Formic acid above the Jungfraujoch during 1985–2007: observed variability, seasonality, but no long-term background evolution

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Abstract

This paper reports on daytime total vertical column abundances of formic acid (HCOOH) above the Northern mid-latitude, high altitude Jungfraujoch station (Switzerland; 46.5° N, 8.0° E, 3580 m altitude). The columns were derived from the analysis of infrared solar observations regularly performed with high spectral resolution Fourier transform spectrometers during over 1537 days between September 1985 and September 2007. The investigation was based on the spectrometric fitting of five spectral intervals, one encompassing the HCOOH ν_6 band Q branch at 1105 cm^{-1}, and four additional ones allowing to optimally account for critical temperature-sensitive or timely changing interferences by other atmospheric gases, in particular HDO, CCl_2F_2 and CHClF_2. The main results derived from the 22 yr long database indicate that the free tropospheric burden of HCOOH above the Jungfraujoch undergoes important short-term daytime variability, diurnal and seasonal modulations, inter-annual anomalies, but no statistically significant long-term background change at the 1-sigma level.

A major progress in the remote determination of the atmospheric HCOOH columns reported here has resulted from the adoption of new, improved absolute spectral line intensities for the infrared ν_6 band of trans-formic acid, resulting in retrieved free tropospheric loadings being about a factor two smaller than if derived with previous spectroscopic parameters. Implications of this significant change with regard to earlier remote measurements of atmospheric formic acid and comparison with relevant Northern mid-latitude in situ findings will be assessed critically. Sparse HCOOH model predictions will also be evoked.

1 Introduction

Numerous field investigations of volatile organic compounds in the boundary layer and through the free troposphere have been undertaken in situ and remotely during isolated, time-limited local as well as regional studies since the mid-1980s (e.g., local
ground-based background monitoring; measurements during natural and controlled savannah fires; related Atlantic and Pacific ship- and airborne campaigns; stratospheric balloon flights- and space-based solar observations). They indicate that HCOOH is, after acetic acid (CH$_3$COOH), the most abundant global organic carboxylic acid present in the Earth’s atmosphere, in both gaseous and liquid phases (e.g., Talbot et al., 1988, 1990; Puxbaum et al., 1988; Andreae et al., 1988; Arlander et al., 1990; Klemm et al., 1994; Sanhueza et al., 1996; Kesselmeier and Staudt, 1999; Khare et al., 1999; Poisson et al., 2000; Legrand et al., 2003, 2004). These studies further suggest that the main HCOOH sources are of biogenic origin (e.g., direct emissions and indirect production via oxidation and ozonolysis of reactive hydrocarbons released by soils, vegetation, forests, seas, biomass burning, ants) and anthropogenic releases (via wild bush and waste fires, domestic heating, road traffic, . . . ). The dominant atmospheric HCOOH removal occurs through dry and wet deposition, thus contributing along with sulfuric, nitric, acetic and hydrochloric acids to the acidity in the boundary layer, out to far remote areas. Measurement techniques used in the above researches, as well as formic acid photochemistry modeling calculations have been evoked, e.g., by Keene et al. (1989), Pszenny et al. (1989), Reiner et al. (1999), Poisson et al. (2000), Baboukas et al. (2000), von Kuhlmann et al. (2003), Christian et al. (2004), Rinsland et al. (2004, 2006), Remedios et al. (2007). Despite original findings acquired along these field investigations and related chemical studies in the laboratory, there remain numerous gaps in the observational datasets, preventing regional and global background emissions to be satisfactorily quantified and understood, thus difficult to be adequately modeled.

Observed short-term variability of atmospheric HCOOH during many field campaigns has led to estimates of its atmospheric lifetime, ranging from hours in the boundary layer to days in the upper troposphere (e.g., Keene and Galloway, 1988; Hartmann et al., 1991; Hahn et al., 1992; Chebbi and Carlier, 1996). Causes for this variability include heterogeneity in the distribution and release intensities of the sources – both biogenic and anthropogenic – at the ground. The latter depend on local/regional vege-
tation types, orography, and diurnally and seasonally varying environmental conditions such as solar insolation, atmospheric circulation-, temperature- and moisture fields, which also affect dry and wet deposition. Despite time-limited field campaigns, evidence for diurnal as well as seasonal variations also emerged (e.g., Arlander et al., 1990; Talbot et al., 1988, 1995; Khwaja 1995; Kajii et al., 1997; Baboukas et al., 2000; Legrand et al., 2004). The combined short lifetime and the location of the main sources at the ground preclude a HCOOH distribution versus altitude with a small scale-height through the free troposphere.

Within this introductory context, two relevant reviews ought to be highlighted here, namely (i) a comprehensive overview of the fate of biogenic volatile organic compounds in the atmosphere (including HCOOH) by Kesselmeier and Staudt (1999), and (ii) a more specific overview by Khare et al. (1999) on the state of measurements of atmospheric formic and acetic acids in the boundary layer and likely interpretations.

In this paper, we provide further evidence of short-term daytime and day-to-day variability, and evaluate diurnal and seasonal modulations, inter-annual variations, as well as the long-term trend of free tropospheric HCOOH above the high-altitude International Scientific Station of the Jungfraujoch (Switzerland; 46.5° N, 8.0° E, 3580 m altitude). The findings are based on the spectrometric analysis of solar spectra regularly recorded since the mid-1980s with modern Fourier transform infra-red (FTIR) spectrometers. As in past infrared remote sensing investigations (e.g., Goldman et al., 1984; Shephard et al., 2003; Rinsland et al., 2004, 2006, 2007; Remedios et al., 2007; González Abad et al., 2009), the present analysis focuses on the \( \nu_6 \) band Q branch of HCOOH near 1105 cm\(^{-1}\), which is by far the most appropriate absorption feature for studying its atmospheric concentration through ground-, airborne- and space-based observations. Here, however, we adopt a new, improved set of spectroscopic parameters for this HCOOH \( \nu_6 \) band, recently released by Perrin and Vander Auwera (2007), and now implemented in the HITRAN-2008 line parameter compilation (Rothman et al., 2009; see also http://www.hitran.com). This new set returns formic acid loadings being about a factor two lower than if derived with parameters found in earlier HITRAN
versions (i.e., Rothman et al., 2003, 2005).

2 Observational database and ancillary inputs

The database analyzed here is a subset of all solar observations made between 2 and 15 µm (about 5000 to 650 cm\(^{-1}\)) with two Fourier transform spectrometers (FTS) operated at the Jungfraujoch under typical spectral resolutions of 0.003 to 0.006 cm\(^{-1}\) (corresponding to maximum optical path differences of 165 and 82 cm, respectively), allowing to study quasi-simultaneously over two dozen atmospheric gases (Zander et al., 2005, 2008). The retained subset contains 8256 individual spectra recorded during 1537 days between September 1985 and September 2007 with an optical filter specifically covering the 700 to 1400 cm\(^{-1}\) spectral domain, thus encompassing the HCOOH \(\nu_6\) band Q branch at 1105 cm\(^{-1}\). These spectra were all acquired at solar zenith angles (SZA) < 87 degrees, a geometry mitigating target- versus interfering gas absorptions in near-horizontal observations. Notice that the mountainous orography around the Jungfraujoch station prevents this SZA limit to be reached during some periods of the year, either at sunrise or at sunset. Until March 1995, the observations were predominantly made with a home-built FTS (Delbouille and Roland, 1995), while a faster commercial Bruker-Type120 HR FTS was used more consistently onwards. The signal-to-noise (S/N) ratio varied from about 300 (essentially earlier single recordings) to over 2500 (corresponding to “average” spectra, resulting from individual Bruker scans obtained during time intervals of 6 to 30 min, depending on the solar zenith angle rate of change). Near-coincident HCOOH measurements with both instruments were specifically inter-compared to assess the consistency of the data retrieved with either instrument.

The model atmosphere adopted above the 3.58 km Jungfraujoch site to compute synthetic spectra for comparison with the observed ones (see Sect. 4) consisted of 41 layers: the two lower ones were successively 0.32 and 0.70 km thick, followed higher up by 24 layers equally spaced by 1.2 km, and additional ones progressively increasing
in thickness to reach 100 km altitude. The mean physical pressure-temperature characteristics of these layers were derived from midday pressure-temperature profiles, specifically calculated for the location of the Jungfraujoch by the US-National Centers for Environmental Prediction (NCEP, Washington, DC; see http://www.ncep.noaa.gov).

The “a priori” volume mixing ratio (VMR) profile adopted for retrieving the HCOOH target gas decreases quasi-monotonically through the free troposphere above the site (i.e., 440, 300, 190, 100 pptv (parts per trillion or \(10^{-12}\) by volume), respectively at 3.58, 6.0, 8.5, and 11 km altitude, and set to zero at 17 km and above). The slope of this a priori profile is commensurate with the altitude-dependent characteristics derived by Reiner et al. (1999) from in situ measurements of HCOOH between 7 km (215 pptv) and 11.3 km (59 pptv) altitude made in the fall of 1991 during five airplane flights above Southern Germany, only a few latitudinal degrees north of the Jungfraujoch. Supporting HCOOH concentration decreases versus altitude at Northern mid-latitudes have also been reported by Talbot et al. (1996a,b, 1997a,b) and by Singh et al. (2000) during airborne missions, respectively over the Northern Pacific and Atlantic oceans. The adopted slope is further in line with vertical transport investigations based on tropospheric measurements of \(^{222}\)Radon (lifetime of 5.5 days) distributions at Northern mid-latitudes (e.g., Nazarov et al., 1970; Liu et al., 1984). Validation of the adopted HCOOH profile in the upper troposphere and up through the lower stratosphere was assessed by one of us (P. Du) and more recently confirmed by González Abad et al. (2009), based on solar occultation measurements at Northern mid-latitudes with the ACE (Atmospheric Chemistry Experiment) FTS aboard the Canadian SciSat satellite (Bernath et al., 2005). A priori VMR profiles for major interfering gases were compiled from reliable sets of space- (i.e., ATMOS (Atmospheric Trace Molecule Spectroscopy; Irion et al., 2002); HALOE (Halogen Experiment; http://haloe.gats-inc.com); ACE) and stratospheric and stratospheric balloon-based in situ measurements at Northern mid-latitudes (i.e., A. Engel, Goethe University of Frankfurt, private communication, 2004).
3 New HCOOH $\nu_6$ band intensity

The first tentative identification of formic acid in the upper troposphere was reported by Goldman et al. (1984), based on solar spectra recorded from aboard a stratospheric balloon platform above Arizona (USA). They used HCOOH $\nu_6$ band parameters derived by Goldman and Gillis (1984) from the analysis of a laboratory spectrum whose total HCOOH line intensities over the 1000–1200 cm$^{-1}$ interval corresponded to $19.96 \times 10^{-18}$ cm$^{-1}$(mol cm$^{-2}$) at 296 K. Assuming this band intensity, Perrin et al. (1999) undertook a theoretical Hamiltonian analysis of the fundamental $\nu_6$ and the interacting $\nu_8$ bands of HCOOH. They generated a detailed set of line positions and intensities, which significantly improved the quality of fits to observed HCOOH spectra. This set of lines was incorporated in the HITRAN-2004 compilation (Rothman et al., 2005).

Meanwhile, Vander Auwera et al. (2004) raised a concern, namely the fact that several independent spectroscopic studies, – both laboratory as well as theoretical –, had reported integrated intensities of the HCOOH $\nu_6$ band that were higher than the Goldman and Gillis (1984) value by about a factor of 2 (see Table V in Vander Auwera et al., 2007). Using new HCOOH laboratory spectra recorded between 600 and 1900 cm$^{-1}$ with a high resolution (0.002 cm$^{-1}$) Fourier transform spectrometer, at various total sample pressures ranging from 14.3 to 314 Pa, they measured the absolute intensity of 72 lines in the $\nu_6$ band, with an absolute uncertainty estimated to be at most 7%, and performed a detailed Hamiltonian analysis of $\nu_6/\nu_8$ interacting bands. Specifics of the approach – both experimental and analytical –, as well as emerging issues can be found in Vander Auwera et al. (2007). In particular, an integrated HCOOH band intensity over the 1045–1150 cm$^{-1}$ interval was found equal to $38.8 \times 10^{-18}$ cm$^{-1}$(mol cm$^{-2}$) at 296 K, thus nearly twice the value reported by Goldman and Gillis (1984). Perrin and Vander Auwera (2007) synthesized the original improvements of this research with respect to earlier ones, pointing out that the new band intensity refers to the trans-form of HCOOH, as natural formic acid only contains about
0.1% of the HCOOH cis-form (Hocking, 1976). They further compiled an improved line list for fitting HCOOH in the 9-μm region, containing individual positions and intensities of almost 50000 lines between 940.20 and 1244.41 cm$^{-1}$. All air-broadening line half-widths were set to 0.101 cm$^{-1}$ atm$^{-1}$ at 296 K (a value assessed experimentally by Notholt et al., 1991), with a temperature dependence exponent $n=0.75$, and a self-broadening parameter $\gamma=0.32$ cm$^{-1}$ atm$^{-1}$ at 296 K derived in Vander Auwera et al. (2007). This new HCOOH database, merged with the line parameters of the HITRAN-2004 compilation for all other atmospheric molecules, has been used here prior to publication. At the time of this writing it has been formally incorporated in the most recent HITRAN-2008 compilation (Rothman et al., 2009), thus available to the scientific community. The factor 2 needs to be kept in mind when relating the HCOOH results reported here with earlier infrared remote sensing findings, e.g., by Goldman et al. (1984), Shephard et al. (2003), Rinsland et al. (2004, 2006, 2007), Remedios et al. (2007), some of which will be discussed in Sect. 6 and in Appendix A. Incidentally, this factor 2 also explains the discrepancy observed by Christian et al. (2004) when inter-comparing HCOOH emissions of biomass-burning material conducted in the laboratory, simultaneously with a proton transfer reaction-mass spectrometer (PTR-MS) and an open path-Fourier transform infrared (OP-FTIR) instrument. The OP-FTIR findings were consistently about twice larger than those derived with the PTR-MS technique; with the new parameters, the results would have been comparable.

4 Spectrometric analysis approach

The analysis of the 8256 solar spectra retained here was performed with the SFIT2 retrieval algorithm, using the spectroscopic parameters compiled by Perrin and Vander Auwera (2007) evoked in Sect. 3. The main features of the SFIT2 code have been described by Pougatchev et al. (1995) and Rinsland et al. (1998). It incorporates a semi-empirical implementation approach of the optimal estimation formalism developed by Rodgers (1976), as well as an error evaluation discussed by Connor et al. (1995). The
SFIT2 Version 3.91 used here specifically allows to simultaneously (i) derive information on the VMR variation versus altitude of up to 3 target gases, (ii) uniformly scale the a priori profiles of up to 8 interfering gases, and (iii) further deal with the absorption characteristics of up to 50 other ones according to prescribed, most realistic VMR profiles.

In order to properly account for a critical interference by a temperature-sensitive HDO line at 1104.8759 cm$^{-1}$ ($E''=701.6$ cm$^{-1}$; a concern already stressed by Perrin et al., 1999), as well as time evolving continuum-type absorptions by CCl$_2$F$_2$ and CHClF$_2$ (e.g., Meier et al., 2004), a variant of the simultaneous multiple-windows approach used by Rinsland et al. (2004) was adopted here. In a first run, four pre-selected micro-windows were fitted together in each retained spectrum, namely two for HDO (1206.18 to 1206.67 cm$^{-1}$ and 1208.58 to 1208.98 cm$^{-1}$; lines with $E''=683.5$ cm$^{-1}$, thus very close to the temperature dependence of the interfering HDO line), one for CCl$_2$F$_2$ (922.50 to 923.60 cm$^{-1}$) and one for CHClF$_2$ (828.85 to 829.25 cm$^{-1}$) by scaling their a priori VMR profiles. These spectrum-specific retrieved profiles were then used as such in a second fitting run encompassing the 1104.65 to 1105.60 cm$^{-1}$ interval, in which we retrieved the distribution of the 2 most important interfering O$_3$ isotopomers ($^{16}$O$_3$ and $^{16}$O$^{16}$O$^{18}$O) and scaled the a priori VMR profile of HCOOH characterized in Sect. 2. Additional discrete absorptions by gases such as H$_2$O, CH$_4$, CO$_2$ and N$_2$O were also accounted for in both runs by scaling assigned, most realistic VMR profiles. A signal-to-noise ratio of 500 was set in all fitting sequences, commensurate with the noise level of most analyzed spectra. Inspection of extra-atmospheric solar observations by the shuttle-based ATMOS-FTIR instrument (Farmer and Norton, 1989) showed that no solar lines perturb any of the 5 spectral intervals used here.

Frame B of Fig. 1 reproduces a fit to a typical spectrum observed at the Jungfraujoch on 8 June 2005, with residuals (Frame C) resulting from the combined absorption characteristics of the 6 most important gases identified in Frame A, as well as contributions from second order ones listed in the previous paragraph. Individual inspection of the fittings to all spectra analyzed here, led to rejecting 616 of them, based on objec-
positive criteria such as local fringing, no fitting convergence, excessive root-mean square fitting residuals with respect to the spectral S/N ratio, and those HCOOH columns in the \(10^{13}\) mol/cm\(^2\) range which were affected by fitting uncertainties exceeding the retrieved columns. The database discussed hereafter thus includes 7640 individual HCOOH total column measurements encompassing 1501 days. They were normalized to the mean local pressure observed at the Jungfraujoch during the past two decades (i.e. 654 mb) versus the specific daily NCEP-reported pressures.

Key observational and analytical sources of uncertainty are reported in Table 1, showing that the largest errors are linked to the adopted a priori HCOOH and HDO profiles. For both species, the uncertainties have been split into a random and a systematic component, the former relating more specifically to short-term, altitude-dependent variability, and the latter to an overall uncertainty, e.g., the mean slope of the profiles. This distinction is worth being made, considering that the root square sum of the total random errors refers to individual HCOOH retrievals, thus reducing by the square root of the number of measurements intervening in averaged spectra as well as in daily and monthly mean column loadings. The reported uncertainties result from evaluations or simulations according to specifics given in the last column of Table 1. Overall, they are commensurate with those reported in a similar research by Rinsland et al. (2004) based on Kitt Peak (AZ, USA) solar observations evoked in Appendix A. One exception however, namely the new HCOOH spectroscopic \(\nu_6\) band intensity adopted here being only affected by a systematic uncertainty of less than 7% (Vander Auwera and Perrin, 2007), as compared to 12% for the older band intensity used by Rinsland et al. (2004), which is further affected by the systematic factor 2 discussed in Sect. 3.

5 Results

The retained individual pressure-normalized HCOOH column abundances are reproduced in Fig. 2a, while b shows the daily arithmetic means and related one-sigma standard deviations resulting from the individual values involved, on average 5 per day,
with extremes ranging from 1 to 56. This wide range of day-to-day HCOOH observations results from the fact that (1) two dozen atmospheric gases are being monitored at the Jungfraujoch using alternatively one among 4 wide band-pass optical filters encompassing altogether the spectral interval 2 to 15 µm, and (2) on many days, solar observations are restricted to periods of breaks in partly cloudy skies. Both frames of Fig. 2 reveal striking features, namely frequent sub-daily and day-to-day variability, as well as a clear seasonal modulation which is definitely characterized by low HCOOH columns occurring predominantly during the months of November-December-January (abbreviated by N-D-J; similar abbreviations will be adopted hereafter for other month groupings). All N-D-J daily mean columns are identified in Fig. 2b by the open circles with central crosses. After discarding some of these “crossed” data points, namely those from November 1991 to January 1995 (slightly biased high due to degradation of the home-made FTIR instrumental line shape, which affects the correct fitting of the strong interfering ozone lines; see Fig. 1), as well as a few abnormally high outliers (i.e., 21 January 1996, 16 November 2006 and 14 January 2007), a linear adjustment to the remaining 269 N-D-J daily mean columns returns a best fit HCOOH mean daytime “background” value (continuous line in Fig. 2b) and a 1-σ standard deviation (dashed lines) of \((0.38\pm0.20)\times10^{15} \text{ molec./cm}^2\). This value also applies to Fig. 2a when considering the individual measurements made during the same retained days. The present database therefore suggests that the N-D-J mean HCOOH daytime background loading above the Jungfraujoch has not changed statistically during the past two decades.

The short-term, sub-daily HCOOH variability can be appreciated by the magnitude of the 1-σ standard deviation bars in Fig. 2b, with extreme individual values often diverging by over ±50% around the means. Day-to-day variations of similar magnitude have also been observed frequently, as confirmed by the extent of the standard deviation bars associated to many monthly mean HCOOH columns displayed on a one-year time base in Fig. 3. A few obvious monthly mean outliers are identified by triangular symbols, namely from left to right for January 07, June 93, August 95, August 94, and September 1989. The thick curve in Fig. 3 corresponds to a running mean fit to the
black filled symbols only, with a 15-day step and a 2-month wide integration time (each step includes between 27 and 43 monthly mean data points). This fit reveals a clear yearly modulation of the mean daytime HCOOH loading, with a broad maximum during June-July-August (J-J-A) averaging $1.70 \pm 0.50 \times 10^{15}$ molec./cm$^2$, and a minimum of $0.42 \pm 0.18 \times 10^{15}$ molec./cm$^2$ (thin continuous line) for all N-D-J data, which is commensurate with the more selective value derived in Fig. 2. The shaded area visualizes the 1-$\sigma$ standard deviation associated to the running mean curve and reflects primarily the inter-annual variability noticeable in Fig. 2. If, as suggested by Talbot et al. (1997a), we consider as “enhancement” those HCOOH columns exceeding the N-D-J background level plus twice its standard deviation (thin dashed line in Fig. 3), then the observed columns from about mid-April to the end of August indicate a statistically significant seasonal modulation at the one sigma level.

To allow some comparison of our total HCOOH column measurements with other findings frequently reported as local mole fractions, we have reproduced on the right side of Fig. 3 a scale that allows converting an observed total HCOOH column abundance read off the left side scale of Fig. 3 into a mean VMR concentration (expressed in pptv) over the 3.58 to 10.6 km altitude range. It was determined by ratioing the total number of HCOOH molecules derived over that altitude interval against the total air molecules over the same range. The upper 10.6 km limit of the range corresponds to the top height of the 7th layer of the adopted model atmosphere (see Sect. 2), which is nearest to the mean tropopause height ($\sim$11.3 km) above the Jungfraujoch during the past two decades. Notice that the conversion from HCOOH columns to mean VMRs carries the systematic uncertainties reported in Table 1.

Table 2 lists the arithmetic monthly mean HCOOH daytime columns and corresponding mean VMRs, as well as their relative uncertainties and number of years involved in each monthly mean, derived from the filled data points of Fig. 3. They are commensurate with the 2-month wide running means reproduced by the thick curve in that figure. At the one-sigma level, they can be considered as representative of the month-specific climatology of the total columns and mean free tropospheric VMRs of HCOOH above
the Jungfraujoch from mid-1987 to mid-2007, returning a maximum in July and a minimum in December. The mean annual loading for that period is provided in the last row of Table 2.

Attention has also concerned the HCOOH diurnal variation above the Jungfraujoch. This is illustrated in Fig. 4, which displays all the individual vertical columns versus solar zenith angle derived from observations made with the Bruker instrument between 1995 and 2007, i.e., all open black circles of Fig. 2a (a total of 5158 data points gathered during 1219 days). Only the Bruker data were considered here, owing to their overall improved temporal regularity. Because of the large short-term variability superimposed on seasonal and inter-annual modulations, a vertical logarithmic scale was adopted to better appraise lower column loadings. While there is indication of an overall qualitative sunrise to sunset total HCOOH column modulation peaking around mid-day, a more quantitative assessment requires evaluating specific data subsets, as solar culmination occurs at heights varying over the year. This is highlighted in Fig. 4, where the filled cyan circles correspond to all N-D-J data points, with solar culmination occurring at about 62° zenith angle. First order fittings, separately to the AM and PM cyan points are featured by the continuous yellow lines. Extrapolated to 90° SZA to the left for “sunrise” and to the right for “sunset”, they show that the mean N-D-J loading of HCOOH above the Jungfraujoch increases by a factor 2, from $0.26 \times 10^{15}$ molec./cm$^2$ at sunrise to $0.51 \times 10^{15}$ molec./cm$^2$ at culmination, then decreases towards $0.32 \times 10^{15}$ molec./cm$^2$ at sunset. The black lines correspond to similar AM and PM fittings to all J-J-A data points, while the red lines refer to all May data points only (featured for a comparison with Mauna Loa observations discussed in Appendix A). Specific AM increases and PM decreases regarding these subsets, along with other first-order, bi-month evaluations, are summarized in Table 3.
6 Discussion

All observational field campaigns referred to in this paper consistently report large short-term variability of atmospheric HCOOH either in ground-, sea- or airborne based in situ air samplings, as well as in slant column remote-sensing spectrometric observations from the ground or from space. Likely causes include regional sources diversity and strength variations as well as specific meteorological conditions affecting local emissions and dry/wet deposition evoked in the introductory section. In this context, Fig. 2 shows that the Jungfraujoch HCOOH database makes no exception, despite the fact that, with respect to local in situ concentration measurements, short-term variability is further smoothed in spectrometric remote sensing observations, as they integrate information over long optical slant paths through the atmosphere.

The difficulty of identifying prevailing processes responsible for observed variability of short lifetime trace constituents (gases as well as aerosols) in a complex mountainous terrain such as the surrounding Jungfraujoch area has been documented and discussed by Baltensperger et al. (1997) and Forrer et al. (2000). These authors stressed the key role of local thermal convection during spring and summer, resulting in upslope winds that carry aerosols and gases from boundary layer valleys and catchment slopes to the Jungfraujoch site and higher up. This enhanced thermal effect, combined with the progressive agricultural soil treatment activity and natural biogenic vegetation metabolisms (e.g., Kesselmeier and Staudt, 1999) is in line with the observed regular increase of the free tropospheric HCOOH loading during March to June (see Fig. 3). The S-O-N decrease is clearly associated with the progressive slowing of these processes. Overall, the observed seasonal modulation of the HCOOH loading is also in phase with the annual solar insolation cycle and with the moisture content in the atmosphere, but off-phase with the observed modulation of CO$_2$ whose minimum is an indicator of high vegetation photosynthesis activity during springtime. While special attention has been given to correct for the spectral interference by the HDO line at 1104.8759 cm$^{-1}$ (see Sect. 4), the moisture content above the Jungfraujoch (showing
a minimum during J-F-M and maximum in J-A) is likely to affect the variability of formic acid which is highly soluble, but no clear correlation has been established statistically between them. It is also worth noting that the annual variation of the mean HCOOH loading above the Jungfraujoch is in excellent phase with the local aerosol concentration reported by Baltensperger et al. (1997) for the period 1988 to 1996.

In Table 4 we compare findings performed here with results from other field investigations identified in the literature, with emphasis on free tropospheric HCOOH measurements at Northern mid-latitudes. Details on these comparisons and related comments are provided in Appendix A.

The large variability in the tropospheric HCOOH loading is ubiquitously confirmed by the standard deviations associated with most Northern mid-latitude field measurements referred to in Table 4, independently of their observational approach, their temporal extent and geographic location, and the specificity of the surrounding milieu (e.g., marine versus continental; boundary layer versus free troposphere). Considering the magnitudes of these standard deviations, it is encouraging to notice that there is commensuration, within a factor 2, among most mean values reported in Table 4, including timely relevant Jungfraujoch findings provided in bold (between parentheses in column 3).

Few, isolated comparisons between HCOOH observations and model calculations have shown that the latter generally predict HCOOH concentrations significantly lower than measured. For example, Baboukas et al. (2000) report a factor 9 difference between boundary layer observations during a 63° N–39° S Atlantic ship cruise and the 3-D global tropospheric chemistry-transport MOGUNTIA model (e.g., Kanakidou and Crutzen, 1999; Poisson et al., 2000). Similarly, mean free tropospheric concentrations of HCOOH derived from infrared solar spectra observed at Kitt Peak Observatory (2.09 km a.s.l., Southern Arizona, USA) by Rinsland et al. (2004) were an order of magnitude higher than related calculations with the MATCH-MPIC 3-D model (Von Kuhlmann et al., 2003); this latter difference remains large even after halving it as a consequence of the new HCOOH spectroscopic parameters produced by Perrin and Van-
der Auwera (2007). According to Von Kuhlmann et al. (2003), the most likely reason for the large differences between HCOOH field observations and model calculations is that the latter only consider HCOOH production from the first order ozonolysis of alkenes (mainly isoprene) totaling 17 Tg/yr, and an additional 14 Tg/yr from biomass burning. Even more recent model runs incorporating some direct emissions of carboxylic acids resulting from biomass burning (i.e., Ito et al., 2007) have not yet approached reasonable commensuration with field investigations through the troposphere. The modelers are aware of other potential HCOOH sources, e.g., direct emissions from forests and land vegetation, and contributions from anthropogenic releases by domestic heating and road traffic, but their related optimal model implementation demands more reliable evaluations of season-related emission rates versus environmental specifics of regional areas.

7 Summary and conclusions

The spectrometric analysis of 8256 solar spectra recorded regularly at the Jungfraujoch during 1537 clear-sky days between September 1985 to September 2007 with very high spectral resolution Fourier transform infrared spectrometers has allowed to characterize various aspects of the free tropospheric loading of formic acid (HCOOH) above the mountainous Swiss Alpine plateau. The investigation used the new, more accurate set of spectroscopic parameters for the HCOOH ν_6 band Q branch at 1105 cm^{-1} (Perrin and Vander Auwera, 2007), which resulted in retrieved formic acid total column abundances nearly a factor 2 smaller (1.97±0.08 from a representative test performed here) and definitely more accurate than if retrieved with the older parameters (see Sect. 3). Adoption of a two-steps retrieval approach involving fittings of five micro-windows helped to optimally account for all major temperature-sensitive interferences such as HD_2O and O_3, as well as continuum-type, long-term evolving absorptions by CCl_2F_2 and CHClF_2 (see Sect. 4). The main sources of uncertainty, both random and systematic, have been discussed at the end of Sect. 4, and quantified in Table 1, with...
the total random and systematic root square sums (RSS), respectively evaluated at 13 and 18% maximum.

Overall, the individual HCOOH column abundances of the 22-yr long database (Fig. 2a) compacted into daily- and monthly means (respectively Figs. 2b and 3) have unambiguously confirmed the occurrence of significant sub-day, diurnal-, and day-to-day variability all along the reported period. Causes for such short-term variability have been evoked in the introductory Sect. 1.

For the first time, the seasonal modulation of the free tropospheric mean daytime HCOOH loading above a Northern mid-latitude site has been assessed statistically at the 1-σ confidence level (Fig. 3), showing a broad maximum during June-July-August and a November-December-January minimum, with a mean maximum to minimum ratio of 4. At the same confidence level, the shaded area in Fig. 3 reflects primarily the observed inter-annual variability noticeable in Fig. 2. A month-specific climatology of the mean daytime HCOOH loading above the Jungfraujoch for the period 1985 to 2007 is quantified in Table 2. The seasonal modulation, characterized by a regular tropospheric HCOOH increase from March to June and a relatively fast decrease during September and October, is in phase with various ground-level biogenic vegetation metabolisms described, e.g., by Kesselmeier and Staudt (1999), which are largely dependent on solar insolation, local temperature and related atmospheric dynamics, as well as specific regional soil coverage. The influence of the vegetation photosynthesis is corroborated here by the observed off-phase loading of CO₂ above the Jungfraujoch versus the seasonal HCOOH modulation. Thermal convection resulting from solar irradiation appears to dominate local releases of source gases at the season-changing ground cover, followed by their transport up to and above the Jungfraujoch site, where they mix and dilute with trade winds.

The lack of a long-term November-December-January change (see Sect. 6) supports the idea that the background formic acid content in the remote troposphere above the Alpine plateau is essentially of natural (direct or indirect) recurring biogenic origin, in line with findings by Legrand et al. (2003). This view is further supported by the
research reported by Glasius et al. (2001), who found that the $^{14}$C content measured in boundary layer carboxylic acids at various remote sites through Western Europe is indicative of a natural biogenic contribution in excess of 80% to the atmospheric HCOOH loading.

While the first evidence for the presence of HCOOH in the upper troposphere was based on remote solar FTIR observations from aboard a stratospheric balloon platform (Goldman et al., 1984), subsequent studies have largely relied on local in situ ground-, sea- and airborne measurements. However, numerous recent Northern mid-latitude investigations listed in Table 4 have re-actualized the infrared remote sensing approach, based on either the solar occultation- or the thermal limb emission mode. In particular, the first global distribution of upper tropospheric formic acid at sunrise and sunset by the ACE-FTS space instrument (Bernath et al., 2005; González Abad et al., 2009) is definitely a “first”, likely to help understanding and modeling the HCOOH distribution on a global scale. In line with the comprehensive laboratory intercomparison exercise of biomass-burning emissions (Christian et al., 2004) evoked in Sect. 3, we believe that a concerted effort should be undertaken to identify measurement techniques and observational platforms, to optimally achieve complementarity, temporal continuity and spatial coverage, with focus on accuracy, detection sensitivity, and long-term stability. Observation priorities of sets of key species and documentation of ancillary environmental parameters should be further identified in concert with modeling groups.

The potential of modern, fast, wide band-pass FTIR spectrometers using the sun as source of radiation, is their ability to regularly record solar spectra over the entire middle-infrared spectral domain from (∼2 to 15 µm) under daytime clear sky conditions, with very high spectral resolution and large signal-to-noise ratios. This remote sensing technique, which was pioneered at Kitt Peak during the 1970s by Brault (1978), progressively supplanted slow and bulky grating spectrometers used till then for solar observations made at the Jungfraujoch since the early 1950 (Migeotte et al., 1956; Delbouille and Roland, 1995). At the onset of the 1980s, the Jungfraujoch solar observation strategy became increasingly dictated by the need to support the Montreal- and
the emerging Kyoto Protocols, thus focusing on the monitoring of stratospheric ozone and ozone-depleting inorganic gases of the NO\textsubscript{y} and Cl\textsubscript{y} families, as well as all related ground-released organic natural- and anthropogenic greenhouse gases with significant absorption strengths in the thermal infrared (e.g., Mahieu et al., 2000, 2007; Zander et al., 2005). In addition to these Protocol-related target gases, the middle-infrared domain further contains characteristic absorption signatures of many other organic trace constituents (Zander et al., 2008) among which we chose to deal here with HCOOH, because of its indirect influence on the oxidizing capacity of the troposphere, as well as its critical, ubiquitous contribution to the boundary layer acidity.

An effort dealing with correlations between HCOOH and other species (i.e., CO, HCN, C\textsubscript{2}H\textsubscript{2}, C\textsubscript{2}H\textsubscript{6}, H\textsubscript{2}CO, CH\textsubscript{3}OH ) also monitored at the Jungfraujoch, is currently in progress. Specifics regarding these databases and preliminary correlations can be obtained by contacting Emmanuel.Mahieu@ulg.ac.be.

Considering the original HCOOH-related findings reported in this work, and found to be commensurate to within a factor 2 with numerous other Northern mid-latitude field observations, it would be very appropriate to see a modeling effort focusing on this zonal part of the Earth. A particular attention should even focus on the Western European Continent, incorporating adjusted emission sources reported by Legrand et al. (2003) for summer 2000 over France. Top-down versus bottom-up modeling outcomes may also shed light on the possible occurrence of indirect production of formic acid within the free troposphere.

**Appendix A**

In the following paragraphs, we briefly discuss findings derived here versus results from other field campaigns reported in the literature, with emphasis on free tropospheric HCOOH evaluations at Northern mid-latitudes. Relevant numerical comparisons are provided in column 3 of Table 4. Where relevant, boundary layer HCOOH concentrations are also evoked.
Because of observational- and analytical similarity, we first focus on the long-term HCOOH investigation reported by Rinsland et al. (2004). This research was based on the analysis of solar spectra recorded between June 1980 and September 2002 at the Kitt Peak National Solar Observatory (31.9° N, 111.6° E, 2.09 km altitude), a mountain top site located in the semi-arid Sonora desert in South-West Arizona, USA. It resulted in a retained set of 116 daily mean constant VMRs of HCOOH through the free troposphere out to 10 km altitude, much sparser than the 1501 days available here over a similar time period (22 yr). Due to large variability in these daily means, Rinsland et al. (2004) reported quarterly (3 months) mean VMR values, namely (356±179) pptv for J-F-M, (∼680±300) pptv for A-M-J, a maximum of (792±323) pptv during J-A-S, and a minimum of (313±175) pptv in O-N-D, with the uncertainties corresponding to statistical means of the intervening daily averages. The magnitude of the quarterly uncertainties prevented a seasonal cycle to be assessed statistically, thus the cautious wording “evidence for a seasonal cycle” in the title of their paper (Rinsland et al. 2004; see their Fig. 4). Compared with our present findings and other Northern mid-latitude ones (i.e., Norton et al., 1992; Talbot et al., 1997a,b; Reiner et al., 1999; Singh et al., 2000; Remedios et al., 2007; Gonzalès Abad et al., 2009), the mean Kitt Peak values appear exceedingly high, more representative of remote boundary layer concentrations documented in various reviews (e.g., Kesselmeier and Staudt, 1999; Khare et al., 1999; Poisson et al., 2000). However, lowering them by 2 in accordance with the new improved spectroscopic parameters of Perrin and Vander Auwera (2007), brings them closer to expectation as shown in Table 4. Despite this absolute halving, they still remain high compared to the mean Jungfraujoch values. This discrepancy may result from the fact that Kitt Peak is 15 degrees further south and a full 1.5 km lower in altitude than the Jungfraujoch, thus experiencing enhanced convective transport of warmer boundary layer air masses which carry formic acid and various precursor gases up to and above the site.

Another noticeable difference deals with the relative 3-month maximum (J-A-S) over minimum (O-N-D) concentration ratio found equal to 2.5 at Kitt Peak, which is lower...
than the max. (J-J-A) to min. (N-D-J) ratio of 4.0 found for the Jungfraujoch. If we strictly consider the quarterly periods adopted for Kitt Peak, then the Jungfraujoch J-A-S over the O-N-D mean VMRs derived from Table 2 return a ratio of 3.0, which is in reasonable agreement when accounting for the systematic uncertainties reported for both sites, namely 28% for Kitt Peak and 18% for the Jungfraujoch. Rinsland et al. (2004) pointed to the sparseness of the Kitt Peak database, which prevented quantifying diurnal- or inter-annual variations, nor a long-term trend. Whether the reported occurrence of both the maximum and the minimum HCOOH loading above Kitt Peak during the two adjacent last quarters of the year is real or the result of insufficient internal consistency of the database remains an open question.

Earlier HCOOH investigations, e.g., Arlander et al. (1990) along a sea cruise in the Pacific- and Indian Oceans, Moody et al. (1991) during a campaign at Amsterdam Island, and Baboukas et al. (2000) based on a N-S Atlantic sea cruise, suggested some seasonal variation with maxima occurring during local summer and minima in winter, but their significance was statistically weak, because of time limited campaigns.

It is worth evoking here the analysis of an ice core extracted at the remote Col du Dôme (4250 m, French Alps) by Legrand et al. (2003), who report records of carboxylic acids covering the period from 1925 to 1995. Formate-related findings showed a seasonal cycle, with somewhat “chaotic” mean summer/winter contrasts, i.e., 3 from 1930 to 1940, 1.7 between 1950 and 1975, 7 from 1980 to 1990, and a ratio dropping towards 4 during the last years of their time base. This latter ratio is commensurate with the mean seasonal peak-to-peak modulation of HCOOH derived from our Fig. 3 for the past two decades, i.e., 4. The authors attributed the changes in the max./min. seasonal contrast to the evolving magnitude of summer versus winter thermal convection of air masses from the boundary layer to the free troposphere. The Col du Dôme record also showed that 1985 to 1995 summertime concentrations of formate were close to values determined between 1925 and 1950, which led Legrand et al. (2003) to conclude that the more recent impact of industry-related anthropogenic contributions to atmospheric HCOOH remains weak. This conclusion is corroborated by Glasius et al. (2001) who
found that the $^{14}\text{C}$ content measured in boundary layer carboxylic acids at remote sites through Western Europe is indicative of a natural biogenic contribution well in excess of 80% to the total HCOOH loading.

Incidentally, Legrand et al. (2004) also reported on another multiple gas in situ study, namely at the Antarctic station Dumont d’Urville (DdU; 60° S, 140° E, 140 m a.s.l.) between 1997 to 2002, with mean HCOOH mixing ratios of about 70 pptv during local winter (May to August; polar night), increasing to $\sim$200 pptv during local summer (November to February). Little influence of air masses origin suggested this seasonal variation to be characteristic of the Antarctic Ocean biogenic emissions. Considering the “extreme” marine environment, nearly free of vegetation and industrial activity, it is relevant to notice that the local HCOOH concentrations reported by Legrand et al. (2004) indicate (1) a mean DdU summer/winter ratio of 3, which is in line with the mean values reported for both Northern mid-latitude Jungfraujoch and Kitt Peak continental sites during the past 2 decades, and (2) daytime concentrations some 30 pptv higher than night-time ($\sim$50 pptv) values during a summer test, thus in line with mean J-J-A sunrise to daytime concentrations found above the Jungfraujoch and documented in Tables 2 and 3.

How do the HCOOH findings derived in situ by Norton (1992) during a short 1–21 May 1988 campaign at the Mauna Loa Observatory (MLO; 19.5° N, 155.6° W) compare with those reported here? MLO is located on the Hawaii Island, in the middle of the Pacific Ocean, at an altitude of 3400 m (similar to that of the Jungfraujoch) on the northern slope of the still active Mauna Loa volcano. Average local HCOOH day- and night-time concentrations and standard deviations were reported equal to 452±268 pptv and 63±34 pptv, respectively. Owing to the MLO altitude and geographic location, Norton et al. (1992) concluded that the daytime values are characteristic of local, sub-tropical marine air masses modified by contact with the island ground and vegetation along their way to the site, whereas the night-time ones are representative of descending air from the free troposphere. While the latter are in line with those reported by Talbot et al. (1997a,b) for marine air masses above 2 km during both PEM-West A and B cam-
campaigns as well as with other free tropospheric measurements referred to in Table 4, the MLO findings in advected marine air during daytime are substantially higher than those found below 2 km during both PEM-W campaigns. The daytime MLO values are also much higher than observed by Arlander et al. (1990), i.e. an average HCOOH concentration of about 150 pptv during the last 3000 km of their 7° N longitudinal leg cruise between Singapore and Hawaii.

The local MLO day- to night-time ratio of 7 contrasts significantly with the mean Jungfraujoch culmination/sunrise ratio of 1.5 for the month of May, specifically derived from the red lines in Fig. 4 and reported in Table 3. This large diurnal ratio difference may result from the fact that MLO is a sub-tropical island site experiencing stronger daytime thermal convection, with marine background air masses advecting then convecting efficiently (a kind of “chimney effect”) towards the site over warm ground and dense local growing vegetation. Also relevant to this high MLO day/night ratio of HCOOH are findings reported by Arlander et al. (1990) during an April–July 1987 cruise transect between 50° N to 40° S in the Pacific and Indian Oceans, near 195° W longitude. They observed broad maxima for both formic and acetic acids centred at 15–20° N, and suggested they may result from increased biogenic emissions of olefinic precursors emitted by marine microbiota, then volatilized to the atmosphere and subsequently photo-oxidized to form organic acids (Keene and Galloway, 1988). A similar maximum was observed around 20° N by Baboukas et al. (2000) during a 30° W Atlantic transect from 63° N to 39° S between 11 October to 2 November 1996. Such marine features may have influenced HCOOH daytime concentrations at MLO. When only considering those mean daytime HCOOH concentrations strictly associated with the night-time data retained by Norton (1992; his Fig. 3), the day-to night ratio reduces from 7 to 5, which still remains relatively high. Although based on limited datasets, HCOOH diurnal ratios reported in the consulted literature, namely about 2 by Arlander et al. (1990), 2.5 by Khwaja (1995), 2.5 by Kajii et al. (1997), 2.2 by Baboukas et al. (2000), and 1.7 by Legrand et al. (2004), are in closer agreement with the mean Jungfraujoch ratio of 1.5, than with the Mauna Loa one of 7.
Further documented in Table 4 are airborne investigations of HCOOH in the free troposphere, based on in situ measurements at different northern mid-latitude locations. In particular, between 27 September and 2 October 1991, Reiner et al. (1999) performed five flights over Southern Germany, only a few degrees North of the Jungfraujoch. They measured HCOOH concentrations in the upper troposphere, showing a regular decrease from mean 215±100 pptv at 7 km to 60±40 pptv at 11.3 km altitude (this slope was accounted for in the confection of the HCOOH a priori profile adopted in the present retrieval process; see Sect. 2). The mean tropospheric VMR concentration read off the thick curve of Fig. 3 for late September is equal to 95±38 pptv, while a more specific value of 115±18 pptv was derived from 17 spectra recorded on 3 and 4 October 1991. If we reasonably assume that the Reiner et al. (1999; their Fig. 2c) slope of HCOOH applies further down through the troposphere, then their corresponding mean VMR concentration would be larger than the mean Jungfraujoch one by almost a factor 2, but still commensurate within the standard deviations of both data sets.

Limited but original measurements of HCOOH concentrations in the upper troposphere have been reported recently by Remedios et al. (2007). They were based on the spectrometric analysis of the HCOOH $\nu_6$ band observed in thermal limb emission spectra recorded during night-time on 8 May 1998 with the cryogenically cooled MIPAS-B2 FTIR balloon spectrometer (Friedl-Vallon et al., 2004). Altitude-specific concentrations reported by Remedios et al. (2007) using the old $\nu_6$ band intensities of HCOOH were equal to 320±120 pptv and 200±80 pptv, respectively at 7.5 and 10.4 km altitude. As can be seen in Table 4, dividing these values by 2 to account for the new spectroscopy (Perrin and Vander Auwera, 2007) brings them in reasonable agreement with the values found by Reiner et al. (1999), and commensurate with our mean tropospheric concentration of 114±19 pptv derived from Jungfraujoch observations made in the very early morning of 8 May 1998. Historically speaking, Goldman et al. (1984) reported the first tentative identification of HCOOH in the free troposphere, namely, 600 pptv at 8 km and 400 pptv at 10 km, based on the analysis of solar occultation spectra recorded from aboard a stratospheric balloon instrument above Arizona (USA). Even halving their re-
ported values to account for the new spectroscopy (Perrin and Vander Auwera, 2007) still leaves them quite high in comparison with the Reiner et al. (1999) and the Remedios et al. (2007) values; one of the causes certainly lies in the poor knowledge, at that time, of many interfering gases and their related spectroscopy.

The goal of the SONEX airborne field campaign of 13 October to 12 November 1997 (Singh et al., 2000; Jaeglé et al., 2000) was to evaluate the impact of aircraft emissions on the HO<sub>x</sub>-NO<sub>x</sub>-O<sub>3</sub> chemistry in the troposphere, between 2 and 12 km altitude, with focus on the North Atlantic flight corridor area between 40° to 65° N latitude. Among many oxygenated organic species concurrently studied, mean HCOOH concentrations were found equal to about 120±65 pptv, 45±30 pptv, and 39±25 pptv, respectively in the lower (2–6 km) and upper (6–12 km) troposphere, and in the lowermost stratosphere (see Table 4). Overall, these values agree well with the mean tropospheric VMR above the Jungfraujoch read off the thick curve of Fig. 3 for late October, namely 62±22 pptv. This comparison is a sound one, as Fuelberg et al. (2000) found that the SONEX campaign occurred during a year-to-year climatologically average situation. However, a look at Fig. 2 shows that 1997 was a year with an overall relatively low HCOOH loading above the Jungfraujoch. This is substantiated by observations specifically made at the Jungfraujoch during the SONEX period (i.e., 30 measurements encompassing 12 days between 19 October and 11 November 1997), returning a mean VMR of 39±17 pptv. While this lower value remains commensurate with the SONEX findings when considering the respective uncertainties, one can’t exclude that some SONEX soundings may have been made in air masses affected by biogenic emissions originating from the Eastern rim of the North American subcontinent. Based on all SONEX measurements made in the upper troposphere (8 to 12 km), Jaeglé et al. (2000) classified the sounded air masses into six categories according to their origin, namely local background, stratospheric influence, cirrus clouds, and convection of tropical marine-, midlatitude marine-, and continental air masses. Among these categories, the extreme median concentrations of HCOOH ranged from 37 pptv in local background air to 21 pptv in convected continental air masses, with an
occurrence-weighted average equal to 33.2 pptv. Noteworthy is the fact that the first
three categories returned a median HCOOH concentration of 35±2 pptv in about 80%
of the measurements, while the categories dealing with convection air masses showed
HCOOH medians of only 23±2 pptv. Whether this difference is indicative of the pres-
ence of a HCOOH source in the upper troposphere is worth of further consideration.

During two airborne “Pacific Exploratory Missions” (PEM) over the western Pacific
basin, east of the Asian rim, namely PEM-West A (from 16 September to 21 Octo-
ber 1991) and PEM-West B (from 7 February to 14 March 1994), HCOOH was one of
the numerous compounds simultaneously investigated in situ between 0.3 and 12.5 km
altitude. The timing of the campaigns was chosen to characterize the eastward Asian
outflow of material to the North Pacific troposphere, which is lowest during fall and high-
est during springtime, when the high-pressure oceanic system is optimally displaced
from West to East (Prospero et al., 1985). Overall findings derived from the PEM
missions have been reported by Talbot et al. (1996a, 1997a,b) according to air mass
origin (i.e., continental south (0 to 20° N), continental north (20 to 60° N) and marine
(air masses that were over the Pacific for at least 5 days)), and over specific altitude
layers (i.e., <2 km, 2–7 km, and 7–12 km). Relevant HCOOH data collected over the
20° to 60° latitude band are listed in Table 4. Contrary to the expected continental air-
mass outflow evoked above, the HCOOH concentrations were lower during PEM-West
B (late winter) compared to PEM-West A (early fall). Talbot et al. (1997b) related the
observed “seasonal” difference of monocarboxylic acids to depressed biogenic vegeta-
tion metabolisms during PEM-West B, as evidenced by a 2–3% increase in concurrently
measured CO₂ concentrations. While the PEM campaigns were occurring east of the
Asian continent, the Jungfraujoch is located east of the Atlantic Ocean, both un-
der the influence of dominant westerly winds. It is therefore likely that, on average, the
HCOOH loading above the Jungfraujoch predominantly reflects free tropospheric ma-
ine air masses, occasionally “polluted” on their way towards the Alpine plateau. This
was already suggested by Hartmann et al. (1989) who found that HCOOH mixing ratios
measured during three airplane flights over Western Germany averaged 170±60 pptv.
in westerly marine air flows, substantially lower than those found in intrusion flows from other directions. The mean HCOOH concentrations read off the thick curve in Fig. 3 for the periods of PEM-West-A and -B, namely $85 \pm 38$ pptv and $75 \pm 20$ pptv, are in better agreement with mean concentrations found in characteristic marine air masses during these campaigns (see Table 4).

A first near global distribution of formic acid between 5 km altitude and the tropopause has been reported recently by González Abad et al. (2009). It is based on remote solar occultation observations made at sunrise and sunset between February 2004 and September 2007 with the ACE-FTS equipment operating aboard the Canadian SCISAT satellite (Bernath et al., 2005). The instrument scanning speed provides a vertical resolution of 2 to 3 km in targeted concentration profiles, but only during a few minutes at local sunrises and sunsets. The HCOOH retrieval analysis (Boone et al., 2005) is based on the fitting of the $\nu_6$ band Q-branch and the updated spectroscopic parameters (Perrin and Vander Auwera, 2007) as adopted here. The HCOOH profiles determined by ACE over the continental USA have been inter-compared with near time-coincident solar occultation observations made over New Mexico (USA) by the MKIV-FTIR balloon instrument (Toon, 1991), with profiles agreeing to within their respective one standard deviation. Overall also, both inter-compared datasets confirm the decrease of the HCOOH concentration with increasing altitude, thus in line with earlier airplane observations (e.g., Reiner et al., 1999; Singh et al., 2000; Talbot et al., 1996a, 1997b) and with the slope adopted for our HCOOH a priori profile (see Sect. 2). The pole-to-pole ACE data for the period December 2004 to September 2007 have been reported as average VMR profiles of HCOOH in 10° wide latitudinal bins integrated over all longitudes (see Table 1 at http://www.atmos-chem-phys-discuss.net/9/12465/2009/acpd-9-12465-2009-supplement.pdf). Mean numerical values found in the 40° to 50° North latitude zone (which encompasses the Jungfraujoch location), show discrete concentrations of 157, 95, 75 and 41 pptv, respectively at 5, 7, 9 and 11 km altitude, which are in good agreement with a yearly mean combined sunrise-sunset free tropospheric concentration of $89 \pm 32$ pptv derived from our Table 3.
Acknowledgements. This work is part of an overall atmospheric monitoring effort conducted at the University of Liège and primarily financed by the Belgian Federal Science Policy Office, the Belgian Fonds National de la Recherche Scientifique, the Belgian Communauté Française, and the European Commission, all in Brussels. The affiliation of the Jungfraujoch site to the International “Network for the Detection of Atmospheric Composition Change” (NDACC, formerly NDSC (Network for the Detection of Stratospheric Change); see http://www.ndacc.org) since 1989 has significantly boosted the implication of the University of Liège group in monitoring Earth’s atmospheric composition changes. We thank all colleagues and collaborators who have contributed to the decades long production of the database used here, and to the operation and upgrading of the Liège laboratory at the Jungfraujoch. Thanks are also extended to colleagues from the Royal Observatory of Belgium and from the Belgian Institute for Space Aeronomy for their participation to intensive observational campaigns. Acknowledgements further go to the Swiss Jungfraubahnen for their continuous maintenance of and year-round access to the Jungfraujoch Station, as well as to the HFSJG Directorate for the logistic management of the numerous researches conducted at the Jungfraujoch site. J. VDA acknowledges financial support from the Belgian Fonds de la Recherche Scientifique (contracts FRFC and IISN), and the Actions de Recherches Concertées of the Communauté Française de Belgique. Analysis at the NASA Langley Research Center was supported by NASA’s Upper Atmospheric Chemistry and Modeling Program (ACMAP) and Upper Atmospheric Research Program (UARP).

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Formic acid above the Jungfraujoch during 1985–2007

R. Zander et al.


Talbot, R. W., Bradshaw, J. D., Sandholm, S. T., Smyth, S., Blake, D. R., Blake, N. J.,


Table 1. Impact of major sources of random (R) and systematic (S) uncertainties on typical individual HCOOH column retrievals above the Jungfraujoch.

<table>
<thead>
<tr>
<th>Sources of uncertainty</th>
<th>R or S</th>
<th>Max. Error (%)</th>
<th>Remark</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spectra quality</td>
<td>R</td>
<td>4</td>
<td>zero offset and S/N</td>
</tr>
<tr>
<td>Pressure-Temperature profile</td>
<td>R</td>
<td>4</td>
<td>±4 K against noon NCEP profile</td>
</tr>
<tr>
<td>HCOOH a priori profile</td>
<td>R</td>
<td>7</td>
<td>±50% variability around a priori</td>
</tr>
<tr>
<td>HDO a priori profile</td>
<td>R</td>
<td>6</td>
<td>±50% variability around a priori</td>
</tr>
<tr>
<td>Other interferences</td>
<td>R</td>
<td>5</td>
<td>other gases, i.e. O, H₂O, CHClF₂</td>
</tr>
<tr>
<td><strong>RSS</strong> Total Random</td>
<td></td>
<td><strong>&lt;13</strong></td>
<td></td>
</tr>
<tr>
<td>HCOOH spectroscopy</td>
<td>S</td>
<td>&lt;7</td>
<td>band intensity; see Sect. 3</td>
</tr>
<tr>
<td>HDO spectroscopy</td>
<td>S</td>
<td>6</td>
<td>HITRAN-2004</td>
</tr>
<tr>
<td>O₃ spectroscopy</td>
<td>S</td>
<td>3</td>
<td>HITRAN-2004</td>
</tr>
<tr>
<td>HCOOH a priori profile</td>
<td>S</td>
<td>12</td>
<td>constant versus sloped profile</td>
</tr>
<tr>
<td>HDO a priori profile</td>
<td>S</td>
<td>8</td>
<td>factor 2 change in a priori slope</td>
</tr>
<tr>
<td>ILS and forward model</td>
<td>S</td>
<td>5</td>
<td>retrieval algorithm-related</td>
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<tr>
<td><strong>RSS</strong> Total Systematic</td>
<td></td>
<td><strong>&lt;18</strong></td>
<td></td>
</tr>
</tbody>
</table>

(*) RSS stands for square root of the sum of the squares
Table 2. Month-specific climatology of the mean daytime HCOOH loading above the Jungfraujoch for the 1985 to 2007 period.

<table>
<thead>
<tr>
<th>Month+ Annual Mean</th>
<th>Nr. of Years</th>
<th>Mean Col. ( \times 10^{15} ) molec./cm(^2)</th>
<th>Mean VMR in pptv</th>
<th>Std. Dev. in %</th>
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</thead>
<tbody>
<tr>
<td>Jan</td>
<td>17</td>
<td>0.455</td>
<td>50.1</td>
<td>46.7</td>
</tr>
<tr>
<td>Feb</td>
<td>18</td>
<td>0.679</td>
<td>74.7</td>
<td>28.5</td>
</tr>
<tr>
<td>Mar</td>
<td>20</td>
<td>0.895</td>
<td>98.5</td>
<td>29.4</td>
</tr>
<tr>
<td>Apr</td>
<td>15</td>
<td>1.146</td>
<td>126.0</td>
<td>29.8</td>
</tr>
<tr>
<td>May</td>
<td>18</td>
<td>1.283</td>
<td>141.1</td>
<td>26.0</td>
</tr>
<tr>
<td>Jun</td>
<td>19</td>
<td>1.697</td>
<td>186.7</td>
<td>27.1</td>
</tr>
<tr>
<td>Jul</td>
<td>15</td>
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<td>108.1</td>
<td>47.2</td>
</tr>
<tr>
<td>Oct</td>
<td>20</td>
<td>0.639</td>
<td>70.3</td>
<td>34.6</td>
</tr>
<tr>
<td>Nov</td>
<td>17</td>
<td>0.441</td>
<td>48.1</td>
<td>32.8</td>
</tr>
<tr>
<td>Dec</td>
<td>18</td>
<td>0.360</td>
<td>39.6</td>
<td>46.1</td>
</tr>
<tr>
<td><strong>Annual Mean</strong></td>
<td><strong>17.5</strong></td>
<td><strong>1.01</strong></td>
<td><strong>110.9</strong></td>
<td><strong>34.4</strong></td>
</tr>
</tbody>
</table>
**Table 3.** Mean AM increases and PM decreases of the HCOOH loading above the Jungfraujoch based on the 1995–2007 observations with the Bruker instrument.

<table>
<thead>
<tr>
<th>Months</th>
<th>All columns in $\times 10^{15}$ molec./cm$^2$ – In brackets, related mean VMRs in pptv</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&quot;Sunrise&quot; ← Increase → Culmination ← Decrease → &quot;Sunset&quot;</td>
</tr>
<tr>
<td>Nov-Dec-Jan</td>
<td>0.26 [29] 0.25 [27] 0.51 [56] 0.19 [21] 0.32 [35]</td>
</tr>
<tr>
<td>Feb-Mar</td>
<td>0.58 [64] 0.20 [22] 0.78 [86] −0.09 [−10] 0.87$^{(1)}$ [96]</td>
</tr>
<tr>
<td>Apr-May</td>
<td>0.90 [99] 0.39 [44] 1.29 [142] 0.34 [37] 0.95 [104]</td>
</tr>
<tr>
<td>Jun-Jul-Aug</td>
<td>1.50 [165] 0.40 [44] 1.90 [209] 0.54 [59] 1.36 [160]</td>
</tr>
<tr>
<td>Sept-Oct</td>
<td>0.50 [55] 0.40 [44] 0.90 [99] 0.13 [14] 0.77 [85]</td>
</tr>
<tr>
<td>Year Mean</td>
<td>0.77 [85] 0.33 [36] 1.10 [121] 0.25 [35] 0.85$^{(1)}$ [93]</td>
</tr>
<tr>
<td>May only</td>
<td>0.94 [103] 0.48 [53] 1.42 [156] 0.45 [48] 0.97 [107]</td>
</tr>
</tbody>
</table>

$^{(1)}$ the Feb–Mar sunset value is significantly biased high due to missing low sun observations evoked in Sect. 2; it has therefore been excluded from the yearly mean.
Table 4. Timely comparison of local free tropospheric HCOOH concentrations (in pptv) at Northern mid-latitudes with relevant mean values above the Jungfraujoch.

<table>
<thead>
<tr>
<th>Location and Observation Period</th>
<th>Measurement specifics and Altitude or Range</th>
<th>Typical HCOOH Concentrations in pptv (1)</th>
<th>References for Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jungfraujoch, Swiss; 3.58 km altitude; 46.5° N; Sep 1985 to Sep 2007</td>
<td>Daytime IR remote sensing 3.58–10.6 km (free troposphere)</td>
<td>J-J-A max. mean: 182±55</td>
<td>This work</td>
</tr>
<tr>
<td>Kitt Peak, AZ-USA; 2.09 km altitude; 31.9° N; Jun 1990 to Oct 2002</td>
<td>Daytime IR remote sensing 2.09–10 km (free troposphere)</td>
<td>Published values divided by 2 J-A-S max. mean: 386±162 (161) O-N-D min. mean: 156±88 (83)</td>
<td>Rinsland et al., 2004</td>
</tr>
<tr>
<td>South Germany; 48–53° N; 5 flights; 27 Sep–2 Oct 1991</td>
<td>Airborne in situ, daytime – 7 km altitude – 11.3 km altitude</td>
<td>215±100 (95–115) 60±40</td>
<td>Reiner et al., 1999(2)</td>
</tr>
<tr>
<td>Western Mediterranean; Night-time IR limb emission Balloon flight; May 8, 1998</td>
<td>– 7.5 km altitude – 10.4 km altitude</td>
<td>Published values divided by 2 160±61 (114) 100±38</td>
<td>Remedios et al., 2007(2)</td>
</tr>
<tr>
<td>ACE-FTS Scisat; partim. 40–50° N, longitude-integrated; Dec 2004–Sept 2007</td>
<td>Space-based IR remote SR+SS</td>
<td>157±60 95±30 (89)</td>
<td>González Abad et al., 2009(2)</td>
</tr>
</tbody>
</table>

(1) values between parentheses refer to timely Jungfraujoch findings discussed in Appendix A.
(2) some reported uncertainties have been estimated from Figures in last column related references.
Fig. 1. Typical fitting example to an observed spectrum (respectively crosses and continuous line in Frame (B) extending from 1104.65 to 1105.60 cm\(^{-1}\), recorded at the Jungfraujoch on 8 June 2005, with a spectral resolution of 0.004 cm\(^{-1}\). The resulting residuals (observed minus calculated signals) are displayed in Frame (C), with a corresponding root mean squares fitting residual equal to 0.101%, which is consistent with the displayed spectrum’s S/N ratio of 1285. Frame (A) shows the individual discrete absorption characteristics (shifted vertically for clarity) of the HCOOH target gas and the 5 most important interfering ones which, complemented by contributions from other second order interfering gases, correspond to the fit in Frame (B).
Fig. 2. Frame (A) reproduces all individual total HCOOH column abundances derived in this work, while Frame (B) shows the daily arithmetic means and their related standard deviations. In both frames, the red- and black symbols correspond to columns derived from observations made, respectively with the home-built FTIR spectrometer and with the commercial Bruker-120 HR instrument. In Frame (B), the crossed circles identify daily mean columns observed during the November-December-January months, which have been used selectively to determine a “background” mean daytime loading (continuous line) and a standard deviation (dashed lines) of HCOOH above the Jungfraujoch during that period of the year. For details, see the beginning of Sect. 5.
Fig. 3. Monthly mean column abundances and associated standard deviation bars displayed on a one-year time base, with 5 obvious monthly mean outliers identified by triangular symbols. The thick curve corresponds to a running mean fit to the black filled symbols only, with a 15-day step and a 2-month wide integration time. The shaded area visualizes the $1-\sigma$ standard deviation associated to the running mean curve and reflects primarily the observed inter-annual variability noticeable in Fig. 2. The dashed line, which corresponds to the mean HCOOH loading of November-December-January (thin continuous line) plus twice its standard deviation, visualizes the presence of a seasonal modulation in the 1985–2007 database. The right side scale allows converting an observed total HCOOH column abundance read off the left side scale into a mean VMR concentration (expressed in parts per trillion by volume, i.e., pptv) over the 3.58 to 10.6 km altitude range. For details, see Sects. 5, 2 and 3.
Fig. 4. Illustration of the diurnal variation of HCOOH above the Jungfraujoch, based on the individual Bruker FTIR observations between 1995 and 2007, with overall AM increases and PM decreases evaluated separately. First order fittings to all N-D-J data (filled cyan points) are highlighted by the two yellow lines. The black and red lines correspond to similar fittings, respectively to all J-J-A and May observations. Related AM increases and PM decreases are given in Table 3. The extreme high HCOOH columns observed on 15 August 1995 and on 14 January 2007 (respectively 25 black- and 7 pink filled circles) were excluded from the fittings.