

# Rapid transport of East Asian pollution to the deep tropics

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**Abstract.** Anthropogenic emissions from East Asia have increased over recent decades. **These increases have led to changes in atmospheric composition as far afield as North America under the prevailing westerly winds.** Here we show that, during Northern Hemisphere (NH) winter, pollution originating in East Asia also directly affects atmospheric composition in the deep tropics. We present observations of marked intra-seasonal variability in the anthropogenic tracer perchloroethene (C<sub>2</sub>Cl<sub>4</sub>) collected at two locations in Borneo (117.84°E, 4.98°N and 118.00°E, 4.22°N) during the NH winter of 2008/2009. We use the NAME trajectory model to show that the observed enhancements in C<sub>2</sub>Cl<sub>4</sub> mixing ratio are caused by rapid meridional transport, in the form of ‘cold surges’, from the relatively polluted East Asian land mass. In these events air masses can move **from ~35°N to Borneo** in 4 days. We then present data from the Monitoring Atmospheric Composition and Climate reanalysis which suggests that air masses high in C<sub>2</sub>Cl<sub>4</sub> may also contain levels of the pollutants carbon monoxide and ozone that are approximately double the typical ‘background’ levels in Borneo. **In addition to strengthening the meridional transport from the north, cold surges can enhance convection in Southeast Asia, and further trajectory calculations indicate that the polluted air masses can subsequently be lifted to the tropical upper troposphere.** This suggests a potentially important connection between mid-latitude pollution sources and the very low stratosphere.

## 1 Introduction

The rapid growth of East Asian economies over recent decades has led to enhanced emissions of various pollutants, including ozone (O<sub>3</sub>) precursors (e.g. Granier et al., 2011) and halocarbons (e.g. Wan et al., 2009). The emitted pollutants are known to be transported eastward by the prevailing mid-latitude winds, and plumes have been observed at the edge of East Asia (e.g. Akimoto et al., 1996; Tanimoto et al., 2008) and over the Pacific (e.g. Liu et al., 2003; Hudman et al., 2004). East Asian pollution has also been shown to affect atmospheric composition further afield, leading to increased O<sub>3</sub> in both western North America during Northern Hemisphere (NH) spring (Cooper et al., 2010) and Hawaii during NH autumn (Lin et al., 2014). Southward transport of polluted air masses originating in East Asia, associated with the Northeast Monsoon which occurs during NH winter, has also been reported (e.g. Liu et al., 2003; Pochanart et al., 2003; Wang et al., 2003).

Here we describe an additional impact of East Asian pollution and show that particularly strong meridional transport events within the Northeast Monsoon, or ‘cold surges’, are able to transport polluted air masses to remote parts of equatorial Southeast Asia. These cold surges are typically caused by a southeasterly movement of the Siberian High pressure system, and are associated with movement of cold air masses towards Southern China and a strengthening of the northeasterly monsoon winds in the South China Sea (Zhang et al., 1997; Chan and Li, 2004; Chang et al., 2004). Cold surges are also known to increase convective activity generally in equatorial Southeast Asia, with enhancements near the northwest coast of Borneo often highlighted (e.g. Slingo, 1998; Compo et al., 1999; Chang et al., 2005). However, their influence on atmospheric composition in the region has yet to be demonstrated.

It is also interesting to consider the possibility of pollutant transport via convection. For example, while the detailed mechanisms for transport into the stratosphere are the subject of current debate (e.g. Park et al., 2009; Bergman et al., 2013), strong uplift of polluted air masses has already been demonstrated during the Asian (NH) Summer Monsoon (Lawrence and Lelieveld, 2010; Randel et al., 2010). Such a process may also be important in the context of our study because Southeast Asian air masses are preferentially lifted towards the stratosphere during NH winter (e.g. Levine et al., 2007; Aschmann et al., 2009), and because of the link between cold surges and convection outlined above.

Our analysis is based on observations of an anthropogenic halocarbon, perchloroethene (C<sub>2</sub>Cl<sub>4</sub>), collected in Borneo during the winter of 2008/2009 (Sect. 2). We conduct simulations with a trajectory model to confirm the influence of East Asian pollution on our measurements (Sect. 3). We then investigate the wider air quality implications of these transport events by studying data from the Monitoring Atmospheric Composition and Climate (MACC) reanalysis (Sect. 4). In Sect. 5 we use further trajectory calculations to investigate whether the polluted air masses are lifted towards the upper troposphere once they have reached the deep tropics. Finally we discuss the implications of our results (Sect. 6).

## 2 Observations

55  $C_2Cl_4$  is an excellent marker of air masses recently subjected to industrial emissions for several rea-  
sons. First, the sources of  $C_2Cl_4$  are predominantly anthropogenic, and largely aseasonal. They  
span a range of commercial and industrial activities, which include dry-cleaning and metal de-  
greasing (McCulloch et al., 1999). Global emissions, and background mixing ratios, have declined  
over the past 20 years or so (McCulloch et al., 1999; Simpson et al., 2004, and data available at  
60 <http://agage.eas.gatech.edu/data.htm>). Nevertheless, there is evidence for continued and significant  
 $C_2Cl_4$  emissions from East Asia over this period (e.g. Barletta et al., 2006, 2009; Shao et al., 2011).  
Second,  $C_2Cl_4$  has a straightforward atmospheric chemistry **which is dominated by reaction with the  
hydroxyl radical** (Singh et al., 1996). Under typical tropospheric conditions its lifetime is  $\sim 3$  months  
(Montzka and Reimann, 2011). This lifetime is both long enough for large-scale transport and short  
65 enough for measurable inhomogeneities to exist. Finally, it appears that East Asian  $C_2Cl_4$  emis-  
sions are similarly distributed to the anthropogenic portion of carbon monoxide (CO, a key ozone-  
precursor) emissions in the region (e.g. Wang et al., 2003; de Gouw et al., 2004; Shao et al., 2011).

**Few continuous measurements of atmospheric composition have been reported in Southeast Asia  
to date.** To fill this gap we have used University of Cambridge  $\mu$ -Dirac instruments (Gostlow et al.,  
70 2010) to measure a suite of halocarbons at a number of sites in the region (see Pyle et al., 2011;  
Robinson et al., 2014). Here we focus on  $C_2Cl_4$  observations collected at two locations in Sabah,  
Malaysian Borneo. The sites, the WMO Global Atmospheric Watch station at Bukit Atur (117.84°E,  
4.98°N) and a Global Satria facility near Tawau (118.00°E, 4.22°N), are  $\sim 85$  km apart. **The data we  
consider here were collected during one NH winter, between 22 November 2008 and 28 February  
75 2009, and have been averaged over 3 h periods (Fig. 1a). There are clear variations in the data at  
weekly scales, which occur concurrently at the two sites (coefficient of determination,  $r^2 = 0.85$   
in Fig. 1b). This coherence is indicative of changes in composition occurring over relatively large  
scales, and suggests very local emissions of  $C_2Cl_4$  do not have an important influence on the data.  
Our measurements are also characterised by a number of abrupt changes between ‘background’  
80 mixing ratios ( $\sim 1.0$ – $1.5$  ppt), and ‘polluted’ mixing ratios ( $\sim 2.0$ – $3.0$  ppt). We will show in Sect. 3  
that changes in Northeast Monsoon circulation are the primary cause of these transitions. **We have  
observed similar features in measurements collected in Borneo during the two subsequent winters  
(2009/2010 and 2010/2011) and are therefore confident these changes are typical of the region.****

Fig. 1 shows **that** there are some differences in the absolute mixing ratio observed at the two  
85 sites. Instrument performance is generally good for this compound (Gostlow et al., 2010) **although  
Robinson et al. (2014) have raised** the possibility of calibration uncertainty leading to part of this  
difference. **In this study we focus on the variations observed concurrently at both locations, and  
hence the absolute mixing ratios are of lesser importance.**

### 3 Transport pathways

90 We use the Met Office’s Lagrangian atmospheric dispersion model, NAME (Jones et al., 2007), to interpret our  $C_2Cl_4$  measurements. Trajectories are calculated using three-dimensional meteorological fields produced by the **Met Office’s Numerical Weather Prediction tool, the Unified Model (UM)**. **These fields have a horizontal grid resolution of  $0.5625^\circ$  longitude by  $0.375^\circ$  latitude, 31 vertical levels below  $\sim 19$  km, and are available at 3 h intervals. Vertical velocities are obtained from the UM**  
95 **and available at grid nodes. The sub-grid-scale processes of convection (currently available only in forward mode, see Meneguz and Thomson, 2014a, b) and turbulence (available in both forward and backward modes, see Webster et al., 2003; Morrison and Webster, 2005) are parameterised in NAME.**

Initially we use NAME to calculate batches of **inert** backward trajectories started at each measurement site within an altitude range of 0–100 m. 33000 trajectories were started throughout each  
100 3 h period for which measurements were available at a particular location. They ran for 12 days, and every 15 minutes the location of all trajectories within the lowest 18 km of the model atmosphere was recorded on a grid with the same horizontal resolution as the meteorological fields.

The result of the above calculation is that a map of time-integrated and column-integrated trajectory (or ‘particle’) density exists for each 3 h period. Fig. 2 shows composites of these maps when they are grouped according to the 3 h mean  $C_2Cl_4$  mixing ratio at Tawau (maps for Bukit Atur are very similar, and not shown). These maps indicate that air masses containing the lowest mixing ratios ( $< 0.8$  ppt) often passed to the south of the Philippines during transport from the unpolluted subtropical Pacific. **As a larger fraction of trajectories travel from mid-latitudes, passing through**  
105 **the Philippines, the  $C_2Cl_4$  mixing ratio at Tawau increases. Beyond the Philippines, there are two diverging origins.** Air masses containing  $\sim 1.0$ – $1.4$  ppt of  $C_2Cl_4$  are largely transported from the Pacific Ocean. In contrast, air masses containing  $> 1.6$  ppt of  $C_2Cl_4$  appear to be increasingly affected by an anti-cyclonic circulation, originating from the Asian land mass and containing higher levels of anthropogenic pollution. This difference is qualitatively consistent with the idea of cold surges  
115 leading to elevated  $C_2Cl_4$  in Borneo.

In a further, more quantitative analysis we examine the relationship between the 3 h mean  $C_2Cl_4$  mixing ratios and the fraction of trajectory mass (or equivalently of trajectory residence time) north of  $35^\circ N$  in the corresponding individual air history maps. This fraction increases linearly as the  $C_2Cl_4$  mixing ratio **increases** (for both sites  $r^2 > 0.7$ , **graph not shown**). Other latitude thresholds  
120 (between  $25$ – $45^\circ N$ ) were tested and found to yield similar relationships. This reinforces the argument made above: cold surges are able to move polluted air from northern mid-latitudes rapidly (**over  $\sim 4$  days in the case study we consider in Sect. 4**) to equatorial Southeast Asia.

#### 4 Wider air quality implications

Thus far we have focused on  $C_2Cl_4$ , an industrial pollutant observed in relatively small quantities. It is clearly of interest to consider how the pollution transport pathway identified in Sect. 3 influences air quality in Southeast Asia more generally. However, the continuous air quality measurements collected in Northern Borneo that we are aware of appear to be affected by local sources of pollution. As such they are unlikely to reflect the regional mechanism suggested by the coherent variations in  $C_2Cl_4$  observed at our two measurement sites (Sect. 2) and by our trajectory calculations. In this section we therefore use data available from the MACC Reanalysis (Inness et al., 2013, downloaded from [http://apps.ecmwf.int/datasets/data/macc\\_reanalysis/](http://apps.ecmwf.int/datasets/data/macc_reanalysis/)). This dataset is created using satellite observations, emission inventories and chemical transport model calculations. For the two pollutants we consider in this section, CO and  $O_3$ , Inness et al. (2013) report small negative biases of  $\sim 10\%$  and  $\sim 20\%$  respectively when the MACC data are compared to available independent observations in the tropical troposphere. We concentrate here on a two week period containing a particularly strong surge event (10–23 January 2009, shaded in Fig. 1a), during which there are abrupt changes in our data and the highest  $C_2Cl_4$  mixing ratios occur.

To begin, we extract time series for CO and  $O_3$  (Fig. 3) in the grid cell nearest to Bukit Atur in the MACC reanalysis. Our  $C_2Cl_4$  measurements from Bukit Atur are also plotted for comparison. We obtained MACC data at 925 hPa (rather than 1000 hPa) because Bukit Atur, at an elevation of 426 m, often appears to be more representative of the free troposphere (Pike et al., 2010). In Fig. 3 the mixing ratios of  $C_2Cl_4$ , CO and  $O_3$  all follow the same pattern, with a sustained increase to maximum values on 15 January, followed by a decline back to more typical ‘background’ mixing ratios in the subsequent days. This suggests that the air masses arriving in Borneo with high levels of  $C_2Cl_4$  also contain a range of other pollutants and significantly impact air quality in this part of the tropics.

Next, in Fig. 4 we contrast daily mean maps of CO and  $O_3$  from the MACC reanalysis (again at 925 hPa) for the two days shaded in Fig. 3: 15 January 2009, when concentrations of  $C_2Cl_4$  were relatively high, and 20 January 2009, when concentrations of  $C_2Cl_4$  were relatively low. Corresponding daily mean air history maps are also presented. In these maps the mean horizontal locations of the back trajectories after 2 and 4 days are marked to highlight the strength of the cold surge event. All together, the six maps demonstrate that our measurement sites in Northern Borneo sit near the edge of a steep pollution gradient associated with contrasting polluted air masses from East Asia and cleaner air masses from the tropical Pacific. On days when air masses are moved rapidly by cold surges from East Asia towards Borneo, such as 15 January (top row of Fig. 4), air quality is significantly reduced. According to the MACC reanalysis mixing ratios of CO and  $O_3$  can reach, respectively,  $\sim 150$  ppb and  $\sim 40$  ppb. Our high  $C_2Cl_4$  measurements are excellent markers of this pollution. By contrast, modelled levels of CO and  $O_3$  and measured levels of  $C_2Cl_4$  are more repre-

sentative of the local background (approximately half of the polluted levels) on days when the winds  
160 blow from the Pacific, such as 20 January (bottom row of Fig. 4).

## 5 Uplift of polluted air masses

As noted in the introduction, cold surges are known to affect the characteristics of convection in Southeast Asia (e.g. Chang et al., 2005). A further NAME calculation was performed to assess whether air masses that move rapidly from mid-latitudes to the tropics may subsequently ascend.  
165 Previous work suggests that NAME is a useful tool for analysing vertical transport of relatively short-lived compounds in regions of tropical convection (Ashfold et al., 2012). In this case, such transport could lift East Asian pollution to the tropical upper troposphere. Forward trajectories ( $3000 \text{ h}^{-1}$ ) were released continuously in a kilometer deep surface box over East Asia, covering  $100\text{--}140^\circ\text{E}$ ,  $30\text{--}50^\circ\text{N}$ . These trajectories were not subject to any chemical losses and their travel was recorded  
170 for 12 days. The aim was not to simulate any particular pollutant, but to consider in a simple manner the transport of air masses originating in the polluted East Asian mid-latitudes.

To assess whether the mid-latitude air masses are lifted in the tropics, the shading in Fig. 5a shows the density of particles (i.e. trajectories), between 600–400 hPa (i.e. the mid-troposphere) and between  $0\text{--}10^\circ\text{N}$  during January 2009 for Southeast Asian longitudes. Also plotted are CO contours  
175 from the MACC reanalysis at 500 hPa in the same time–longitude space. Within the figure there are periods, notably around 12–13 January near the time of a strong cold surge (Figs. 3 and 4), when peaks in simulated mid-latitude air masses coincide with enhanced CO mixing ratios. Similar regions of agreement are found at the same time in a corresponding plot for  $\sim 200 \text{ hPa}$  (Fig. 5b). These features indicate that East Asian mid-latitude pollution is capable of influencing atmospheric  
180 composition through much of the depth of the Southeast Asian tropical troposphere.

Our argument is supported further by Fig. 6a and b, which show, respectively, longitude–altitude and latitude–altitude slices through the Southeast Asian atmosphere on 15 January 2009 (our selected ‘cold surge’ day in Fig. 4). Again, the density of the mid-latitude tracer is indicated with shading. The plots demonstrate that the air masses originating in mid-latitudes can be lifted above 200 hPa  
185 within the tropics. Evidence for this type of vertical transport is weaker in similar plots for days without cold surge activity (20 January, for example, Fig. 6c and d). An additional analysis of the trajectory timescales shows that it can take fewer than 10 days for air masses to travel from the East Asian boundary layer to the tropical upper troposphere (i.e. above 200 hPa). This is a sufficiently short time for this process to be important for even relatively short-lived pollutants.

## 190 6 Summary and discussion

We analyse observations of  $\text{C}_2\text{Cl}_4$ , an excellent tracer for anthropogenic pollution, collected at two sites in Borneo during the NH winter of 2008/2009. Backward trajectories calculated using NAME

show that measurements of high  $C_2Cl_4$  mixing ratios were associated with rapid ‘cold surge’ transport towards the equator from polluted mid-latitude East Asia. Data available from the MACC re-analysis demonstrate that the polluted air masses may contain levels of CO and O<sub>3</sub> that are approximately double the typical ‘background’ levels in Borneo. Steep gradients in atmospheric composition have been identified before in this region (Hamilton et al., 2008), but the gradients identified here appear to be mobile (Fig. 4) and capable of periodically reducing air quality significantly in Southeast Asia. Once in the deep tropics, we show that the polluted air masses can be lifted out of the boundary layer into the mid- and upper troposphere. This process may be enhanced by the more vigorous regional convection typically induced by the cold surge circulation, as considered before within the framework of Borneo Vortex events (Braesicke et al., 2012). **Our trajectory calculations suggest that in total, transport from the East Asian boundary layer to the tropical upper troposphere (above 200 hPa) can occur in fewer than 10 days.** The tropical upper troposphere is a key gateway to the lower stratosphere (e.g. Fueglistaler et al., 2009). The route we have identified may therefore transport polluted air from NH mid-latitudes into the tropical upper troposphere and thence to the lower stratosphere during NH winter. **To illustrate the potential importance of this route, air masses enriched in  $C_2Cl_4$  may also contain high levels of various other relatively short-lived chlorine containing gases (e.g. Wang et al., 2014) that could have a negative impact on stratospheric ozone. The significance of this transport pathway therefore needs to be investigated further.**

Pollutant emissions from East Asia have increased over recent decades, and they are unlikely to decline significantly in the next 20 years or so (e.g. Zhao et al., 2014). At the same time, rising greenhouse gas levels could lead to changes in the regional climate. There remains, however, significant uncertainty in how the various large-scale processes that influence East and Southeast Asian climate variability will change this century (Christensen et al., 2013). Any climatic changes are likely to influence the importance of the pollution transport pathway we have identified. To illustrate, the El Niño–Southern Oscillation is known to moderate the strength of cold surge activity (e.g. Zhang et al., 1997). In order to **better quantify** the future impact of East Asian pollution on the tropical atmosphere there is a need to increase our understanding of both past multi-decadal variations (e.g. Huang et al., 2011), and possible future changes (e.g. Park et al., 2011), in cold surge frequency.

Observational evidence from other parts of Southeast Asia will be needed to assess more fully the influence these transport events have on regional atmospheric composition. One step in this direction is afforded by a new program of long-term measurements of a suite of compounds (including the halocarbons measured by  $\mu$ -Dirac) at the University of Malaya’s Bachok Marine Research Station on the east coast of Peninsular Malaysia ([http://www.ioes.um.edu.my/research\\_facilities.html](http://www.ioes.um.edu.my/research_facilities.html)). Levels of pollution associated with the prevailing northeasterly flow through the South China Sea may be more severe in, for example, Peninsular Malaysia, than those in Borneo. There are clearly possible implications for human health, though the effect of pollutants such as O<sub>3</sub> on tropical forests and

230 crops is yet to be well understood (e.g. Ainsworth et al., 2012). Further long-term measurements  
will also facilitate more detailed investigation of the influences of the climatic variations discussed  
above.

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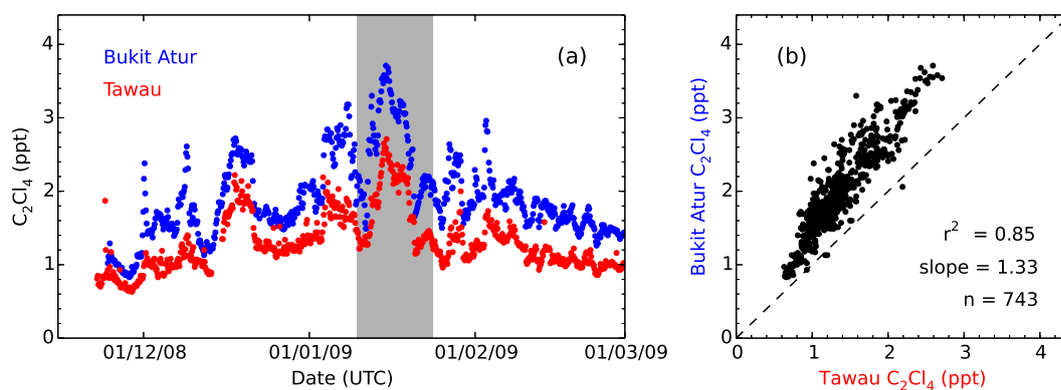
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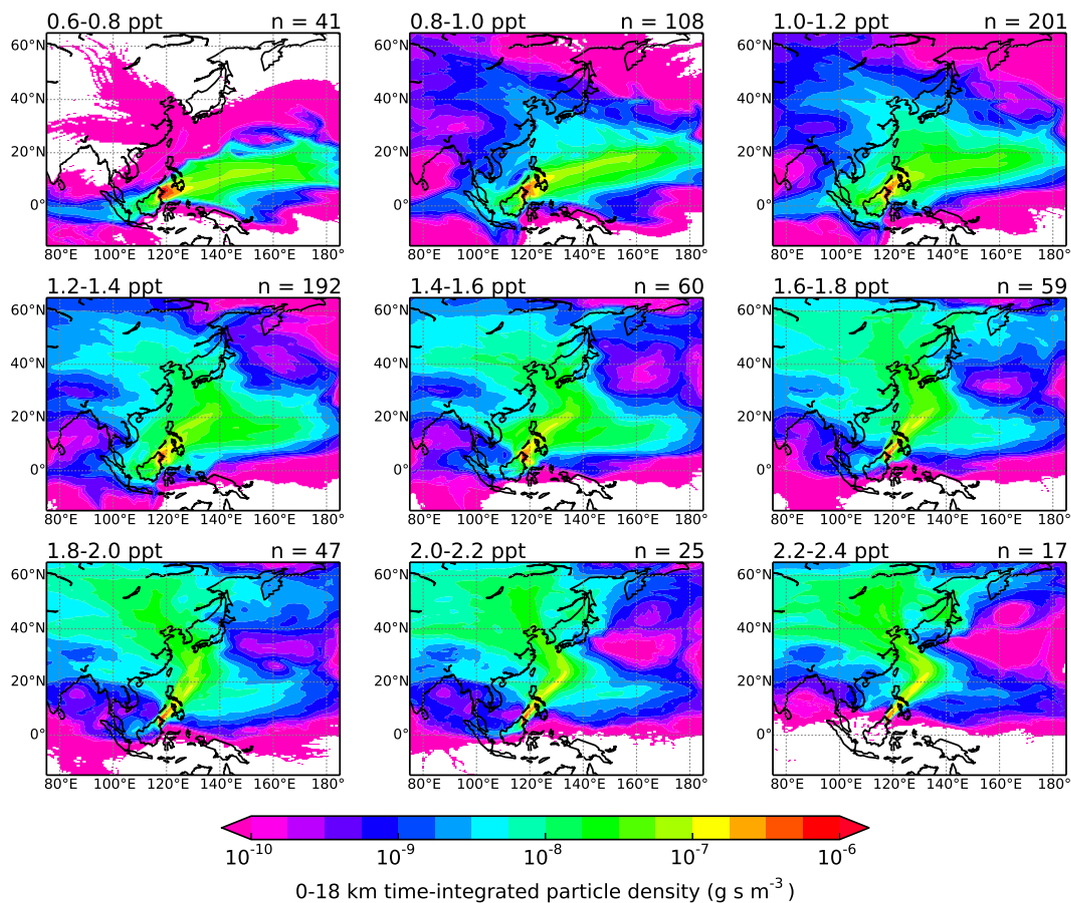
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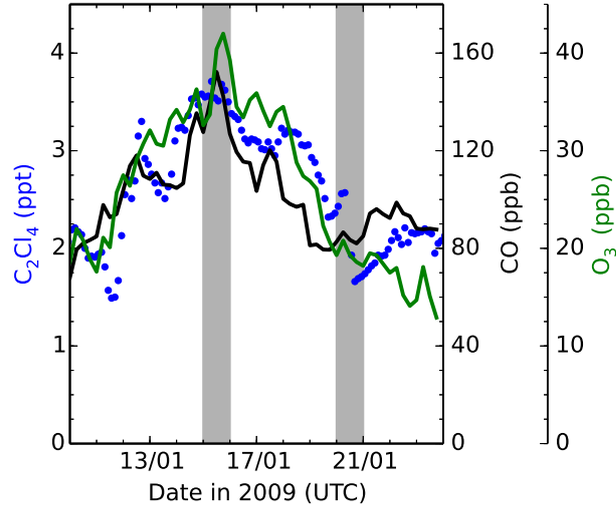
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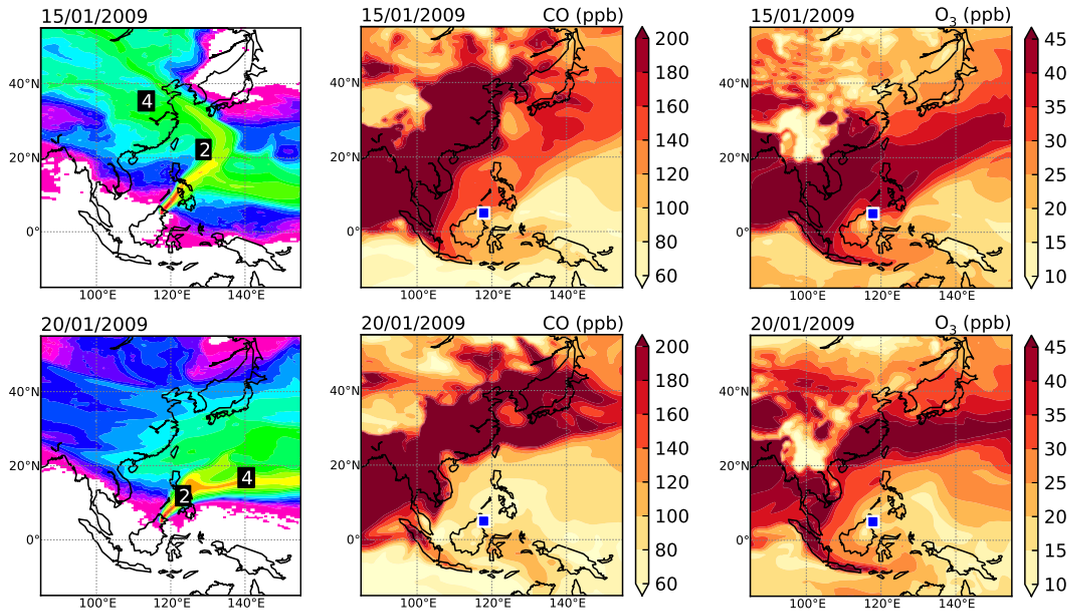
**Figure 1.** (a) shows a time series of 3 h mean  $C_2Cl_4$  measurements collected at Bukit Atur (blue) and Tawau (red) during the winter of 2008/2009. The period analysed in more detail in Section 4 is shaded grey. (b) shows the correlation between the measurements at the two sites. The coefficient of determination ( $r^2$ ), the gradient of the regression line (slope) and the number of 3 h periods for which data exist at both measurement locations ( $n$ ) are noted. In constructing this figure two outlying 3 h mean measurements from Tawau ( $>6$  ppt) have been discounted. Both high mixing ratios are likely to be due to a very local  $C_2Cl_4$  source.



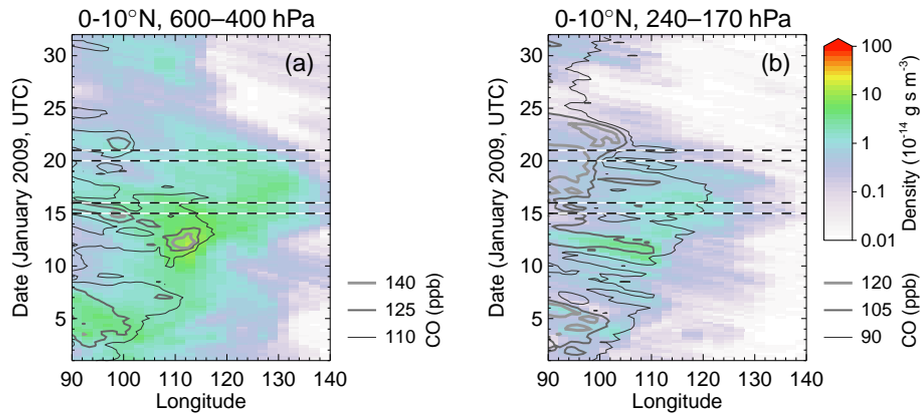
**Figure 2.** Composite air history maps, for each  $\text{C}_2\text{Cl}_4$  mixing ratio interval of 0.2 ppt at Tawau. The number of 3 h periods,  $n$ , contributing to each composite is noted above each panel. A mixing ratio greater than 2.4 ppt was observed in eleven 3 h windows; these periods are not accounted for in the figure.



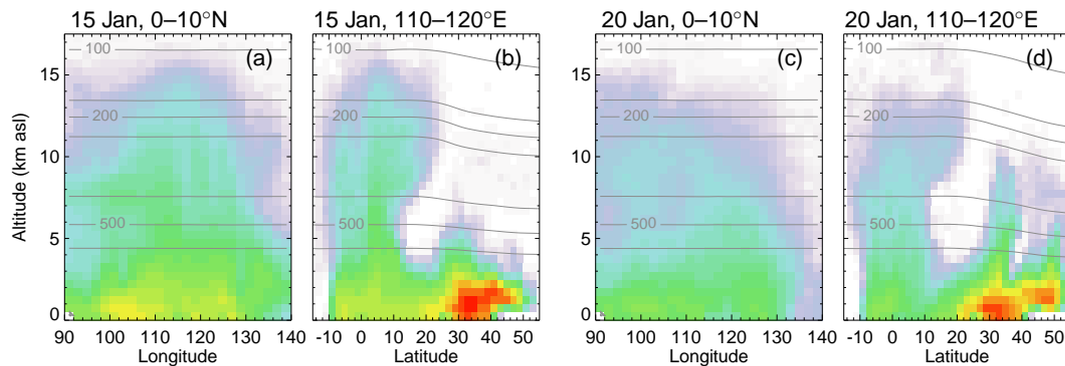
**Figure 3.** Time series of 3 h mean  $C_2Cl_4$  measurements from Bukit Atur (blue) and 6 hourly CO (black line) and  $O_3$  (green line) at  $118^\circ E$ ,  $5^\circ N$ , 925 hPa in the MACC reanalysis. The days analysed in more detail in Fig. 4 are shaded grey.



**Figure 4.** Daily mean air history maps calculated using NAME (left), CO at 925 hPa from the MACC reanalysis (center) and  $O_3$  at 925 hPa from the MACC reanalysis (right) for 15 January 2009 (top) and 20 January 2009 (bottom). The colour scale for the air history maps is the same as in Fig. 2, and the mean horizontal location of the trajectories after 2 and 4 days is marked. In the CO and  $O_3$  panels the location of Bukit Atur is marked with a blue square.



**Figure 5.** Time-integrated particle (i.e. trajectory) density, as a function of longitude and time through January 2009, resulting from a mid-latitude particle source in NAME. (a) shows particle density between 600–400 hPa averaged between 0–10°N. (b) shows particle density between 240–170 hPa, also averaged between 0–10°N. The contours show CO from the MACC reanalysis at, respectively, 500 hPa and 200 hPa, for the same spatial and temporal dimensions.



**Figure 6.** Time-integrated particle (i.e. trajectory) density resulting from a mid-latitude particle source in NAME, on 15 and 20 January 2009 (marked with dashed lines in Fig. 5). The colour scale is the same as in Fig. 5. (a) shows a longitude–altitude cross-section for 15 January 2009, averaged over 0–10°N. (b) shows a latitude–altitude cross-section, also for 15 January 2009, and averaged over 110–120°E. (c) and (d) show the same for 20 January 2009. The altitude scale is km above sea level (asl). Pressure contours (in hPa) from NAME's driving meteorological data are marked with grey lines (contours without labels bound the pressure ranges used in Fig. 5).