Tropospheric ozone production related to West African city emissions during the 2006 wet season AMMA campaign

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Abstract. During African Monsoon Multidisciplinary Analyses (AMMA) airborne measurements of ozone, CO and nitrogen oxides (NOx) were collected by French and German Falcon aircraft near three cities in West Africa (Cotonou, Niamey and Ouagadougou). They have been analysed to identify the good conditions to observe ozone plumes related to city emissions during the monsoon season. Results show that an O3 increase of 40–50 ppbv above the summer average concentration took place during two specific events: one near Cotonou on the coast of the Gulf of Guinea, and the other near Niamey in the Sahel region. In both cases a high level of NOx (3–5 ppbv) is related to the ozone production. Air mass transport simulations with FLEXPART and a tracer simulation with the BOLAM mesoscale model shows that Southern Hemisphere biomass burning emissions are always at higher altitude (>3 km) compared to the city emissions. In Niamey and Ouagadougou, the daily variability of ozone and CO correlates with the FLEXPART analysis showing the role of air mass stagnation near the city for 1–2 days and advection of emissions from the vegetated areas. Absence of ozone enhancements for high CO values can be explained by the occurrence of deep convection near the city. In the Sahel region, convection must be accounted for to understand the small number of observed ozone plumes but also to explain the high level of NOx in the 3–5 ppbv range, due to increasing soil emissions after rainfall. To verify that daily ozone production can reach 20 ppbv day−1 for the NOx and CO conditions encountered near West African cities, a simulation of the CTTyCAT Lagrangian model was conducted using the observed average chemical composition reported by other aircraft during AMMA. Such ozone production is possible for NOx levels up to 5 ppb showing that West African cities are potentially significant sources of tropospheric ozone.

1 Introduction

According to UNEP (United Nations Environment Programme), Africa will have the fastest population growth rate in the world between 2000 and 2050, twice the rate of any other region during that time. Given their relatively low industrial development, air pollution in west African countries is not as widespread as in other regions of the world. However, in most populous cities, long-term exposure to poor-air quality has been found to be a severe health issue for the population. Africa is also known to contribute to ozone formation at the global scale mainly because of large amounts of biomass burning (Andreae and Merlet, 2001; Thouret et al., 2009), biogenic emissions (soil and vegetation) (Guenther et al., 2006; Stewart et al., 2008) and lightning NOx (Schumann and Huntrieser, 2007). Anthropogenic emissions related to fossil fuel combustion are generally considered as a regional problem (Aghedo et al., 2007). For example large cities like Lagos (6°35’N, 3°2’E) along the coast of the Guinea Gulf are known to influence air quality and ozone production at local scale (Baumbach et al., 1995; Minga et al., 2010). Most recent emission data bases still
underestimate this contribution and are needed for accurate modelling of aerosol and trace gas distributions in this region (Lioussse et al., 2010).

Although many papers have discussed the link between tropospheric ozone total column variability over West Africa and tropical Atlantic with biomass burning emissions (Thouret et al., 2009; Sauvage et al., 2007b; Thompson et al., 2000), there are few papers about city plume studies in this region. Minga et al. (2010) showed that during the dry season very large amounts of O\textsubscript{3} can be found near Cotonou (2°26′E, 6°60′N) due to the Lagos industrial area. Hopkins et al. (2009) also discussed the impact of the city of Lagos on ozone precursor emission using the data of the UK BAe-146 aircraft recorded during a circular flight around the city in July 2006. The emission levels for NO\textsubscript{x} and CO were found to be high enough to characterise the Lagos area as a polluted megacity, but the fate of these emissions for photochemical ozone production is not discussed in that study. Thus ozone production due to African megacities remains poorly documented especially during the wet season.

In this paper, our objective is to analyse aircraft data collected near three major cities: Cotonou, 2°26′E, 6°60′N, Niamey, 2°05′E, 13°32′N and Ouagadougou, 1°32′W, 12°22′N, focusing on ozone production and the respective impact of anthropogenic emission and Southern Hemisphere (SH) biomass burning. They have been analysed to identify the good conditions to observe ozone plumes related to city emissions during the monsoon season. Aircraft data has the great advantage of providing vertical profiles of O\textsubscript{3} and its precursors, limited to NO\textsubscript{x} and CO observations in this work. The aircraft were deployed during the African Monsoon Multidisciplinary Analyses (AMMA) campaign in July–August 2006 aimed at studying the role of convection on the atmospheric composition (Reeves et al., 2010). During the wet season, photochemical O\textsubscript{3} production in the lower troposphere is limited by frequent fast vertical mixing (Chatfield and Delany, 1990), but during this season there is a better altitudinal decoupling of the respective influence of emissions from biomass burning and from African cities on the regional ozone production. Thouret et al. (2009) showed that the advection of biomass burning from the SH during the wet season is seen at higher altitudes (3–5 km) than during the dry season (1–3 km).

The paper is structured as follows. In Sect. 2, we introduce the meteorological context and differences between Cotonou and Niamey/Ouagadougou. In Sect. 3, the aircraft observations are presented and discussed. Different model results are used in Sect. 4 (i) to demonstrate that we are able to separate the O\textsubscript{3} plumes from city emissions and those related to biomass burning, (ii) to discuss the good meteorological conditions for O\textsubscript{3} plume formation and (iii) to assess an upper limit of the O\textsubscript{3} photochemical production in the city plumes considering the chemical observations available for this campaign. In Sects. 4.1 and 4.2, the air mass transport is studied using Lagrangian model simulations (FLEXPART), to identify whether air masses stayed long enough under the influence of urban emissions to increase O\textsubscript{3} concentrations or if they were influenced by other emissions (Southern Hemisphere, tropical forest). In Sect. 4.3, the vertical structure of ozone in Cotonou is discussed using a tracer simulation from the mesoscale model (BOLAM) to show that biomass burning emissions can be distinguished from the city emissions. The photochemical O\textsubscript{3} production is discussed in Sect. 4.4 using the chemical box model CTTeYCAT. Finally, we discuss the representativeness of the limited number of aircraft measurements in Sect. 5.

2 Meteorological context and characteristics of the selected cities

The synoptic situation over West Africa during the summer monsoon is characterised by the convergence of the Harmattan flux blowing from the Sahara desert to the gulf of Guinea and the monsoon fluxes advecting inland the maritime air masses. The main averaged characteristics of these two flows are shown in Fig. 1 for the 0–1 km and 3–4 km altitude layers during August 2006. The wind data are from the ECMWF reanalysis with assimilated AMMA soundings (Agusti-Panareda et al., 2009). Below 1 km the humid and relatively cold monsoon flow penetrated inland reaching Ouagadougou (Burkina Faso) and Niamey (Niger). Above 3 km, the most striking features are the African Easterly Jet (AEJ), located north of 10°N, with wind speed over 15 m s\textsuperscript{-1} (AEJ-N) and a similar wind flow south of the equator with wind speed up to 9 m s\textsuperscript{-1} (AEJ-S) (Parker et al., 2005; Tompkins et al., 2005). The AEJ-S interacted with the monsoon flow and played an important role in the interhemispheric transport of biomass burning pollution, while the AEJ-N limits the northward extent of such a transport (Mari et al., 2008). The role of the monsoon circulation on the regional ozone and CO distribution is discussed in Reeves et al. (2010), and in this paper we only consider the meteorological factors important for O\textsubscript{3} production resulting from anthropogenic emissions.

According to the weather patterns in the lower troposphere, we distinguish two groups of cities: (i) Lagos or Cotonou mainly influenced by SH emissions and the Atlantic boundary layer chemical composition, (ii) Niamey and Ouagadougou, named after the Sahelian cities which are mainly influenced by the northernmost part of the monsoon flow advecting biogenic emissions from forested regions to the south. This is why we analyse separately the Cotonou observations and the Niamey/Ouagadougou data set. Even though Niamey and Ouagadougou are similar with respect to the weather patterns, we expect different emission factors for these two cities considering their different population sizes (1.2 million for Ouagadougou and 700 000 for Niamey). The RETRO anthropogenic emission inventory at 0.5′ resolution and with a monthly time resolution for year
Fig. 1. Wind velocity (m s$^{-1}$) and vectors from ECMWF reanalysis averaged over August 2006 for the 3–4 km (bottom), and 0–1 km (top) tropospheric layers.

2000 (http://retro.enes.org/) indeed shows CO fluxes three time more important in Ouagadougou than in Niamey (Table 1). During the wet season, numerous mesoscale convective systems (MCSs) can develop over West Africa with a strong impact on vertical mixing of O$_3$ and O$_3$ precursors (Reeves et al., 2010; Thouret et al., 2009). This is known to impact the ozone distribution over the whole troposphere, but also to control the timing of the low level ozone production. Therefore, vertical mixing needs to be taken in to account when discussing the data set.

### Table 1. CO fluxes in µg m$^{-2}$ s$^{-1}$ for the 0.5 × 0.5° grid cell containing each of the 4 West African cities where aircraft data are available. Data are from the RETRO inventory (see http://retro.enes.org/).

<table>
<thead>
<tr>
<th>City</th>
<th>Longitude</th>
<th>Latitude</th>
<th>CO flux</th>
</tr>
</thead>
<tbody>
<tr>
<td>LAGOS</td>
<td>3.396</td>
<td>6.453</td>
<td>5.5935</td>
</tr>
<tr>
<td>COTONOU</td>
<td>2.433</td>
<td>6.367</td>
<td>0.9672</td>
</tr>
<tr>
<td>NIAMEY</td>
<td>2.083</td>
<td>13.667</td>
<td>0.09242</td>
</tr>
<tr>
<td>OUAGA</td>
<td>−1.533</td>
<td>12.370</td>
<td>0.3324</td>
</tr>
</tbody>
</table>

3 Aircraft data analysis

3.1 The data set

The measurements used in this paper were recorded by the French and German Falcon aircraft (named FF20 and DF20 hereafter) when they flew at altitudes less than 3 km just after takeoff or before landing in one of the West African cities. The FF20 data were obtained near Cotonou (2 profiles) and near Niamey (11 profiles), while the DF20 data were collected near Ouagadougou (18 profiles). The chemical species discussed in this paper are mainly O$_3$, CO and NO$_x$ measurements. The specific humidity is also used to described the Planetary Boundary Layer (PBL) structure, especially its diurnal evolution between takeoff and landing profiles. The FF20 and DF20 instruments are respectively described in Ancellet et al. (2009) and in Baehr et al. (2003). The NO$_x$ instrument on the FF20 experienced problems with an ozone leak during the AMMA campaign and NO$_x$ was only measured in Cotonou (2 profiles) and near Niamey on 19 and 20 August (4 profiles). The FF20 CO measurements required frequent zero monitoring and only 8 out of the 11 profiles are available near Niamey because the zero monitoring occurred during the aircraft landing. The DF20 did not measure NO$_x$ concentration, but NO. Since it is important to make the Ouagadougou data set comparable to the Niamey data, we have estimated the NO$_x$ from the measured NO and O$_3$ concentrations assuming that (i) the NO$_x$ partitioning is only determined by the NO$_2$ photolysis and NO + O$_3$ reaction (ii) the measured NO$_2$ photolysis coefficient is of the order of 0.015 s$^{-1}$. There is a 20–30% uncertainty in the NO$_x$ values but good enough to discuss the occurrence of polluted layers with high NO$_x$ values.

The dates and times of the available profiles near Niamey and Ouagadougou are summarised in Table 2. In both cities, 6–7 O$_3$ profiles were obtained after 14:00 UT, i.e. when an O$_3$ plume was more likely to be detected considering the strong diurnal cycle controlling the ozone concentrations in the PBL. This corresponds in both cases to 4–5 different days. So the two datasets are quite comparable.
Table 2. List of the flights near Niamey (Niam.) and Ouagadougou (Ouag.).

<table>
<thead>
<tr>
<th>Day (UTC)</th>
<th>City</th>
<th>Takeoff UTC</th>
<th>Landing UTC</th>
<th>Species measured</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 August</td>
<td>Ouag.</td>
<td>1130</td>
<td></td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>4 August</td>
<td>Ouag.</td>
<td>0845</td>
<td>1215</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>6 August</td>
<td>Ouag.</td>
<td>0930</td>
<td>1245</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>7 August</td>
<td>Ouag.</td>
<td>1220</td>
<td>1515</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>11 August</td>
<td>Ouag.</td>
<td>1450</td>
<td>1750</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>11 August</td>
<td>Niame.</td>
<td>1530</td>
<td></td>
<td>CO$^2$</td>
</tr>
<tr>
<td>13 August</td>
<td>Ouag.</td>
<td>1040</td>
<td>1400</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>13 August</td>
<td>Niame.</td>
<td>1215</td>
<td>1530</td>
<td>O$_3$, CO$^1$</td>
</tr>
<tr>
<td>14 August</td>
<td>Niame.</td>
<td>0630</td>
<td>0930</td>
<td>O$_3$</td>
</tr>
<tr>
<td>15 August</td>
<td>Ouag.</td>
<td>0915</td>
<td>1230</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>15 August</td>
<td>Niame.</td>
<td>1230</td>
<td>1545</td>
<td>O$_3^1$, CO$^1$</td>
</tr>
<tr>
<td>15 August</td>
<td>Ouag.</td>
<td>1430</td>
<td>1700</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>16 August</td>
<td>Niame.</td>
<td>1500</td>
<td>1700</td>
<td>O$_3$, CO$^1$</td>
</tr>
<tr>
<td>16 August</td>
<td>Ouag.</td>
<td>1415</td>
<td>1700</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
<tr>
<td>17 August</td>
<td>Niame.</td>
<td>0930</td>
<td>1245</td>
<td>O$_3$, CO$^1$</td>
</tr>
<tr>
<td>19 August</td>
<td>Niame.</td>
<td>0915</td>
<td>1630</td>
<td>O$_3^2$, NO$_x$, CO$^2$</td>
</tr>
<tr>
<td>20 August</td>
<td>Niame.</td>
<td>1400</td>
<td>1745</td>
<td>O$_3$, NO$_x$, CO</td>
</tr>
</tbody>
</table>

$^1$ measured during takeoff only, $^2$ measured during landing only.

3.2 Analysis of the Cotonou observations: a city on the southern coast

Aircraft observations of O$_3$ allow the analysis of the potential impact of anthropogenic pollutants on photochemical ozone production, provided that one can use vertical profiles east and west of the city of Cotonou. The aircraft data also offer the possibility to address the question of ozone production more precisely by using the vertical profiles of CO and NO$_x$. On 19 August 2006, the FF20 aircraft made a landing at 12:00 UT and a takeoff at 13:45 UT in Cotonou. The aircraft position around the city of Cotonou is shown on Fig. 2a when it was exploring the 0–3 km altitude range. Considering the small time difference between the two corresponding vertical profiles, the comparison provides a good estimate of the local scale horizontal gradients: southwest-northeast gradient below 3 km and northwest-southeast gradient above 3 km. A dropsonde launched when flying over Cotonou at 10:41 UT (Fig. 2a) provides the observed wind vertical profile and the boundary layer structure can be examined using the water mixing ratio. The most remarkable feature looking at the difference between the east and west ozone profiles is the significant O$_3$ increase (+50 ppbv) in the 1–3 km altitude range with mixing ratios reaching 80 ppbv East of Cotonou (Fig. 2b). The water vapour mixing ratio profiles from the aircraft and the dropsonde show that the monsoon layer height extended up to 2.5 km and that it did not change very much during the time period 11:00 UT to 14:00 UT over the spatial domain corresponding to the aircraft measurements. Therefore the southwest-northeast ozone gradient is

![Fig. 2](image-url)
not related to a sharp discontinuity of the boundary layer height as observed sometimes near a coastline. Notice also that below 1 km and above 3 km, the change in the ozone vertical distribution is completely different. The large ozone values above 3 km are now observed in the whole area above the coastline and are known to be related to advection of biomass burning from the SH (Mari et al., 2008). The effect of these emissions on O$_3$ are already seen above 2.5 km in an ozone sonde profile taken on 17 August (see Sect. 5). When the aircraft approached the city from the north, O$_3$ mixing ratios at altitudes below 1 km for the east profile were as low as the 1- to 3-km O$_3$ values in the west profile (20 ppbv). In contrast, O$_3$ at altitudes below 1 km in the west profile increased to values near 50 ppbv when the aircraft came closer to Cotonou. Thus, the black line in Fig. 2a represents the limit between the areas with high (East of the line) and low (West of the line) O$_3$ concentrations. The low ozone values near Cotonou below 750 hPa are also discussed in Reeves et al. (2010) showing the role of the weak ozone production over the ocean and the strong ozone sink over the vegetation area inland.

Looking now at the corresponding CO and NO$_x$ vertical profiles (Fig. 2c) for the east sector in the 1–3 km range, they also indicate larger CO (up to 250 ppbv corresponding to a 100-ppbv increase) and NO$_x$ (up to 5 ppbv corresponding to a 3-ppbv increase). Notice that the shorter lived compound, NO$_x$, followed more closely the structure of the boundary layer with a sharp gradient at 2.5 km. The CO vertical profile is more comparable to the ozone profile. Thus, the ozone increase across the line of Fig. 2a corresponds to a concentration build-up longer than the daily time-scale of the boundary layer height evolution. Above 3 km, the weak northwest-southeast CO gradient is consistent with the impact of the biomass burning emissions. Below 1 km, the relative increase in NO$_x$ up to 2.5 ppbv as the aircraft approached Cotonou again (see Fig. 2a) suggests that the 50 ppbv ozone mixing ratios seen in the lowest altitude range of the west profile are indeed related to the emissions from cities located on the coast. Notice also that the wind profile measured by the dropsonde shows the southwesterly direction of the monsoon flow explaining the southwest to northeast direction of the limit between the polluted and unpolluted air in Fig. 2a. There was also a layer of weak wind between 1 and 2.5 km suggesting that polluted city plumes stayed near Cotonou in this altitude range.

To address more explicitly the question of the respective influence of the emissions from the cities along the coast, namely Cotonou and Lagos, and biomass burning products advected aloft at altitudes higher than 3 km, a more detailed analysis of the transport mechanisms is required. Before discussing this part of our work, it is important to study similar measurements near the two other cities less influenced by the SH emissions.

### 3.3 Analysis of the Niamey and Ouagadougou observations: Sahelian cities

All the CO, O$_3$ and NO$_x$ vertical profiles measured by the aircraft are shown in Fig. 3 for the two cities. In Niamey the CO vertical profiles show a daytime PBL extending generally up to 1.2 km (Fig. 3c). The same feature was obtained using the water vapour vertical profiles (not shown). Although the water vapour profiles indicate that the PBL vertical structure does not change very much from one day to the next, there is a significant variability in the CO mixing ratio ranging from 100 ppbv to 400 ppbv in the 0.5 to 1.5 km altitude range. The occurrence of low values indicates that city emissions were not large enough to maintain a constant CO level around 200 ppbv as was observed near the coast. Nevertheless, freshly polluted plumes were detected by the aircraft near Niamey as shown by the occurrence of high CO values in this altitude range. There were only 2 days with NO$_x$ vertical profiles (19 and 20 August) and they also show plumes with high NO$_x$ values in the 0.5–1.5 km altitude range during landing (Fig. 3e). NO$_x$ mixing ratios were less than in Cotonou but reached 3 ppbv in the plumes.

In Ouagadougou, the CO and NO$_x$ (Fig. 3d and f) profiles show a slightly lower PBL top (<1 km). The CO variability within the PBL is less than in the Niamey data set with mixing ratios ranging between 100 and 250 ppbv. This is a good indication that no well defined polluted plumes advected away from the city at altitudes above 1 km were encountered by the aircraft. The mean NO$_x$ mixing ratios were of the order of 1 ppbv near 1 km, but values as large as 4–5 ppbv were observed near the ground, especially on 16 August. The 1 ppbv NO$_x$ level is similar to the BAe-146 aircraft average NO$_x$ vertical profile observed during the AMMA campaign in the 12°–16° N latitude band (Saunois et al., 2009). Such a NO$_x$ mixing ratio range (0.5–1.5 ppbv) is measured outside city plumes and it is related primarily to NO emission from recently wetted soil (Stewart et al., 2008).

In Niamey, the ozone values were generally lower than 35 ppbv in the free troposphere above the PBL as discussed in Reeves et al. (2010) and the day to day variability is low at this altitude (Fig. 3a). Within the PBL or near its maximum altitude, O$_3$ mixing ratios reaching 50 ppbv were observed in 3 profiles, especially on 16 August when O$_3$ > 70 ppbv. This is very different from Cotonou where there was a positive ozone vertical gradient. This suggests that photochemical O$_3$ production occurred within the PBL in Niamey and not above the 1.5 km because of the latitudinal decrease of ozone precursors due to biomass burning emissions. The day to day variability of the O$_3$ and CO mixing ratios (Fig. 4a) shows that the largest probability of sampling the Niamey plume, i.e. with the largest CO values, was during the period 15 August to 17 August. More high O$_3$ episodes were not observed probably because very specific conditions are required such as air mass stagnation around the city or significant addition of NO$_x$ emissions from wetted bare soil as
discussed by Saunois et al. (2009) using a regional model and observations. The role of the transport in the observed day to day variability is discussed more precisely in Sect. 4.2.

In Ouagadougou, the vertical ozone profiles (Fig. 3b) show a small variability around 30 ppbv at all altitudes. This is consistent with less ozone produced by polluted plumes at altitudes < 1 km. The slightly positive O₃ vertical gradient is different from the Niamey data set and can be related to the more efficient ozone sink over the more vegetated situation in Ouagadougou. Notice also that according to Table 3, the DF20 flights often took place during the days with active convection increasing vertical mixing and preventing ozone

Fig. 3. Vertical profiles of O₃ – top panels (a), (b) – CO – middle panels (c), (d) – and NOₓ – bottom panels (e), (f) – mixing ratio measured by the FF20 near Niamey (left) and the DF20 near Ouagadougou (right).
Table 3. List of Mesoscale Convective System (MCS) derived from Meteosat satellite observations near Niamey and Ouagadougou.

<table>
<thead>
<tr>
<th>Day</th>
<th>Position</th>
<th>Comments on MCS evolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>31 July</td>
<td>2.5° W, 15° N</td>
<td>Travelling over Ouagadougou at 16:30 UT and growing during the day</td>
</tr>
<tr>
<td>4 August</td>
<td>2.5° W, 14° N</td>
<td>North of Ouagadougou in the morning, decays after 09:00 UT but cloudy in the afternoon</td>
</tr>
<tr>
<td>6 August</td>
<td>2° E, 13° N</td>
<td>Large MCS travelling west over Ouagadougou after 12:00 UT</td>
</tr>
<tr>
<td>7 August</td>
<td>2° E, 13° N</td>
<td>A patchy MCS develops over Ouagadougou after 15:00 UT</td>
</tr>
<tr>
<td>11 August</td>
<td>3° E, 13° N</td>
<td>Large MCS over Niamey in the morning and Ouagadougou after 12:00 UT</td>
</tr>
<tr>
<td>14 August</td>
<td>2° E, 14° N</td>
<td>MCS arriving over Niamey at 12:00 UT and decays during the day</td>
</tr>
<tr>
<td>16 August</td>
<td>2° W, 12° N</td>
<td>Small system developing over Ouagadougou during the day</td>
</tr>
<tr>
<td>17 August</td>
<td>2° E, 15° N</td>
<td>Small system forming North of Niamey, decays during the day but cloudy in Niamey until 14:00 UT</td>
</tr>
</tbody>
</table>

Production (Chatfield and Delany, 1990; Reeves et al., 2010). The daily variability of the O\textsubscript{3} or CO mixing ratios (Fig. 4b) confirms that there were no days with conditions favourable to photochemical O\textsubscript{3} production near Ouagadougou, even on 16 August when large NO\textsubscript{x} and CO are encountered. Subsequent O\textsubscript{3} production could have still occurred downwind.

To explain the Niamey O\textsubscript{3} production on 16 August 2006, we unfortunately do not have NO\textsubscript{x} mixing ratio measured by the FF20. However, the NO\textsubscript{x} measured on 19 and 20 August (Fig. 3e) shows that values of the order of 3 ppbv can be found in the afternoon in the upper part of the PBL. Since CO is not very high on 19 and 20 August (Fig. 4a), we expect even higher NO\textsubscript{x} values for the days corresponding to the polluted plume crossing (15 to 17 August). Notice that NO\textsubscript{x} can be as large as 5 ppbv in Ouagadougou on 16 August, showing that high NO\textsubscript{x} can be also recorded in Sahelian cities. Since Stewart et al. (2008) shows that a significant fraction of the 5 ppbv NO\textsubscript{x} level found in West Africa can be attributed to the wetted soil emissions (up to 1.5 ppbv), it is important to consider also the variability of the convective activity. This was at a maximum in the first half of August (Table 3) and slowed down at least over western Niger only after 14 August. So indeed soil NO\textsubscript{x} emissions must be considered during the period 15 August to 17 August in addition to the emissions from the city of Niamey. After 14 August, reductions in local cloudiness and precipitation could have also maintained high levels of NO\textsubscript{x}, HCHO, NO\textsubscript{2} photolysis and production of HO\textsubscript{x} radicals. This could explain the observed ozone increase of 30 ppbv in the polluted plume on 16 August. To explain the low ozone values observed again on 17 August in Niamey, one can notice that new convective activity on this day (Table 3) limited further O\textsubscript{3} plume formation even though the aircraft still measured high CO mixing ratios. Vertical mixing during convective episodes will limit the lifetime of ozone plumes forming near the top of the PBL. The influence of convection and related mixing on surface O\textsubscript{3} in Senegal is discussed in Grant et al. (2008). The impact of NO\textsubscript{x} removal processes on O\textsubscript{3} photochemistry is discussed in Sanderson et al. (2008). The role of the synoptic scale transport must be also accounted for to explain the ozone variability during the 15 August to 17 August period. This will be analyzed in the Sect. 4.

To support the role of the efficient O\textsubscript{3} sink over the vegetation area south of Niamey, one can use all the O\textsubscript{3} aircraft
measurements performed on-board another French aircraft (ATR-42) which often flew in the 11.5° to 14.5° latitude band at low altitude level around Niamey during the same time period (11 to 19 August 2006). The average meridional cross-section (Fig. 5) clearly shows the low O3 mixing ratios (<30 ppb) south of Niamey at altitudes below 1 km. Higher values observed near and to the north of Niamey illustrate the combined role of city and soil NOx emission in the O3 budget as discussed in Reeves et al. (2010).

To summarise, the Sahelian cities also exhibit the potential for ozone production in the city plumes. Differences in anthropogenic emissions between Ouagadougou and Niamey should lead to more O3 pollution episodes in the former. This does not seem to be the case, so factors other than the magnitude of emissions are needed to explain the differences in O3 concentration:

- The occurrence in Niamey of a non-convective period lasting at least 2 or 3 days to reduce vertical mixing and wet removal of NOx reservoir species in NOy but following very active convection and related precipitation necessary to increase NOx soil emissions.

- Significant vegetation cover around Ouagadougou increasing local ozone surface dry deposition.

Different large scale transport mechanisms may also be responsible for the daily variability observed in both places and for the difference of the ozone enhancement in Niamey and Ouagadougou. In the following section we address these issues.

4 Model results

The first object of this section is to provide a characterisation of air mass transport using the FLEXPART Lagrangian particle dispersion model. Other modelling work performed in this study were aimed at demonstrating that: (1) regional ozone production near the coast is not related to the SH biomass burning at altitudes < 3 km, (2) the chemical composition of the air masses influenced by the city emissions is consistent with O3 mixing ratios reaching 70–80 ppbv in a polluted plume staying for two days in the area influenced by these emissions. To answer the first question, a simulation using a mesoscale model was needed to reproduce the detailed dynamical processes of the monsoon layer. The advantage of using a mesoscale model to address the complex interaction of emissions and dynamics in the monsoon layer has been demonstrated by several studies (Vizy and Cook, 2002; Sauvage et al., 2007a; Delon et al., 2008). The second question has been addressed, albeit in a fairly simplistic manner, using runs with a chemical box model initialised with observations. A similar approach was used in the Minga et al. (2010) work with a chemical box model (0D model) to study the ozone production downwind of Lagos and to study ozone production in biomass burning plumes (Real et al., 2010)

4.1 Transport of the air masses in Cotonou (19 August)

The FLEXPART model version 6 (Stohl et al., 1998, 2002) driven by 6-hourly ECMWF re-analyses (T213L91) interleaved with operational forecasts every 3 h, was run for 5 days in a backward mode by releasing 2000 particles in 2 boxes corresponding to the altitude range 2–2.5 km and 3.5–4 km. The horizontal extent of the box was 1° × 1° around the city of Cotonou. FLEXPART used ECMWF analyses with 91 model levels and a resolution of 0.5°. The initial version of the model included the computation of potential vorticity (PV) for each air parcel. The fraction of particles with PV > 2 PVU (1 PVu = 10−6 K kg−1 m2s−1) was calculated for each time step to estimate the probability of transport from or into the stratosphere. Previous studies have shown that a fraction larger than 20% encountered during a 5-days time period corresponds to a significant influence of the stratosphere-troposphere exchange (STE) for the air mass at the release time (Colette and Ancellet, 2006). We modified the FLEXPART model to introduce the calculation of the fraction of particles originating below an altitude of 3 km, for two areas corresponding to different emissions of ozone precursors and CO: the SH (latitude < 0° N, longitude < 40° E) for the biomass burning emission, a 1° latitude band along the coast line of the Gulf of Guinea (latitude ∈ [6° N, 7° N]) for the anthropogenic emissions. The upper troposphere-lower stratosphere (UTLS) fraction was defined as the particles coming from altitudes above 8 km and latitudes > 8° N. These fractions are shown as a function of time in Fig. 6. For the layer with a release between 2–2.5 km, i.e. corresponding
Fig. 6. Fraction of particles being in 3 different regions (see text for the definition of the regions) at each time step of backward FLEXPART runs for 2 releases above Cotonou in the altitude range 2.0–2.5 km (solid line) and 3.5–4.0 km (dotted line). The UTLS fraction (>8 km) is always negligible.

to the layer with the southwest-northeast positive ozone gradient, the fraction of particles remaining in the coastal region is high for at least 3 days, while the transport from the SH remains weak (fraction < 10%), even after 5 days. For the layer above 3 km, local emissions near the coast have little influence compared to biomass burning emissions from the SH, even though the fraction from the SH remains of the order of 10%. The UTLS fraction is always very small ruling out the downward transport of stratospheric ozone, but showing also the limited role of the upper troposphere lightning NOx sources (Huntrieser et al., 2008) in the NOx concentrations below 3 km. Therefore, the FLEXPART simulation shows that the low level southwest-northeast positive ozone gradient is largely related to local emissions near the coast.

4.2 Transport of air masses in Niamey and Ouagadougou

FLEXPART was also run in a backward mode for two boxes located above Niamey and Ouagadougou corresponding to the altitude range 0.5–1.5 km. We again release 2000 particles in the 500 m x 1° x 1° boxes. The fraction at a given time step in two specific regions were also calculated: the 1° latitude band around Niamey or Ouagadougou and the area with more vegetation, i.e. at latitudes < 10°. A large fraction in the first region (named city fraction hereafter) points to the influence of local anthropogenic emissions and also possibly soil NOx emissions provided that the stagnation took place after a convective period. A large fraction in the second region (named southern region fraction hereafter) is useful to determine how advection of biogenic emissions from the forest may change the ozone production (i.e. by increasing isoprene or terpene plant emissions and by changing the HOx radical chemistry). Enhancements in ozone production in a city plume at mid-latitudes have been previously observed when a plume mixed with background air strongly influenced by biogenic species (Derognat et al., 2003). The daily variability of the FLEXPART fraction of particles staying in the 1° latitude band around the city and the fraction of particles influenced by the land emission south of 10° N are reported in Fig. 7. The city fraction is given 24 h before the observations as it is the minimum time period to obtain a measurable ozone enhancement due to city emissions. The southern region fraction is given 48 h before the observations as a longer time period reduces the lifetime of the ozone precursors from biogenic emission.

For Niamey, results show that there are 3 days with city fractions > 20%: 11, 15, and 16 August. The southern region fraction becomes significant after August 14 (>30%) while 16 August is the only day with a stagnation of the air mass near Niamey following advection from the southern region, i.e. the optimum air mass transport conditions for O3 production. This is consistent with the aircraft observations showing the largest ozone enhancements on 16 August in Niamey.

For Ouagadougou, 11 August is the only day with a fraction > 20% in the city latitude band 24 h before the measurements. But, it is also the day when the northward transport was smallest (<10%) and when there was a strong MCS with frequent overcast skies and mixing during the day. The influence of advection from latitudes < 10° is often very high and even dominates the air mass composition on 4 and 6 August. This makes the Ouagadougou vertical profiles more representative of photochemical conditions found above the forest and so less impacted by local city emissions. These findings are consistent with the absence of elevated ozone values and moderate NOx mixing ratios in the DF20 flights compared to the Niamey FF20 flights.

While it is not shown in Fig. 7, the UT air mass fraction is also calculated with the same conditions used in the Cotonou simulation. Results are similar with a negligible influence of transport from the upper troposphere in both cities at altitudes near 1 km.

In Sect. 3.3, we discussed the role of surface ozone deposition and convection in the analysis of the differences between Niamey and Ouagadougou observations, but the FLEXPART analysis also shows that specific synoptic scale transport conditions are also needed to explain the occurrence of ozone rich plumes near Niamey, i.e. the stagnation of air for more than 24 h near the city after advection from the forest region in order to increase the influence of biogenic emissions.
Fig. 7. Daily variability of local stagnation and northward transport from the forest using backward FLEXPART runs with releases at observation time $t_0$ and in the altitude range 0.5–1.5 km above Niamey (right column) and Ouagadougou (left column). Top row (a,b): is the fraction of particles remaining within the 1° latitude band around the city at $t_0–1$ day. Bottom row (c,d): is the fraction of particles coming from latitudes $<10^\circ$N at $t_0–2$ days.

4.3 BOLAM model results

4.3.1 Mesoscale model description and simulations set-up

BOLAM (BOlogna Limited Area Model) is a limited-area meteorological model based on primitive equations in the hydrostatic approximation. Prognostic variables (horizontal wind components, potential temperature, specific humidity and surface pressure) are defined on hybrid coordinates and are distributed on a non-uniformly spaced Lorenz grid. It includes a microphysical scheme that has five prognostic variables (cloud water, cloud ice, rain, snow and graupel), as derived from the one proposed by Schultz (1995). Deep convection is parameterized with the scheme of Kain-Fritsch (Kain, 2004). Vertical diffusion is modelled using the mixing-length assumption and the explicit prediction of turbulent kinetic energy (Zampieri et al., 2005). Further details of the model are provided in Malguzzi et al. (2006).

Here, the mesoscale model BOLAM was used to simulate the transport of southern hemispheric biomass burning and regional anthropogenic emissions over West Africa. To take into account the different spatial and temporal scales involved in the transport, two simulations were performed: a three month run for biomass burning emissions and a shorter and higher resolution run for anthropogenic emissions. For both simulations ECMWF AMMA re-analyses have been used as initial and boundary conditions (Agusti-Panareda et al., 2009).

1. Local anthropogenic emissions simulation (hereafter 8 km-anthro) characteristics are as follows: the simulation covered a period of 5 days, starting on 00:00 UTC of 15 August 2006. The horizontal grid resolution was $0.07^\circ \times 0.07^\circ$ (around $8 \times 8$ km) and 60 vertical levels used with 21 levels below 3 km and the lowest level at 65 m. The domain’s limits were longitude = $[-2.8, 7.6]$ and latitude = $[1.1, 11.5]$. The tracer field was initialised to zero and CO fluxes from the RETRO
inventory (Table 1) were used to inject anthropogenic emissions into the lowest model layer during the whole simulation. High values for emissions are reported for Lagos, while the overall area encompassing Cotonou and Lagos reaches more than 6.5 µg m$^{-2}$ s$^{-1}$). The CO tracer underwent exponential decay with a lifetime of 20 days imposed to match the lifetime of CO in the troposphere (Mauzerall et al., 1998).

2. The biomass burning emissions simulation (hereafter 24 km-bb) characteristics are as follows: the simulation started on 15 June and ended on 31 August 2006. Meteorological fields were re-initialised on 1 July and 1 August. The horizontal grid resolution was $0.216^\circ \times 0.216^\circ$ (around 24 × 24 km) and 38 vertical levels were used. The domain included both West Africa and the area of SH wild fires, domain's limits were longitude = [−22, 40] and latitude = [−20, 29]. The tracer was injected up to 1 km altitude and 5-day averaged CO emissions from the biomass burning emission inventory described in Lioussé et al. (2010), were used for the horizontal distribution of the tracer. An exponential decay was imposed on the CO tracer with a lifetime of 20 days. Even if an exponential decay was imposed and CO emissions were used, the biomass burning tracer is reported in arbitrary units instead of ppbv of CO because the simulation is too long to neglect chemistry.

To discuss the capability of the model to reproduce the local transport near Cotonou, Fig. 8 shows the vertical profile of wind speed, wind direction and relative humidity measured by the dropsonde launched at 10:41 UTC by the FF20 and calculated in the 8 km-anthro simulation at 10:00, 12:00 and 14:00 UTC on 19 August above Cotonou. Measurements show the monsoon layer with a uniform southwesterly wind from ground up to 1.2 km and wind speed with a maximum of 10 m s$^{-1}$ at 0.5 km and decreasing toward 0 m s$^{-1}$ at 1.2 km. BOLAM (black line, 10:00 UTC) reproduces such a layer, underestimating the maximum wind speed. Above the monsoon layer, measurements report a counter-clockwise rotation of 360° of the wind between 1.2 and 2.5 km and then again a southwesterly wind from 2.5 km upward. BOLAM model is able to reproduce the vertical variation of the wind direction and the overall vertical structure of the wind, even if it underestimates wind speed above 2 km and, due to the finite vertical resolution, is not able to catch the sharp increase in wind speed at 2.1 km.

Regarding relative humidity, both the measurements and BOLAM show a boundary layer extending up to 1.9 km with a nearly saturated atmosphere and a sharp decrease toward a value of 30% at 2.5 km. The model shows also very little change in the vertical structure humidity profile as shown by the sonde and aircraft measurements reported in Fig. 2.

Figure 9 shows the time evolution of wind speed and wind direction from 8 km-anthro and ECMWF AMMA reanalyses, averaged between 0 and 1 km and between 1.5 and 3 km over Cotonou. Within the monsoon layer (upper panel) BOLAM shows a high correlation with analyses concerning both wind speed and wind direction. Regarding the 1.5–3 km layer, where the highest differences in ozone mixing ratio were found, there is a good agreement between simulation
and analyses during the whole simulation except for the last day, when the clockwise rotation of the wind is slower in the simulation than in the analyses. This is of small importance for the transport of anthropogenic tracer because the last day of simulation is also characterized by weak winds (<2 m s\(^{-1}\)). Simulated wind fields averaged over the whole simulated period (not shown) also show a reasonable agreement with analyses in both altitude ranges within the monsoon layer.

### 4.3.2 Tracer simulation results

Figure 10 report the horizontal distribution and the vertical cross section of anthropogenic tracer concentration from 8 km-anthro simulation. The horizontal cross section shows that the anthropogenic tracer is transported from the emission regions along the coast (pink squares in Fig. 10). Highest concentrations are related to Lagos emissions. The vertical cross section of tracer concentration shows high values at the ground near Cotonou, where there are some emissions of CO, and a minimum around 0.9 km due to the southwesterly advection of air with low tracer concentrations from the ocean. Above the minimum and up to 2.8 km, a tracer layer with maximum around 2.4 km is visible, in agreement with O\(_3\) and CO profiles influenced by Cotonou city plume (west profiles in Fig. 2). The boundary of the area influenced by anthropogenic emissions is indeed located near Cotonou at the different altitudes below 3 km, supporting the fact that the horizontal gradient at these altitudes are related to anthropogenic emissions in the Lagos/Cotonou area.

In Fig. 11 left panel, the 900 hPa horizontal distribution of biomass burning tracer, from the 24km-bb simulation, shows that around Cotonou (red asterisk in Fig. 11) the tracer remains confined far from the coast. In Fig. 11 right panel, the vertical structure of the tracer over Cotonou (red asterisk) shows that a layer of high concentrations is present from 2.5 km upward. This structure agrees with the observations and also the work of Thouret et al. (2009) that reported frequent ozone enhancements above Cotonou between 3 and 5 km in August 2006 (see Sect. 5). At lower altitudes, the tracer remains confined far from the coast in the simulation in agreement with the observations above the ocean. Nevertheless regional simulations can fail in reproducing the position and strength of the Southern African Easterly Jet and/or the re-circulation above the continent around the equator. This may lead to an uncertainty in the transport of biomass emissions from the SH (Williams et al., 2010; Real et al., 2010).

The BOLAM results then show that the observed ozone local anomaly near Cotonou can be explained by the advection of polluted plumes from the Lagos area and that advection of biomass burning emissions is not related to the occurrence of this anomaly.
4.4 Modelling of the ozone production

In order to estimate if the O$_3$ difference > 40 ppbv between the background composition and the city plume can be produced by anthropogenic emissions in Cotonou/Lagos and Niamey, the CiTTyCAT (Cambridge Tropospheric Trajectory Model of Chemistry and Transport) chemistry box model was used. It is a photochemical model including 90 chemical species (see Evans et al., 2000 for details) and has been previously used to simulate the chemical evolution in the biomass burning plume originating from central Africa and advected towards West Africa in the lower-mid troposphere in 2006 (Real et al., 2010). FLEXPART simulations (see Sects. 4.1 and 4.2) showed that air masses containing the high O$_3$ plume had remained in the city region for at least 2 days. Meteorological conditions 24 h before the flight were generally characterised by low winds and no MCS, so it is a reasonable approximation to neglect mixing and wet deposition, in this case. Dry deposition is also assumed to negligible in this case since the plumes were located above 500 m. Therefore, a 2-day stationary run was conducted using diurnal cycles of temperature and relative humidity corresponding to ground based observations collected during AMMA at this location. Near the coast, the temperature varied between 23° and 27° with relative humidity rangeing from 80 % to 100 %.

Since pollutant concentrations directly emitted over the source region are unknown, FF20 observations of O$_3$ precursors were used to initialize the model. Unfortunately, only CO and NO$_x$ were measured in this plume. Volatile Organic Compound (VOC) initial concentrations were estimated from the mean ratio of VOC/CO measured during the whole campaign by the BAe-146 aircraft at lower altitudes (<3 km), by scaling the VOC to the CO measured in
the high O$_3$ plume by the FF20. To initialise NO$_y$, a typical ratio NO$_y$/NO$_x$ of 2 was used. This is the mean ratio measured during the campaign by the DF20 when encountering high NO$_x$ plumes. NO$_2$ concentrations (NO$_y$ - NO$_x$) was then partitioned between PAN and HNO$_3$, with HNO$_3$ and PAN being respectively 75% and 25% of NO$_x$. This partitioning corresponds to the values observed at low altitudes during the whole campaign by the BAe-146 aircraft. Ozone concentrations measured by the aircraft in the cleanest air were used to initialise the model simulations. Concentrations used in this reference polluted run are detailed in Table 4.

Maximum values of NO$_2$ and O$_3$ photolysis rates are respectively $9.1 \times 10^{-3}$ s$^{-1}$ and $4.3 \times 10^{-3}$ s$^{-1}$ around 12:00 UTC. These values are typical of photolysis rates near the equator, without any clouds (Liao et al., 1999). The model climatological data base for clouds was not used since it is not very well suited for the very complex cloud structure observed in Africa during the monsoon season. This assumption is also justified because no MCS were observed over Cotonou during the two days corresponding to the simulation. What may be missing is the aerosol impact on photolysis rates, as Fig. 11 shows that a biomass burning aerosol layer is present in the free troposphere above the city plume. Our estimation of the O$_3$ production is therefore an upper limit since photolysis rate may be slightly overestimated.

Considering uncertainties in the O$_3$ precursor concentrations estimated in this reference run, sensitivity tests were conducted by multiplying or dividing NO$_x$ and VOC concentrations with a factor 2 and 4. This is also justified by the fact that the aircraft measurements are representative of a processed plume, and not a very fresh plume. When the NO$_x$ concentrations were modified, the NO$_y$ concentrations were also changed using the same scaling factor between these two species. Daily net photochemical O$_3$ production rates obtained from these runs are shown in Fig. 12 for the different NO$_x$ and VOC values. A value of 20 ppbv O$_3$ per day, is estimated for a NO$_x$ range between 5 and 6 ppbv, already for the standard VOC concentrations. Higher VOC concentrations are needed to achieve a production rate as high as 25 ppbv per day for the same NO$_x$ range. The dependance on NO$_x$ concentrations also increases as VOC concentrations increase. For the reference polluted concentrations and NO$_x$ near 4 ppbv, the O$_3$ regime NO$_x$ limited since changing NO$_x$ concentrations by 2 leads to an O$_3$ net production change twice more effective than varying VOC concentrations by 2. Therefore, ozone production rates approaching the enhancements that were observed have been estimated, suggesting that anthropogenic plumes in this region are likely to be very photochemically active due to their high NO$_x$ concentrations.

### Table 4. Chemical initialisation of the box model reference run. Observations used to define the chosen concentrations are indicated.

<table>
<thead>
<tr>
<th>Species</th>
<th>Origin</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
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<td>238 ppbv</td>
</tr>
<tr>
<td>H$_2$O$_2$</td>
<td>FF20 clean air mass</td>
<td>2360 pptv</td>
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<tr>
<td>O$_3$</td>
<td>FF20 clean air mass</td>
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<td>CH$_4$</td>
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<tr>
<td>HCHO</td>
<td>BAe-146</td>
<td>815 pptv</td>
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<td>DF20 NO$_y$/NO$_x$ ratio</td>
<td>2 to 16 ppbv</td>
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<td>25% of NO$_y$-NO$_x$</td>
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<tr>
<td>HNO$_3$</td>
<td>75% of NO$_y$-NO$_x$</td>
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<td>MeCHO</td>
<td>BAe-146</td>
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</tr>
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</table>

### 5 Discussion of data representativeness

Since only one flight was analysed over Cotonou, it is difficult to estimate how often O$_3$ is produced downwind from the city emissions. However, we can compare our data with a more extensive study of ozone vertical profiles in Cotonou by using the 24 ozone sondes record presented in Thouret et al. (2009), corresponding to the entire 2006 wet season.
The 2006 summer mean ozone profile and the data for the sonde launched on 17 August 2006, (i.e. two days before the aircraft measurements) have a similar structure showing a positive ozone vertical gradient, but with ozone mixing ratios exceeding 50 ppbv only at altitude above 3 km (Fig. 13a). The fact that the mid-troposphere O₃ maximum is still visible in the mean is consistent with the SH biomass burning origin of the O₃ plume (Mari et al., 2008). Out of a total of 24 profiles, 8 were clearly influenced by convection (low O₃ values in the upper troposphere) and 7 show a 3- to 5-km O₃ maximum linked to advection of biomass burning emissions. Ozone mixing ratios larger than 50 ppbv at altitudes below 2.5 km are not very common in this dataset because the Cotonou ozone sonde launch site is located west of the city centre and there is a mean southwesterly flow near the coast (Fig. 1). When the wind direction shifts to an easterly flow, as for 30 June, 10:00 UT sounding, a significant O₃ increase can be observed in the monsoon layer below 3 km (Fig. 13a) in the balloon data set. So when convection does not limit the O₃ production (30 % of the profiles) and provided O₃ is measured downwind of the city, ozone values exceeding 50 ppbv can be encountered near Cotonou.

To discuss the representativeness of the Niamey aircraft data, one can use the DMI ozone sonde record obtained in Niamey from 26 July to 25 August. The O₃ profiles are shown in two different figures for convective (Fig. 13b) and non-convective days (Fig. 13c) using the MCS positions and developments near Niamey as indicated by the hourly maps of the Meteosat observations available on the AMMA web site. Days with convection represented 40 % of the 27 profiles. The O₃ variability is larger for the days without convection and there are 3 days (10 % of all observations) with O₃ > 45 ppbv in the 0–2 km altitude range. The Niamey aircraft data resemble more the O₃ vertical profiles of the non-convective days. For these days, the fraction of low O₃ values (<30 ppbv) is of the order of 50 %. It probably corresponds to the fraction of days with recent advection from the regions south of Niamey where O₃ is reduced by deposition over the vegetation (see Fig. 7). Using the fraction of days without convection (60 %) and assuming less than 50 % of the profiles with a stagnation over Niamey, puts an upper limit of 30 % on the number of days where O₃ enhancements.

6 Conclusions

Aircraft observations obtained near 3 cities in the altitude range 0–3 km have been examined for evidence of ozone enhancements related to anthropogenic emissions in West Africa during the wet monsoon season. Ozone mixing ratios outside the city plumes can be as low as 20 ppbv in air masses advected at low altitudes from vegetated areas or from the ocean. Our aircraft observations showed two episodes with O₃ mixing ratios exceeding 70 ppbv in well defined plumes near the top of the PBL. To identify the good meteorological
conditions necessary for the formation of these O3 plumes, air mass transport was calculated with the FLEXPART model and the BOLAM mesoscale model. The BOLAM model was run with two different tracers for anthropogenic emission and biomass burning emissions to separate the ozone enhancement due to the transport of SH emissions and due the emissions from cities located near the observations. Both FLEXPART and BOLAM simulations show that the biomass burning plume remained above 3 km while the city emissions controlled the chemical composition in the 0.5–2.5 km altitude range. In Cotonou, it is also consistent with the NOx profile showing a maximum of 5 ppbv near 2 km. In Niamey NOx can only be estimated from measurements obtained at the end of the campaign, showing mixing ratios in the 3–5 ppbv range. Wetted bare soil NOx emissions in the Niamey latitude band during convective activity (1 to 14 August) can explain up to 1.5 ppbv of the observed NOx according to (Delon et al., 2008) and (Stewart et al., 2008). The remaining amount of NOx (1.5 ppbv) must be related to the anthropogenic emissions in Niamey.

In Niamey and Ouagadougou, the daily variability of ozone and CO can be explained with the results of the FLEXPART analysis showing a correlation of ozone and CO enhancements for air mass stagnation near the city for 1–2 days and advection of emissions from the vegetated areas. A lack of ozone enhancements is always correlated with the occurrence of deep convection near the city identified on the METEOSAT satellite images. This explains the differences between Niamey and Ouagadougou where no ozone enhancements were observed even for elevated CO values. The larger vegetated area around Ouagadougou also implies a larger O3 surface deposition.

To verify that photochemical ozone production can reach 20 ppbv per day for typical NOx and CO conditions encountered near West African cities, simulations were carried out with the CIITyCA Lagrangian model and using observed average PBL chemical composition data reported by other aircraft during AMMA. Such a production is possible with NOx mixing ratios reaching 5 ppbv.

The limited number of observations makes it difficult to derive general conclusions about the role of West African cities to the O3 budget during the wet season. In fact, the main contribution of the paper is to identify the conditions needed to observe ozone enhancements related to anthropogenic city emissions. A more extensive ozone data set is available for Cotonou and Niamey when using ozonesondes. The analysis of the sonde data confirms that the very specific conditions established using the aircraft observations are also valid for the ozonesonde analysis, namely the role of convection. The analysis of the statistics of these conditions is not within the scope of this paper but could be based in future work on BOLAM simulations using tracers provided that the capability of the model to reproduce the convection variability is established. Despite the very specific conditions needed to observe ozone enhancements during the wet season, this study show that west african cities can be a substantial contributor in the ozone production balancing the efficient ozone surface deposition at the regional scale.

Appendix A

Photochemical ozone production

Many atmospheric chemistry textbooks describe the ozone photochemistry. In this appendix, the main features are very briefly discussed. Ozone can be formed in the atmosphere by chemical reaction involving NOx and non methane hydrocarbons (NMHC). The complex role of the NMHC chemistry is often simplified by considering the following cycle leading to ozone production where NMHC is abbreviated by RH:

\[
\begin{align*}
2(NO_2 + h\nu (< 430\text{nm}) &\rightarrow NO + O_3^3P) \\
2(O^3P + O_2 + M &\rightarrow O_3 + M) \\
RH + OH &\rightarrow R + HO_2 \\
R + O_2 + M &\rightarrow RO_2 + M \\
RO_2 + NO &\rightarrow RO + NO_2 \\
RO + O_2 &\rightarrow R'CHO + HO_2 \\
HO_2 + NO &\rightarrow OH + NO_2
\end{align*}
\]

In absence of RH oxidation by OH, O3 is converted back to O2 through its reaction with NO and through its photolysis, then leading to a null cycle. Without NOx, ozone reacts with HO2 to recycle OH and ozone is chemically destroyed. This is why NOx and RH are considered as ozone precursors. The NO2 lifetime in the atmosphere depends on the conversion of NO2 to other form of reactive nitrogen like HNO3, its major oxidation product:

\[
\begin{align*}
NO_2 + OH + M &\rightarrow HNO_3 + M \\
HNO_3 + h\nu (< 320\text{nm}) &\rightarrow NO_2 + OH
\end{align*}
\]

The loss of the total reactive nitrogen, often called NOy, is then important for the net O3 production rate as it implies a true loss of NOx. The NOy loss is mainly related to wet and dry deposition.

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