The Pagami Creek smoke plume after long-range transport to the upper troposphere over Europe – aerosol properties and black carbon mixing state

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Abstract. During the CONCERT 2011 field experiment with the DLR research aircraft Falcon, an enhanced aerosol layer with particle linear depolarization ratios of 6–8 % at 532 nm was observed at altitudes above 10 km over northeast Germany on 16 September 2011. Dispersion simulations with HYSPILT suggest that the elevated aerosol layer originated from the Pagami Creek forest fire in Minnesota, USA, which caused pyro-convective uplift of particles and gases. The 3–4 day-old smoke plume had high total refractory black carbon (rBC) mass concentrations of 0.03–0.35 µg m\(^{-3}\) at standard temperature and pressure (STP) with rBC mass equivalent diameter predominantly smaller than 130 nm. Assuming a core-shell particle structure, the BC cores exhibit very thick (median: 105–136 nm) BC-free coatings. A large fraction of the BC-containing particles disintegrated into a BC-free fragment and a BC fragment while passing through the laser beam of the Single Particle Soot Photometer (SP2). In this study, the disintegration is a result of very thick coatings around the BC cores. This is in contrast to a previous study in a forest-fire plume, where it was hypothesized to be a result of BC cores being attached to a BC-free particle. For the high-altitude forest-fire aerosol layer observed in this study, increased mass specific light-absorption cross sections of BC can be expected due to the very thin coatings around the BC cores, while this would not be the case for the attached-type morphology. We estimate the BC mass import from the Pagami Creek forest fire into the upper troposphere/lower stratosphere (UTLS) region (best estimate: 25 Mg rBC). A comparison to black carbon emission rates from aviation underlines the importance of pyro-convection on the BC load in the UTLS region. Our study provides detailed information on the microphysics and the mixing state of BC in the forest-fire aerosol layer in the upper troposphere that can be used to better understand and investigate the radiative impact of such upper tropospheric aerosol layers.

1 Introduction

Light-absorbing aerosol species, such as black carbon (BC), contribute to a large part to the positive radiative forcing of aerosols (Jacobson, 2001). Recent modeling studies suggest that BC is the second strongest source to current global warming after CO\(_2\) (Ramanathan and Carmichael, 2008) because it is the dominant solar-radiation-absorbing component in the visible spectrum in the atmosphere. A comprehensive scientific assessment of the role of BC in the climate system is given in Bond et al. (2013). The terminology used in the literature for carbonaceous particles from combustion emissions is often ambiguous (Petzold et al., 2013). Here we...
use the term “BC” exclusively for the most refractory and light-absorbing component of carbonaceous combustion particles, which is essentially pure carbon, not including organic carbon, brown carbon or inorganic aerosol components. The term “BC-containing” particle is used for referring to combustion particles with some BC content. A particle is classified as a “BC-free” particle, if the Single Particle Soot Photometer (SP2) does not detect any BC in the particle, which does not exclude the presence of a small amount of BC below the lower detection limit of the SP2 (∼0.5 fg BC in a particle). Furthermore, we use the more specific term refractory black carbon (rBC; see Petzold et al., 2013), whenever we refer to quantitative measurements of the rBC mass (or quantities inferred from rBC mass measurements) with the SP2 (Schwarz et al., 2006).

Besides BC, other light-absorbing materials such as brown carbon, a light-absorbing organic carbon (OC), can contribute significantly to light absorption (e.g. Andreae and Gelencser, 2006; Moosmüller et al., 2009). The largest source of primary carbonaceous fine particles in the global troposphere is anthropogenically- and naturally-occurring biomass burning defined as the sum of all open natural and anthropogenic combustion processes using non-fossilized vegetative or organic fuels (e.g. Akagi et al., 2011). Biomass burning has a major impact on the radiation budget of the atmosphere (Knorr et al., 2012). Diehl et al. (2012) estimate the global BC emissions from biomass burning as 1.8–7.0 Tg per year, while OC emissions range from 14–57 Tg per year. Biomass-burning emissions can be transported to the free troposphere and even into the stratosphere by, for example, pyro-convection, as it was observed for a North American forest fire by, for example, Jost et al. (2004).1 There elevated aerosol layers may form. Biomass-burning aerosol layers are often transported over long distances on hemispheric scales (e.g., Damoah et al., 2004; Mattis et al., 2008; Ramanathan and Feng, 2009; Petzold et al., 2007). In this study we define an elevated aerosol layer as a vertically extended layer situated above the planetary boundary layer, containing aerosol particles. Such long-range transport of biomass-burning emissions was previously measured and reported by, for example, Fiebig et al. (2003), Petzold et al. (2007), Real et al. (2009), Quennehen et al. (2011), Weinzierl et al. (2011) and Quennehen et al. (2012). Barnaba et al. (2011) estimate for the 6 year period, 2002–2007, that up to 35 % (maxima in April and in summer) of the aerosol optical thickness of the European fine particle fraction \(D_{\text{opt}} < 1 \mu m\) is attributable to wildfire emissions, including long-range transport from North America. Based on ten years of ground-based multi-wavelength Raman lidar (light detection and ranging) observations, Mattis et al. (2008) observed free-tropospheric aerosol layers above the continental boundary layer of central Europe in 43 % of their measurements, with a maximum from June to August. In 19 % of their measurements, they found forest-fire smoke advected from Canada and anthropogenic pollution advected from urban areas of the USA and Canada.

During transport, the biomass-burning plumes can mix with other aerosols (e.g., Ramanathan and Carmichael, 2008; Tesche et al., 2009; Petzold et al., 2011; Weinzierl et al., 2011). Depending on the composition and altitude, and albedo of the underlying surface, these layers can impact the radiative budget and may have cooling or warming effects on different parts of the atmosphere. In particular, BC in elevated aerosol layers absorbs incoming solar radiation effectively and transforms it into thermal radiation (Moosmüller et al., 2009). In this context, the mixing state of BC-containing particles is important because coatings of light-scattering material can enhance the absorption of the BC core due to the so-called “lensing effect” and can thus amplify the heating potential of BC-containing aerosol particles on the surrounding air (Bauer et al., 2010; Shiraiwa et al., 2010). Enhancements of the absorption by at least 30 % (Schwarz et al., 2008b) up to 100 % have been calculated for thickly coated particles with an optical particle diameter to core diameter \(D_{\text{opt}}/D_{\text{BC}}\) ratio of 2 (Shiraiwa et al., 2010). However, Cappa et al. (2012) report a significant difference in the coatings’ absorption enhancement in ambient observations versus laboratory measurements and model results. Thus, the understanding of the absorption enhancement and the radiative forcing by mixed atmospheric BC is still incomplete and needs further investigation. Aside from this idealized internal shell-core mixture of particles, more complex structures of BC particles and BC-free materials exist in nature (e.g., Sedlacek et al., 2012). Semi-external mixtures such as particles attached to each other or near-surface BC cores can be observed in the atmosphere as well (Mishchenko et al., 2004; Liu and Mishchenko, 2007). More information about particle mixing is given in Buseck and Posfai (1999), Posfai et al. (2003), and Posfai et al. (2004).

Although complicated mixing structures are observed in nature, most modeling studies assume homogeneously mixed or idealized core-shell morphologies as a basis for radiative transfer calculations. Only some recent model developments implement the particle mixing state (e.g., Aquila et al., 2011). In addition to a changed impact on the radiative aerosol properties, the structure of mixed particles, for example, internal mixing of a BC core with organic or inorganic aerosol components, might have an influence on the activation of BC particles to form cloud droplets in warmer regimes, possibly leading to an increased BC wet deposition and a related decrease in BC residence times. Furthermore, the ice-nucleation efficiency of the particle and thus the associated indirect climate effect of these aerosols may be impacted (Hoos and Möhler, 2012).

Although recently more studies focus on airborne in situ measurements of the vertical distribution of BC in the

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1In this study, the general term “biomass-burning aerosol” is used if utilized in referenced studies while “forest-fire aerosol” is used for the case treated in this study.
troposphere (e.g., McMeeking et al., 2010; Schwarz et al., 2010), so far, processes concerning light-absorbing aerosol layers and their impact on the radiation budget are only insufficiently quantified and characterized. One reason for this deficiency is the sparseness of airborne in situ measurements in elevated aerosol layers. In this study, we present airborne in situ data of elevated aerosol layers observed with the Deutsches Zentrum für Luft- und Raumfahrt (DLR) research aircraft Falcon-20E in the framework of Contrail and Cirrus Experiment 2011 (CONCERT 2011) over central Europe in September 2011 (Voigt et al., 2014). In particular, we focus on the investigation of an elevated forest-fire aerosol layer observed on 16 September 2011 over northeastern Germany, originating from the Pagami Creek forest-fire in North America.

The present study is organized as follows: in Sect. 2, we give an overview of the CONCERT research flights and describe the instrument calibration and data analysis procedures with focus on the SP2. In Sect. 3, we show the results of our measurements in the Pagami Creek forest fire including a detailed analysis of microphysical and optical aerosol properties, vertical and horizontal extension of the Pagami Creek plume and use dispersion simulations with HYSPLIT for source attribution. In Sect. 4, we compare the observations made on 16 September 2011 with other measurements in elevated aerosol layers during CONCERT 2011 and with the literature. Furthermore, we discuss the morphology and potential disintegration of the observed rBC particles on the basis of our SP2 measurements. Finally, we present an estimate on the black carbon mass import into the UTLS by the Pagami Creek fire and compare it with the BC import into the UTLS by aviation.

2 Material and methods

2.1 Research flights and instrumentation

The main focus of the CONCERT 2011 mission, the follow-up mission of CONCERT 2008 (Voigt et al., 2010), was the measurement of microphysical and optical properties of contrail cirrus (Voigt et al., 2011; Schumann et al., 2011a) and natural cirrus in order to investigate their climate impact (Kübbeler et al., 2011; Schumann et al., 2013; Jeßberger et al., 2013). Here we focus on the serendipitous observation of an enhanced aerosol layer in the upper troposphere/lower stratosphere (UTLS) region, originating from the Pagami Creek forest fire in Minnesota, USA. During CONCERT 2011, the DLR Falcon research aircraft was based in Oberpfaffenhofen, Germany, and performed 11 research flights between 7 September and 30 September 2011. Most of the flights were conducted over Germany, with additional landings in Norway and Italy. Instruments from DLR and the Universities of Mainz and Clermont-Ferrand were operated onboard the DLR Falcon. The following basic aerosol parameters were measured: particle size distribution, rBC mass concentration, rBC core size distribution, mixing state of rBC-free and rBC-containing particles, light scattering coefficient and light absorption coefficient. In addition, instruments for trace gases were deployed on the Falcon. Most of them are described in Voigt et al. (2010) and the references therein. In the following, we only describe the instruments used in this study.

In the past decades BC measurements in the atmosphere were conducted predominantly with filter-based instruments. In this study, the more recently introduced Single Particle Soot Photometer (SP2; Stephens et al., 2003) was applied to obtain information about the rBC mass concentration as well as the mixing state and coating thickness of rBC-containing particles. The SP2 determines the optical size and the rBC mass of single rBC-containing particles, using light-scattering and laser-induced incandescence methods, respectively (Schwarz et al., 2006). More details about the functional principles, calibration and data analysis approaches of the SP2 will be summarized in Sect. 2.2.

To obtain information on the entire aerosol size distribution, measurements from five condensation particle counters (two modified TSI Condensation Particle Counters (CPC; models 3010 and 3760A) and three Condensation Particle Size Analyzers (CPSA; built by DLR) set to different lower cut-off diameters (4 nm, 10 nm, 14 nm)) are combined with optical sizing measurements from the SP2 and a wing-mounted Forward Scattering Spectrometer Probe (FSSP-300). One of the CPCs was operated on 16 September behind a thermal denuder, heating the aerosol to 255°C to measure non-volatile particles. Table 1 summarizes the aerosol parameters and the size ranges covered by the different aerosol instruments used in this study.

2.2 Instrument calibration and data analysis

In the SP2, a concentric-nozzle jet system is used to direct the aerosol sample flow through a high-intensity continuous-wave intra-cavity laser beam (λ = 1064 nm) and four different detectors are used to characterize individual particles with 100 % detection efficiency within the respective detection limits (see Table 1). The response curves of four detectors for each particle are recorded for 44 μs with a temporal resolution of 0.2 μs.

A broadband incandescence detector (BID) and a narrowband incandescence detector (NID), both standard detectors implemented in the SP2 by the manufacturer, are used to detect the thermal radiation emitted by light-absorbing and refractory particles heated to incandescence in the 300–800 nm and 630–800 nm wavelengths bands, respectively. The peak intensity of the thermal radiation is typically reached at the vaporization temperature of the most refractory particle component. The incandescence signals from atmospheric aerosol particles normally originate from rBC and the peak height of the incandescence signal is proportional to the rBC mass.
Table 1. Size ranges of the aerosol instrumentation onboard the Falcon aircraft considered in this study during the flights on 16 September 2011. The first column displays the name of the instrument, the second column the measured property, and the third column the respective size range. CPC, Condensation Particle Counter; CPSA, Condensation Particle Size Analyzer; SP2, Single Particle Soot Photometer; LSD, light scattering detector; BID, broadband incandescence detector; NID, narrowband incandescence detector; PSD, position sensitive detector; FSSP 300, Forward Scattering Spectrometer Probe.

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Property</th>
<th>Size/mass range</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPC/CPSA</td>
<td>integral number concentration(^1) of ultrafine particles</td>
<td>0.004 µm &lt; (D_{\text{opt}}) &lt; (D_{\text{cutoff}})</td>
</tr>
<tr>
<td>SP2</td>
<td>LSD</td>
<td></td>
</tr>
<tr>
<td></td>
<td>single particle optical diameter of rBC-free particles</td>
<td>0.14 µm &lt; (D_{\text{opt}}) &lt; 0.29 µm</td>
</tr>
<tr>
<td></td>
<td>ambient number size distribution of rBC-free particles</td>
<td>0.14 µm &lt; (D_{\text{opt}}) &lt; 0.29 µm</td>
</tr>
<tr>
<td></td>
<td>BID, NID</td>
<td></td>
</tr>
<tr>
<td></td>
<td>single particle rBC mass</td>
<td>0.5 fg &lt; (m_{\text{rBC}}) &lt; 80.3 fg</td>
</tr>
<tr>
<td></td>
<td>ambient rBC mass size distribution</td>
<td>0.08 µm &lt; (D_{\text{rBC}}) &lt; 0.44 µm</td>
</tr>
<tr>
<td></td>
<td>LSD, BID, NID, PSD</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Leading-edge-only (LEO) fit:</td>
<td></td>
</tr>
<tr>
<td></td>
<td>single particle optical diameter of rBC-free and rBC-containing particles</td>
<td></td>
</tr>
<tr>
<td></td>
<td>coating thickness of rBC-containing particles</td>
<td>(\sim 0.15 \mu m &lt; D_{\text{opt}} &lt; \sim 0.55 \mu m) (cut-offs dependent on rBC fraction)</td>
</tr>
<tr>
<td>FSSP-300</td>
<td>ambient size distribution(^2)</td>
<td>nominal size range</td>
</tr>
<tr>
<td>Falcon standard</td>
<td>(T, \rho, \text{RH}, \text{3-D-wind velocity})</td>
<td>0.3 µm &lt; (D_{\text{opt}}) &lt; 30 µm</td>
</tr>
<tr>
<td>instrumentation</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^1\) Aerosol in situ instruments operated inside the cabin of the Falcon behind an isokinetic inlet. The inlet system has an upper cut-off diameter \(D_{\text{cutoff}}\) where 50% of the particles are carried into the isokinetic inlet, depending on air speed and outside pressure. For typical Falcon speed, the cut-off decreases from roughly 2.5 µm at ground level to roughly 1.5 µm at 10 km altitude (Fiebig, 2001; Wendisch et al., 2004).

\(^2\) Aerosol in situ instruments sampling at wing stations.
sensitive detector (PSD), a modified multi-element LSD, which makes it possible to determine the scattering cross-section of rBC-free as well as rBC-containing particles at any time (or position) in the laser beam. The approach for the time-resolved scattering cross-section analysis applied here is largely based on Gao’s method, described in detail in Laborde et al. (2012a). The scattering cross-section of the unperturbed particle is obtained from the LSD signal in the leading edge of the laser beam (up to 3% of maximal laser intensity) before the onset of coating evaporation. This is commonly referred to as the leading-edge-only (LEO) fit method (Gao et al., 2007). The LEO-fit extends the upper detection limit for the optical sizing of rBC-free particles to larger diameters, in our case up to around 550 nm (Table 1). Similar to the standard optical sizing (see above), the optical diameter of rBC-free particles is inferred from the scattering cross-section measured with the LEO-fit approach assuming a homogeneous sphere and a refractive index of 1.59 + 0.00i (see below). The uncertainty of the optical sizing associated with uncertainties of the assumed refractive index is about 5% (±10 nm) for a 200 nm particle and about 8% (±50 nm) for a 600 nm particle when varying the refractive index from 1.59 + 0.00i to 1.50 + 0.00i or 1.70 + 0.00i. Inferring the optical size of rBC-containing particles from their scattering cross-section is not straightforward as the particles are inhomogeneous and typically not spherical. We use the approach described in Schwarz et al. (2008b) and Laborde et al. (2012b). Briefly, a concentric-spheres core-shell morphology is assumed for the Mie calculations. The core size is constrained by the rBC core mass equivalent diameter, $D_{BC}$, from the incandescence measurement. To obtain the total optical diameter, $D_{opt}$, of the unperturbed particle, including core and shell, the diameter of the total particle is varied until the calculated scattering cross-section matches the measured scattering cross-section. The coating thickness ($\Delta R_{coat}$) of a BC-containing particle is defined as the difference between the radii of the unperturbed particle and the BC core, i.e., $\Delta R_{coat} = 0.5(D_{opt} - D_{BC})$. This provides information about the mixing state of individual BC-containing particles. In this study, refractive indices of 2.26 + 0.00i and 1.59 + 0.00i are used for the rBC core and the coating, respectively, in the Mie calculations. The particular value for the refractive index of the BC core was measured by Moteki et al. (2010) for thermally denuded ambient BC particles using an APM (Aerosol Particle Mass Analyzer) and an SP2.

The reliability of the SP2 has been studied extensively by inter-comparisons of six different SP2s at the Aerosol Interaction and Dynamics in the Atmosphere (AIDA) chamber in November 2011. Laborde et al. (2012b) report a reproducibility of 10% for the rBC mass and 17% for the coating thicknesses for this inter-comparison study. However, due to calibration uncertainties the absolute uncertainty may be higher. Kondo et al. (2011b) compared rBC mass concentrations measured with the SP2 and a filter-based absorption photometer, continuous soot-monitoring system (COSMOS) and found that the methods agree to within about 10% on average. Besides, Laborde et al. (2013) made a comparison of the rBC measurements by the SP2 with a collocated elemental carbon (EC) measurement by a thermal-optical EC instrument (Sunset ECOC analyzer), revealing agreement well within the uncertainty of either method. The SP2 was prepared for the research flights, following the instructions given in Laborde et al. (2012b). For this study we estimate a total uncertainty of 15% for the rBC mass, which includes the described reproducibility and calibration uncertainties (Laborde et al., 2012b) combined with uncertainties due to airborne in situ measurements.
The aerosol inlet system on the DLR Falcon research aircraft has been extensively tested and successfully used for years. For example, Fiebig (2001) investigated the inlet system on the Falcon and Schumann et al. (2011b) show the cut-off of the Falcon aircraft inlet and the effect on the derived particle size distribution.

To obtain particle number size distributions in the size range of ~0.15–30 µm, measurements from the SP2 and the FSSP-300 are combined. To ensure a correct signal-to-diameter assignment for the optical particle counters such as the FSSP-300 ($\lambda = 633$ nm), an appropriate refractive index has to be applied for the bulk aerosol. Problems with this procedure are explained in more detail in Weinzierl et al. (2009), for example. Real parts of the refractive index at visible wavelengths of several types of organic aerosols and biomass-burning organic aerosol are reported to be between 1.50 and 1.63 and between 1.56 and 1.64, respectively (Kondo et al., 2011a, and references therein). However, the FSSP-300 cannot distinguish between BC-free and BC-containing particles. Thus, the particles in the considered FSSP-300 size range are assumed to be non-absorbing (the imaginary part of the refractive index is zero), with real parts of the bulk refractive index ranging between 1.59 and 1.60. That increases the uncertainty of the FSSP-300 data compared to the SP2 size distribution, which distinguishes between BC-containing and BC-free particles. However, due to a low fraction of BC-containing particles in the data set presented later (8–13 %) we expect only minor effects of the refractive index assumption on the inferred particle size distribution. The combination of the FSSP-300 and the SP2 data allows us to derive the particle number size distribution in the size range between about 0.14 µm and 30 µm (Table 1). To obtain information on the size distribution in the Aitken mode ($D_{\text{opt}} < 0.14$ µm) data from the CPC/CPSA is added. The number concentration of Aitken mode particles in the size range between 0.01 µm (lower CPC cut-off) and 0.14 µm (lower SP2 cut-off) is determined as the differential of particles in the accumulation mode, i.e., measured with the SP2 and the FSSP-300 and the integral number concentration determined by the CPC/CPSA. This method is only valid if nucleation mode particles can be excluded. This is ensured by the measurement of particles in the size range 10–14 nm, showing low particle number concentrations (< 260 cm$^{-3}$). The shape of the particle number size distribution in the Aitken mode range is approximated by applying a log-normal fit, assuming that a single log-normal mode represents all particles in the Aitken mode (Weinzierl et al., 2011) with the particular minimum between Aitken and accumulation mode being operationally defined as 0.14 µm. Figure 1b sketches the procedure for fitting the number size distributions. The parameters for a log-normal fit of mode $j$ are the mode number concentration ($N_j$), the geometric standard deviation ($\text{GSD}_j$), and the count median diameter ($\text{CMD}_j$). The CMD for the Aitken mode, $\text{CMD}_1$, is set constant to 0.07 µm, and $\text{GSD}_1$ is estimated to provide the best fit to the residual between the measured accumulation mode and the log-normal fit to the accumulation mode in the lower end of the measurement range (of SP2/FSSP-300) and to match values regularly found for forest-fire aerosol layers (see Sect. 4.2). In addition to one log-normal distribution used to describe the size distribution in the Aitken mode, one or two log-normal distributions are applied to parameterize the particle size distribution in the accumulation mode and the coarse mode, i.e., the measured size range of the SP2 and
FSSP-300. Because no considerable number fraction of particles was measured in the coarse mode, no separate coarse mode fit was applied.

Unless stated otherwise, the data presented in this study refer to standard temperature and pressure (STP; 273.15 K, 1013.25 hPa), corresponding to mixing ratios. STP concentration data can be converted to ambient conditions by applying a factor \( f_{\text{STP}} \) indicated in Table 2.

To determine the origin and the dispersion of the investigated aerosol, we use the Hybrid Single Particle Lagrangian Integrated Trajectory Model HYSPLIT (Draxler and Hess, 1997, 1998). Input for dispersion calculations is the location of the forest fire and the altitude range of the emission. For the HYSPLIT dispersion runs the following assumptions are made: the dispersion is calculated for 0.5° grid cells for 3 h time steps. All particles are set to a diameter of 1 µm and a particle density of 1 g cm\(^{-3}\). The particle deposition velocity is assumed to be 0.1 mm s\(^{-1}\) for all particles. Wet deposition is not considered for this qualitative approach. HYSPLIT is initialized with meteorological data from the European Centre for Medium-Range Weather Forecasts (ECMWF) and the Global Forecast System (GFS). In this study only the results for the ECMWF data are presented because no relevant differences are observed for the different meteorological data sets.

### 3 Results from the measurements in the Pagami Creek forest-fire plume

On 16 September 2011, two flights were performed with the DLR Falcon. For the first flight (#110916a, 10:48–13:20 UTC), the Falcon flew from Oberpfaffenhofen (EDMO; 48°05′ N, 11°16′ E, 593 m a.s.l.) to Neubrandenburg-Trollenhagen (ETNU; 53°36′ N, 13°18′ E, 69 m a.s.l.), both in Germany. The second flight (#110916b, 14:54–17:48 UTC) was from Neubrandenburg-Trollenhagen back to Oberpfaffenhofen. The scope of these two flights was to sample contrails from commercial aircraft. To achieve this, the aircraft climbed to more than 10 km altitude and stayed between 10.0 km and 11.8 km for roughly 105 min and 95 min during flights #110916a and #110916b, respectively. Figure 2 shows the time series of rBC mass concentration for both flights with elevated rBC mass concentrations at high altitudes (> 10 km a.s.l.) starting at about 11:45 UTC. Averaged 10 s rBC mass concentrations were observed as high as 0.67 µg m\(^{-3}\) (STP). For comparison, maximum rBC mass concentrations in the polluted boundary layer over Munich, averaged for 10 s, showed similar values with roughly 0.70 µg m\(^{-3}\) (STP) on that day. The observed aerosol layer was visible by eye as distinct broad greyish-brownish aerosol layer (Fig. 3) and covered a large area over northeastern Germany. Figure 4 gives an overview of the horizontal distribution of the aerosol layer during the two flights on 16 September 2011 by means of color-coded rBC mass concentrations (size range: 80–440 nm) along the flight track. To emphasize upper tropospheric BC, we show only values for rBC mass concentrations above 9 km a.s.l. In the following we explicitly investigate the chemical, microphysical and optical properties of this vast aerosol layer and we will show that this layer originated from the Pagami Creek fire in North America. In this study we therefore refer to the aerosol layer observed over Germany on 16 September 2011 as Pagami Creek fire aerosol layer or Pagami Creek fire plume. We are aware that the observed layer does not contain the entire emissions of the Pagami Creek fire and smaller layers and plumes might have been transported along other trajectories at other altitudes, which we did not observe and which are consequently not presented in this study.

### 3.1 Microphysical properties of the observed aerosol layer

To derive particle number size distributions and BC mass distributions and to infer BC particle mixing state, we subdivide the measurements in the Pagami Creek fire aerosol layer into sequences with rather homogeneous rBC mass concentrations. Altogether, 14 time sequences are chosen, as shown in Fig. 2, including sequences with maximum concentrations as well as less concentrated sequences within this heterogeneous aerosol layer. Aircraft exhaust and contrail sequences, which are characterized by short and distinct increases of particle and trace gas concentrations, are excluded, using NO\(_3\) and SO\(_2\) as aircraft plume markers.

#### 3.1.1 rBC mass size distributions

Figure 5 shows the median measured rBC mass size distribution and log-normal fits for the measured rBC mass size distributions of the Pagami Creek fire plume of all described
Table 2. rBC characteristics and characteristics of the total particle number size distributions in the Pagami Creek fire plume. Shown are the minimum, median, and maximum values of rBC mass concentrations, the parameters of the log-normal fit to the rBC mass size distribution, the parameters of the log-normal fit to the total number size distribution, and STP-ambient conversion factors of 14 sequences during the forest-fire aerosol layer measurements. Values in italics are extrapolated from total number concentration values. In comparison to the rBC mass concentration in the Pagami Creek fire within the SP2 size range (0.03–0.29 µg m\(^{-3}\)) the free tropospheric background sequence has a mean rBC mass concentration of < 0.004 µg m\(^{-3}\). GSD is the geometric standard deviation and CMD, the count median diameter.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Sequence with rBC mass concentration</th>
<th>minimum</th>
<th>median</th>
<th>maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>rBC mass concentration (in SP2 size range) (STP)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(µg m(^{-3}))</td>
<td>0.03</td>
<td>0.12</td>
<td>0.29</td>
<td></td>
</tr>
<tr>
<td>Parameters of log-normal fit to rBC mass size distribution as function of (D_{rBC}) (STP)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rBC total mass concentration (µg m(^{-3}))</td>
<td>0.03</td>
<td>0.16</td>
<td>0.35</td>
<td></td>
</tr>
<tr>
<td>GSD</td>
<td>–</td>
<td>1.75</td>
<td>1.70</td>
<td>1.70</td>
</tr>
<tr>
<td>CMD</td>
<td>nm</td>
<td>121</td>
<td>144</td>
<td>150</td>
</tr>
<tr>
<td>Parameters of log-normal fit to total particle number size distribution as a function of (D_{opt}) (STP)</td>
<td></td>
<td></td>
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<tr>
<td>(N_1) cm(^{-3})</td>
<td>1077</td>
<td>1517</td>
<td>760</td>
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<tr>
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<td>1.36</td>
<td>1.30</td>
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<tr>
<td>CMD(_1)</td>
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<td>70</td>
</tr>
<tr>
<td>(N_2) cm(^{-3})</td>
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<td>1.41</td>
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<tr>
<td>CMD(_2)</td>
<td>nm</td>
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<td>251</td>
<td>281</td>
</tr>
<tr>
<td>(N_3) cm(^{-3})</td>
<td>1</td>
<td>–</td>
<td>–</td>
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<td>3.93</td>
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</table>

sequences. Error bars represent 15 % uncertainty. The total rBC mass concentration was estimated with adding the extrapolated tails (log-normal fit to the measurements) of the rBC mass size distribution to the rBC measurement of the SP2 integrated over the whole detection range (see Sect. 2.2 for details). The tails of the size distribution consistently contributed ∼ 20 % to the total estimated rBC mass for each of the 14 plume sequences. Due to a higher fraction of fresh and small rBC-particles and possibly even an extra mode of small rBC particles not captured by the log-normal fitting, the determined rBC mass fraction might be lower in the boundary layer, implying a potential systematic uncertainty for the comparison of the determined total rBC mass of the Pagami Creek fire plume and the boundary layer. However, with increasing age the mode of small rBC particles shifts to larger sizes, therefore we expect the log-normal fit to capture the total mass fairly well. The mass median diameter (MMD), which is equal to the modal diameter for a log-normal size distribution, of all mass size distributions of the Pagami Creek fire plume is in the size range of 120 to 160 nm. The total rBC mass concentrations for the 14 sequences (0.03–0.35 µg m\(^{-3}\); STP) reveal a variability factor of approximately 10. A UTLS background sequence at the same altitude as the Pagami Creek fire plume shows rBC average total mass concentrations < 0.01 µg m\(^{-3}\) (STP), implying the Pagami Creek fire plume contains a factor of 3–34 higher rBC mass concentrations than the background air. Parameters of the log-normal fit to the rBC mass size distribution (STP) and the total rBC mass concentrations derived from the log-normal fit (STP) are shown in Table 2.

3.1.2 Total particle number size distributions

Figure 6 shows the total particle number size distributions of the Pagami Creek fire plume in comparison with the UTLS background aerosol sequence. The presented minimum, median, and maximum size distributions represent the range of observations for the 14 considered sequences (Fig. 2) in the Pagami Creek fire plume. The parameters of the fitted total number size distributions (Fig. 6) are shown in Table 2. Additionally, the factors used to convert the data from STP to ambient conditions are given in Table 2. For the Aitken mode (\(j = 1\)), i.e., particles < 0.14 µm (Table 2), the number
Figure 4. Flight tracks for flight #110916a (left) and #110916b (right) on 16 September 2011 with averaged (10 s) rBC mass concentrations (STP) color-coded along the flight track. For clarity, only data for altitudes above 9 km a.s.l. and rBC mass concentrations > 0.01 µg m\(^{-3}\) are shown.

3.1.3 Coating thickness

Besides the size distribution information of aerosol layers, the mixing state of individual particles plays an important role in atmospheric processes. As described in Sect. 2.2, the coating thickness of individual particles inferred from the SP2 measurements delivers information about the mixing state of individual particles. Figure 7a shows normalized histograms of the observed coating thickness of all BC-containing particles with BC cores in two size ranges \((D_{rBC} = 140–160 \text{ nm}, \ D_{rBC} = 180–220 \text{ nm})\) for the 14 different sequences in the Pagami Creek fire plume. These two size ranges are chosen as examples to test if the coating thickness \(\Delta R_{\text{coat}}\) varies with changing BC core size. The coating thickness of individual aerosol particles from the Pagami Creek fire aerosol layer ranges from \(\Delta R_{\text{coat}} = 20 \text{ nm}\) up to the upper detection limit of \(\sim 180 \text{ nm}\), in both BC core size ranges shown, with a flat maximum between \(\sim 80 \text{ nm}\) and 160 nm (saturated signals originating from particles with coatings thicker than 180 nm were also frequently observed). The small number of BC cores with coatings below 20 nm observed within the Pagami Creek fire plume is likely from UTLS background particles and/or aircraft emissions. The observed coating thickness distributions are very similar for all 14 sequences, indicating very homogeneous BC particle properties across the different plume areas.

For comparison, the BC particles observed in the boundary layer in the regions of Neubrandenburg and Munich are thinly coated and show median coating thicknesses of < 20 nm with the majority (75 %) showing coating thicknesses of < 65 nm. Thus, the particles in the observed Pagami Creek fire aerosol layer exhibit much larger coating thicknesses than younger aerosol particle populations such as the planetary boundary layer (PBL) aerosol. The negative coating thickness values measured for some particles (Fig. 7a) are caused by random noise and potential systematic biases resulting from the assumptions made about the refractive indices and particle shape. The systematic uncertainty for the coating thickness is assessed to be < ±20 nm.

The raster graphic in Fig. 7b illustrates the relationship between the BC core size and the coating thickness integrated over all Pagami Creek fire plume sequences. The color-coding and the black line represent the number of particles in each pixel and the median coating thickness, respectively. The grey lines represent the detection limits of this method (see figure caption for explanation). Coating...
Figure 5. Median rBC mass size distribution (red dots) out of the 14 measurement sequences (see Fig. 2) in the Pagami Creek fire plume. Log-normal fits (solid lines) are shown for all sequences for the measured size range of 80–250 nm. The grey shaded area represents the range between maximum and minimum log-normal fit. The mean total rBC mass concentration (accounting for the data outside the SP2 range) is 0.03–0.35 µg m\(^{-3}\) (STP) for the 14 sequences in the forest-fire layer, indicated by the numbering. Error bars represent 15% uncertainty (see Sect. 2.2). For comparison the rBC mass of a free tropospheric (FT) background sequence (blue) is shown.

Figure 6. Number size distributions (STP) for background aerosol (blue markers), minimum (orange markers), median (red markers), maximum (dark red markers), and range (light grey area) of measurements in the Pagami Creek fire plume. Number size distributions are extrapolated by log-normal fits for the minimum, median, maximum, and background measurements (orange, red, dark red, and blue solid lines), and compared to forest-fire plumes (dashed lines) from LACE 98 (purple) as ambient values, ICARTT ITOP (International Consortium for Atmospheric Research on Transport and Transformation, Intercontinental Transport of Pollution; light green), SAMUM-2 (Saharan Mineral Dust Experiment; dark green), and SAFARI 2000 (Southern African Regional Science Initiative; black). Normalized values from SAFARI 2000, showing only the accumulation mode due to a lack of measurements for particle sizes < 100 nm, are scaled to similar magnitudes.

3.2 Estimates on the dimension and classification of the observed aerosol layer

Figure 8 shows vertical profiles of rBC mass concentration for both flights on 16 September 2011 with 100 m vertical resolution, including the median and 10th/25th/75th/90th percentiles for the specific size range 80–440 nm. The vertical profiles are averaged over the entire flight for altitudes > 2 km. For altitudes < 2 km the profiles are separately shown for the Falcon and the ground-based lidar in Leipzig also observed the Pagami Creek fire aerosol layer on 16 September 2011 in the UTLS. Figure 9 presents lidar observations performed in Leipzig, Germany (51.3° N, 12.4° E), between 08:36 UTC and 10:30 UTC on 16 September 2011. In the time series of the 532 nm range-corrected backscatter signal (Fig. 9a), the Pagami Creek fire plume was observed at altitudes between 8 km and
A humid layer was present below 1.5 km altitude, which causes the vertically striped pattern in the signal of the forest-fire plume in the UTLS due to partial attenuation of the laser light by thin clouds or haze. The base of the heterogeneously-layered Pagami Creek fire plume slowly descended with time. In the course of the late morning around 10:30 UTC the layer diminished above Leipzig. The time-resolved lidar profiles strongly confirm our observations, showing the maximum of the plume between 10.0 km and 11.5 km a.s.l. and an upper limit at roughly 12 km. At the time of the Falcon flyover (15:43 UTC) in Leipzig, neither the DLR Falcon nor the lidar observed signatures of the plume. In agreement with the lidar and in situ observations, the spaceborne lidar CALIPSO (not shown) observed the Pagami Creek fire plume on 16 September 2011 at about 10–12 km altitude over Europe.

The vertical profiles of particle backscatter coefficient $\beta_{\text{par}}$ and particle linear depolarization ratio $\delta_{\text{par}}$ are presented in Fig. 9b based on the averaged 532 nm elastic backscatter signal of the time period from 08:36–09:25 UTC. Sequences strongly affected by the low-level clouds are removed from the analysis to increase the signal-to-noise ratio. Because no Raman-scattering information is available, the lidar data analysis is performed using the Klett–Fernald method (Klett, 1981; Fernald, 1984). This requires the assumption of the extinction-to-backscatter ratio (lidar ratio). Because earlier studies (Tesche et al., 2011; Groß et al., 2013) show that a lidar ratio of 70 sr is reasonable for biomass-burning aerosol, we use this value for the analysis of the lidar data. Uncertainties in the retrieved optical properties are due to noise and the assumption of the lidar ratio (Rocadenbosch et al., 2010). Particle linear depolarization ratios of around 6–8% at 532 nm are observed throughout the Pagami Creek fire plume between 8 km and 12 km altitude. This is comparable to findings by Groß et al. (2013) who presented particle linear depolarization ratios of $7 \pm 2$% for aged (4–6 day-old) biomass-burning aerosol from their own measurements and summarize that values between 2% and 8% are reported in the literature for fresh to aged pure biomass-burning aerosol. Müller et al. (2007) report that particle linear depolarization ratios $< 5$% can be expected for biomass-burning aerosol particles, probably due to their spherical shape. However, in agreement with our results, Müller et al. (2007) observed particle linear depolarization ratios of up to 10% at 532 nm wavelength during the Lindenberg Aerosol Characterization Experiment 1998 (LACE 98), which they explained by soil dust particles possibly lifted together with the smoke particles at the source of the fire. Weinzierl et al. (2011) report particle linear depolarization ratios of 6–16% (median: 13%) at 532 nm wavelength for biomass-burning aerosol mixed with mineral dust from the Sahara measured during the SAMUM-2 field experiment in the Cape Verde region. Other aerosol types can be excluded as source of the observed layer with ice clouds and mineral dust having higher and sea salt and sulphates having lower particle linear depolarization ratios (Groß et al., 2013). A mixture of ash and sulphate of volcanic origin could also explain the observed depolarization ratios besides biomass-burning aerosol. However, volcanic eruptions did not occur in an appropriate spatiotemporal window. In addition, enhanced BC is normally not observed in volcanic ash layers (Weinzierl et al., 2012). The lidar in Leipzig only probed the edges of the plume (Fig. 4), such that the observed particle backscatter coefficient $\beta_{\text{par}}$ (up to 1 Mm$^{-1}$ sr$^{-1}$) above Leipzig is most likely lower than that found in the central part of the Pagami Creek fire aerosol layer. Using the observed particle backscatter coefficient, the optical thickness (532 nm) of the Pagami Creek fire plume is estimated to be at least 0.15–0.20, which is in...


Figure 8. Vertical profiles (STP) of flight #110916a (left) and #110916b (right) on 16 September 2011 for rBC mass concentration measured with the SP2 as median (black lines), 25th/75th percentile (dark grey shading), and 10th/90th percentile (light grey shading) for 100 m vertical intervals. Below 2000 m the data set is split into the PBL of Oberpfaffenhofen (593 m a.s.l.; dashed line) and Neubrandenburg (69 m a.s.l.; dotted line). Values are representative for the approximate size range of 80–440 nm for the rBC particles.

Figure 9. (a) Time-height cross-section of the 532 nm range-corrected backscatter signal of the lidar system in Leipzig and (b) vertical profiles of particle backscatter coefficient $\beta_{\text{par}}$ (black curve) and particle depolarization ratio $\delta_{\text{par}}$ with error bars (red curve) for the time interval 08:36–09:25 UTC. The optical thickness (at 532 nm) of the observed aerosol layer is derived to be 0.15–0.20.

the range of previous findings for biomass-burning aerosol layers, but higher than the median, of Weinzierl et al. (2011).

3.3 Identification of the Pagami Creek forest fire as the source of the observed aerosol layer

Figure 10a shows a true-color image of the Pagami Creek fire located in Minnesota, USA (approximate location 47°52′N, 91°30′W; US Forest Service), taken by the Moderate Resolution Imaging Spectroradiometer (MODIS) on 12 September 2011 at 18:50 UTC. This fire was caused by lightning on 18 August 2011. Before 12 September the fire did not evolve significantly and thus caused only minor pollution according to the MODIS images. On 12 September 2011 the fire grew rapidly exhibiting a thick plume, which is transported eastwards (Fig. 10a). The Pagami Creek fire burned a total area of roughly 375 km$^2$. A cloud-top 11.0 µm brightness temperature of roughly −60°C, corresponding to altitudes > 11 km, was observed with MODIS, which is comparable to deep thunderstorm clouds and is indeed colder than at any level up to 16 km in the radiosonde observation (RAOB) at International Falls, Minnesota, that day. Thus, this brightness temperature minimum certainly indicates cloud top altitudes up to the cold point above 12 km altitude. Radar echo tops on the cloud anvil reach $\sim$13.6 km at 19:54 UTC caused by pyro-convection forming this pyrocumulonimbus (pyroCb) cloud as a result of the extensive forest fire (Fromm et al., 2010). Similar observations of brightness temperatures for pyro-convective clouds are discussed in Damoah et al. (2006). Additional pyro-convection events are observed and described in Andreae et al. (2004) and Fromm et al. (2005). MODIS imagery suggests that the bulk emission and the pyro-convection event were limited from roughly 17:00 UTC on 12 September to 04:15 UTC on 13 September. In fact, Advanced Very High Resolution Radiometer (AVHRR) imagery (not shown) suggests that pyro-convective lofting shut down at sometime between 20:41 UTC and 22:41 UTC on 12 September.
To show that the aerosol plumes observed by the Falcon over Germany on 16 September originated from the Pagami Creek fire, we conduct HYSPLIT dispersion calculations starting at the location of the Pagami Creek fire. Based on the described observations, we assume that the Pagami Creek fire released emissions for 12 h between 17:00 UTC on 12 September 2011 and 05:00 UTC on 13 September 2011 as a maximum estimate. The release location is retrieved from satellite images and the release altitude is set from ground to 11 km a.g.l., assuming pyro-convection (e.g., Fromm et al., 2005; Damoah et al., 2006) and a homogeneous vertical distribution. Results are presented in Fig. 10b–f, which show the column-integrated (6–14 km a.s.l.) particle number concentration (arbitrary units; a.u.) in the time from 20:00–23:00 UTC between 12 September and 16 September 2011. The calculations underline that the Pagami Creek fire plume was transported through the area and at the altitude where the aerosol layer has been measured by the Falcon and the lidar in Leipzig. According to those simulations the Pagami Creek fire plume should have reached Germany on 15 September 2011 (Fig. 10e). A detailed analysis of the simulation results suggests that the smoke layer was north of Leipzig on 15 September, explaining why the lidar in Leipzig did not detect the Pagami Creek fire plume on that day. Figure 10g–i show zoomed-in dispersion calculations of column-integrated (10–12 km a.s.l.) particle number concentrations on 16 September 2011 for central Europe for the time between 08:00–11:00 UTC (g), 11:00–14:00 UTC (h) and 14:00–17:00 UTC (i). Figure 10h and i additionally include the Falcon flight tracks of flight #110916a (10:48–13:20 UTC) and #110916b (14:54–17:48 UTC), respectively. Comparing the location of the measured plume and the plume location simulated by the dispersion calculations strongly suggest that the Pagami Creek fire is most likely the source of the observed aerosol layer. This is also supported by observations of CO, a tracer for biomass burning, by the Infrared Atmospheric Sounding Interferometer (IASI) and by data of the GOME-2 Absorbing Aerosol In-
dering, by the Infrared Atmospheric Sounding Interferometer supported by observations of CO, a tracer for biomass burn-

4 Discussion

4.1 Comparison of the measurements in the Pagami Creek fire plume with other measurements during CONCERT 2011

Although the localization and observation of elevated aerosol layers was not a scientific objective of the CONCERT 2011 field experiment, in 9 of the 11 flights elevated aerosol layers, including the Pagami Creek fire plume, were observed. From about 32.3 h total flight time we flew by chance roughly 3.2 h (~10 % of the flight time) within elevated aerosol layers (volcanic ash layers, which we intentionally measured, are not counted here). Table 3 gives an overview of all CONC-
ERT 2011 flights, including the altitude of the detected aerosol layers. As can be seen from Table 3, most of the aerosol layers were found between 2 km and 5 km altitude with rBC mass concentrations of 2–425 ng m\(^{-3}\) (size range: 80 \(< D_{\text{rBC}} < 440\) nm). Figure 11 shows the median vertical profile of the rBC mass concentration and the rBC mass mixing ratio for the entire CONCERT 2011 field experiment, i.e., for all 11 research flights, with a 500 m vertical resolution and a temporal resolution of 10 s. This temporal resolution is used with consideration of the trade-off between reducing the noise from insufficient integration time and retaining sufficient horizontal resolution of about 2 km. The rBC mass mixing ratio (MMR) is calculated from the mass concentration \(m\) at standard conditions and the density of dry air at standard conditions (\(\rho_0\) = 1.29 kg m\(^{-3}\)): MMR \([\text{ng kg}^{-1}] = m_{\text{STP}} \left[\text{ng m}^{-3}\right] / \rho_0 \left[\text{kg m}^{-3}\right]\). The median vertical distribution can be used as an approximation of the typical vertical profile of BC above central Europe during autumn 2011. Below roughly 2 km altitude the vertical profile is dominated by the influence of the boundary layer: we see the highest MMR, a small spread between 10th/90th percentiles and a decreasing rBC MMR with increasing altitude. Above 2 km the signal is dominated by the wide spread of the percentile values, showing high variability in rBC mixing ratios above the boundary layer. This variability can be attributed to the presence of aerosol layers as a source of heterogeneous aerosol loadings in the free troposphere. The highest variability is found above 10 km altitude (Figs. 8 and 11). The greatly increased 75th and 90th percentile values at this altitude are related to the high loadings detected in the Pagami Creek fire plume.

4.2 Comparison with previous studies from literature

4.2.1 rBC mass concentrations and rBC size distributions

In Sect. 3 we showed that the Pagami Creek fire plume, observed in the altitude range of 9–12 km on 16 September 2011, reached rBC mass concentrations between 0.03 \(\mu\)g m\(^{-3}\) and 0.35 \(\mu\)g m\(^{-3}\) (STP), which are
comparable with the STP rBC mass concentration in the PBL over Neubrandenburg-Trollenhagen (0.11 µg m\(^{-3}\) and 0.05 µg m\(^{-3}\); STP) and Oberpaffenhofer (0.38 µg m\(^{-3}\) and 0.96 µg m\(^{-3}\); STP) on that day but far larger than the background concentration (< 0.01 µg m\(^{-3}\); STP) at the same altitude as the plume. Oberpaffenhofer is located near the metropolitan area of Munich. Therefore these values are representative of a polluted PBL. To put the results of the observed Pagami Creek fire plumes in context to the literature and to other typically expected BC-containing aerosol types, several papers are chosen: biomass-burning plumes from Wyoming (Pratt et al., 2011), biomass burning from North America and Asia (Kondo et al., 2011a), biomass-burning and anthropogenic emissions from California (Sahu et al., 2012), biomass-burning and urban emissions from Texas (Schwarz et al., 2008a) and urban emissions from California (Metcalf et al., 2012). In 12–24 h-old biomass-burning plumes at altitudes from ground to roughly 5 km Kondo et al. (2011a) measured the highest rBC mass concentration in smoldering burning events (0.27 µg m\(^{-3}\)–1.24 µg m\(^{-3}\); STP). Sahu et al. (2012) observed rBC mass concentrations of 0.50 ± 0.17 µg m\(^{-3}\) (STP), while Pratt et al. (2011) reported total rBC mass concentrations of up to 2 µg m\(^{-3}\) in fresh biomass-burning plumes (~2–120 min old) in altitudes up to 3.5 km a.g.l. These concentrations are a factor of 5–60 larger than those found for the aged Pagami Creek fire plume (3–4 day old) in this study. However, the measurements by Pratt et al. (2011) were taken at altitudes lower than 3.5 km whereas our measurements were taken above 10 km. These differences, however, may be attributed at least partly to the dilution caused by turbulent mixing and wet removal processes. Besides the emissions of forest fires, preexisting BC particles originating from other sources such as traffic and other anthropogenic combustion processes may contribute to the BC load in forest-fire plumes. However, due to the intensity and the remote location of the fire, this fraction is assumed to be small so that the described characteristics can be considered to be representative of the forest-fire particles.

Besides the rBC mass concentration, the mass size distribution is an important aspect for BC in the atmosphere. As can be seen from Fig. 5, the MMD in the Pagami Creek fire plume is between 120 nm and 160 nm. These MMDs are smaller than observed by Kondo et al. (2011a) who found MMDs between 177 nm and 197 nm in fresh

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Figure 10. Evolution and dispersion of the Pagami Creek fire plume calculated with HYPLIT. (a) MODIS picture of Pagami Creek fire in Minnesota, USA on 12 September 2011, 18:50 UTC (NASA images courtesy Jeff Schmaltz, MODIS Rapid Response Team at NASA GSFC. Caption by Michon Scott.). (b–f) Daily evolution of the plume vertically integrated from 6–14 km from 12 September 2011 through to 16 September 2011 for time period 20–23 UTC on each day; (g–i) dispersion of the plume over Germany vertically integrated from 10–12 km for the 16 September 2011, (g): 08:00–11:00 UTC, (h): 11:00–14:00 UTC and (i): 14:00–17:00 UTC and locations of Oberpaffenhofer (OP; star) and Leipzig (triangle). Green dashed lines in (h) and (i) represent the sequences where the Pagami Creek fire plume was measured.
Table 3. List of research flights during CONCERT 2011. The lower and upper boundary* of the detected aerosol layers is given for each flight. The Pagami Creek fire plume is written in bold type. OP, Oberpfaffenhofen; NB, Neubrandenburg-Trollenhagen; TRA, Temporary Restricted Area.

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* The boundaries are derived from the in situ measurements, representing the minimal vertical depth of the layers.
organic material are also reported by Kondo et al. (2011a) for fresh biomass-burning plumes with mass fractions of 0.3–34 % in fresh biomass-burning plumes (age: 2–3 days, origin: Asia). MMDs of roughly 160–180 nm for urban emissions and roughly 200–220 nm for biomass-burning aerosol, while Metcalf et al. (2012) observed modes of roughly 120–140 nm for boundary layer aerosol and roughly 160 nm for free-tropospheric aerosol during airborne measurements in the Los Angeles Basin. Moteki et al. (2012) and Oshima et al. (2012) describe the wet removal of rBC during transport. A size-dependent wet removal (Moteki et al., 2012) can shift the MMD of aged BC layers towards smaller particle sizes and might cause differences in observed MMDs for BC particle populations with varying exposure to wet removal conditions during the transport as potential reason for the observed differences. They observed, for instance, a decrease from 127 nm to 97 nm with decreasing transport efficiency for the count median diameter.

Pratt et al. (2011) observed with their SP2 average BC mass fractions of 0.3–34 % in fresh biomass-burning plumes with the largest part of the mass (60–90 %) being nonrefractory organic material. Comparably high mass fractions of organic material are also reported by Kondo et al. (2011a) who find an organic fraction of 56–92 % and a BC mass fraction of 1.7–4.9 %. The values from Kondo et al. (2011a) are comparable to findings from Sahu et al. (2012) presenting <5 % BC mass fractions. The higher amount of OC compared to BC can be explained by emission factors, which are 0.2 g BC and 7.8 g OC per kg burned mass for boreal coniferous forests (McMeeking, 2008; Wiedinmyer et al., 2011).

In our study, the volume fraction for rBC-containing particles derived from the SP2 for the 14 considered sequences is 2.5–3.9 % in the described size range. No measurements of organic material were performed but we have to assume that in our case OC is also a major constituent in terms of mass fraction.

### 4.2.2 Total particle number size distributions

The comparison of the total aerosol number size distributions, including BC-free particles (representing the largest fraction, and a minor fraction of BC-containing particles), of the Pagami Creek fire plume with measurements in clean UTLS “background” air (Fig. 6) exhibits one striking feature: the Aitken and accumulation modes give roughly comparable contributions to particle number concentration in the Pagami Creek fire cases, while the accumulation mode has a very minor contribution in the background case. On 16 September 2011, no particles larger than 400 nm were present in the background air, while particles of 1–2 µm in diameter were detected in the Pagami Creek fire plume. In all Pagami Creek fire plume cases the accumulation mode is strongly enhanced compared to the background sample. Deviating from the monomodal structure of the background aerosol, the particles larger than 0.14 µm can be attributed to the presence of forest-fire aerosol. Our results of number size distributions of biomass-burning plumes as well as measurements from the SAMUM-2 field experiment at 0.8–4.2 km altitude (Weinzierl et al., 2011), the SAFARI 2000 field experiment at roughly 2–5 km altitude (Haywood et al., 2003), the LACE 98 field experiment at 3–5 km altitude (Fiebig et al., 2003) and the ICARTT-ITOP field experiment at 3–9 km altitude (Petzold et al., 2007) show that in all biomass-burning aerosol cases, an enhanced accumulation mode is present. Especially in the ICARTT-ITOP case where a 6–9 day-old Canadian forest-fire aerosol layer over Europe was investigated, the bimodal structure of the size distribution is very pronounced and similar CMDs are found as in our case. By approximating STP conditions, results from LACE 98 (Fiebig, 2001) exhibit a bimodal structure with similar CMDs as well as with total number concentration about 2–9 times higher than in the Pagami Creek fire plume (not shown). Compared to our results, the SAMUM-2 case (3–6 day old) shows a similar CMD for particles > 0.14 µm but the mode for particles < 0.14 µm is not as enhanced. In addition, Weinzierl et al. (2011) reported a significant coarse mode fraction for data collected during the SAMUM-2 field experiment attributable to mineral dust. In contrast we observed no significant amount of aerosol particles larger 1 µm.
Figure 12. SP2 raw signals for selective particles with LSD signal (black) and BID signal (blue) and the scattering cross-section (red) for (a) – an rBC-free particle, (b) – a coated rBC particle and (c) – a disintegrating rBC particle. Schematics for the particle–laser interaction for each raw signal plot are shown on the left. For the disintegrating rBC particle in (c) several initial structures are possible: a very thickly coated rBC particle (top); a thickly coated particle with an eccentrically located rBC core (middle); an rBC particle located at the surface of an rBC-free particle (bottom). For depiction purposes the size of the schematically-shown particles (left) is enlarged compared to the laser diameter. The scattering cross-section was obtained by dividing the scattering signal by the laser intensity profile. The incandescence is indicated by yellow stars.

Results from SAFARI 2000 show only the accumulation mode since no measurements or extrapolations for particles < 100 nm were performed. Furthermore, the SAFARI 2000 size distribution was scaled to quantities found in our and other studies, since in Haywood et al. (2003) the size distribution is given as a normalized distribution. In summary, our observations are similar to biomass-burning number size distributions observed previously, especially for ICARTT-ITOP and LACE 98, indicating similar origins, source material, environmental and burning conditions or transportation. The aerosol layers often show similar modes in the accumulation mode size range. Thus, a dominant accumulation mode and a relatively large modal diameter (∼200–400 nm) seem to be common features of forest-fire aerosols.

4.2.3 Coating thickness

The coating thickness of BC-containing particles in biomass-burning plumes has previously been reported in the literature. Akagi et al. (2012) observed for a chaparral fire that 85% of the rBC cores may have been thickly coated 4 h after emission, i.e., with shell/core diameter ratios ($D_{opt}/D_{rBC}$) > ∼2 based on the time delay approach. Kondo et al. (2011a) observed rBC particles in biomass-burning plumes with shell/core ratios of $D_{opt}/D_{rBC} = 1.3–1.6$ in biomass-burning plumes, based on the LEO-fit approach. These values are somewhat lower than our observations (Sect. 3.1), which may be related to differences in the properties of the freshly emitted particles or continued mass acquisition of the BC particles during the long-range transport in the Pagami Creek fire plume. The coatings reported for
the BC particles from forest fires in all three studies discussed here are substantial. This is in contrast to BC particles emitted from diesel engines and wood burning for residential heating, which emit particles with very little and moderate coatings, respectively, and also the median coating thickness observed in aged aerosols typically remains clearly below 100 nm (Laborde et al., 2013, and references therein).

4.3 Morphology and disintegration of black carbon particles

The interaction of BC-containing particles with solar radiation depends on their morphology and mixing state with light-scattering aerosol components (Kahnt et al., 2012). The coating thickness values reported in the previous section were inferred assuming a concentric-spheres, core-shell morphology for the BC core and a purely scattering coating. However, the BC cores of combustion particles are fractal-like or compact aggregates and the BC can be mixed with light-scattering material in different ways, including, e.g., surface contact of the BC with the scattering components, full immersion of the BC in the scattering component or immersion of the scattering components in the BC aggregate (Scarmato et al., 2013). Previous laboratory and field experiments have shown that the SP2 signals of BC-containing particles can sometimes give indirect evidence of how the BC core is mixed with the scattering components beyond just quantifying the coating thickness (Sedlacek et al., 2012; Moteki and Kondo, 2007).

Figure 12 schematically shows the particle–laser interaction of different particle types and the corresponding SP2 signals including the time-resolved scattering cross-section as inferred from the raw LSD signal, according to the method described in Laborde et al. (2012a). For BC-free particles (Fig. 12a) the LSD signal usually follows an almost Gaussian shape. The LSD signal is directly proportional to the laser intensity profile along the flight path of the particle through the laser beam, because the size and also the scattering cross-section of a non-absorbing particle remains unchanged during laser transition. The maximum of the LSD signal from BC-free particles occurs at the center of the laser beam (here at time 26 µs; Fig. 12a).

In contrast, the size of a BC-containing particle is reduced during laser transition, first through evaporation of non-refractory coating material followed by evaporation of BC, which has a higher boiling point. Figure 12b shows typical SP2 signals of BC-containing particles with a medium-thick coating. The LSD responds with an increasing signal when the particle enters the laser beam and scatters the laser light with a constant scattering cross-section. Traveling further into the laser beam, the BC core absorbs the laser radiation and heats up. As a consequence, the coating evaporates and causes a decrease in the scattering cross-section. This often results in a first local maximum of the LSD signal (here at time 21 µs; Fig. 12b). After complete evaporation of the coating, the scattering cross-section of the particle stabilizes for a short time and the LSD signal raises again, because the laser intensity increases still (time range 22–23 µs; Fig. 12b). The BC core heats up further, until it reaches the boiling point of BC. The thermal radiation, emitted by the particle and detected by the incandescence detectors, now reaches its maximum (here at time 24 µs; Fig. 12b), and the BC core evaporates, resulting in a fast decrease of the BC mass and thus also the scattering cross-section of the residual particle, which is reflected in a fast decrease of the BID and LSD signals, respectively. Consequently, the LSD signal can have a second local maximum around the peak of the BID signal (here at time 23 µs; Fig. 12b). This second local maximum is typically higher than the first local maximum of the LSD signal for BC particles with a thin or medium coating, such that the global maximum of the LSD signal occurs at about the same time as the peak of the BID signal. For thickly coated BC particles, the first local maximum of the LSD signal is typically higher than the second local maximum, such that the global maximum occurs before the peak of the BID signal. Moteki and Kondo (2007) have shown that the time lag between the peaks of the LSD and the BID signals gives crude information about the mixing state of BC-containing particles, where values > ~ 2 µs are associated with thick coatings, while values < ~ 2 µs are associated with thin to medium coatings.

However, particles that disintegrate in the laser beam of the SP2 have also been reported in the literature (Moteki and Kondo, 2007; Sedlacek et al., 2012). Disintegration occurs before the BC core reaches its boiling point and results in a BC-free fragment, which remains stable after disintegration, and a BC fragment that eventually evaporates. Figure 12c shows the SP2 signals of such a disintegrating BC-containing particle. Disintegration can be identified by the fact that the LSD signal does not fully disappear after evaporation of the BC core. More precisely, the scattering cross-section stabilizes again after the peak BID signal, as the BC-free fragment does not evaporate any further (here at time > 24 µs; Fig. 12c). Furthermore, the stable scattering cross-section in both the leading and trailing edges of the laser beam is a key feature that distinguishes disintegrating particles from coincidence of a BC-containing and a BC-free particle (the position dependent correction for the laser beam intensity would fail for coincident particles, that are, while simultaneously being in the laser beam, not exactly in the same position; this would cause an apparent time dependence of the inferred leading and/or trailing edge scattering cross section for coincident particles). The BC-free fragment of a disintegrated particle, which behaves as a BC-free particle after its separation, causes another local maximum of the LSD signal at the center of the laser beam (here at time 28 µs; Fig. 12c). This local maximum of the LSD signal from the BC-free fragment can be the global maximum if the BC-free fragment is a substantial portion of the original particle. In this case the time lag between the peaks of the LSD and the BID signals
becomes negative as the peak of the BID signal normally occurs before the center of the laser beam.

Sedlacek et al. (2012) have reported, for the first time, the identification of disintegrating particles through negative lag times. However, in our study, only a minor fraction of all disintegrating particles could be identified via a negative lag time because most of the disintegrating particles either exhibited a positive lag time or because the peak of the LSD signal, and with that, also the lag time, were not defined due to LSD detector saturation. Therefore, the time-dependent (position-dependent) scattering cross-section, as inferred from the raw LSD signal, according to the method described in Laborde et al. (2012a), was used here to identify disintegrating particles. Specifically, the scattering cross-section measured in the trailing edge of the laser beam (at the position with 3% of the maximum laser power) was used to determine for all BC-containing particles whether a detectable BC-free fragment (producing an LSD signal above background noise) remained left in the trailing edge of the laser beam. The scattering cross-section measured in the leading edge of the laser beam (at the position with 3% of the maximum laser power) was used to determine the size of the undisturbed particles, in order to further investigate the influence of particle size on disintegration. The relation of the scattering cross-section at the leading edge and at the trailing edge of the laser beam for BC-containing particles is shown Fig. 13a for a sequence measured in the Pagami Creek fire plume (11:36–11:46 UTC, 16 September 2011). The trailing edge LSD signal indicates that about 50% of the BC-containing particles in the plume with particle size of about $D_{\text{opt}} = 300$ nm disintegrated in the laser beam, leaving a detectable BC-free fragment. The fraction of disintegrating particles increased with increasing particle size, so that > 90% of the BC particles disintegrated at large sizes ($D_{\text{opt}} > \sim 480$ nm). Conversely, the majority of the smaller particles ($D_{\text{opt}} < \sim 300$ nm) showed no disintegration characteristics. However it cannot be distinguished if they actually evaporated completely or if potential BC-free fragments just remained below the detection limit of the LSD. Overall ~40–50% of all detected BC-containing particles were identified as disintegrating particles with the trailing edge LSD-signal method. Data points shown in Fig. 13a remaining on or above the 1:1 line are attributable to the coincidence of a BC-free and a BC-containing particle, marked by the grey pattern. However, coincidence accounts for only a very low fraction of the data points.

Only 3.5–6.8% (median: 4.6%) of all BC particles, corresponding to roughly 5–20% of all identified disintegrating BC particles, exhibited a negative lag time in this study. This shows that the lag-time method only provides a lower limit for the total fraction of disintegrating particles. The success rate of the lag-time method for the identification of disintegrating particles depends on particle and instrument properties such as, e.g., the particle size and the saturation level of the LSD.

Sedlacek et al. (2012) observed disintegrating particles during a ground-based field campaign in summer 2011. More than 60% of BC-containing particles in the forest-fire aerosol plume from several wildfires east of Lake Winnipeg and into western Ontario on 30 July were disintegrating BC particles, exhibiting negative lag times in the SP2. Our observations show for the first time the occurrence of disintegrating BC particles in high-altitude BC layers (40–50% of all detected BC-containing particles, according to the trailing-edge LSD signal method).

Sedlacek et al. (2012) suggested that disintegrating particles are associated with a particular particle morphology,
where the BC is located near or at the surface of the BC-containing particles. Moteki and Kondo (2007) also reported the occurrence of disintegrating particles in laboratory experiments with graphic cores that were coated with glycerol or oleic acid. The number fraction of disintegrating particles was mainly a function of overall particle diameter. No disintegration was observed for small sizes, while almost all larger particles disintegrated. The threshold particle diameter for the occurrence of disintegration was 450–600 nm depending on the coating type. Moteki and Kondo (2007) speculated that disintegration might be associated with eccentrically positioned BC cores in their case, while near-surface BC morphology was not expected based on the process applied to coat the BC cores.

The specific BC particle morphology, i.e., core-shell versus near-surface BC, is relevant because the light-absorption efficiency enhancement effect by light-scattering coatings internally mixed with BC depends on the coating volume fraction as well as the particle morphology. Coatings with $D_{\text{opt}}/D_{\text{BC}}$-ratios of 2 can already enhance the light-absorption by the BC cores by 100 % (Schwarz et al., 2008b; Shiraiwa et al., 2010), while the relation between the coating thickness and the enhancement factor is not linear and saturation occurs above a certain threshold. The high $D_{\text{opt}}/D_{\text{BC}}$ ratios observed in the Pagami Creek fire plume imply that substantial light-absorption enhancement effects are expected, though large uncertainties of this effect remain (e.g., Cappa et al., 2012). Sketches of three conceivable morphologies of disintegrating BC-containing particles are shown on the left side of Fig. 12c: a concentric core-shell, an eccentric core-shell or a near-surface BC/attached BC morphology. Light-scattering matter simply attached to a BC core causes very little enhancement of the light-absorption efficiency (Liu and Mishchenko, 2007; Scarnato et al., 2013). In contrast, a substantial light-absorption efficiency enhancement can be expected for coated BC-containing particles with a concentrically or eccentrically positioned core.

We note here that disintegrating particles might also complicate the interpretation of the Soot-Particle Aerosol Mass Spectrometer (SP-AMS; Onasch et al., 2012) measurements because the BC-free fragment would not be evaporated by the laser. This should be further investigated with specific laboratory experiments involving biomass-burning aerosol and the SP-AMS.

A process that can produce near-surface BC morphologies is coagulation of almost bare BC aggregates with BC-free particles. Condensation of secondary organic or inorganic aerosol components on BC particles can either result in particles with core-shell morphology (concentric or eccentric) or with near-surface BC morphology. Adachi et al. (2010) proposed a mechanism producing near-surface BC particles via condensation processes, which involves condensation of organic material onto nascent BC that effectively preserves the aggregate of primary BC particles in an un-collapsed form, followed by further mass acquisition via condensation, preferentially onto the more organic portion of the particle. Based on our observations in the aged Pagami Creek fire plume, disintegration was associated with large BC-containing particles (overall optical diameter $> 300$ nm; see above). These large particles generally had a very low rBC mass fraction since the majority of the rBC cores were $< 130$ nm in diameter. They can be the result of extensive condensation of secondary aerosol components onto the nascent BC aggregates or of coagulation of BC-particles with large BC-free particles during transport. Indeed, the initial plume was highly concentrated such that both processes are likely to occur during aging of the BC particles in the plume. Thus, it is in principle possible that both BC particle types with near-surface BC or core-shell morphology are present in the aged plume. However, the observed number fraction of disintegrating particles is for the most part a function of overall particle size, indicating that disintegration was possibly just the result of very thick coatings, as reported in the laboratory study by Moteki and Kondo (2007). Figure 13b shows the fraction of disintegrating particles to all rBC-containing particles as a function of their coating thickness for the same sequence as shown in Fig. 13a (11:36–11:46 UTC, 16 September 2011). The increasing fraction of disintegrating particles with respect to all rBC-containing particles as a function of increasing coating thickness supports the assumption that thick coatings trigger the disintegration. At a coating thickness of about 120 nm 50 % of the rBC-containing particles produce an LSD signal above background noise in the trailing edge of the laser beam. For particles with coating thicknesses $> 200$ nm, more than 90 % of the rBC-containing particles disintegrate. Moteki et al. (2014) showed with controlled laboratory experiments that attached-type rBC particles can be identified by insignificant evaporation before incandescence onset. The dominant fraction of the disintegrating rBC-containing particles observed in the Pagami Creek fire plume exhibited substantial coating evaporation, such as the example particle shown in Fig. 12c, which is clear evidence that those rBC-containing particles had a coated-type morphology.

The specific features of the particles in the aged Pagami Creek fire plume observed in this study are the enhanced size of all particles (Fig. 6) and the enhanced coating thickness of BC-containing particles (Fig. 7a) compared to the observations outside the plume. Both can be attributed to extensive coagulation and/or condensation in the forest-fire plume. The very thick coatings of BC particles in the forest-fire plume will influence their light-absorption properties. Furthermore, they will also influence their interaction with clouds. They are expected to be good cloud condensation nuclei through the mere size effect as well as the additional hygroscopic water uptake by the coating (e.g., McMeeking et al., 2011; Laborde et al., 2013; Liu et al., 2013). Furthermore, the mixing state of the BC with organic and inorganic matter might have an influence on the ice-nucleation efficiency of the particles and is thus relevant for the indirect aerosol effect (Hooge and Möhler, 2012).
It is important to consider large particle size when assessing the radiative impact of such plumes by direct light-scattering of solar radiation. When assessing the radiative impact via absorption of solar radiation, it is recommended, based on the above discussion about the morphology of BC particles, to consider the light-absorption enhancement effect that is expected to occur for thickly coated BC particles with core-shell morphology.

The first two reports of disintegrating BC particles in ambient aerosols are from forest-fire plumes (Sedlacek et al., 2012 and this study). The question if disintegrating particles are uniquely or predominantly associated with biomass-burning aerosol remains to be addressed. A UTLS background sequence on 16 September 2011 showed an insignificant amount of disintegrating particles (< 1%). The fact that biomass-burning BC particles often exhibit thicker coatings compared to BC particles from, e.g., diesel engines, even allowing for coating acquisition during atmospheric aging (e.g., Laborde et al., 2013), might be a reason for the preferential occurrence of disintegrating particles in biomass-burning aerosol.

4.4 Black carbon import into the UTLS by the Pagami Creek fire in comparison to aviation

In this section we present two estimates for determining the primary emissions from the Pagami Creek fire (first estimate) and the import of BC from the fire to the UTLS region (second estimate).

First, we estimate the total OC and BC emissions of the Pagami Creek fire (Fig. 14a) based on the burned area, the wood mass per area and the emission factor of boreal needle-leaf forests based on the values presented in Wiedinmyer et al. (2011). These consist of BC and OC emission factors for boreal coniferous forests (0.2 g per kg biomass burned and 7.8 g per kg biomass burned, respectively; McMeeking, 2008) and fuel loadings for North American boreal forests (25 kg m\(^{-2}\); Akagi et al., 2011, and references therein) are combined with the burned area (~350 km\(^2\); US Forest Service), resulting in roughly \(2 \times 10^3\) Mg BC and \(68 \times 10^3\) Mg OC emitted from the Pagami Creek fire. This estimate depends strongly on the assumptions made for the burned area and the applied emission factors and must be considered with caution, implying large uncertainties. A large fraction of these primary emissions will fall out quickly by, e.g., wet deposition (see Sect. 4.1), will be transported to altitudes below 8 km (see Fig. 3; Mattis et al., 2008; Weinzierl et al., 2011) or will not even enter the free troposphere. The fraction of the emissions entering the upper troposphere is more relevant in terms of its impact on the radiation budget due to longer life time and the possibility of long-range transport.

To assess the BC import of the Pagami Creek fire to the UTLS directly, we create a second estimate. Assuming a homogeneously distributed aerosol layer, we can approximately derive the BC mass import from the Pagami Creek fire event into the UTLS region above Europe (Fig. 14b) from our measurements and dispersion model simulations. The area of the aerosol layer is deduced from the HYPLIT dispersion calculation by sizing the plume area for 15 September 2011 (20:00–23:00 UTC) by assuming a plume concentration threshold (0.1 a.u.; see Fig. 10) and a reasonable uncertainty (area: \(3 \pm 2\) \(\times 10^3\) km\(^2\); Sect. 3.3). This plume area is representative of the plume observed 12 h later above Europe based on observations from IASI (Sect. 3.3) and GOME (Sect. 3.3). The depth of the aerosol layer is deduced from the lidar observations (depth: 2 \pm 1 km; Sect. 3.1). Thus, the volume of the plume is derived as 0.1–1.5 \(\times 10^6\) km\(^3\) (best estimate: \(6.0 \times 10^6\) km\(^3\)). Furthermore, we use the rBC mass concentration from our measurements at the edge of the plume (7–84 ng m\(^{-3}\); median of measurement sequences: 42 ng m\(^{-3}\); at ambient conditions). From these data we obtain a best estimate for the BC total mass import into the high-altitude aerosol layer above Europe of 25 Mg BC, ranging from 1–126 Mg BC. This estimate depends strongly on the plume area at the time of measurements, which in turn depends on the assumed duration of the emission in the dispersion calculations. Nevertheless, we show in Sect. 3.3 that the duration of the emissions used in the simulation is in good agreement with the observations of the duration of the fire. The local inhomogeneities observed in both the simulation (Fig. 10) and the measurements (Fig. 2) cause large uncertainties in plume area and average concentration. However, the uncertainties of this estimate are smaller than the uncertainties for the theoretical estimate on the direct total emissions of the Pagami Creek fire described above. This is because the input parameters are based on direct measurements and calculations related to the specific event rather than generalized emission factors used in the first estimate. As the dispersion calculation suggests that the measurements took place in the less concentrated tail of the plume, the maximum
value of 126 Mg BC might still underestimate the imported BC mass. Nonetheless, it is highly likely that the real BC mass of the Pagami Creek fire plume is within the presented span of 1–126 Mg BC.

Comparing our best estimate of the direct emissions at the source with the best estimate of the BC mass in the plume we observed after 4 days on 16 September 2011, this aerosol layer above Europe contains less than 7% of the total BC mass emitted by the fire.

These estimates show that singular forest-fire events can be a relevant source for substantial BC loadings in the UTLS region on time scales of at least several days. Aside from forest fires, another (permanent) source of BC in the UTLS region is emissions from aircraft (Fig. 14c). Lee et al. (2010) estimate the soot (defined as BC + OC) emissions from aviation to 16 Mg d\(^{-1}\). The BC/OC ratio for aviation-induced soot is roughly 3.5 (Bond et al., 2004, and references therein). Thus, the estimate of Lee et al. (2010) can be converted to roughly 12 Mg d\(^{-1}\) of BC. Bond et al. (2013) estimate the aviation-induced BC to be 55 Mg d\(^{-1}\). The BC emissions from aviation in the UTLS region are even smaller, taking into account that the total aviation-induced emissions contain also emissions during take-off and landing at lower altitudes. Nevertheless, the best guess of the BC mass imported from the Pagami Creek fire to the UTLS region over Europe after several days of transport is of the same magnitude as the daily globally averaged BC mass emissions from aviation. The comparison of 1–126 Mg BC in the Pagami Creek fire plume in the UTLS region over Europe to 12–55 Mg d\(^{-1}\) BC related to aviation shows the relevance of such singular forest-fire events as a source for the import of black carbon into the atmosphere, keeping in mind that the BC size distribution of aviation-induced BC shows high concentrations in the smaller size ranges in contrast to forest-fire BC in the UTLS region. As this result is based on several assumptions, the given values should be considered with caution but do illustrate the potential relevance of forest fires as a source for BC in the UTLS region. A future increase in forest-fire activity due to a changing climate might be of great importance to the BC import into the upper troposphere.

5 Summary and conclusion

In the framework of the CONCERT 2011 field experiment we observed numerous elevated aerosol layers over Europe with the DLR research aircraft Falcon. This study focused on the Pagami Creek forest-fire aerosol layer detected accidentally on 16 September 2011 between 10 km and 12 km altitude over northeast Germany. From our measurements with the SP2, we derived the median vertical profile of rBC mass concentration for the entire CONCERT 2011 field experiment, representing an estimate for the typical black carbon mass-mixing ratios in central Europe in September 2011. We discussed the Pagami Creek fire plume with respect to this average situation. To obtain information about the aerosol layer characteristics, refractory black carbon (rBC) mass size distributions and total aerosol particle number size distributions were calculated for different parts of the Pagami Creek fire plume over Europe. The mode parameters of the applied log-normal fits were compared to size distributions of biomass-burning aerosol from other studies. Determining the impact of those elevated aerosol layers on the radiation budget it is essential for investigating the mixing state of the particles. Therefore, the coating thickness of individual aerosol particles was calculated, assuming ideal spherical BC particles coated with a non-absorbing material. Additionally, we provided information about the mixing structure of individual particles, more precisely, whether a BC-containing particle disintegrated in the SP2 laser beam. This effect was recently found in ground-based measurements of forest-fire aerosol and has now been confirmed for the first time for airborne in situ measurements.

In summary, the major findings of this study were as follows:

- The aerosol layer observed on 16 September 2011 is attributed to the Pagami Creek fire in Minnesota, USA, supported among other observations by the measured lidar linear depolarization ratio (532 nm) of 6–8%, indicating a forest-fire source.
- The total rBC mass concentration of the aged (3–4 day-old) Pagami Creek fire plume ranged from 0.03–0.35 µg m\(^{-3}\) (STP), comparable to rBC mass concentrations typical for moderately-polluted planetary boundary layers. The MMDs of the rBC size distribution ranged between 120 nm and 160 nm.
- In contrast to a background air sample showing a monomodal number size distribution, the Pagami Creek fire plume showed a bimodal number size distribution with a pronounced accumulation and Aitken mode.
- The rBC particles in the Pagami Creek fire plume showed coating thicknesses of roughly 50–220 nm with maxima of the histograms between ~80 nm and 160 nm (medians of 14 considered sequences: 105–136 nm), whereas fresh PBL aerosol exhibited coatings predominantly <65 nm (median: <20 nm). These coatings may modify the BC radiative forcing, the BC hygroscopicity, and the efficiency to act as cloud condensation or ice nuclei. Resulting from small rBC diameters and large coating thicknesses, we observed shell/core ratios of up to 7.4 with median shell/core diameter ratios of \(D_{\text{opt}} / D_{\text{rBC}} = 3.7\), 2.8 and 2.4 at \(D_{\text{rBC}} = 90\) nm, 130 nm and 170 nm, respectively.
- We showed that the majority of the rBC-containing particles in the Pagami Creek fire plume disintegrated in the SP2 laser beam. Generally, this instrument-specific
process will not occur without an intense laser beam but it is an indication of either very thickly coated rBC cores or rBC cores located near/attached to the surface of rBC-free particles. In this study, the dominant fraction of the disintegrating particles was of the thickly-coated type, for which a light-absorption enhancement can be expected. The question if disintegrating particles are uniquely or predominantly associated with biomass-burning aerosol remains to be addressed. The fact that biomass-burning BC particles often exhibit thick coatings might be a reason for preferential occurrence of disintegrating particles in biomass-burning aerosol. When assessing the radiative impact via absorption of solar radiation, it is recommended to consider the light-absorption enhancement effect that is expected to occur for thickly-coated BC particles with core-shell morphology.

- We presented a rough estimate of the BC mass import due to the Pagami Creek fire to the UTLS region above Europe (best estimate: 25 Mg BC) and compared it with the daily BC emissions of global aviation (12–55 Mg BC) thereby highlighting the relevance of forest fires as a potential BC source for the upper free troposphere.

In conclusion, the microphysical properties of the observed Pagami Creek fire plume obtained within this case study clarify the relevance of pyro-convection as a particle source and as a cause for long-range transport of forest-fire aerosol layers in the UTLS region. Consequently, this study enables us to assess the impact of this light-absorbing aerosol layer on the radiation budget using radiation transfer calculations. Additional in situ measurements of aerosol layers similar to the presented layer and additional studies of the processes of aerosol transport by pyro-convection to the UTLS region are necessary to globalize the estimate of BC mass import into the upper troposphere and to reduce uncertainties of model estimates.

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