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Overview: oxidant and particle photochemical processes above a south-east Asian tropical rainforest (the OP3 project): introduction, rationale, location characteristics and tools

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Abstract. In April–July 2008, intensive measurements were made of atmospheric composition and chemistry in Sabah, Malaysia, as part of the “Oxidant and particle photochemical processes above a South-East Asian tropical rainforest” (OP3) project. Fluxes and concentrations of trace gases and particles were made from and above the rainforest canopy at the Bukit Atur Global Atmosphere Watch station and at the nearby Sabahmas oil palm plantation, using both ground-based and airborne measurements. Here, the measurement and modelling strategies used, the characteristics of the sites and an overview of data obtained are described. Composition measurements show that the rainforest...
site was not significantly impacted by anthropogenic pollution, and this is confirmed by satellite retrievals of NO$_2$ and HCHO. The dominant modulators of atmospheric chemistry at the rainforest site were therefore emissions of BVOCs and soil emissions of reactive nitrogen oxides. At the observed BVOC:NO$_x$ volume mixing ratio ($\sim$100 pptv/pptv), current chemical models suggest that daytime maximum OH concentrations should be $\sim 10^5$ radicals cm$^{-3}$, but observed OH concentrations were an order of magnitude greater than this. We confirm, therefore, previous measurements that suggest that an unexplained source of OH must exist above tropical rainforest and we continue to interrogate the data to find explanations for this.

1 Introduction

Tropical and equatorial forests account for over half of the World’s forests (1.8 billion ha) and act as a massive source of matter and energy to the lower atmosphere. They exhibit some of the most dynamic yet poorly understood biogeochemical behaviour on Earth. This behaviour is driven by solar radiation and is largely mediated by its transformation into latent and sensible heat, with the concomitant uptake of carbon by photosynthesis and the associated emission of reactive, less-reactive and un-reactive trace gases, water vapour and energy into the atmosphere. Simultaneously, ozone and other trace gases, aerosol particles, and momentum, are lost to the forest surface. A further important consequence of the large solar radiation flux in the tropics is the very vigorous convective uplift that occurs, which results in the rapid movement of chemical species emitted at or near ground level into the free troposphere, as shown, for example, in Surinam (Andreae et al., 2001). Hence reactive trace gas emissions from the surface in the tropics may take part in chemical processes at greater distances and at higher altitudes from their sources than might otherwise occur.

Globally, tropical and equatorial forests are estimated to account for almost half of all biogenic reactive volatile organic compound (VOC) emissions into the atmosphere (Guenther et al., 1995, 2006: global total 1150 Tg C/y, estimate for tropical forests $\sim$500 Tg C/y). These compounds are believed to play a major role in mediating the chemistry of the atmosphere, yet their roles in controlling chemical budgets and processes in the atmosphere on the local, regional and global scales are poorly understood, with considerable and surprising gaps and uncertainties in knowledge remaining (e.g. Lelieveld et al., 2008). In addition, it is possible that biological primary and secondary organic particles play a pivotal role in the formation of cloud condensation nuclei (CCN) and thus control precipitation patterns in forested regions (Barth et al., 2005).

Most previous work on the interactions between tropical forests and atmospheric composition has been carried out in Amazonia (e.g. the LBA project: Andreae et al., 2002; Avisar et al., 2002), with less in Africa (e.g. the AMMA project: Redelsperger et al., 2006) and very little in SE Asia. Unlike the LBA and AMMA domains, which are contiguous continental regions, the complex mosaic of tropical seas and islands that exists in SE Asia makes the likely atmospheric chemistry occurring there somewhat different to that elsewhere. Structurally and floristically, the lowland dipterocarp forest of SE Asia is very different to the rainforest of Amazonia, and it is not known what differences this may cause in the speciation and rates of emission of VOCs and hence in atmospheric composition and chemistry. Furthermore, there is strong evidence that transport from the boundary layer in this “maritime continent” region into the upper troposphere, and possibly subsequently into the stratosphere, is particularly efficient (Fueglistaler and Haynes, 2005), so that the region’s importance to global atmospheric processes may be disproportionately large.

In common with the other tropical forest regions, SE Asia is undergoing very rapid, and in some cases catastrophic, rates of land use change. For example, in Malaysia, the area of total land cover dedicated to oil palm plantations has increased from $\sim$1% in 1974 to $\sim$13% (FAO, 2005; MPOC, 2008). In spite of attempts to implement policies to conserve rainforest, logging of this dwindling resource continues at a rapid rate, and natural forests are being replaced by crop monocultures.

The multi-national OP3 (“Oxidant and particle photochemical processes above a south-east Asian tropical rainforest”) project had the goal of better understanding the interactions that exist between natural forests, atmospheric composition and the Earth’s climate system. The project had the specific objectives of (a) understanding how emissions of reactive trace gases from a tropical rainforest mediate the regional scale production and processing of oxidants and particles, and (b) better understanding the impacts of these processes on local, regional and global scale atmospheric composition, chemistry and climate. By very closely coupling ground-based and airborne measurements of surface fluxes and atmospheric composition of reactive trace gases and particles with modelling studies of chemical processes, the project aimed to address the following questions:

1. What are the rates of transfer of organic compounds emitted from the tropical forest?
2. How are these organic compounds chemically processed immediately after release?
3. To what extent do the regional organic emissions contribute to the atmospheric aerosol in the region, and what are the effects of the aerosol? What is the composition of the organic fraction of the aerosol?
4. What are the effects of these biogenic emissions on global chemistry and climate?
The OH radical initiates the oxidative degradation of biogenically emitted VOCs, and its concentration defines the rate of production of secondary products. A consistent and important finding from field studies conducted in forested environments, characterised by high emissions of isoprene and low levels of NOx, is the significant underestimation of OH by models (Lelieveld et al., 2008; Ren et al., 2008; Butler et al., 2008; Carslaw et al., 2001; Martinez et al., 2008; Tan et al., 2001; Kubistin et al., 2009). These model underestimations scale with isoprene concentration and indicate a current inability to correctly describe isoprene oxidation. The OP3 project provided an excellent opportunity to confirm these findings and to seek an explanation. Atmospheric chemistry models, constrained to measured isoprene emission rates, predict dramatic reductions in ambient OH concentrations in forested areas, in contrast to observations and, as a consequence, predict unrealistically high concentrations of other trace gas constituents (Guenther et al., 2008). Simultaneous measurements of the OH concentration, isoprene concentration and fluxes and isoprene oxidation products were made during OP3, together with many species that control the rate of production and destruction of OH, providing a stringent set of model constraints to investigate in detail any modelled/measured discrepancies for OH.

Similarly, current models suggest that secondary organic aerosol (SOA) in the tropics is dominated by biogenic aerosol (e.g. Kanakidou et al., 2005), but the measurement database is sparse. Emerging first measurements by aerosol mass spectrometry indicate that sub-micrometre organic aerosol concentrations are at the lower end of the model estimates, with median concentrations of around 1 μg m⁻³ observed in subtropical West Africa and Amazonia (Capes et al., 2009). Despite recent progress, our picture of the formation processes of biogenic SOA (BSOA) is still far from complete (Hallquist et al., 2009). Again, the suite of measurements during OP3 was designed to improve our understanding of the levels, composition and formation processes of BSOA in the SE Asian domain.

As described below, the focus of activity was the Global Atmosphere Watch (GAW) station at Bukit Atur, Sabah, Malaysia, on the island of Borneo (4°58′49.33″N, 117°50′39.05″E, 426 m a.s.l.) (http://gaw.empa.ch/gawsis/). Two field campaigns were held during the periods 7 April–4 May 2008 (OP3-I) and 23 June–23 July 2008 (OP3-III). During OP3-III, the UK’s largest atmospheric science research aircraft, a converted BAe 146–301, was based at Kota Kinabalu International Airport during the period 8–23 July 2008 and operated for over 60 h over northern Borneo. Between these two forest campaigns, a sub-set of instruments were deployed in an oil palm plantation, where measurements were made during the period 11 May–20 June 2008 (OP3-II).

In this paper, the land use, vegetation and climate characteristics of the ground-based measurement sites are described, together with an overview of the chemical climatology of the region. The measurement and modelling tools used in the project are also described, as are some preliminary conclusions.

2 Climate, weather, land use and vegetation of Sabah

2.1 Equatorial climate and forest formations

The equatorial tropics are characterised by rain throughout the year, i.e., an absence of marked seasonal droughts. This climatic regime covers: (i) Malaysia, Papua New Guinea and much of Indonesia within tropical monsoon Asia, (ii) coastal regions of Liberia, Nigeria and Cameroon, and central Congo in Africa, and (iii) western Amazonia and a belt extending from the western Caribbean coast to the Pacific coast in Ecuador in tropical America (McGregor and Nieuwolt, 1998; Walsh, 1996). The equatorial tropics can be further classified into tropical superwet, tropical wet and tropical wet seasonal, using a perhumidity index, based on a cumulative annual score of the number of months with >200 mm (+2 index value), 100–199 mm (+1), 50–99 mm (−1) and <50 mm (−2) of rainfall (Walsh, 1996). The extent of tropical superwet and wet climates (with a perhumidity index >10) shown in Walsh (1996) is similar to the extent of tropical climates of Asia and America lacking marked dry seasons shown in McGregor and Nieuwolt (1998).

With the exception of the central Congo and the western Caribbean, there is a good correspondence of regions with a tropical wet or superwet climate and the extent of tropical lowland evergreen broadleaf rainforest. This forest formation dominates within the majority of the Asian, West African and American wet/superwet zone of the equatorial tropics (Whitmore, 1998), and is the most common forest formation in the tropics as a whole (Schmitt et al., 2008). However, in areas locally above 750–1200 m altitude, lowland evergreen broadleaf rainforest grades into lower montane and then upper montane forest. Such areas of mountain forests are noted particularly in the wet/superwet zones of the Asian tropics. Within low-lying areas of the wet and superwet zone, peat swamp, freshwater swamp and heath forest are also present. In areas of podzolic sands, limestone or ultrabasic rocks other forest formations are developed locally.

The key exception to the link between climate and the extent of lowland evergreen rainforest is found within central Congo and the western Caribbean, where tropical semi-evergreen rainforest dominates in areas classified as tropical wet. This semi-evergreen forest formation is important throughout the surrounding seasonal tropics of continental tropical Asia, north-east Australia, and eastern and southern Amazonia (Whitmore, 1998).

Within the equatorial tropics, the dominant lowland evergreen broadleaf rainforest is characterised by a lofty (45 m or taller) and dense canopy with a large number of different tree species occurring together. Usually over two thirds of the upper canopy comprises tree species not contributing
more than 1% to the total number (Whitmore, 1998). The soils typically associated with the occurrence of this forest formation in tropical Asia are the Ultisol group, and within western Amazonia the Oxisol group (Baillie, 1996; Chappell et al., 2007).

2.2 Local climate and forest formations

The majority of the island of Borneo (total area 743 330 km²) has a superwet climate (Walsh, 1996) and the most extensive forest formation is lowland evergreen broadleaf rainforest, occupying some 257 000 km² (Schmitt et al., 2008). A similar situation is observed at the 76 115 km² regional scale of the state of Sabah, Malaysian Borneo. Within Borneo Island, and elsewhere within equatorial Asia, the lowland evergreen rainforest typically has a tree family dominance of Dipterocarpaceae (Whitmore, 1984). Most of the state of Sabah was once covered with rainforest (Schmitt et al., 2008), particularly such mixed dipterocarp forest. Currently, some 47% (36 049 km²) of the state lies within Permanent Forest Estate (PFE; Fig. 1a). Most of this PFE (74%) is maintained under a selective harvesting system (PFE Production Forest), while the remaining 26% is classified as PFE Protection Forest. Within eastern Sabah, most of the cleared lands are now used for the cultivation of oil palm trees (Fig. 1b). Some commercial timber plantations are also present within Sabah.

The Bukit Atur GAW tower used for OP3 sampling is located within the PFE Production Forest of the Ulu Segama – Malua Forest Reserve, but is less than 5 km east of the 438 km² area of PFE Protection Forest known as the Danum Valley Conservation Area (DVCA). The Ulu Segama – Malua Forest Reserve is 2411 km² in area and is divided into annual timber harvesting coupes. The GAW tower lies at the centre of the 22.6 km² “Coupe 88” which was subjected to selective timber harvesting in 1988. An average of 96 m³ of timber ha⁻¹ was cut by both tractor and high-lead harvesting (Tangki and Chappell, 2008). The central area of this coupe was subsequently rehabilitated by enrichment planting (Moura-Costa, 1996). As part of a study covering a 225 km² area Tangki and Chappell (2008) calculated an average tree biomass of 172 t ha⁻¹ for Coupe 88, (using five inventory plots surveyed in March 1997), and demonstrated a strong correlation ($r^2=0.76$) between such coupe-averaged values and Landsat-5 TM band 4 (near-infra red) radiance. Tangki (2008) demonstrated that Dipterocarpaceae were the most abundant tree family recorded for the 225 km² area as a whole (comprising of only lowland evergreen broadleaf rainforest), but Euphobiaceae had become more abundant in Coupe 88 following timber harvesting.

Within the immediate vicinity of the Bukit Atur GAW tower, we calculate the average perhumidity index to be 22 (using the daily rainfall data for the period 1996–2008 from the Danum Valley Field Centre (DVFC), located 8 km from Bukit Atur at 4°58’ N, 117°48’ E, 100 m a.s.l.). Hence Bukit Atur is classified as having a superwet climate. The index did however range from 9 (1997) to 23 (1999) as a result of the 4–5 year cycle in the rainfall caused by El Niño South Oscillation phenomena (Chappell et al., 2001). During the OP3 campaign year of 2008, the perhumidity index was 22, and therefore identical to the longer-term average.

The 23-year mean rainfall (1986–2008 inclusive) for the DVFC rain gauge is 2840 mm (±438 mm standard deviation). The wettest month is typically January with 310 mm precipitation. April is typically the driest month with 155 mm on average, but it also has the most variable rainfall total with a coefficient of variation (CV) of 69% against
a mean monthly CV of 46%. Indeed, the three-month period from February to April has the least predictable rainfall totals, with CV values all over 57%. This variability in rainfall totals may relate to the period being at the change from the northeast monsoon (approximately November–April) to the southwest monsoon (approximately May–October: Bidin and Chappell, 2006).

Rainfall in 2008 totalled 3220 mm, the fifth wettest year in the 23-year record with 113% of normal rainfall. The magnitude of the seasonal variations in rainfall was the smallest on record with a CV of 23%, against the long-term average monthly CV of 47%. This lack of marked seasonality was also shown in the number of days with rainfall. A total of 257 rain-days, the largest number in the 23-year record, were observed in 2008 against an average of 226 rain-days.

2.3 Campaign meteorology

Most of the OP3 measurements were undertaken within the four-month period of April to July 2008. This period was 124% more wet than normal, with 1045 mm of rainfall. Notably, the driest month according to the longer-term record, April, received 170% of the normal rainfall at 263 mm. The April–July 2008 period was also cooler, with a mean temperature of 27.1 °C, which was 99% of the norm for April–July 2001–2008.

A clear diurnal cycle in the rainfall is observed, even within the records of the relatively short OP3 campaign period of April–July (Fig. 2a). The presence of a late afternoon peak in rainfall at DVFC, which is more pronounced when several years of data are summarised, as in Bidin and Chappell (2006), results from the diurnal development of convective rainfall cells, which is consistent with LIDAR observations (peak rainfall typically observed around 15:00 LT) and measurements of radiation and heat fluxes at the site (Pearson et al., 2010; Helfter et al., 2010). The predominance of rainfall delivery by convective events also results in an extreme localisation of the rainfall field. Within a 5 km² region encompassing the summit of Bukit Atur, Bidin and Chappell (2003) demonstrated that inter-gauge correlation in annual rainfall totals fell to 0.90 over distances of only 1.1 km, which is short even by comparison with other convective systems.

Figure 2 also shows median temperatures at various heights at Bukit Atur for the combined periods of OP3-I and OP3-III, measured with aspirated thermocouples. The overall median temperature at 30 m was 25.1 ± 1.6 °C. This is similar to the long-term surface temperature data from DVFC, taking into account a typical temperature gradient with altitude. During the campaign, the atmospheric stability varied from strongly stable at night to strongly unstable during the middle of the day, as would be expected for a convective region with low wind speeds.

When the boundary layer was stable, the rainforest canopy was decoupled from the overlying atmosphere, resulting very frequently in nocturnal radiation fogs. LIDAR data (not shown) demonstrate this, and also suggest a day-time (10:00–18:00 LT) mixing height of ∼800 m. The strong
daytime turbulence, along with weak winds, makes chemical box-modelling a reasonable strategy for interpretation of daytime atmospheric composition (Sect. 5, below), but understanding measurements made at night, when the atmosphere near the surface is strongly stratified, requires more careful consideration of vertical mixing (Pugh et al., 2010a).

The bottom panel of Fig. 2 shows the diurnal cycle in photosynthetically active radiation (PAR), measured at canopy-top height. There is a tail of dull days (shown by the 5th percentile) but, on the whole, the period of the campaign was bright. The warm temperatures and bright sunshine produced substantial emissions of biogenic volatile organic compounds (BVOCs) from the forest (Sect. 4.2.2, below).

Turning from vertical mixing/convective effects, to take a more horizontal/advective view, backwards air mass trajectories were analysed in order to characterise the origins of chemical species measured at Bukit Atur during the measurement campaigns. They were calculated by the British Atmospheric Data Centre (BADC) Web Trajectory Service using European Centre for Medium-Range Weather Forecasts (ECMWF) wind fields. A series of trajectories was calculated, with one trajectory for every hour during OP3-I and OP3-III. These were then analysed to give an ensemble representation of air mass residency time as a function of location for the whole of each campaign (Ashbaugh et al., 1985).

Back trajectories are calculated for the 24 h before arrival at Bukit Atur, with a time resolution of 30 min, and a final pressure altitude of 950 hPa. Back trajectories that touch the ground have been removed. Figure 3a and b show air mass residency time on a $0.1^\circ \times 0.1^\circ$ grid for all back trajectories from the first and third campaign periods, respectively.

The first campaign period (OP3-I) was influenced by air masses from most directions, in contrast to OP3-III when the air was predominantly from the south. The southern air in the third period can be split into two main areas of origin: the SE air from the sea with a minimal fetch over land, and the SW air which is exclusively over land. This can be used to identify and compare periods of marine and terrain influenced air. It is possible to extend this analysis by using only trajectories from periods when a certain measurement is elevated, giving a coloured probability distribution of the source of the measured quantity. A subsequent paper will use these techniques to present a more in-depth analysis of chemical origins in the future.

2.4 Land-cover classification and VOC emissions modelling

Biogenic VOC emissions are highly sensitive to land-cover characteristics and can vary over several orders of magnitude across different landscapes. This is partly due to variability in total biomass density but is greatly enhanced by variability among different plant species, especially for compounds such as isoprene that are emitted by less than a third of all plant species. This presents a daunting challenge for attempts to characterize regional BVOC emissions, especially in highly diverse tropical forests.

Guenther et al. (1995) estimated global biogenic VOC emissions using the highest resolution ($0.5^\circ$) and most detailed global map of land-cover and land-use (Olson, 1992) available at that time. Figure 4 (top left panel) illustrates the Olson (1992) classification of Borneo landscapes, which include six unmanaged land-cover types, dominated by tropical rainforest (50%), marsh/swamp (14%) and tropical montane (7%) and three managed land-cover types dominated by re-growing woods (9%) and paddy rice (7%). The associated isoprene emission factor map shown in Fig. 4 (top right panel) characterizes the gross features of Borneo but does not represent the full diversity of landscapes in this region. A major limitation of the Olson global ecosystem approach is that, for example, all re-growing woods are lumped together and observations from North American and European forests...
were used to assign a single isoprene emission factor to all occurrences of this land-cover type across the globe. The Guenther et al. (1995) estimate of annual isoprene emission from Borneo is about 10 Tg of carbon, which is 2% of the estimated global total.

The availability of satellite observations has greatly improved quantitative estimates of eco-region distributions and other land-cover variables, including leaf area indices (LAI) and plant functional type (PFT) cover fractions. In addition, the development of a global geo-referenced eco-region map by Olson et al. (2001) represents an additional major advance for biogenic emission modeling. This high resolution digital map is the product of over 1000 biogeographers, taxonomists, conservation biologists and ecologists from around the world. Each of the 867 eco-regions represented on this map has relatively uniform species composition and is accompanied by an online database (http://www.nationalgeographic.com/wildworld/) that includes a description of the dominant plant species. Figure 4 (lower left) shows that this database divides Borneo into seven eco-regions with Borneo lowland rainforests covering just over half of the total land area, Borneo montane rainforests and Sundaland heath forests together comprise about 25%, and the remaining four ecoregions (Kinabalu montane alpine meadows, Borneo peat swamp forests, Borneo freshwater swamp forests, Sunda Shelf mangroves) each make up 1 to 8% of the total. While the total area associated with broad types (e.g. tropical forest, montane forest) agree reasonably well with the Olson (1992) database, the details differ considerably. Figure 4 (bottom right) shows that land-cover results in considerable differences between the isoprene emission factor distribution of Guenther et al. (1995) and the MEGAN model (Guenther et al., 2006) which uses the Olson et al. (2001) eco-region map and satellite derived PFT (e.g. crop, broadleaf tree, shrub) cover fractions. In addition, MEGAN uses plant species composition estimates from the Olson et al. (2001) global terrestrial eco-region database and the Leff et al. (2004) crop species distribution database. Although the spatial pattern is quite different, the annual isoprene emission for Borneo differs by less than 5% when the MEGAN land-cover is used in place of Guenther et al. (1995) land-cover data. This good agreement, however, is the result of major offsetting differences between these two land-cover databases. The Guenther et al. (1995) foliar density and LAI estimates are about 50% higher, resulting in about 25% more isoprene, but the emission factors are about 25% lower. The result is a very similar annual isoprene emission but for different reasons.

The MEGAN framework can be used to estimate regional to global biogenic VOC emissions but the accuracy of the results is dependent on the availability of representative measurements of individual ecoregions. A lack of BVOC
measured emissions from Borneo resulted in the assignment of MEGAN version 2.1 emission factors to the ecoregions of Borneo that were based on observations from other tropical regions. Improved estimates for future versions of MEGAN and other models are highly dependent on the availability of observations characterizing the dominant plant functional types within major global ecoregions. In fact, our initial analysis of emissions from Bukit Atur show that the default base emission rates in MEGAN prior to the OP3 observations are a factor of four too high for this forest ecoregion. We also confirm that, unusually for a species classified as crop in MEGAN, oil palm is an intense isoprene emitter.

3 Measurement strategy and methods

The overall measurement strategy for OP3 was to perform integrated measurements from the forest floor, through the forest canopy, above the canopy and then up-scaled to the regional scale using airborne measurements, with clear linkages between measurements made at the different sites at the different scales.

3.1 Ground based measurements

Ground based measurements were centred on the 100 m tower at the Bukit Atur GAW station (4°58′49.33″ N, 117°50′39.05″ E, 426 m a.s.l.) (http://gaw.empa.ch/gawsis/). An “in-canopy” sampling site was established 2 km E of the Bukit Atur tower (4°58′49.10″ N, 117°51′19.12″ E) and a further “oil palm plantation” sampling site was established at the Sabahmas oil palm estate, 70 km NE of Bukit Atur (5°14′58.69″ N, 118°27′15.76″ E).

3.1.1 Forest floor (soil) NO\(_x\) flux measurements

The fluxes of NO\(_x\) from forest soil were measured using a continuous automated dynamic chamber system at the “in-canopy site”. Seven spatially distributed chambers were used in order to represent the spatial variation inherent in soil NO\(_x\) emissions at each site. The chamber construction and operation is described in Pilegaard (2001) while the flux calculation method was modified from Conrad (1994), with further details available in Dorsey and Gallagher (2010). In addition, fluxes of nitrous oxide (N\(_2\)O) and methane (CH\(_4\)) were measured at the “in-canopy site”, at a near-by undisturbed primary forest site, at a heavily disturbed road-side site and from the soils at the oil palm plantation (see below), using static soil chambers (Siong et al., 2009; Ryder et al., 2010).

3.1.2 Within-canopy concentration profiles

At the “in-canopy site”, measurements were made of the vertical gradients (from the ground to 32 m height) of the concentrations of ozone, NO\(_x\), volatile organic compounds, particle size distribution and composition, as well as temperature, relative humidity, radiation and turbulence. Four platforms were strapped against an emergent tree (Canarium decumum), at 8, 16, 24, 32 m. Each platform supported sonic anemometers, inlets to a gradient system for O\(_3\)/NO/NO\(_2\) and fine thermocouples. In addition, an automated winch system continuously raised and lowered a temperature/humidity probe, a PAR sensor, an optical particle spectrometer (GRIMM 1.08) and an inlet leading to a PTR-MS. This gave vertical gradient measurements between 2 and 28 m (Ryder et al., 2010). In a clearing near the in-canopy site, LIDAR measurements were made of wind speed and direction and of aerosol backscatter throughout the boundary layer.

3.1.3 Concentration and flux measurements at Bukit Atur

The largest number of ground-based measurements were made at the Bukit Atur GAW station, which routinely records CO\(_2\) and O\(_3\) mixing ratios, and various aerosol parameters. The station consists of a main building with four air-conditioned laboratories at the base of a 100 m tower, all located in a large (~150×50 m) clearing on the top of a hill and surrounded by forest. The surrounding forest canopy extends ~10 m above the base of the tower. Four mobile sea-container laboratories were deployed around the base of the tower to provide extra instrument accommodation. Electrical power was provided by generators, located 2 km E of the station. Pollution events, attributable to individual vehicles arriving on site, the generators, or to small leaks of reactive compounds on site, were identified by elevated concentrations of the oxides of nitrogen and by wind direction analysis, and are excluded from subsequent data analysis. Table 1 summarises the measurements made at Bukit Atur: the critical measurements included:

- eddy covariance and gradient flux measurements of trace gases and particles;
- speciated concentration measurements of trace gases and particles;
- measurements of aerosol size-dependent hygroscopicity and critical supersaturations for cloud growth;
- concentration measurements of OH, HO\(_2\), and the sum of hydroperoxy and organic peroxy radicals;
- OH reactivity measurements (the rate at which OH is removed from the atmosphere);
- characteristics of boundary layer turbulence and mixing.
3.1.4 Concentration and flux measurements at the Sabahmas oil palm plantation

The Sabahmas oil palm plantation measurement site used during OP3-II was located in a 33 ha flat area of oil palm (*Elaeis guineensis × Elaeis oleifera* hybrid, progeny “Guthrie”) trees. The trees were of uniform age (12 years) and height (12 m). The site comprised a 15 m tower and an 8 m canopy access platform. Instruments were housed in a hut at the base of the platform. The analytical methods used were the same as at Bukit Atur, including the measurements of aerosol sub-micrometre composition, fluxes of aerosol, BVOCs, ozone and CO$_2$, although the suite of measurements made was not as comprehensive. In particular, measurements of OH and other radicals and of the oxides of nitrogen were not made.

3.1.5 Airborne measurements

Airborne measurements (see Table 1) were made during OP3-III using the Natural Environment Research Council/UK Meteorological Office’s BAe 146–301 Facility for Airborne Atmospheric Measurements (FAAM) aircraft (Lewis et al., 2007), deployed to Kota Kinabalu airport, less than 30 min flying time from Bukit Atur. In general, the same type of flight plan was executed on each flight, with one profile up and one down, interrupted by straight and level runs at altitudes of 100–250, 1500, 3000 and 6000 m above ground over the rainforest (centred on Bukit Atur), over an extensive and homogeneous agro-industrial oil palm landscape surrounding and including the Sabahmas oil palm plantation estate, and in transects between the two sites. Flights were made morning and afternoon giving typically four stacked profiles, allowing a picture to be built up of the concentrations of trace gases and particles during the daytime. Two flights were also made over the ocean up- and down-wind of Sabah.

In order to link the ground-based and airborne measurements, the aircraft flew past the Bukit Atur measurement station on more than ten occasions, at the same height above sea-level as the base of the GAW tower and at approximately 500 m horizontal separation. Table 2 summarises the flights made during the OP3 deployment and Fig. 5 shows the geographical extent of the flights over Sabah.

4 The chemical climatology of Sabah and Bukit Atur

4.1 Satellite observations

Satellite observations of key tropospheric trace gases allow the surface and aircraft trace gas measurements above the rainforest surrounding Bukit Atur to be put in a wider context relative to Borneo and the larger south-east Asian region. We focus on formaldehyde (HCHO) and nitrogen dioxide (NO$_2$), which are good indicators of emissions and photochemical activity, to examine the atmospheric chemistry over Borneo during the OP3 campaigns in 2008. Global background concentrations of HCHO are determined by the balance between the source (from the oxidation of methane) and the OH sink. Concentrations are typically much larger over continents due to additional sources from the oxidation of biogenic and anthropogenic VOCs, and from biomass burning (either directly released or from the oxidation of co-emitted VOCs) (Palmer et al., 2003; Fu et al., 2007). Anthropogenic activities, biomass burning and soil emissions are the main source
Table 1. Overview of the measurements made in OP3.

| Species                                      | Method/Analytical Technique | Ground† | Air Temporal Resolution | Detection Limit | Measurement Uncertainty | Reference                          |
|----------------------------------------------|----------------------------|---------|-------------------------|-----------------|-------------------------|------------------------------------|
| NMHC, including isoprene and oxygenates      | Disjunct eddy covariance flux measurement with continuous flow and analysis by PTR-MS | √/BA, OP | Fluxes: 30 min          | Fluxes: < 0.05 mg m⁻² h⁻¹ | Fluxes: Precision = ~ ±30% Mixing ratios: ~7 s Mixing ratios: 10–100 pptv | (Langford et al., 2009) |
|                                             |                           |         |                         |                 |                         |                                    |
| NMHC, including isoprene and oxygenates      | Automated PTR-MS gradient  | √/IC    | Gradient every 7 min    | 10–100 pptv     |                         | (Karl et al., 2007)                |
|                                             |                           |         |                         |                 |                         |                                    |
| NMHC, including isoprene, monoterpenes and oxygenates | Dual Channel Gas Chromatograph with Flame ionisation detectors (DC-GC-FID) | √/BA    | 1 h ground variable air  | 1 pptv          | variable, typically around 10% | (Lewis et al., 2007; Lewis et al., 2005) |
|                                             |                           |         |                         |                 |                         |                                    |
| Terpenoids, alcohols, aldehydes              | GC-PID, portable mass spectrometer | √/BA    |                         |                 |                         |                                    |
| CO₂/H₂O flux                                 | Eddy-covariance flux using infrared gas analyser Li-Cor 7000/7500 | √/BA, OP | 30 min fluxes           |                 |                         | (Aubinet et al., 2000)            |
| NO₂ flux                                    | Eddy covariance using laser induced fluorescence | √/BA    | 30 min fluxes           |                 |                         | (Farmer et al., 2006)             |
| HNO₃, HCl, HNO₂, NH₃, SO₂, NH₄⁺, NO₃⁻, Cl⁻, SO₄²⁻ | Wet effluent denuder & steam jet aerosol collector, online IC (GRAEGOR) | √/BA    | 1 h                     |                 |                         | (Thomas et al., 2009)             |
| Turbulence, sensible heat flux               | Eddy-covariance using sonic anemometry | √/BA    | 30 min fluxes           |                 |                         |                                    |
| Soil NO flux                                 | Dynamic auto-chamber using NO analyser | √/IC    |                         |                 |                         | (Pilegaard et al., 2006)          |
| Species          | Method/Analytical Technique                                                                 | Ground† | Air Temporal Resolution | Detection Limit       | Measurement Uncertainty | Reference                  |
|------------------|-------------------------------------------------------------------------------------------|---------|-------------------------|------------------------|--------------------------|----------------------------|
| Soil \( \text{N}_2\text{O}/\text{CH}_4 \) flux | Static soil chamber with off-line GC analysis                                               | ✓IC, OP | 1 h                     | (OH) \(2.4 \times 10^5\) molecule cm\(^{-3}\) (3 min av.) (HO\(_2\)) \(3.8 \times 10^6\) molecule cm\(^{-3}\) (3 min av.) | 44 % (OH) 50 % (HO\(_2\)) \((2\sigma)\) | (Whalley et al., 2010a, b) |
| \( \text{OH}, \text{HO}_2 \) | FAGE (Fluorescence Assay by Gas Expansion) laser-induced fluorescence                     | ✓BA     | 10 s                    | (OH) \(2.3 \times 10^6\) molecule cm\(^{-3}\) (1 min av.) (HO\(_2\)) \(2.0 \times 10^6\) molecule cm\(^{-3}\) (1 min av.) | 28 % (OH & HO\(_2\)) \((2\sigma)\) |                           |
| \( \text{OH}, \text{HO}_2 \) | FAGE (Fluorescence Assay by Gas Expansion) laser-induced fluorescence                     | ✓        | 60 s                    | (OH) \(2.3 \times 10^6\) molecule cm\(^{-3}\) (1 min av.) (HO\(_2\)) \(2.0 \times 10^6\) molecule cm\(^{-3}\) (1 min av.) | 28 % (OH & HO\(_2\)) \((2\sigma)\) |                           |
| OH Reactivity    | FAGE                                                                                      | ✓BA     | 1 min                   | 0.4 pptv (10 min)      | 22 % \((2\sigma)\)       | (Ingham et al., 2009)      |
| \( \Sigma \text{RO}_2 + \text{HO}_2, \text{HO}_2 \) | PERCA (PERoxy Radical Chemical Amplifier), dual inlet                                     | ✓BA     | 1 min                   | 1.5 pptv 500 pptv 60 pptv 130 pptv 4 ppbv 130 pptv | 38% \((1\sigma)\)         | (Fleming et al., 2006)     |
| \( \text{NO}_3, \text{CHO}_2, \text{NO}_2, \text{HONO}, \text{O}_3, \text{CHOCHO} \) | Differential Optical Absorption Spectroscopy                                               | ✓BA     | 10 min                  | 2 pptv 480 pptv 80 pptv 150 pptv 4.6 ppbv 150 pptv | 0–90° SZA 4 ppbv 130 pptv 4 ppbv 130 pptv | (Plane and Saiz-Lopez, 2006) |
| \( \text{NO}_2, \text{HCHO}, \text{CHOCHO} \) | MAX-DOAS                                                                                  | ✓BA     | 1 min                   | 14% and 13% 0–90° SZA 4 ppbv 130 pptv 4 ppbv 130 pptv | 14% and 13% 0–90° SZA     | (Bohn et al., 2008; Edwards and Monks, 2003; Volz-Thomas, et al., 1996) |
| \( \text{O}_3 \) | UV absorption                                                                              | ✓BA     | 1 s                     | 0.6 ppbv               | 0–90° SZA 4 ppbv 130 pptv 4 ppbv 130 pptv | 10%±3.4 ppbv \((\pm1\sigma)\) | (Heard et al., 2006)       |
| Species | Method/Analytical Technique | Ground† | Air Temporal Resolution | Detection Limit | Measurement Uncertainty | Reference |
|---------|----------------------------|---------|-------------------------|-----------------|------------------------|-----------|
| O$_3$ flux eddy correlation | Dry chemiluminescence | ✓/BA, OP | 30 min fluxes from 0.05 s | 0.1 ppbv | | (Güsten et al., 1990; Gusten and Heinrich, 1996) |
| O$_3$/NO/NO$_2$ gradient | Chemiluminescence (O$_3$), thermal converter | ✓/BA/IC | 15 min | | | |
| NO, NO$_2$, ΣNO$_2$, ΣNO$_2$-HNO$_3$ | NO/O$_3$ chemiluminescence detectors Photochemical converter + above Gold tube/CO converter + above Gold tube converter and denuder | ✓ | 10 min | 3 pptv for NO, 7 pptv for NO$_2$ | 15% for NO and 20% for NO$_2$ at 50 pptv | Pike et al., 2009 |
| NO, NO$_2$, ΣNO$_2$, ΣNO$_2$-HNO$_3$ | NO/O$_3$ chemiluminescence detectors Photochemical converter + above Gold tube/CO converter + above Gold tube converter and denuder | ✓ | 10 s | 3 pptv for NO and 15 pptv for NO$_2$ | 8% for NO at 1 ppbv and 9% for NO$_2$ at 1 ppbv | (Brough et al., 2003; Stewart, et al., 2008) |
| NO$_2$ | Laser-induced fluorescence (LIF) | ✓/BA | 1 Hz | 3.6 pptv/60 s | 20% | (Dari-Salisburgo et al., 2009) |
| ΣPNs | Thermal-dissociation LIF | ✓/BA | 1 Hz | 13 pptv/60 s | 40% | (Dari-Salisburgo et al., 2010) |
| ΣANs | Thermal-dissociation LIF | ✓/BA | 1 Hz | 13 pptv/60 s | 40% | (Aruffo et al., 2010) |
| NO$_2$ flux | LIF-Eddy covariance | ✓/BA | 10 Hz | 11 pptv/60s | 20% | (Dari-Salisburgo et al., 2010) |
Table 1. Continued.

| Species                                                                 | Method/Analytical Technique                  | Ground† | Air Temporal Resolution | Detection Limit | Measurement Uncertainty | Reference                          |
|------------------------------------------------------------------------|---------------------------------------------|---------|-------------------------|-----------------|-------------------------|-----------------------------------|
| H$_2$O$_2$, ROOH                                                       | Dual-channel fluorometric detector           | √       | 1 min                   | 50 pptv         | 14% (1σ)                | (Penkett et al., 1995)            |
| H$_2$O vapour                                                         | Dew point hygrometer                        | √/BA    | √                       |                 |                         |                                   |
| H$_2$O flux                                                            | Eddy-covariance using UV Absorption          | √/BA    | 30 min fluxes 100 Hz    | 8% (q-dependent) |                         | (Coe et al., 1995)                |
| CH$_2$O                                                                | Fluorometric detection (Hantzsh reaction)    | √/BA    | 1 min                   | 100 pptv        | 17% (2σ)                | (Still et al., 2006)              |
| Speciated aldehydes, ketones and alcohols NMHC, including isoprene and oxygenates | GC/GC detection, PTR-MS                      | √       | 15 s                    | 50–120 pptv     | 13–16% (1σ)             |                                   |
| >C7 NMHC (e.g. terpenes)                                              | Adsorbent tubes & GC/TOF-MS                 | √       | Variable                | 50 pptv         | ±(5%+20 pptv)           | (Capes et al., 2009)              |
| CO                                                                    | Chemiluminescence                            | √/BA    | √                       |                 |                         | (Gerbig et al., 1999)            |
| PAN                                                                   | GC/ECD (electron capture detection)          | √       | 90 s                    | 5 pptv          | 5%                      | (Whalley et al., 2004)           |
| PAN                                                                   | GC/ECD (electron capture detection)          | √/BA    | 10 min                  | PAN:15 pptv, PPAN, MPAN:25 pptv | 20% (2σ) | (Harrison et al., 2006) |
| Alkyl nitrates, organic N                                             | GC and negative ion CI GC/MS                 | √/BA    | √                       | G: 1 h A: variable | 0.005 pptv | 13% (2σ) | (Reeves et al., 2007; Worton et al., 2008) |
| Reactive halocarbons                                                  | GC/MS                                        | √/BA    | √                       | G: 1 h A: variable | 0.005 pptv | 15% (2σ) | (Worton et al., 2008) |
| Halocarbons                                                           | GC/ECD                                       | √/BA    | ∼ 15 min                | ∼ 0.5 pptv      | 5–10%                  | (Gostlow et al., 2009)           |
| Species                        | Method/Analytical Technique | Ground† | Air Temporal Resolution | Detection Limit | Measurement Uncertainty | Reference |
|-------------------------------|----------------------------|---------|-------------------------|-----------------|-------------------------|-----------|
| Aerosol number concentration | CPCs total particle number concentration (>3 nm); | ✓/BA | ✓ | 30 min (ground) 1 s (aircraft) | N/A | N/A |
| Aerosol size distribution    | SMPS ground: 3 < \(D_{md}\) < 700 nm Air: 3 < \(D_{md}\) < 350 nm | ✓/BA/IC | ✓ | 30 min (ground) 1 min (aircraft) | N/A | N/A |
| Optical and aerodynamic particle counters (0.3/0.1–20/3µm) | ✓/BA/IC | ✓ | 30 min (ground) 1 s (aircraft) in-canopy gradients (2–32 m) | N/A | N/A |
| Ultra-fine particle fluxes    | Eddy Correlation \(D_p > 3\ nm, >10\ nm, independent CPCs | ✓/BA | Fluxes 30 min. Raw 0.3 s. | < 0.01 particle/cm\(^3\) | ±10% @ 3 × 10\(^5\) particles/cm\(^3\). | (Buzorius et al., 1998) |
| Aerosol size segregated chemical composition | Aerodyne Aerosol Mass Spectrometer (40 nm < \(D_{vad}\) < 0.8 µm), non-refractory \(SO_2^-\), \(NO_3^-\), \(NH_4^+\), organic species | ✓/BA, OP | ✓ | 30 min (ground) 10 s (profile) 30 s (SLR) | Ground: 3 ng m\(^{-3}\) (NO\(_3^-\)), \(SO_2^-\) and 11 ng m\(^{-3}\) (NH\(_4^+\)), 10 min average, high res mode Air: 3 ng m\(^{-3}\) (NO\(_3^-\)), \(SO_2^-\) and 30 ng m\(^{-3}\) (NH\(_4^+\)) 30 s average | 15% | (Nemitz et al., 2008) |
| Size segregated chemically speciated aerosol mass fluxes | Eddy-covariance using Aerodyne Aerosol Mass Spectrometer, (40 < \(D_{vad}\) < 800 nm), non-refractory \(SO_2^-\), \(NO_3^-\), \(NH_4^+\), organic species | ✓/BA, OP | 30-min fluxes | 10 ng m\(^{-2}\) s\(^{-1}\) | ±10 ng m\(^{-2}\) s\(^{-1}\) | (Nemitz et al., 2008) |
Table 1. Continued.

| Species                          | Method/Analytical Technique | Ground† | Air       | Temporal Resolution | Detection Limit | Measurement Uncertainty | Reference                  |
|----------------------------------|-----------------------------|---------|-----------|--------------------|-----------------|--------------------------|----------------------------|
| Size-segregated bioaerosol       | WIBS                        | √BA, IC |           |                    |                 |                          | (Kaye et al., 2005)         |
| Genetic bioaerosol               | Genetic analysis of filters  | √BA, IC, OP | 12-h     |                    |                 |                          |                            |
| Cloud liquid water               | Gerber PVM-100              | ✓       | 0.1 s     | 0.005 g m⁻³         |                 |                          | (Gerber, 1991)              |
| Aerosol Scattering properties   | 3 wavelength nephelometer   | ✓       | 2 mm⁻¹    | 5%                 |                 |                          | (Osborne et al., 2008)      |
| Cloud Condensation Nuclei Activity | CCN spectrometer (single) | ✓ (two) | 1 h (ground) |                    |                 |                          | (Roberts and Nenes, 2005)   |
| Aerosol hygroscopicity          | Hygroscopic Tandem Differential Mobility Analyser | ✓BA    | 1 h       |                    |                 |                          | (Gysel et al., 2007)        |
| Meteorological parameters (Wind speed & direction, solar radiation, PAR, precipitation, wetness, pressure, temperature, RH, turbulence, sensible heat flux) | Standard meteorological sensors (aspirated thermocouples, Vaisala WXT) | ✓BA, IC, OP | 30 min temperature gradients | | | | |
| Boundary Layer Height           | LIDAR                       | ✓BA     | 1 s       | 20–60 m            |                 |                          | (Pearson et al., 2009)      |

† – BA=Bukti Atur site; IC=In-canopy site; OP=oil-palm site (OP3-II).

of variability in the tropospheric NO₂ columns (Martin et al., 2002, 2003).

Figure 6a and b shows HCHO and NO₂ column distributions over Borneo during April–July 2008, quantified using UV spectroscopic measurements taken by the nadir-viewing data from SCIAMACHY (HCHO; De Smedt et al., 2008) and OMI (NO₂; ATBD-OMI-02) instruments respectively. NO₂ is a strong absorber at UV wavelengths and, because the OMI instrument provides daily coverage, the NO₂ column data can be averaged onto a fine resolution grid of 1.0° × 1.0° (longitude × latitude). In contrast, we average the HCHO columns onto a coarser 2.5° × 2.0° grid (to improve the signal-to-noise ratio), since it is a much weaker absorber and because global coverage is only achieved by SCIAMACHY at the Equator every 6 days. Scenes with fractional cloud coverage >40% are excluded from our analysis.

Figure 6b shows that NO₂ measurements from OMI identify anthropogenic NOₓ signals from Bangkok, Jakarta, Surabaya and Singapore. Enhanced HCHO also seen over these cities indicates intense photochemical activity, likely associated with polluted conditions. Over Indochina in April, elevated HCHO and NO₂ columns are loosely correlated.
Table 2. Overview of aircraft flights made in OP3.

| Flight | Flight description | Date in 2008 | Take off-Landing Times (UTC) |
|--------|--------------------|--------------|-----------------------------|
| B384   | Survey flight (Bukit Atur, oil palm, SW of Kota Kinabalu) | 9 July       | 02:13–05:32                 |
| B385   | Bukit Atur (ground sampling site) and SW of Kota Kinabalu | 10 July      | 01:10–05:04, 07:04–10:41    |
| B386   | Bukit Atur         | 12 July      | 01:02–04:48, 06:10–9:59     |
| B387   | Maliau Basin, S of Sabah, N of Bukit Atur and W of oil palm plantation | 13 July      | 00:54–06:01, 06:30–10:10    |
| B388   | Bukit Atur and oil palm | 15 July    | 00:54–04:43, 06:01–09:33   |
| B389   | Ocean flight SE of Sabah | 16 July    | 00:49–04:31, 05:39–09:40   |
| B390   | CASCADE (Marine flight, W and NW of Sabah) | 18 July     | 22:57–03:14                |
| B391   | SW of Bukit Atur, oil palm | 19 July    | 00:54–05:06, 06:15–09:48   |
| B392   | Mainly oil palm plantation | 21 July     | 00:48–04:46, 05:53–09:46   |
| B393   | Ocean flight (NW and SE of Sabah) | 22 July    | 00:43–05:10                |

Local Time is (UTC+8).

with AATSR fire-counts (not shown) (Arino et al., 2005) and hence are probably due to fire emissions (Fu et al., 2007). During May, enhanced HCHO columns over the Gulf of Thailand may reflect outflow from fires occurring in Sumatra and from the Sibu area of eastern Borneo. However, during the OP3 campaigns, fire activity over Malaysian Borneo was minimal and the levels of HCHO and NO$_2$ were generally low, with their column amounts typically approaching their background values of $\sim5 \times 10^{15}$ molecules cm$^{-2}$ and $\sim5 \times 10^{14}$ molecules cm$^{-2}$, respectively. These low values also suggest that during the OP3 measurement period the HCHO source from biogenic VOC oxidation was weak. We observe a slight HCHO enhancement during April over Danum Valley (Fig. 6a), but it is difficult to assign source attribution.

In Fig. 6c we show the monthly-mean 12-year time series of (continuous) HCHO and (non-continuous) NO$_2$ columns, retrieved from the GOME and SCIAMACHY instruments (De Smedt et al., 2008; Martin et al., 2002, 2003), together with the total number of fire-counts detected by the ATSR/AATSR instruments (Arino et al., 2001, 2005). High correlations between HCHO and NO$_2$ columns and the fire-count data suggest that biomass burning over Borneo drives observed variability of these trace gases. We find that anomalously high HCHO and NO$_2$ columns were due to intense burning periods associated with strong El Nino conditions, indicated here by the Multivariate El Nino Southern Oscillation (ENSO) Index (Wolter and Timlin, 1998), as expected. For example, during 1997/98 an increased number of forest and peat fires during the warmer and drier El Nino conditions (Levine, 1999) led to extremely high trace gas column concentrations.

Our recent analysis shows that spatial correlations between the 12-year HCHO and NO$_2$ data, and the associated assimilated meteorological data, vegetation activity and fire-counts were strongest in southernmost Borneo and not over the Bukit Atur (Danum Valley) region (Barkley et al., 2009). The low trace gas columns observed over this region during 2008 are consistent with our understanding of past variability.
4.2 In-situ observations

4.2.1 Airborne measurements

The in-situ observations made during the OP3 project were centred on sites representative of natural rainforest and oil palm plantations (see above), with aircraft flights over each landscape and over the up- and down-wind oceans (see above). Figure 7 shows aircraft measurements of six species taken on flights near to the Bukit Atur GAW station and the adjacent oil palm plantations. Data are shown from flights (a) below 500 m above ground level, representative of the boundary layer and (b) above 2500 m, representative of the free troposphere. Also indicated on the plots is the approximate boundary between the natural rainforest and the palm oil plantations.

Within the boundary layer (Fig. 7a), the most striking concentration difference between the rainforest and plantation is for isoprene. Concentrations over the rainforest were typically 1000–3000 pptv, with concentrations over the plantation significantly higher (2–5 times higher; 5000–10 000 pptv). This is consistent with the higher emission rate of isoprene from oil palm trees (*Elaeis guineensis*) compared to most rainforest tree species (Wilkinson et al., 2006). The plantation landscape also contains associated agro-industrial activities (e.g. road traffic, oil palm processing plants and nitrogenous fertiliser application) and as a result, NO\textsubscript{x} concentrations were also higher compared to the rainforest. However, whereas the higher isoprene concentrations observed tended to be in the west of the oil palm plantation area, higher NO\textsubscript{x} levels were concentrated in the north and east of the plantation area, where the majority of plantation processing plants were observed to be situated. In this area, typical NO\textsubscript{x} levels were 1000–1500 pptv, with up to >2000 pptv observed when flying through or close to the plume of a processing plant. Further south in the plantation area, closer to the boundary with the rainforest, NO\textsubscript{x} was much lower, with levels typically <300 pptv, which is similar to levels that are observed over the rainforest.
CO shows a similar behaviour to NO\textsubscript{x}, with higher levels (100–110 ppbv) observed over the northern and eastern plantation area compared to further south and over the rainforest (70–90 ppbv). Levels of sulphate aerosol show little difference between the two land use areas, with both exhibiting aerosol loading of 0.25–1 µg m\textsuperscript{−3}.

The result of the increased levels of primary pollutants over the oil palm plantation compared to the rainforest can be seen in the observed ozone, a species that is not directly emitted into the atmosphere but produced photochemically. Ozone was significantly higher (20–25 ppbv) over the eastern area of oil palm plantation, corresponding to the area where NO\textsubscript{x} was generally higher. Over the western area of oil palm plantation, where NO\textsubscript{x} levels are lower but isoprene levels are still high, and over the natural rainforest, where both NO\textsubscript{x} and isoprene are lower, ozone is observed to be 8–15 ppbv. This demonstrates the need for increased NO\textsubscript{x} emissions, as well as isoprene, in order for significant extra ozone production to occur (Hewitt et al., 2009). More organic particulate was observed over the oil palm plantation than the rainforest, with average loadings of 0.67 µg m\textsuperscript{−3} and 0.48 µg m\textsuperscript{−3} respectively, which is consistent with our ground based measurements (see below) and could be due to enhanced production of secondary organic material, although primary emissions from the processing plants and human settlements may also be contributing.

In the free troposphere (Fig. 7b), very little isoprene or sulphate aerosol was observed over either the rainforest or plantation and levels of CO were similar over both areas (~50–60 ppbv). NO\textsubscript{x} levels were higher in the eastern plantation region (200–300 pptv) compared to the rainforest (<50 pptv); however this did not lead to increased ozone in the free troposphere over the plantation. This is likely to be due to the very low levels of isoprene and other biogenic VOCs, which do not have a long enough lifetime to escape the boundary layer. Organic aerosol was also very low over both plantation and rainforest in the free troposphere, with an average organic loading of 0.03 µg m\textsuperscript{−3}.

As described above, some flights were also undertaken over the ocean, both upwind and downwind of Sabah. Levels of isoprene and NO\textsubscript{x} were very low in both cases and there was no obvious enhancement in ozone in the upwind flights. These data were collected on only two days; however our initial interpretation of this is that Sabah is not a large exporter of primary or processed pollutants to the wider region.

### 4.2.2 Bukit Atur measurements

Data were taken within the rainforest at the Bukit Atur GAW station during two four-week intensive measurement periods (OP3-I and OP3-III). The GAW station has made ozone measurements since January 2007 and these data are shown in
Fig. 8. Ground level ozone concentrations measured at the Bukit Atur GAW site 2007–2008. Data are shown as hourly points with a 24 h running average.

Fig. 8. Gaps in the data are due to lightning strikes rendering the instrument inoperable. However the plot shows that the levels of ozone observed during the OP3 intensive periods in 2008 are typical for the site.

Campaign daytime (10:00–14:00) average isoprene and total monoterpene fluxes measured at 75 m on the GAW tower were 1.2 and 0.3 mg m\(^{-2}\) h\(^{-1}\) respectively during OP3-I and 2.1 and 0.49 mg m\(^{-2}\) h\(^{-1}\) during OP3-III (Langford et al., 2010). These values were comparable to measurements made above regions of the Congo during the EXPRESSO field campaigns (Greenberg et al., 1999; Klinger et al., 1998; Serca et al., 2001) and are somewhat smaller than values reported above regions of the Amazon (Karl et al., 2007; Kuhn et al., 2007). The typical monoterpene to isoprene ratio was between 0.2–0.25, which is within the upper range of ratios reported in other tropical regions (Guenther et al., 2008). The fluxes of other biogenic VOCs, including methanol, acetone and acetaldehyde, were very much smaller, accounting for just 2% of the total measured VOC emissions.

Figure 9a shows hourly averaged median diurnal cycles for NO, NO\(_2\), O\(_3\), isoprene and sulphate and organic aerosol mass concentration, taken at Bukit Atur during June and July 2008. Only data from the six days when flights took place over the station were used in the averaging for a direct comparison between the ground-based and aircraft data in the boundary layer and free troposphere over the rainforest. Note that since Bukit Atur is on a ridge, some of the boundary layer data collected by the aircraft over the surrounding forest were at the same or lower altitude (above sea level) as the GAW tower.

For NO, a peak of ~100 pptv was observed between 07:00 and 08:00 local time, which corresponds to the onset of NO\(_2\) photolysis. This was followed by a drop to ~40 pptv by 11:00, with NO then remaining fairly constant throughout the day, until a further drop to ~20 pptv occurred at around 17:00. NO then remained constant throughout the night. NO\(_2\) exhibits a clear diurnal cycle peaking at around midnight at 300 pptv, with a minimum of ~150 pptv in mid-afternoon. These values are comparable to NO\(_2\) measurements made in Amazonia (e.g. Ganzeveld et al., 2008) and over West Africa during the AMMA experiments (e.g. Saunois et al., 2009).

Boundary layer aircraft measurements of NO and NO\(_2\) were in reasonable agreement with the ground based measurements, although airborne NO levels were consistently lower. This, along with the non-zero night-time levels, suggests a local NO source that was diluted throughout the boundary layer. NO and NO\(_2\) levels in the free troposphere were much lower, being <10 pptv and ~20 pptv, respectively. A full description of ground level NO\(_x\) data and discussion of the chemistry is given by Pike et al. (2009) and Pugh et al. (2010b).

Isoprene was the most abundant terpenoid compound observed and showed a clear diurnal cycle with very low (<40 pptv) levels at night and a peak of ~2000 pptv in mid afternoon. This was caused by the dependence of isoprene emission from trees on temperature and light (Guenther et al., 1993). Boundary layer isoprene measurements from the aircraft were in good agreement with the ground-based Bukit Atur measurements, and free tropospheric isoprene levels were, as expected, very low. Limonene, alpha-pinene and beta-pinene were the most abundant monoterpene compounds observed. Relatively high concentrations of 2-ethyl hexanol and some aldehydes were also seen in samples taken from within the forest canopy in the vicinity of Bukit Atur.

The concentrations of NO\(_2\), peroxy nitrate (RO\(_2\)NO\(_2\)) and alkyl nitrate (RONO\(_2\)) were measured at Bukit Atur with a laser induced fluorescence system (Aruffo et al., 2009). The diurnal cycle of RO\(_2\)NO\(_2\) followed the isoprene diurnal cycle, with maximum concentrations in the afternoon.
RO$_2$NO$_2$ compounds are produced in the oxidation of isoprene and other terpenoids, this result provides evidence for the oxidation of BVOC in the vicinity of the forest canopy. Additional evidence for the active photochemical processing of isoprene is provided by the increase in concentrations of its oxidation products, methyl vinyl ketone and methacrolein, observed in the afternoons.

The concentrations of HO$_2$+RO$_2$ were measured at Bukit Atur using PEROxy Radical Chemical Amplification (PERCA) technology (Karunaharan et al., 2009) over 20 days in OP3-I and 29 days in OP3-III. Mean midday mixing ratio of 33.5±3.3 and 35.5±3.5 parts per trillion (pptv) were calculated from the continuous measurements of the sum of the inorganic and organic peroxy radicals from OP3-I and OP3-III respectively. The mean net ozone production rate can then be calculated using:

$$N(\text{O}_3) = k_p[\text{NO}][\text{HO}_2 + \text{RO}_2] - (f \cdot j(O_1D) + k_1[\text{OH}] + k_2[\text{HO}_2])[\text{O}_3]$$

where $k_p$ is a combined rate coefficient for the oxidation of NO to NO$_2$ by all peroxy radicals and $f \cdot j(O_1D)[O_3]$ represents the fraction of ozone photolysed to yield O($D_1$) atoms and then OH (with $f$ being the proportion of O($D_1$) atoms which react with H$_2$O to give OH rather than being collisionally deactivated). The rate constants $k_1$ and $k_2$ are from the ozone loss reactions:

$$\text{OH} + \text{O}_3 = \text{HO}_2 + \text{O}_2 \quad (R1)$$

$$\text{HO}_2 + \text{O}_3 = \text{OH} + 2\text{O}_2 \quad (R2)$$

The net formation of ozone was found to be strongly dependent on the concentration of isoprene, having a linear sensitivity (e.g., $d\ln(P(\text{O}_3))/d\ln(\text{isoprene})=1$). Since HO$_2$+RO$_2$ are produced during the oxidation of isoprene and other terpenoids, taken together with the observations of MVK and methacrolein formation, this result provides evidence for the oxidation of BVOC in the vicinity of the forest canopy and of the role of isoprene in the formation of ozone in the near-canopy environment during the daytime.

Evidence of BVOC reactivity also comes from differential optical absorption spectroscopy (DOAS) measurements. DOAS was used at Bukit Atur to make the first measurements in a tropical rainforest of glyoxal (CHOCHO), one of the major products of isoprene oxidation by OH. Mixing ratios much higher than in other rural locations were observed, with a maximum of 1.6 ppbv and average around 600 pptv. HCHO was also measured, peaking at 4.5 ppbv and with an average around 3 ppbv. Both species were closely correlated to isoprene (which should be their major source in this environment), and peaked in the early afternoon. A box model indicates that the production efficiency of glyoxal from isoprene is ∼10%. Although glyoxal was significantly scavenged during rain showers, it is mostly removed by reaction with OH and photolysis (lifetime at midday ∼2.3 h) rather than uptake on background aerosol in this environment, because, as discussed earlier, aerosol mass loadings were low.

There was a slight diurnal trend in ground-based O$_3$ measurements, with levels of ~5 ppbv at 06:00, rising to ~8 ppbv in mid afternoon. Note that the very small amplitude in the diurnal pattern is typical of a hill-top site (Fowler et al., 2008) and contrasts with that observed at the in-canopy site and over the low-lying palm plantations (see below). These values are comparable to above canopy measurements made above regions of Amazonia (Karl et al., 2007). Boundary layer O$_3$ levels also show an increase throughout the day, but at higher levels (10–13 ppbv), probably because the boundary layer O$_3$ is less depleted by deposition to the canopy (Matsuda et al., 2006; Cros et al., 2000). These boundary layer values are slightly less than those measured over Amazonia and West Africa during the GABRIEL and AMMA campaigns (Ganzveld et al., 2008; Saunois et al., 2009). There is little vertical structure in the O$_3$ mixing ratio at higher heights above the rainforest, with levels in the free troposphere being very similar to those in the boundary layer. Ozone vertical profiles were therefore more like those observed during the monsoon over northern Australia, than those observed during monsoon break periods (Heyes et al., 2009).

Concentrations of OH display a clear diurnal cycle, peaking at solar noon, with significant concentrations observed: up to $8.7 \times 10^6$ molecules cm$^{-3}$ (60 min average) was recorded on one day. During the GABRIEL campaign above Amazonia, average midday boundary layer OH concentrations were $5.75 \times 10^6$ molecules cm$^{-3}$ (Martinez et al., 2008). Although $j(O^1D)$ levels and humidity were high, the relatively low O$_3$ concentrations limit primary OH production. The measured OH reactivity was very high, up to 87 s$^{-1}$ (corresponding to a lifetime for OH approaching 10 ms). According to the measurements of reactive hydrocarbons (above), this reactivity is dominated by reaction with isoprene, indicating that significant OH sources must be present – in addition to primary production following ozone photolysis – in order to maintain the elevated levels of OH levels recorded (Whalley et al., 2010b). The OH diurnal profile was asymmetric (i.e., lower post-noon than the equivalent time pre-noon). This may be due to its main measured sink, isoprene, being present at higher concentrations in the afternoon, but the asymmetric profile could also result from other factors that have yet to be identified. OH concentrations up to $6 \times 10^5$ molecule cm$^{-3}$ (60 min average) were also recorded on several nights, consistent with measurements by other groups in forested environments (Tan et al., 2001), with some residual OH observed after $j(O^1D)$ had dropped to zero. Measured concentrations of HO$_2$ and the sum of HO$_2$+RO$_2$ displayed similar profiles.

Boundary layer OH concentrations recorded on the aircraft over Bukit Atur were similar to those measured at ground level, whereas aircraft HO$_2$ concentrations were higher than measured at the ground, a reflection of the elevated NO levels.

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at the surface. A full description of the OH, HO\textsubscript{2} and OH reactivity measurements is given in Whalley et al. (2009, 2010a) and Edwards et al. (2009, 2010).

The non-refractory sub-micrometre aerosol as measured by aerosol mass spectrometry at the Bukit Atur ground site was mostly composed of organics, sulphate, and ammonium, with low levels of nitrate and chloride. Average loadings were 0.74, 0.61, 0.21, 0.04 and 0.01 µg m\textsuperscript{-3}, respectively, giving a total aerosol loading of 1.61 µg m\textsuperscript{-3}. The sulphate was observed to be largely neutralised to ammonium sulphate. Organics showed a diurnal profile peaking around 21:00. The ensemble organic spectrum had a m/z=44 to total organics ratio of 0.15%, with prominent peaks at m/z=55 and 53, features which would be considered typical of a partially oxidised organic aerosol (e.g., Hallquist et al., 2009). The sulphate showed no distinct diurnal profile, suggesting it was regional in nature, potentially from the anthropogenic background, volcanoes or oxidised dimethyl sulphate originating from marine phytoplankton (Dacey and Wakeham, 1986). Of all the gas and aerosol species measured at Bukit Atur, sulphate aerosol most clearly showed the influence of regional-scale transport rather than local sources and sinks. Both nitrate and chloride showed strong nocturnal peaks that could be caused by partitioning from the gas phase as the temperature falls and humidity increases at night.

Slightly larger aerosol concentrations were measured from the aircraft than the ground, both over the rainforest and oil palm, with mean total loadings in the boundary layer of 1.66 and 1.78 µg m\textsuperscript{-3}, respectively. Aerosol loadings were much lower in the free troposphere, with an average total of 0.22 µg m\textsuperscript{-3}. Free-tropospheric aerosol showed a larger sulphate contribution of 69%, compared with 43% in the boundary layer.

The water uptake ability of the atmospheric aerosol at Bukit Atur was probed in both the sub- and super-saturated regimes with the use of a Hygroscopicity Tandem Differential Mobility Analyser (HTDMA; Cubison et al., 2005; Grydeland et al., 2007) and Cloud Condensation Nucleus counter (CCNc; Roberts et al., 2005), respectively. Aerosol hygroscopic growth factors (GF) at 90% relative humidity (RH) for particles of dry diameter (D\textsubscript{0}) from 27 to 217 nm showed the aerosol to be externally mixed, with a mean growth factor (GF\textsubscript{D0,90%}) ranging between 1.3 and 1.6 with increasing dry diameter and multiple growth factor modes at each dry size evident. A Scanning Mobility Particle Analyser (SMPS; TSI 3080) was operated upstream of the CCNc, to supply quasi-monodisperse aerosol in the size range 30 to 220 nm. Critical supersaturations (S\textsubscript{c}) derived from fraction activated CCN spectra, showed a diurnal variation in supersaturation between 0.2 and 0.8% required to activate particles of a given D\textsubscript{0} primarily in the dry diameter range 50 to 120 nm.

### 4.2.3 Oil palm plantation measurements

Campaign daytime average (10:00–14:00) isoprene fluxes measured at 15 m at the Sabahmas oil palm plantation site were 8.6 mg m\textsuperscript{-2} h\textsuperscript{-1} during OP3-II (Misztal et al., 2010a). This is four to seven times greater than the isoprene flux measured at Bukit Atur (see above). The fluxes of other biogenic VOCs, including oxygenated species and more exotic compounds (e.g., estragole: Misztal et al., 2010b), were smaller, contributing ~15% of the total measured VOC emissions. Net fluxes of monoterpenes were very small (<1% of total VOC flux).

Figure 9b shows diurnal cycles of data taken at the ground-based site within the Sabahmas oil palm plantation. Species plotted are the same as for the Bukit Atur rainforest ground site, with the exception of NO and NO\textsubscript{2} which were not measured at ground level at Sabahmas. Boundary layer NO and NO\textsubscript{2} levels show slightly higher levels compared to the rainforest but the scatter on the data (indicated by the 1σ error bars) is much larger. This is indicative of the high NO\textsubscript{x} concentrations observed in certain areas of the plantation landscape, as described above. Free troposphere NO and NO\textsubscript{2} levels over the plantation were again much lower and very similar to those observed over the rainforest.

Ground based isoprene data show a similar diurnal cycle to the rainforest, with near zero levels at night, but with an earlier peak at around 12:00. Isoprene concentrations are much higher in the plantation, with peak daytime levels of ~16 000 pptv. Isoprene concentrations in the boundary layer over the plantation show a very similar diurnal profile, but with a significantly lower peak of ~5000 pptv. The ground-based measurements were made at the top of the oil palm canopy, and hence this difference may be explained by dilution. Also, since the oil palm plantations are generally situated in low-lying, flat land, the aircraft measurements were always made at a significant height above the ground level measurements, unlike at Bukit Atur where the aircraft was able to fly past the base of the GAW tower, located on a hill.

Measurements of O\textsubscript{3} at the ground in the plantation exhibit a very different diurnal cycle compared to the rainforest, with near zero night-time concentrations and a peak of ~5 ppbv at 11:00–12:00. This is followed by a steady decrease throughout the day until O\textsubscript{3} reaches close to zero at around 23:00. This general pattern is typical of that for a low-lying site, where O\textsubscript{3} lost by reaction with NO emitted from fertilised plantation soils and deposition at night is not replenished by mixing from aloft, due to the more ready establishment of a stable boundary layer compared with elevated, more windy, sites (Fowler et al., 2008). Data taken in the boundary layer above the plantation show a similar diurnal cycle to the rainforest, with an increase throughout the day to a peak of ~15 ppbv. This mean value is 2 to 3 ppbv higher than the rainforest levels, but as indicated by the 1σ error bars there is scatter in this value, which reflects the variability in the NO\textsubscript{x} and its impact on O\textsubscript{3} concentrations. Increased
deposition to the oil palm trees, reaction with NO\textsubscript{X} from local sources, and reaction with isoprene, are three possible contributing factors for the low concentrations and earlier peak of day-time O\textsubscript{3} observed on the ground in the plantation.

The total aerosol loading, as measured by the AMS, at the oil palm site was on average 1.6 µg m\textsuperscript{-3}, similar to the aerosol loading measured by AMS at the forest site (Nemitz et al., 2009, 2010). The mass was dominated by organic aerosol, which made up on average 65% of the total mass, with sulphate contributing 20% of the total mass. The organic aerosol exhibits a clear diurnal trend with two maxima, occurring at approximately 08:00 and 15:00 local time. The ensemble mass spectrum measured at the oil palm resembles the classic oxygenated organic aerosol spectrum (e.g. Hallquist et al., 2009), dominated by mass fragment m/z 44, resulting from carboxylic acid functionality. An examination of mass fragment-specific diurnal profiles indicates that fragments thought to be associated with less oxygenated organic aerosol make a smaller contribution to the afternoon maximum in the organic mass relative to those masses arising from oxygenated fragments, e.g. m/z 44.

Sulphate aerosol shows a slightly different trend in mass to the total organics, with a broader maximum during the daytime, although there was large day-to-day variability. Both the nitrate and chloride fractions showed an obvious maximum, coincident with the morning organic aerosol maximum, but both lacked the increase in mass observed in the organic fraction during the afternoon.

5 Modelling and assessment

As stated in the introduction above, a fundamental aim of the OP3 project was to understand the role played by biogenic emissions from the tropical rainforest: what is the nature of the emissions, how are they transformed and how do these processes then impact local, regional and hemispheric scale atmospheric chemistry; what will be the implications of global change for these processes and their impacts?

To address these questions, the OP3 project employed a range of models, varying in their spatial scale of interest and chemical complexity, as shown in Table 3. Thus, a model of transport and chemistry through and above the forest canopy has been used to address the canopy measurements, to study vertical transport of gas and aerosol species, their emission and deposition, and chemical transformation (Ryder et al., 2009). A range of chemical box models (some run along air parcel trajectories) has been employed to study local chemistry, addressing both the ground based and aircraft measurements. These very detailed models aim to inform the chemical mechanisms used in our regional and global models that, in turn, are used to study the impact of the biogenic emissions at the regional and wider scales. They are also used to investigate transport and variability.

Understanding the fast photochemistry processes (mainly involving HO\textsubscript{2} and RO\textsubscript{2}) is carried out using constrained box models. These use the observed concentrations of species providing HO\textsubscript{2} sources, sinks and inter-conversion pathways, to calculate the concentrations of OH, HO\textsubscript{2} and other short lived species such as the sum of peroxy radicals (RO\textsubscript{2}), for comparison with observations. Insight from these calculations allows our understanding of the complex organic chemistry occurring in this region to be evaluated, the level of chemical complexity required to capture these processes to be quantified and the impact of the approximations made in the chemical transport models to be evaluated.

The constrained-box-model focus is split between the ground-based and aircraft data. One investigation has used the Master Chemical Mechanism, a near-explicit mechanism that considers the degradation of a suite of hydrocarbons in the atmosphere (Jenkin et al., 2003; Saunders et al., 2003). For both ground and aircraft data, a zero-dimensional box model is constrained with the observed concentrations of longer-lived species such as hydrocarbons, NO\textsubscript{X}, O\textsubscript{3}, relevant meteorological data and observed photolysis rates; the models predict radical concentrations which can be compared with those measured. The ground-based model is a continuation of a previous modelling framework (e.g. Emmerson et al., 2007) that has been used extensively for similar comparisons (Heard and Pilling, 2003). The aircraft based model uses the Dynamically Simple Model of Atmospheric Chemical Complexity (DSMACC) (Emmerson and Evans, 2009) model and follows previous aircraft based work (Jaegle et al., 2000; Olson et al., 2006). Preliminary results show that for the first (April–May) ground-based campaign,
Table 3. Modelling tools employed in the OP3 project.

| Model name                                      | Brief description                                      | Domain                          | References                  |
|------------------------------------------------|-------------------------------------------------------|---------------------------------|-----------------------------|
| Canopy model                                   | 1-D transport and chemistry                           | Ground to canopy top – six levels| Ryder et al. (2009)         |
| MCM – York                                     | 4500 species, >12 500 reactions. Near-explicit chemistry; long lived gases constrained by observations | Ground-based box                | Emmerson et al. (2007)     |
| Dynamically Simple Model of Atmospheric Chemical Complexity (DSMACC) | Reduced chemistry; long lived gases constrained by observations | Steady state calculations along aircraft flight track | Emmerson and Evans (2009) |
| CiTTyCAT                                       | 200 species, >800 reactions including MIM2 isoprene scheme, Jenkin alphapinene and RACM limonene | Boundary layer: one well-mixed layer in daytime, two at night. | Pugh et al. (2010b)         |
| Cambridge p-TOMCAT chemical transport model    | 63 species, 177 reactions. Simple organic mechanism, including isoprene. Separate bromocarbon scheme. | Global. Variable resolution, usually 3°×3° or 0.5°×0.5° | Cook et al. (2007), Voulgarakis et al. (2009) |
| p-TOMCAT box                                   | As above                                              | Box model                       | Pike et al. (2009)          |

There is reasonable agreement between modelled and measured OH on some days, but an under-prediction on others. The OH loss for the OP3 campaigns was dominated by isoprene chemistry, with smaller contributions from \( \gamma \)-terpinene and limonene. \( \gamma \)-Terpinene becomes much more important at night-time, when isoprene emissions are effectively zero. Modelling of the aircraft data is in a more preliminary state but shows relatively good agreement between the observed and simulated HO\(_2\) with poorer agreement for the OH. Whether this reflects issues with making OH observations from aircraft or with our understanding of the chemistry of isoprene will require further analysis.

Reduced chemical mechanisms, of the kind traditionally employed in global chemical transport models, have also been used in box model mode, to compare against the measured constituent data and to investigate the significance of processes. Thus, the CiTTyCAT model (Wild et al., 1996; Evans et al., 2000; Emmerson et al., 2004; Donovan et al., 2005) has been used to study the chemical regimes probed by the OP3 measurements, both over the rainforest and over palm oil plantations, and to explore how local air quality could be affected should palm oil (a major isoprene-emitter) become an even more dominant crop, replacing the rainforest. Hewitt et al. (2009) show that under these circumstances, what happens to local and regional-scale emissions of NO\(_x\) will be crucial. Without control of NO\(_x\) emissions, the levels of ozone could become unacceptably high, damaging both crop and human health.

We find that CiTTyCAT can represent the broad features of atmospheric composition above a tropical rainforest (Pugh et al., 2010a). In particular, the model can fit daytime NO\(_x\) and O\(_3\) chemistry. Ambient concentration measurements have been used to estimate a net NO flux of 5 µg N m\(^{-2}\) h\(^{-1}\). Deposition of intermediate VOC oxidation products can have a very significant influence on their concentrations and the chemistry of the boundary layer, highlighting the need for further investigation into deposition processes over tropical forests.

Another box model study with a reduced chemical mechanism has focussed on the ground-based measurements of ozone and NO\(_x\) (Pike et al., 2009). This study suggests that the chemical mechanism included in the Cambridge global chemistry transport model, p-TOMCAT (Cook et al., 2007; Voulgarakis et al., 2009a, b) can reproduce the measurements...
if a parametrization of mixing out of and into the boundary layer is included.

The Cambridge p-TOMCAT global model has been used to look at regional-scale chemistry. Figure 10 (Pike et al., 2009) shows surface ozone concentrations from two calculations for April 2008, with the model run at two different spatial resolutions (approx 3°×3° and 0.5°×0.5°, respectively). At the lower resolution (typical of multi-annual integrations), the model significantly overestimates surface ozone over Borneo. Little spatial structure is evident and detailed flight track comparisons (not shown) fail to reproduce much observed structure. In contrast, at the high resolution, the model now captures a significant land – sea difference in surface ozone, driven by deposition over the land. Modelled ozone concentrations are still somewhat higher than observed but are now closer to the observations.

6 Conclusions

The overall aim of the OP3 project was to better understand the interactions that exist between SE Asian rainforests, atmospheric composition and, ultimately, the Earth’s climate system. In pursuing this aim, an extensive field deployment was carried out in 2008, supported by satellite data retrievals and modelling studies. The field deployment, from April to the end of July 2008, comprised extensive composition and flux measurements within and above the rainforest canopy, along with a more limited set of measurements in and above an oil palm plantation. Composition measurements across most of the state of Sabah in northern Borneo, and out to sea on both windward and leeward sides of the island, were provided by flights of the FAAM BAe 146. Remote sensing of land cover has shown that the chosen sites were representative of large areas of Borneo and the rest of insular SE Asia; long-term meteorological and ozone data gathered as part of the Global Atmosphere Watch programme show that weather and ozone mixing ratios during the time period of the field deployment were typical of the long term record.

Composition measurements show that the rainforest site was not impacted significantly by anthropogenic pollution, and this is confirmed by satellite retrievals of NO₂ and HCHO. The dominant modulators of atmospheric chemistry at the rainforest site were therefore emissions of BVOCs and soil emissions of reactive nitrogen oxides. At the observed BVOC:NOX volume mixing ratio (∼100 pptv/pptv), current chemical models suggest that daytime maximum OH concentrations should be ca. 10⁵ radicals cm⁻³, but observed OH concentrations were an order of magnitude greater than this. We confirm, therefore, previous measurements which suggest that an unexplained source of OH must exist above tropical forests (Lelieveld et al., 2008) and continue to interrogate the data to find explanations for this.

Marked differences in BVOC and NOx concentrations, and in BVOC fluxes, were measured in the rainforest and the oil palm plantation. By an accident of the non-linearity of boundary-layer chemistry, ozone mixing ratios over the two landscapes were similar, although plumes of higher ozone were observed in air impacted by emissions from oil-palm processing plants.

The land-sea mosaic of SE Asia makes it difficult for global atmospheric chemistry-and-transport models run at typical resolution to capture the observed atmospheric composition; runs at higher horizontal resolution provide much better agreement with measurements. A missing OH source notwithstanding, simple chemistry schemes, as currently implemented in most global models, can match observed ozone and NOx diurnal patterns, if a parametrization of mixing out of and into the boundary layer is included.

Overall, the SE Asian rainforest has shown itself to have significant differences from the more intensively studied...
Amazon rainforest – in tree speciation and, hence, in BVOC emission potential, and in the apparent lack of seasonality of emissions. It also exhibits significant differences to the oil palm plantations that are replacing it. A striking similarity, however, is the sustained oxidative capacity of the atmosphere above the rainforest in both SE Asia and Amazonia. Our current inability to adequately explain this limits the confidence we can have in our projections of how atmospheric composition and climate may change in the future.

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