synoptical lifting, they may significantly add to the upward transport of gaseous species over the Alpine region. Forrer et al. (1999) showed by trajectory analysis that air masses reaching the JFJ during south föhn episodes from the south and southeast seem to be frequently below the 900 hPa level and are therefore likely to be influenced by sources from this region. The influence of source regions during south föhn events can also



**Fig. 5b.** Frequency distributions including median, mean and number (*N*) of values for PAN, CO and the aerosol surface area concentration from April 1997 to March 1999 for the whole data set and different meteorological situations. For each species, the data set was based on hourly values and divided into 50 bins.

be seen from the present data set, where a median  $NO_x$  mixing ratio of 0.5 ppbv was observed during south föhn events, with 11% of the data being above 2.5 ppbv. This is significantly higher compared to undisturbed FT values with a median  $NO_x$  mixing ratio of 0.11 ppbv. The  $NO_x/NO_y$  ratio was also significantly higher during south föhn events compared to the undisturbed FT. In contrast, only a very low aerosol surface area concentration was observed, which can be explained by wet deposition.

Similar behavior was observed for north föhn events, which occurred with a frequency of  $\sim$ 5% between April 1997 and March 1999. Figures 5a and b show that the  $NO_x/NO_y$  ratio was comparable to south föhn events, in conjunction with low aerosol surface area concentrations. However, the mixing ratios of gaseous species ( $NO_x$ , PAN, total  $NO_y$ ) were lower compared to south föhn events. Explanations for this may be the different pollution levels north (Swiss plateau) and south (Po basin) of the Alps as well



**Fig. 6.** Diurnal variation of seasonal median values of gas and aerosol parameters from April 1997 to March 1999 at the Jungfraujoch Station. The four seasons are defined as summer, June to August, etc. The left panel shows days with thermally induced vertical transport, whereas undisturbed FT data is shown in the right panel. No winter data are shown for thermally induced vertical transport due to the small number of days with convection during winter.

as the orographic situation of the Alps. The Milan area in the Po basin was recognized to form a large emission source for ozone precursors by Prévôt et al. (1997). This is also in line with emission inventories made by the co-operative program for monitoring and evaluation of the long-range transmission of air pollutants in Europe (EMEP) (www.emep.int). In addition, ongoing research using satellite data (www.isac.cnr.it/%7Etrasfene/POLPO\_WEB/HomePage) also points to the Po valley area as a hot spot for NO<sub>2</sub>.

Synoptical lifting (e.g. frontal systems) is another important transport process for  $NO_y$  species, which may be enhanced by Alpine orography. It can be seen from Figs. 5a and b that air rich in  $NO_x$ ,  $NO_y$ , PAN and CO was transported to the JFJ during synoptical lifting. The frequency distributions of these compounds during synoptical lifting are comparable to north föhn events. In contrast to föhn events, the aerosol surface area concentration increased slightly. This may be explained by better vertical mixing and less precipitation during synoptical lifting.

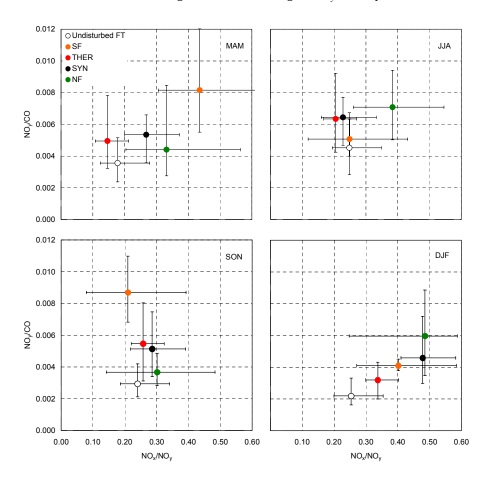


Fig. 7.  $NO_x/NO_y$  versus  $NO_y/CO$  ratios for south föhn (SF), north föhn (NF), thermally induced vertical transport (THER), synoptical lifting (SYN) and the undisturbed FT. Four seasons are shown (MAM = March to May, etc.). The first and third quartiles are indicated by bars.

Thermally induced vertical transport also leads to increased  $NO_y$  and aerosol surface area concentrations (see Figs. 5a and b). Thermally induced vertical transport has been recognized as an important mechanism of upward transport to the JFJ for aerosol particles (Baltensperger et al., 1997; Lugauer et al., 1998, 2000; Nyeki et al., 1998, 2000), as well as for gaseous species (Zellweger et al., 2000).

The identification of thermally induced vertical transport was based on the diurnal variation of NO<sub>v</sub>, the aerosol surface area concentration, and the specific humidity (see Sect. 2.3). This is in contrast to earlier studies (Lugauer et al., 1998, 2000) where thermally induced transport was determined according to the Alpine Weather Statistics (AWS) scheme (Schüepp, 1979; Wanner et al., 1998). Convective AWS days during spring and summer usually exhibit a pronounced diurnal cycle with afternoon peak concentrations for both aerosol and some gaseous parameters. However, a strong diurnal variation was also observed on several nonconvective days according to the AWS classification (e.g. 31 March 1998, AWS Advective type). This might be due to the fact that the AWS classification covers the whole Alpine region and does not account for local wind systems. Figure 6 illustrates diurnal variations of the median NO<sub>y</sub>, O<sub>3</sub>,

and aerosol surface area concentration over the four seasons at the JFJ for days with thermally induced vertical transport and for the undisturbed FT. The aerosol surface area concentration and NO<sub>v</sub> showed a distinct diurnal cycle with afternoon maximum concentrations at ~18:00 CET during days with thermally induced vertical transport. A detailed overview of diurnal cycles is given in Lugauer et al. (1998) and Nyeki et al. (1998) for aerosol parameters. The diurnal cycle of radon daughters was described recently (Lugauer et al., 2000). Ozone showed only a very weak diurnal variation for days with thermally induced vertical transport during spring, and no diurnal variation was observed for the rest of the year. This can be explained by small vertical concentration gradients in combination with dilution by FT air during convective transport. Figure 5a shows that the lowest NO<sub>x</sub>/NO<sub>v</sub> ratios of 0.22 on average were observed during thermally induced upward transport at the JFJ. This supports the conclusion that photochemically well-processed air masses reach the JFJ during spring and summer afternoons. Thus, the NO<sub>x</sub> mixing ratios during these periods were comparable to values of the undisturbed FT, whereas a significant increase of the NO<sub>v</sub> mixing ratios was observed.

### 3.2 Air mass aging

The fact that lower NO<sub>x</sub>/NO<sub>y</sub> ratios were observed during thermally induced upward transport compared to undisturbed FT conditions at the JFJ suggests that the NO<sub>x</sub>/NO<sub>y</sub> ratio is not always a suitable parameter to estimate the processing that has occurred in an air parcel. This is especially the case for remote locations, where deposition (e.g. wet and dry deposition of HNO<sub>3</sub>) and decomposition processes (e.g. thermal decay of PAN) can make the ratio difficult to interpret. The NO<sub>x</sub>/NO<sub>y</sub> ratio accounts for the photochemical processing that has occurred in an air parcel. The photochemical aging of an air parcel may be fast, with NO<sub>x</sub>/NO<sub>y</sub> ratios <0.4 within a few hours (Thornberry et al., 2001). Thus, this ratio may be used to assess the photochemical processing close to sources and on a time scale of less than one day. An alternative parameter to assess the aging process that has occurred in an air parcel is the NO<sub>v</sub>/CO ratio, which accounts for both deposition and dilution effects. The relationship between CO and NO<sub>v</sub> was previously used to assess anthropogenic input to air masses by Parrish et al. (1996). Close to anthropogenic sources, the  $NO_x/CO$  ratio averages  $\sim 0.1$ , whereas values of  $\sim 0.005$  are observed in the upper troposphere (Jaeglé et al., 1998). This is in line with recent findings of Stohl et al. (2002) where the original emission ratio between NO<sub>v</sub> and CO was assumed to be 0.16. This ratio dropped to values below 0.01 for air masses older than four days. However, it should be noted that the NO<sub>v</sub>/CO ratio also shows a seasonal variation, with lower values during the winter months due to a longer lifetime of CO. Figure 7 shows seasonal plots of the NO<sub>x</sub>/NO<sub>y</sub> versus NO<sub>y</sub>/CO ratios for different meteorological situations described in Sect. 2.3 and the undisturbed FT. It can be seen that undisturbed FT conditions are always accompanied by the lowest NO<sub>y</sub>/CO ratios, indicating advanced aging of these air masses. This ratio may therefore be an alternative method to distinguish between disturbed and undisturbed FT conditions at the JFJ.

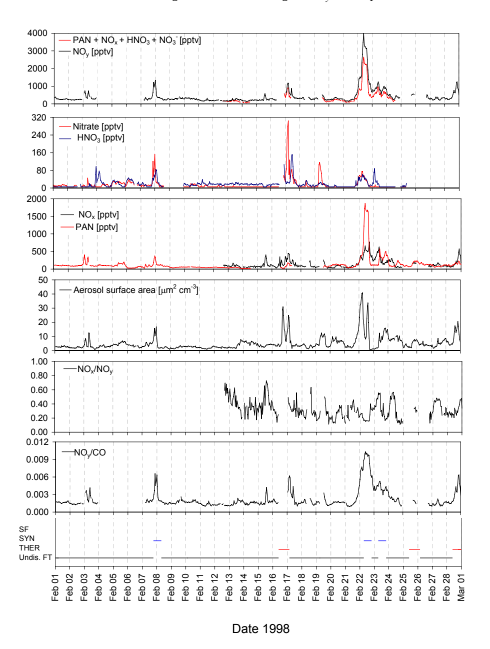
A further illustration of the effect of different meteorological situations on the NO<sub>x</sub>/NO<sub>y</sub> and NO<sub>y</sub>/CO ratios is shown in Figs. 8 and 9, where mixing ratios of individually measured NO<sub>v</sub> species as well as total NO<sub>v</sub>, the NO<sub>x</sub>/NO<sub>v</sub> and the NO<sub>v</sub>/CO ratios and the aerosol surface area concentration are presented for two selected periods in winter and spring 1998. The periods experiencing south föhn, synoptical lifting and thermally induced vertical transport are also shown in Figs. 8 and 9 (no situation with north föhn occurred during these periods). Again the NO<sub>v</sub>/CO ratio increased on average during the above meteorological events. Figure 8 shows mainly undisturbed FT conditions with the exception of a few short periods of thermally induced vertical transport and synoptical lifting. A period of particular interest is during thermally induced vertical transport on 30 March to 1 April (see red highlighted area in Fig. 9). On these days NO<sub>v</sub>/CO ratios exceeded 0.01 during the peak in afternoon concentrations, indicating a "pollution" event. In contrast, NO<sub>x</sub>/NO<sub>y</sub> ratios were very low (below 0.2), which indicates photo-chemically well-processed air masses. The peaks can also be observed in the aerosol surface area concentration, which typically occurs during thermally induced transport processes.

A second interesting period occurred between 3 April and 16 April when several south föhn events were experienced (see yellow highlighted area in Fig. 9). Again, all these events were accompanied by NO<sub>y</sub>/CO ratios exceeding 0.01. In contrast to thermally induced transport, NO<sub>x</sub>/NO<sub>y</sub> ratios also reached high values (approx. 0.7–0.9) during the föhn events. This can be attributed to the high NO<sub>x</sub> concentrations observed during these periods along with efficient wet deposition of HNO<sub>3</sub> and particulate nitrate. For the same argument particle concentrations were also relatively low during these föhn events.

# 3.3 NO<sub>y</sub> partitioning – case studies

Summertime NO<sub>y</sub> speciation at the JFJ has already been discussed by Zellweger et al. (2000). The most abundant NO<sub>y</sub> species was PAN with an average contribution of 36%, followed by NO<sub>x</sub> with 22%, particulate nitrate with 17% and HNO<sub>3</sub> with 7%. The sum of NO<sub>x</sub>, HNO<sub>3</sub>, particulate nitrate and PAN contributed on average 82% to the total NO<sub>y</sub> mixing ratio. However, it should be emphasized that an accurate determination of total NO<sub>y</sub> with individually measured species is difficult due to measurement uncertainties. Furthermore, the presence of reactive nitrogen compounds which are present in significant quantities but which are not measured in a given data set may also lead to an underestimation of total NO<sub>y</sub>.

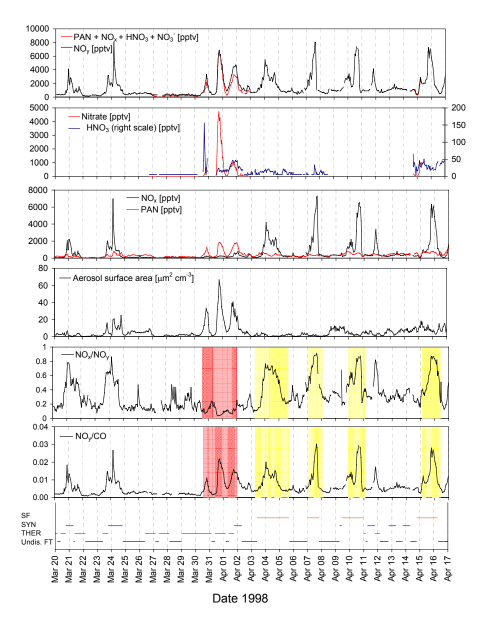
The NO<sub>v</sub> partitioning for selected periods is shown in Figs. 10a-c. During the undisturbed FT period from 12-13 February 1998 (Fig. 10a), the NO<sub>x</sub> fraction was, at 46%, the most abundant NO<sub>v</sub> species, followed by PAN (20%), HNO<sub>3</sub> (9%) and particulate nitrate (3%). An average fraction of 22% could not be identified, and total NO<sub>v</sub> averaged 209 pptv. This period can be considered as representative of clean conditions during the autumn and winter months, since total NO<sub>v</sub> remained almost constant throughout February 1998 with the exception of the periods when synoptical lifting and thermally induced vertical transport was observed. During periods influenced by synoptical lifting (22– 23 February 1998, 07:00–19:00, Fig. 10b), PAN was the most abundant  $NO_v$  species (38%), followed by  $NO_x$  (26%), HNO<sub>3</sub> (1%) and particulate nitrate (1%). The peak concentration of NO<sub>v</sub> was more than 10 times higher during this period compared to the "clean" period, and mainly PAN together with an unidentified fraction was transported to the JFJ. The very low contribution of inorganic nitrate to total NO<sub>v</sub> during synoptical lifting can be explained by wet deposition. The period between 27 March and 2 April 1998 (Fig. 10c) was characterized by clean conditions until the morning of 30 March. During this period, an average NO<sub>v</sub> mixing ratio of 350 pptv was observed with PAN (39%) as



**Fig. 8.** Time series of hourly mean values of  $NO_y$ ,  $HNO_3$ ,  $NO_3^-$ ,  $NO_x$ , PAN, aerosol surface area concentration, the  $NO_x/NO_y$  and the  $NO_y/CO$  ratios together with the sum of the individually measured  $NO_y$  species for February 1998. The occurrence of south föhn (SF), synoptical lifting (SYN), thermally induced vertical transport (THER), and undisturbed FT is also shown.

the most abundant NO<sub>y</sub> species, followed by NO<sub>x</sub> (24%) and inorganic nitrate (4%). Thermally induced vertical transport occurred on the three following days, where very high NO<sub>y</sub> mixing ratios were found. On the first day, a maximum NO<sub>y</sub> concentration of 3.3 ppbv was observed. During this peak event, NO<sub>y</sub> consisted mainly of PAN (42%), followed by particulate nitrate (13%), NO<sub>x</sub> (9%) and HNO<sub>3</sub> (3%). In the following night, the air was replaced by undisturbed FT-influenced air, and low concentrations were observed. A very high injection of NO<sub>y</sub> was then observed on 31 March with a

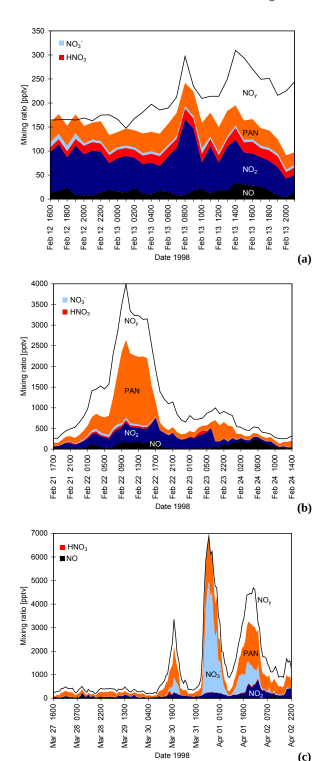
maximum concentration of 6.9 ppbv, which consisted mainly of particulate nitrate (52%), PAN (30%) and NO<sub>x</sub> (5%). HNO<sub>3</sub> was below the detection limit during this episode, indicating that the formation of particulate nitrate was favored due to neutralization of HNO<sub>3</sub>. The NO<sub>y</sub> rich air was again replaced through horizontal advection during the following night. On 1 April, another thermally induced injection was observed. The NO<sub>y</sub> mixing ratio reached 4.7 ppbv, and consisted mainly of PAN (34%), followed by particulate nitrate (20%), NO<sub>x</sub> (10%) and HNO<sub>3</sub> (1%). However, the following



**Fig. 9.** Time series of hourly mean values of  $NO_y$ ,  $HNO_3$ ,  $NO_3^-$ ,  $NO_x$ , PAN, aerosol surface area concentration, the  $NO_x/NO_y$  and the  $NO_y/CO$  ratios together with the sum of individually measured  $NO_y$  species from 20 March to 16 April 1998. The occurrence of south föhn (SF), synoptical lifting (SYN), thermally induced vertical transport (THER), and undisturbed FT is also shown. A period of particular thermally induced vertical transport is highlighted in red, and periods with south föhn events are highlighted in yellow.

advection of clean air was not effective enough to replace the injected air mass completely by FT air, leading to elevated NO<sub>y</sub> mixing ratios during the night of 2 April. This phenomenon was already observed during summer 1997 for convective days with low wind speed at the 500 hPa level (Zellweger et al., 2000). Figure 11 summarizes levels and speciation of NO<sub>y</sub> during selected periods. The NO<sub>y</sub> mixing ratio remained relatively constant during undisturbed FT periods. Highest NO<sub>y</sub> mixing ratios were observed during spring, but the seasonal difference between summer and winter was small. However, differences can be seen in the NO<sub>y</sub>

speciation. The  $NO_x/NO_y$  ratio was highest during winter (0.46) and lowest during spring (0.21). This reflects the lower photochemical activity during the winter months. By similar reasons, PAN/ $NO_y$  was highest during spring (0.39) and lowest during winter (0.20). Meteorological processes also have a strong influence on the level and speciation of  $NO_y$ . The  $NO_y$  speciation for selected periods (thermally induced vertical transport, föhn, frontal systems) is also shown in Fig. 11. The  $NO_y$  mixing ratio reached an average level of 1 to 4 ppbv, a factor 4 to 10 higher compared to undisturbed FT conditions. The  $NO_x/NO_y$  ratio was lowest during episodes with



**Fig. 10.** (a)Average hourly mixing ratios of speciated and total NO<sub>y</sub> at the Jungfraujoch from 12 to 13 February 1998. (b) Average hourly mixing ratios of speciated and total NO<sub>y</sub> at the Jungfraujoch from 21 to 24 February 1998. (c) Average hourly mixing ratios of speciated and total NO<sub>y</sub> at the Jungfraujoch from 27 March to 2 April 1998.

thermally induced vertical transport, accompanied by high  $NO_3^-/NO_y$  and  $PAN/NO_y$  ratios. This again reflects the photochemical transformation of  $NO_x$  during thermally induced upward transport, yielding a photochemically aged air upon arrival at the Jungfraujoch. The low  $NO_x/NO_y$  and the high  $PAN/NO_y$  ratios also suggest that PAN degradation is not a major source of  $NO_x$  at the JFJ. Rather, the Alpine region appears to be a passive transport pathway of PAN (and other  $NO_y$  compounds), which is in line with recent results from FREETEX'98 (Carpenter et al., 2000).

#### 4 Conclusions

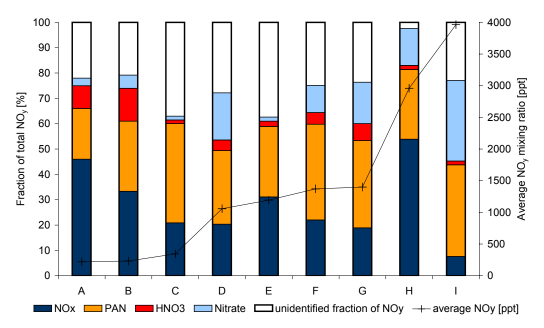
Speciated and total NO<sub>y</sub> were measured at the high-alpine site Jungfraujoch (JFJ) over a two-year period. The NO<sub>y</sub> mixing ratios and partitioning strongly depended on meteorological conditions. As a consequence, filters must be used at Alpine (and presumably other continental) sites to distinguish between undisturbed and disturbed FT conditions. Attempts to develop a filter procedure for the combined European GAW virtual baseline station, comprising the Jungfraujoch (Switzerland), the Sonnblick (Austria) and the Zugspitze (Germany) high-alpine stations, have not been successful so far (Fricke et al., 2000).

This study presents filters based mainly on meteorological parameters for the Jungfraujoch station. The meteorological processes that were identified to influence NO<sub>v</sub> mixing ratios and composition include föhn (both from north and south), synoptical lifting (e.g. fronts) and thermally induced vertical transport. Highest NO<sub>v</sub> mixing ratios were observed during south föhn events. This is in contrast to aerosol parameters, for which thermally induced vertical transport is the major process for upward transport. Seasonal variation in the composition of NO<sub>v</sub> was found, with PAN being dominant during spring and summer, and NO<sub>x</sub> during autumn and winter. The NO<sub>x</sub>/NO<sub>y</sub> ratio reached the lowest values during thermally induced upward transport. As a consequence, the NO<sub>x</sub>/NO<sub>v</sub> ratio is not a suitable parameter to estimate the air mass aging at remote sites. Alternatively, the NO<sub>v</sub>/CO ratio proved to be an interesting approach in assessing the age of an air mass.

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- A: winter, undisturbed FT, 12-13 Feb 1998
- B: summer, undisturbed FT, 27-30 Jul 1997, 0300 0900
- C: spring, undisturbed FT, 27-30 Mar 1998
- D: summer, influenced by thermally induced vertical transport, 27-30 Jul 1997, 1500-2100
- E: winter, influenced by synoptical lifting, 22-24 Feb 1998
- F: summer, influenced by synoptical lifting, 25 Aug 1997, 1900 to 26 Aug 1997, 0700
- G: summer, 16-22 Aug 1997; convective days with low wind speed at the 500 hPa level [Zellweger et al., 2000]
- H: spring, south foehn, 14 Apr 1998, 1400 to 16 Apr 1998, 1600
- I: spring, influenced by thermally induced vertical transport, 30 Mar 2 Apr 1998, 1500-2100

**Fig. 11.** NO<sub>y</sub> levels and speciation for selected periods.

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