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Fluorescence from atmospheric aerosol detected by a lidar indicates biogenic particles in the lowermost stratosphere

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Abstract. With a lidar system that was installed in Lindenberg/Germany, we observed in June 2003 an extended aerosol layer at 13 km altitude in the lowermost stratosphere. This layer created an inelastic backscatter signal that we detected with a water vapour Raman channel, but that was not produced by Raman scattering. Also, we find evidence for inelastic scattering from a smoke plume from a forest fire that we observed in the troposphere. We interpret the unexpected properties of these aerosols as fluorescence induced by the laser beam at organic components of the aerosol particles. Fluorescence from ambient aerosol had not yet been considered detectable by lidar systems. However, organic compounds such as polycyclic aromatic hydrocarbons sticking to the aerosol particles, or bioaerosol such as bacteria, spores or pollen fluoresce when excited with UV-radiation in a way that is detectable by our lidar system. Therefore, we conclude that fluorescence from organic material released by biomass burning creates, inelastic backscatter signals that we measured with our instrument and thus demonstrate a new and powerful way to characterize aerosols by a remote sensing technique. The stratospheric aerosol layer that we have observed in Lindenberg for three consecutive days is likely to be a remnant from Siberian forest fire plumes lifted across the tropopause and transported around the globe.

1 Introduction

Soot released by combustion into the atmosphere absorbs radiation and therefore heats the climate countering the cooling effect generally assigned to anthropogenic aerosols (Chung and Seinfeld, 2002). Aerosol particles are released either at the Earth’s surface from various sources, the crust (dust), the sea (sea salt aerosol) or by combustion processes (soot), or they form in situ by gas to particle conversion, like sulphate aerosol. While particles may enter the stratosphere in the tropics (Brock et al., 1995), the tropopause in the mid latitude efficiently suppresses tropospheric-stratospheric transport (TST) (Holton et al., 1995). However, recent observation of stratospheric aerosol layers by satellite-borne instruments and ground-based lidar suggest that strong thunderstorms are able to inject smoke from forest fires into the stratosphere at high latitudes (Fromm et al., 2000). Latest in situ measurements also proof the presence of forest fire particles in the stratosphere (Jost et al., 2004). Even though these layers are optically thin, they are highly relevant because when aerosols leave the troposphere they escape their most efficient removal mechanism which is wet scavenging and consequently their lifetimes are strongly increased. The implications of particles in the stratosphere on the radiative balance and chemistry crucially depend on their physical and chemical properties. Ground or satellite based sensors may measure the global distribution of the optical depth of aerosols but usually provide little information on the exact type and source of the particles (Kaufman et al., 2002).

Lidar is a powerful tool to investigate atmospheric aerosol since it makes it possible to measure its distribution from near to the ground up to the upper atmosphere with high vertical and temporal resolution. The different lidar techniques applicable to aerosol measurements like multi-wavelength backscatter lidar, depolarisation measurement and Raman techniques allow the determination of the particle’s macro- and microphysical properties to some extent. However, a precise and reliable characterization of aerosol based on lidar data which includes a description of its size distribution and chemical composition is generally hampered by the limited information available from this remote sensing techniques.

Here, we report the observation of a stratospheric aerosol layer by a ground based lidar system. Besides aerosol layers in the middle troposphere and cirrus clouds at about 12 km, Fig. 1 shows a thin layer of particles in the lowermost
stratosphere with a backscatter ratio of about 3–4. This layer
gave rise to a red shifted inelastic backscatter signal that we
detected with our instrument and interpret as laser induced
fluorescence (LIF) created by the aerosol particles. At first
we briefly describe the methods used for this observations
(Sect. 2) and the results obtained from them (Sect. 3). We
will then discuss fluorescence of aerosol as a possible cause
for the detected inelastic signal (Sect. 4) before we draw the
corresponding conclusions in Sect. 5.

2 Instrumentation and water vapour measurements

The Mobile Aerosol Raman Lidar (MARL) of the Alfred
Wegener Institute (AWI) is designed for measurements of
aerosols and water vapour in the stratosphere and upper troposphere (Schäfer et al., 1995). It detects light backscattered
by molecules and condensed matter in the atmosphere from
outgoing laser beams at 532 nm and 355 nm. Inelastic Raman
backscatter from nitrogen is detected at 387 nm and 607 nm
and allows the direct retrieval of the aerosol extinction co-
efficient $\alpha$ (Ansmann et al., 1992). An additional detection
channel at 407 nm with a bandwidth of 5.7 nm detects the vi-
brational Raman scattering from H$_2$O molecules and allows
the determination of the water vapour mixing ratio profiles
up to about 8 to 10 km at night time.

The emitter of the lidar is a Nd:YAG laser with frequency
doubling and tripling, generating laser pulses with about
300 mJ at both 355 nm and 532 nm with a repetition rate of
30 Hz and a pulse duration of 8 ns. The receiver is based on
a Cassegrain telescope with 1.1 m aperture coupled by two
fibre bundles to the 10-channel polychromator. Our system
is able to perform unattended measurements and can be con-
trolled via the internet, thus enabling us to gain a large dataset

![Fig. 1. Lidar backscatter ratio at 532 nm as a function of time and altitude from Lindenberg on 1 June 2003.](image)

even with a minimum of manpower. Continuous data records
spanning several consecutive days were achieved on occasions owing to the high degree of automation of the system.

A lidar signal measured at the wavelength $\lambda$ is proportional
to the total backscatter coefficients at this wavelength $\beta_\lambda$ and
is described by the following equation:

$$P_\lambda(z) = C \frac{O(z)\beta_\lambda(z)T^2(z)}{z^2}$$

(1)

$C$ summarizes all system constants like laser energy and de-
tection efficiency and $T(z)$ is the transmission term that can
be calculated by using Beer-Lambert’s law:

$$T(z) = \exp\left[-\int_0^z dz' \left(\alpha^M(z') + \alpha^A(z')\right)\right]$$

(2)

$\alpha^M$ and $\alpha^A$ refer to the extinction coefficients caused by
molecular and particle scattering, respectively. $O(z)$ de-
scribes the geometric overlap between laser and field of view
of the receiving telescope.

Inelastic scattering can be detected at the vibrational Ra-
man lines of atmospheric gases (Melfi, 1972). The backscatter
coefficient $\beta$ is in good approximation a function of the
number density profile of the corresponding molecule $N(z)$
and the cross section $\sigma_\lambda$ for Raman scattering which we as-
sume here to be independent on temperature. For water
vapour the backscatter coefficient is then simply a function of
the concentration of water molecules and the backscattering
cross section which is constant throughout the atmosphere:

$$\beta_{407}(z) = N_{H_2O}(z) \ast \sigma_{407}^{H_2O}$$

(3)

Introducing this equation into the lidar Eq. (1) yields an ex-
pression describing the water vapour Raman signal measured
by a lidar. In an analogue way the Raman signal from nitrogen that is measured at 387 nm can be described. The ratio of the water vapour and the nitrogen Raman lidar signal is then proportional to the water vapour mixing ratio $w_{H_2O}(z)$:

$$w_{H_2O}(z) = \frac{N_{H_2O}(z) M_{H_2O} w_{N_2}}{N_{N_2}(z) M_{dry}} = C' \frac{P_{407}(z)}{P_{387}(z)}$$

(4)

$O(z)$ is the same for both lidar signals and thus cancelled. $M_{H_2O}$ and $M_{dry}$ are the molecular weights of water and dry air, respectively. The correction for the differences in the remaining transmission terms ($T_{407}(z)/T_{387}(z)$) that was neglected in Eq. (4) is performed as described by Whiteman et al. (1992). The mixing ratio of nitrogen $w_{N_2}$ is constant in the altitude range of interest. However, the calibration constant $C'$ needs to be determined by using another independent measurement of water vapour because the system constants can not be otherwise established.

The German weather service (DWD) provided Radiosonde (Vaisala RS 80) data from four daily launches at the site and these delivered a temperature and humidity profile and allowed the determination of the tropopause height. The water vapour measurement of the RS 80 sondes is based on an improved retrieval algorithm (Leiterer et al., 1997). This data served for calibrating the water vapour measurements of the lidar which is done by determining $C'$ from Eq. (4) using the water vapour mixing ratio $w_{RS}(z)$ determined from the radiosonde profile at an altitude $z_0$. An appropriate fit altitude $z_0$ is usually found in about 3–5 km altitude, where the aerosol load is low and the water vapour is still high. During the campaign in Lindenberg we gathered 37 occasions of coincident lidar and RS water vapour measurements. The calibration constant was determined each time and varied less than 5%.

The water vapour profiles from lidar measurements presented below were calibrated using lidar and radiosonde profiles measured simultaneously shortly before the events under discussion. The stability of the calibration is demonstrated in Fig. 2. On 30 May 2003 a calibration factor was determined at 2.3 km altitude. The differences between the lidar and the RS profiles (Fig. 2b) are low throughout the troposphere and are mainly due to the temporal variability of the water vapour profile. This calibration factor is used to determine the water vapour profile of the consecutive days. A profile calculated this way from 3 June 2004 is shown in Fig. 2c along with a simultaneously measured radiosonde profile. The differences of the two profiles are less than 0.2 g/kg (Fig. 2d). The strong aerosol layer in the boundary layer (<3 km) and the starting cloud formation on its top, indicated by the strong backscatter ratio, do not significantly interfere with the water vapour measurement of the lidar.

Beside water vapour and cirrus clouds another major focus of our measurements is tropospheric aerosol. It was demonstrated during a ship-borne campaign that our system is able to detect and differentiate dust from marine particles in the troposphere (Immler and Schrems, 2003). This is possible due to the fact that the depolarisation and the wavelength dependence of the backscatter coefficient expressed as colour index $C_I$ are different for these two types of aerosols. $C_I$ is defined as by

$$C_I = -\log \left( \frac{\beta_{532}^A}{\beta_{555}^A} \right) / \log \left( \frac{355 \text{ nm}}{532 \text{ nm}} \right)$$

(5)
Fig. 3. Time-altitude plot of lidar backscatter ratio at 532 nm profiles measured in Lindenberg/Germany starting on 1 June 2003, 00:00 UT and ending on 4 June 2003, 17:00 UT. The solid green line indicates the altitude of the lapse rate tropopause determined from radiosonde launches four times a day. The dashed black line indicates the temperature minimum. The upper solid line in red marks the 380 K potential temperature level.

Based on the colour index and the lidar ratio \( S \) defined as the ratio between the aerosol extinction and backscatter coefficient,

\[
S_{\lambda} = \frac{\alpha_{\Lambda}^A}{\beta_{\Lambda}^A}
\]

an estimate of the effective radius of the aerosol particles can be retrieved based on a method described in an earlier publication by Immler and Schrems (2003). Saharan dust has a large and wavelength dependent lidar ratio of about 40 sr at 532 nm and 75 sr at 355 nm and a small colour index of about zero or even below and shows significant depolarisation. Maritim aerosol on the contrary has a lower lidar ratio and does not depolarise. Urban and biomass burning aerosol have different optical properties and can be discriminated from the unpolluted types by its weak depolarisation, larger colour index \( C_{\lambda} \geq 1 \) and a large lidar ratio of about 60–80 sr at 532 nm (Wandinger et al., 2002). However, it is difficult to differentiate between urban and biomass burning aerosol on the basis of elastic backscatter lidar data since the differences in their optical properties are rather small and the variability of this properties due to aging, changes in relative humidity and other factors is relatively large.

3 Observations of inelastic scattering by aerosols

From April to October of 2003 we performed measurements at the Meteorological Observatory in Lindenberg (52.2° N, 14.1° E) near Berlin/Germany with the MARL. During this measuring campaign, strong layers of aerosol in the troposphere were frequently detected, and on several occasions we also observed layers of haze in the lower stratosphere. The most pronounced observation of this kind was made during the first days of June 2003, when a 0.4 to 1 km thick aerosol layer was dwelling at an altitude of about 13 km – about 1 km above the thermal tropopause – for more than 72 h. Figure 1 shows the backscatter ratio \( R \) at 532 nm as a function of time and altitude from 1 June 2003. \( R \) is the total atmospheric backscatter divided by the purely molecular contribution and thus a measure for the content of aerosol particles with \( R > 5 \) above which a thin and steady layer emerges at about 12 km altitude. This layer persisted for the rest of the day, gaining altitude and stayed at 13 km throughout 2 June 2003 and most of 3 June 2003 (Fig. 3).

The tropopause was occasionally split during this period into a cold point tropopause (CPT) defined by the local temperature minimum and a lower lapse rate tropopause (LPT) as defined by the standard WMO definition according to which the tropopause is at the lowest level where the temperature gradient is higher than \(-2\) K/km and the average of the gradient between this level and any level within the next 2 km is still higher than \(-2\) K/km. On the first day of the observation, 1 June 2003, LPT and CPT were at the same height, up to 1 km below the aerosol layer (Fig. 3). Around noon on 2 June 2003 the situation changed and a new temperature minimum appeared in 13.5 km altitude, just above the aerosol layer. The mean temperature in the aerosol layer itself decreased gradually from 213.5 to 209 K during the course of the three days (Fig. 5).
The aerosol layer at about 13 km is clearly above the tropopause but below the level of 380 K potential temperature (Figs. 3 and 6) which marks the border to the “over-world” where air can cross exclusively in the tropics (Holton et al., 1995). Therefore we assign the observed aerosol layer to the lowermost stratosphere. Backward trajectories on the basis of ECMWF data were retrieved using the BADC trajectory service (http://badc.nerc.ac.uk) with vertical advection based on the pressure vertical velocity $\omega$. The results for a 3 June 2003, 00:00 UT shown in Fig. 4 indicate that the air that we sampled in Lindenberg came from North America and has been travelling undisturbed across the Atlantic within five days. Trajectories calculated for other times between 1 June 2003, 12:00 UT and 3 June 2003, 18:00 UT yield very similar results.

The stratospheric aerosol layer was $650\pm300$ m (mean ± std. dev.) thick and the backscatter ratio of the layer varied in the range from 1.5 to 4 at 532 nm. The optical depth at 532 nm determined from the nitrogen Raman signals varied from below 0.01 to 0.05 (0.026±0.013). Within the accuracy of our measurements the values at 355 nm were about the same (0.02±0.01). From these measurements the lidar ratio $S=\alpha/\beta$ can also be determined, yielding fairly high values of $86\pm30$ sr for 355 nm and $100\pm20$ sr for 532 nm. The colour index ($C_I$) of the layer with 1.0±0.2 is rather constant throughout the whole period. The cloud shows a significant depolarisation (Fig. 6) with a clear diurnal cycle, reaching 20% during the day and dropping to 5% at night time (at 355 nm). Using Mie scattering theory an estimate of the effective radius $r_{eff}$ can be retrieved from the colour index. Depending on whether we use the refractive index of water, ice, sulphate or dust (D’Almeida et al., 1991) for the retrieval, we find values between 0.1 and 0.4 µm. The number concentration of the layers is calculated to be 500 to 2000 cm$^{-3}$ (based on an effective radius of 0.1 µm).

The optical properties that we measured are typical for an aged polluted aerosol (Wandinger et al., 2002) and therefore suggest absorbing, solid material containing soot emitted in urban regions or by biomass burning. The measured properties are different from typical values for dust, sea salt or sulphate. Finding aerosol of such properties in the lowermost stratosphere is an intriguing result, in particular with respect to the longevity of this layer. Nevertheless, there is one more remarkable observation: Besides the backscatter ratio and temperature profiles, Fig. 5 shows the signal of the 407 nm channel of our instrument. During the nights of 1 June 2003 and 2 June 2003 (daytime measurements at 407 nm are not available) there is a peak at the altitude of the stratospheric aerosol layer at about 13 km (Fig. 5b and c). This statistically significant signal was present throughout both nights, its strength correlated with the elastic backscatter signal of the aerosol layer.

As was demonstrated in Fig. 2 there is no cross talk from elastic backscatter into the 407 nm channel even in the
presence of low clouds that give rise to very strong elastic backscatter. We have also never observed other unwanted effects in our Raman channel like after-pulsing or strong signal induced noise, that could explain the unusual signal at 13 km altitude (corresponding to a delay of 86 µs after the laser shoots). We therefore came to the conclusion that we detected inelastic scattering originating from the stratospheric aerosol layer. However, it was not clear in the first place what could cause such a strong inelastic signal at that altitude.

The simplest explanation is a high water vapour content in that plume. However, this explanation can be ruled out since the relative humidity above ice according to the calibrated 407 nm signal would be up to several thousand percent given the low temperature at the corresponding altitude. Even though high supersaturations have been observed in the atmosphere, such high values are not conceivable and definitely out of range, in particular in the presence of particles. Also, the radiosonde data does not indicate any enhancement of water vapour in that region (Fig. 6).

In the condensed phase, the Raman lines of H$_2$O are widened to a rather broad band and shifted toward smaller wavelengths. The Raman spectrum of liquid cloud droplets overlaps with the spectrum of water vapour and thus with the transmission spectra of the filter in the 407 nm channel. Therefore, water clouds may interfere with Raman lidar measurements of water vapour (Melfi et al., 1997). However, there is strong evidence against the interpretation that the inelastic signals that we have measured were created by Raman scattering from water. First, the depolarisation shown in Fig. 6a indicates that the particles of the plume are solid and the Raman spectrum of ice does not overlap with the transmission of the filter we use in the 407 nm channel. Second, the Raman signal from condensed water is far to weak to be detected by our system. The size distribution retrieved from the elastic lidar backscatter as discussed above has a Volume density on the order of 10$^3$ cm$^{-3}$. If those particles were liquid water droplets the Raman backscattering coefficient would still be about 3 orders of magnitude below our detection limit even if we apply all the enhancement factors discussed by Melfi et al. (1997). In fact, we do not see any significant inelastic scattering from clouds, neither solid nor liquid, with our instrument. This is confirmed in Fig. 5c where the ice particles of the cirrus cloud at the tropopause, directly below the aerosol layer, do not create a detectable inelastic signal in the 407 nm channel, even though its elastic backscatter with a value of approximately 30 is by more than a factor of 10 higher compared to that of the aerosol layer above. This profile shows clearly, that neither cross talk from strong elastic backscatter, nor inelastic scattering from ice is measured by our system. We therefore came to
the conclusion that the observed inelastic backscatter signal is a specific feature of the stratospheric aerosol layer and is not caused by water, neither liquid nor solid nor in the gas phase.

3.1 Portuguese forest fire plumes

While searching for other occasions where the water vapour Raman channel shows unexpected behaviour, we found a significant discrepancy between water vapour profiles measured with radiosonde and lidar during a number of consecutive days in early August 2003. A fairly strong layer of aerosol was present in these measurements throughout the troposphere with backscatter ratios up to 3 and the water vapour mixing ratio retrieved from the lidar data showed a systematic deviation from the moisture measurements of the radiosonde (Fig. 7b). During this period heavy forest fires were reported in Portugal. The backward trajectories shown in Fig. 8 give a clear indication, that the aerosol we detected in Lindenberg were smoke plumes from these forest fires.

Figure 7 shows on the right side the difference detected between the water vapour mixing ratio retrieved from the lidar data and those from the radiosonde. The error bars indicate the total error (including a 5% uncertainty introduced by the calibration) and show that there is a significant positive bias in the lidar data. The differences between the lidar and the radiosonde data reach 1 g/kg and are therefore much larger than the usual discrepancies (compare with Fig. 2). This bias correlates with the concentration of smoke aerosol that is displayed in green in terms of the elastic backscatter signal measured with the lidar. This correlation suggests, that the aerosol particles itself create the bias by inelastic scattering. The most likely explanation for this finding is laser induced fluorescence from aerosol particles.
4 Discussion

4.1 Fluorescence in the atmosphere

There is a long tradition of using laser induced fluorescence for lidar remote sensing of the atmosphere. Hydroxyl (OH) and metal atoms such as sodium, potassium or iron have been detected by resonant fluorescence in the mesosphere in 85 km altitude and allow the determination of the temperature and wind at these altitudes (Neuber et al., 1988; von Zahn and Höflner, 1996; Brinksma et al., 1998). In the lower atmosphere however, fluorescence by atoms or molecules in the gas phase is severely hampered by collisional quenching of the exited states due to the high pressure. On the other hand, fluorescence of mainly organic substances still allow interesting applications near or at the earth’s surface. For instance, contamination of soils with petroleum products is detected in situ with high sensitivity using fluorescence (Schultze et al., 2004) and fluorescence lidars are used for remotely detecting oil spills, phytoplankton and chlorophyll on the surface of the sea (Cecchi et al., 2003).

The investigation of fluorescence from polycyclic aromatic hydrocarbons adsorbed on suspended particles was pioneered by Allegrini and Omenetto (1979) and the group of Niessner et al. (1991). The fluorescence spectra of Benzo[a]pyrene (BaP, C_{20}H_{12}) in water for example, shows a wide band covering the 407 nm region when excited at 355 nm (Fernández-Sánchez et al., 2003) and therefore could in principal be detected with the set-up used by our lidar. BaP is an ubiquitous product of incomplete combustion and is stable in the atmosphere. BaP and other PAHs adsorbed on soot in a smoke plume are known to persist for many days in the atmosphere (Behymer and Hites, 1988). In polluted air BaP was found in amounts of 200 pg/m$^{-3}$ (Schauer et al., 2004). The presence of BaP and other PAHs in cigarette smoke is not only one of the reasons for the carcinogenic effect of smoking, they are also the cause for the fluorescence exhibited by the smoke of tobacco leaves (Hill et al., 1999). This is one example for a biomass burning aerosol that fluoresces when irradiated with UV light.

A similar fluorescence spectrum such as the one from BaP was measured from bacteria and fungal spores (Hill et al., 1999). The detection and discrimination of biological particles in the atmosphere by their intrinsic fluorescence has received greater recognition in the recent years (Hairston et al., 1997). Besides in-situ detection, the remote detection of disseminated spores by a fluorescence lidar up to a distances of several kilometres have been reported (Christesen et al., 2004).

Another type of particles that shows fluorescence are sulphuric acid droplets which are contaminated with organic compounds. Hegglin et al. (2002) reported that they observed broad fluorescence spectra from single levitated sulphuric acid droplets that contained aliphatic hydrocarbon, alcohols, aldehydes, ketones or organic acids and conclude that ambient aerosol which often contain significant amount of these organics should fluoresce and thereby offer an interesting experimental possibility to identify biogenic material in aqueous droplets.

In considering these observations, we arrived at the conclusion that different types of biogenic material may have the potential to fluoresce at atmospheric conditions when excited with a laser beam like the one used in our lidar. But is it possible to detect such a signal from the lowermost stratosphere with the 407 nm channel?

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Fig. 7. Same as Fig. 2 measured on 5 August 2003, 22:38 UT.

Fig. 8. 3-day-backward trajectories for Lindenberg, 6 August 2003, 00:00 UT at about which time the measurement of Fig. 7 was made. The different colours mark different altitude levels at the trajectories’ end points as indicated in the legend.
4.2 Detecting fluorescence of atmospheric aerosol with a lidar

To estimate the sensitivity of our instrument regarding the detection of fluorescence, we calculated the backscatter coefficient at 407 nm $\beta_{407}(z)$ within the stratospheric aerosol layer. Based on the water vapour calibration discussed earlier, the backscatter coefficient measured with 407 nm channel can be calculated from Eq. (3) using the known scattering cross section $\sigma_{H_2O}$ of the H$_2$O Raman line at 3657 cm$^{-1}$ which is $8 \times 10^{-30}$ cm$^2$sr$^{-1}$ (Melfi et al., 1997). Accordingly, the peak at 13 km (Fig. 6b) corresponds to a backscatter coefficient in the order of $10^{-15}$ cm$^{-2}$sr$^{-1}$.

Unfortunately, little is reported about the cross sections of fluorophores like PAH on aerosol particles. Also, the efficiency of such a process in the atmosphere depends on many parameters including temperature and pressure hampering an assessment of the possibility of detecting the fluorescence from these agents by our lidar. However, measurements of cross sections of the fluorescence of bacteria and pollen are available, a value of $5 \times 10^{-14}$ cm$^2$nm$^{-1}$particle$^{-1}$ for Bacillus globigii (Bg) and $2 \times 10^{-13}$ cm$^2$nm$^{-1}$particle$^{-1}$ for pine pollen when excited with light at 230 nm are reported by Stephens (1999). Given a scattering cross section in this order of magnitude, a concentration of fluorescent particles of about 0.1 cm$^{-3}$ is calculated from the measured backscattered coefficient using Eq. (3). Given the concentration of particles estimated from the elastic backscatter which was about 1000 cm$^{-3}$ this means that one out of $10^4$ particles in the plume should fluoresce in order to explain the inelastic signal that we have measured. The concentration of bacteria in ambient air is 0.001 to 0.01 cm$^{-3}$ (Ariya and Amyot, 2003) and slightly below our limit of detection. An enhancement of bioparticles that may fluoresce is likely to occur in the plume of a forest fire. However, soot particles that fluoresce due to its content of PAH should be a lot more abundant than bioparticles and therefore might play a more important role for creating the fluorescence signal.

Hegglin et al. (2002) reported that the fluorescence signal of sulphuric acid droplets that are contaminated with organic material is 2–3 orders of magnitude higher than the Raman signal from the water stretching vibration that they were trying to measure in the first place. We discussed above, that the Raman signal from aqueous aerosol particles of an amount as inferred from the elastic lidar signals is about 3 orders of magnitude to weak to be detected with our lidar. Accordingly, the fluorescence of organic material could be strong enough to be measurable. The fluorescence of contaminated sulphuric acid droplets was reported to increase as the concentration increases and the aerosol ages (Hegglin et al., 2002). Therefore, in case of a dried and aged layer of particles in the lowermost stratosphere, the efficiency of fluorescence compared to Raman scattering should be particularly large.

This considerations demonstrate, that fluorescence should be taken into account when analysing Raman lidar signals in the presence of polluted air. Even though the information that is currently available is to scarce to give a detailed quantitative analysis, we are confident that fluorescence from organic material is the prime candidate to explain the inelastic scattering that we have detected from aerosol plumes in the troposphere and the lower stratosphere. The powerful laser, the large receiving telescope (1.1 m), and the large bandwidth (5.7 nm) and good performance of the 407 nm channel makes our system particularly suitable for detecting laser induced fluorescence from particles in the atmosphere. The decay time of the fluorescence signals of biologic material is on the order of 100 ns (Niessner et al., 1991). In lidar measurements this delay would be interpreted as an altitude shift of about 15 m. The maximum vertical resolution of 7.5 m of the lidar and the naturally rather diffuse vertical structure of aerosol layers, leads us to the conclusion that the finite lifetime of the excited states of fluorophores has no discernible effect on the lidar measurements.

4.3 Biomass burning aerosol in the stratosphere

According to our investigation, the only agents capable to create the fluorescence that can interfere with the water vapour measurements of the lidar are organic compounds. The only case, where we have found tropospheric aerosol that creates such an interference is smoke from forest fires in Portugal. These facts lead us to the conclusion that the detection of fluorescence indicates aerosol of biogenic origin. Accordingly, the aerosol layer that we have observed in the lowermost stratosphere from 1 June 2003 to 3 June 2003 is must be contaminated with organic pollutants that came from a biomass burning event or something similar.

At that particular time, forest fires were abundant across the northern hemisphere, including an unprecedented 19 million hectares of boreal forest burned in Siberia (Damoah et al., 2004). Backward trajectories trace the air mass that we observed in Lindenberg back to the Pacific region where it obviously experienced convection (Fig. 4). Fromm et al. (2000) have suggested that extreme convection triggered by forest fires may be able to inject aerosol into the stratosphere at high latitudes. The origins of the trajectories in (Fig. 4) in the Pacific are detached from potential continental sources in Siberia and therefore do not support the idea of a direct injection of the forest fire smoke from thunderstorms created by the fires themselves. At that particular time a tropical storm, the Typhoon Chan Hom was active in that region and might have created a TST event that included polluted air from Asia (Fig. 4). A detailed model study of the history of the air mass and the possibility of TST events is beyond the scope of this work. However, it is conceivable that a fraction of the smoke from Siberian forest fires which has affected almost the entire northern hemisphere at that time (Damoah et al., 2004) emerged into the lower stratosphere. The fact that the elastic
optical properties of this aerosol are those of an aerosol containing soot supports this hypothesis.

5 Conclusions

During a measuring campaign in Lindenberg/Germany we have detected inelastic scattering from an aerosol layer in the lowermost stratosphere. We showed that Raman scattering by water is not efficient enough to explain the observation and we are confident that the signal is not an instrumental artefact. Additionally, we presented results from water vapour measurements from August 2003 with our Raman lidar that show that aerosol particles related to Portuguese forest fires exhibit inelastic scattering. In the literature numerous reports describe the fluorescence of particles that are likely to be present in aerosol released by biomass burning. Fluorescence spectra of such particles are detectable by our lidar set-up. All these examples have in common, that organic compounds need to be present to create the fluorescence. Therefore, we conclude that atmospheric aerosol of biogenic origin may fluoresce when excited with a laser and that this fluorescence interfered with the water vapour measurement of the 407 nm Raman channel. The detection of fluorescence from atmospheric aerosol by a lidar has not yet been reported anywhere in literature, however, it is the most plausible explanation for our experimental results.

We interpret the fluorescence that we detected from aerosol layers as an indicator for particles created by biomass burning because both the events that we discussed were strongly linked to forest fires. It is still open which organic compounds could have caused the fluorescence and if particles from other sources may have similar properties. From the observations presented it is not possible to discriminate soot from contaminated sulphate, bioparticles or other possible fluorophores. Future lidar experiments should involve the detection of inelastic scattering at different wavelengths that are off the Raman line of water vapour. Such a set-up could discriminate fluorescence from water vapour Raman scattering without the need for additional measurements and therefore allows a unique identification of fluorescence of aerosol on the one hand and correction or rejection of the Raman lidar water vapour measurement due to fluorescence on the other hand. The measurements of vertically resolved fluorescence spectra of aerosol could become a helpful method for determining the type of particles. Additionally, quantitative laboratory measurements of the efficiencies of laser induced fluorescence created by aerosol particles of different types at different atmospheric conditions are necessary for a more detailed analysis of fluorescence lidar data.

The possibility of detecting fluorescence from atmospheric particles by lidar opens up new opportunities for remote sensing of the distribution and the composition of aerosols in the atmosphere and could therefore overcome some of the shortcomings of current techniques. These data are important prerequisites to assess the climatic impact of these types of anthropogenic aerosol, which is currently still debated (IPCC, 2001). Lidar techniques based on fluorescence detection could become a powerful tool to study the abundance of such aerosol as well as exchange processes between the upper troposphere and the lower stratosphere. Our observation of an aerosol layer lasting for three days above the tropopause, shows an interesting, rarely detected example of such an event in the tropopause region.

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References


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F. Immler et al.: Fluorescing aerosol


