Airborne measurements indicate large methane emissions from the eastern Amazon basin

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[1] Recent results from laboratory, field and remote sensing measurements suggest the presence of large methane emissions from the Amazon basin. Here we present regionally integrative, direct trace gas observations from two sites that confirm the presence of large fluxes of methane in eastern Amazonia. Air samples collected on aircraft near Santarém (2.9°S, 55.0°W) and Manaus (2.6°S, 60.0°W) in eastern and central Amazonia show large enhancements of CH4 that are not seen at the NOAA/ESRL background sites in the tropical Atlantic Ocean. From the surface to about four km, enhancements averaging 34 ppb and up to 200 ppb occur throughout the year and we calculate enhancements averaging 27 mg CH4/m²/day from upwind sources. Citation: Miller, J. B., L. V. Gatti, M. T. S. d’Amelio, A. M. Crotwell, E. J. Dlugokencky, P. Bakwin, P. Artaxo, and P. P. Tans (2007), Airborne measurements indicate large methane emissions from the eastern Amazon basin, Geophys. Res. Lett., 34, L10809, doi:10.1029/2006GL029213.

1. Introduction

[2] Understanding methane growth rate variations and the processes responsible for them is difficult because of the wide variety of methane sources [Cicerone and Oremland, 2006] and the problem of distinguishing individual sources in the atmosphere even when using isotopes [Miller et al., 2002]. Until recently, ESRL measurements have only been made in the tropical marine boundary layer (MBL), where they are largely decoupled from continental emissions. This greatly limits our ability to infer tropical CH4 fluxes [e.g., Bousquet et al., 2006; Houweling et al., 1999].

[3] Retrievals of column-mean CH4 mixing ratio from SCIAMACHY during 2003 and 2004 [Frankenberg et al., 2006] show CH4 abundances above the Amazon basin and other parts of northern South America substantially larger than expected from process-based model estimates of methane emission from wetlands [Walter et al., 2001]. However, more recent estimates of Amazonian wetland emissions [Melack et al., 2004] show more consistency with SCIAMACHY observations [Bergamaschi et al., 2006]. Additionally, recent laboratory results [Keppler et al., 2006] pointing to plants as direct emitters of methane and field measurements from Amazonian upland forests [do Carmo et al., 2006] suggest that sources other than wetlands may also be important contributors to Amazonian methane emissions.

[4] Here, we present a four year record of CH4 vertical profiles from the surface to at least 3600 m (asl), above two sites in the central Amazon, near the cities of Santarém (SAN) and Manaus (MAN) (Figure 1). These data do not have the spatial and temporal density of the satellite columns, but are very accurate and precise and being direct measurements do not suffer from any of the biases of SCIAMACHY, including assumptions about CO2 mixing ratios (to which SCIAMACHY CH4 radiances are scaled) and aerosol contamination. SCIAMACHY also cannot reliably retrieve methane over the oceans, so that its marine – continental gradients are not well defined. We reference our Amazonian measurements to those made at our remote sampling sites at Ascension Island (ASC; 7.9°S, 14.4°W) and Barbados (RPB; 13.2°N, 59.4°W) located in the tropical Atlantic, because these sites represent air entering the Amazon basin through the trade winds. We will use enhancements of CH4 at SAN or MAN relative to background to estimate the surface flux of CH4 between Brazil’s Atlantic coast and the sites.

2. Methods

[5] At SAN and MAN, air was collected with portable sampling systems consisting of separate compressor and flask units [Tans et al., 1996]. These units are loaded onto a light aircraft, and the pilot initiates sample collection at predetermined altitudes. Most flights consisted of one descending and one ascending profile from 3600 m to 300 m. From 2000 to 2003, samples collected in Brazil were sent to the NOAA lab in Boulder, USA, where they were analyzed for CO2, CH4, CO, N2O, SF6, and H2. Measurement repeatability for CH4 is estimated to be better than 0.1% (<2 ppb, 1σ), and all CH4 measurements reported here are on the NOAA2004 scale [Dlugokencky et al., 2005]. Since 2004, a replica of the NOAA analysis system began operating in Brazil at Instituto de Pesquisas Energéticas e Nucleares (IPEN), with precision and accuracy very similar to that at NOAA. Air at ASC and RPB was sampled into 2.2 L glass flasks with Teflon-tipped glass stopcocks and filled to about 1.2 bar [Conway et al., 1994], and shipped to NOAA for analysis of the same suite of gases. All measurements presented here are available via anonymous ftp at ftp.cmdl.noaa.gov/pub/LBA.

[6] At SAN, ascending profiles were made above the Tapajos National Forest, near the “km 67” tower that is located about 10 km to the east of the Tapajos river.
first 4 years, descending profiles were made 30 km to the east of the tower to evaluate possible impacts of fluxes from the river [Richey et al., 2002]. At MAN, ascending profiles were made above the “K34” tower to the northwest of the city of Manaus (population 1.8 million), following descending profiles 50 km to the northeast of Manaus, in order to avoid possible pollution from Manaus at the K34 tower, which can be downwind of the city. At both SAN and MAN, differences between ascending and descending profiles were generally small compared to differences with ASC (Figure S1 of the auxiliary material),1 so for our analysis, we use the vertically denser profiles above the towers at both sites, reserving the alternate profiles for sensitivity tests. Between 2000 and 2003, 11 vertical profiles were also conducted off the Atlantic coast of Brazil, 50 km NE of the city of Fortaleza (FTL, 4.15°S, 38.28°W) to sample air entering the Amazon basin.

3. Results and Discussion

[7] Air entering the Amazon basin is dominated by trade-wind easterlies coming from the tropical Atlantic Ocean, although the relative influence of Northern and Southern Hemisphere air depends upon the seasonally varying latitude of the ITCZ. Thus, the difference between our SAN and MAN measurements and the Atlantic background should be directly related to terrestrial CH₄ fluxes; oceanic CH₄ fluxes are assumed negligible as supported by the small differences between measurements at FTL and ASC (Figure 2). Until 2004, the vast majority of measurements took place during the wet season, but since that time there have been numerous dry season profiles. Large enhancements are evident in both seasons and there are no clear differences in the character of dry and wet season profiles. However, during the dry season, CO can also be significantly enhanced suggesting that elevated CH₄ in the same samples results from biomass burning. The ratio of CO to CH₄ enhancements relative to background ranges from 0.13 to 0.82 mol/mol in the wet season and 1.8 to 6.3 mol/mol during the dry season. A review of emission measurements [Andreae and Merlet, 2001] suggests a ratio of 9 mol/mol for tropical forest burning, roughly consistent with our observations.

[8] Figure 2 shows that CH₄ at SAN and MAN relative to ASC and RPB is almost always enhanced, thus indicating the presence of upwind sources. The largest enhancements are in the convective boundary layer (CBL), but enhancements in the free troposphere can also be seen in numerous profiles, possibly indicating the convective redistribution of methane emitted into the CBL. Because of strong convection one cannot count on surface emissions to be trapped in the CBL, so we do not calculate fluxes using a boundary layer budgeting technique [e.g., Lloyd et al., 2001]. Instead, we use a column integration technique that does not distinguish the CBL and free troposphere.

[9] To apply this technique, we first determine the background CH₄ mixing ratio entering Brazil off the Atlantic Ocean. Because the relative Northern and Southern Hemisphere contributions to the CH₄ background vary, we use co-measured sulfur hexafluoride (SF₆, a purely anthropogenic gas) as a transport tracer. Almost all SF₆ is emitted in the NH, and there are essentially no emissions of SF₆ between the coast and our sites [Olivier et al., 1999]. Thus, all variations seen at our aircraft sites result from varying amounts of Northern and Southern Hemisphere tropical air arriving at our sites. Figure 2 shows that most of the time SF₆ at SAN and MAN is bounded by the time series from ASC and RPB. Using a simple two-end-member mixing model, we then calculate the fractions of air arriving at our Amazonian sites that can be represented by the background sites ASC and RPB, which can then be applied to any other conserved tracer (equations 1 and 2).

\[
ASC_{site} = 1 - RPB_{site} = \frac{SF_6_{site} - SF_6_{RPB}}{SF_6_{ASC} - SF_6_{RPB}}
\]  

\[
X_{bg} = ASC_{SAN} \times X_{ASC} + RPB_{SAN} \times X_{RPB}
\]  

ASC_{site} and RPB_{site} are the respective fractions of air arriving at our Amazonian sites (MAN or SAN), SF₆_{site} is the median SF₆ from the vertical profile site, and SF₆_{(RPB or ASC)} is the SF₆ mixing ratio extracted from a smoothed curve fit [Thoning et al., 1989] of the background data at the same date as the site observations; X refers to the mixing ratio of any co-measured gas. We then calculate the enhancement above background by subtracting X_{bg} from measurements at all altitudes (Figure 3). Measurements of CH₄ at FTL show free troposphere (>1500 m) CH₄ enhancements of 10–20 ppb (Figure S2), indicating that it is not totally justified to use a single MBL-based point, X_{bg}, as a background for the entire 0 to 4 km range. However, the FTL enhancements are small compared to the Amazonian sites and only a few FTL profiles are available for reference, requiring us to use the temporally dense time series at ASC and RPB.

[10] Figure 3 shows the difference between vertical profile measurements of CH₄ at site (SAN or MAN) and the background as calculated in (2) for the dry and wet seasons. There is enhancement in the CBL, while in the free

1Auxiliary materials are available in the HTML. doi:10.1029/2006GL029213.
The differenced profiles can be converted to surface fluxes by integrating the CH$_4$ content from the surface to the top of the profile, and normalizing by the time since the air was at the coast, according to:

$$F_{\text{CH}_4} = \frac{\int_0^4 (\left[\text{CH}_4\right]_{\text{site}} - \left[\text{CH}_4\right]_{\text{bg}}) dz}{t}$$

where $[\text{CH}_4]$ is the concentration of CH$_4$ in mol/m$^3$, which can be determined from vertical profiles of mixing ratio, temperature and pressure (estimated using a lapse rate of 6.5 K/km and a scale height of 7 km). $t$ is the time since the air has been over land, estimated using mean 850 mb windspeed of 10 m/s (www.cdc.noaa.gov/cdc/data.ncep.html) and a mean distance to the coast of 1700 km (it is much less to the northeast and more to the southeast); we derive a mean value for $t$ of 2 days, to which we assign a 50% uncertainty. Uncertainties in SF$_6$ and CH$_4$ values used in equations 1 and 2 are 0.05 ppt and 10 ppb, respectively, and are based on the scatter about smooth curve fits in Figure 2; uncertainty in $[\text{CH}_4]_{\text{site}}$ in equation 3 is just the measurement uncertainty of 2 ppb. Uncertainty in $F_{\text{CH}_4}$ as shown in Figure 4 is estimated by propagating uncertainty from all terms in equations 1–3. The sensitivity of $F$ to possible biases is also tested and discussed below.

[12] Fluxes estimated using SAN data (Figure 4) average 35 ± 23 mg CH$_4$/m$^2$/day and 20 ± 17 mg CH$_4$/m$^2$/day at MAN. The mean uncertainty of each flux determination at SAN and MAN is 21 and 15 mg CH$_4$/m$^2$/day, respectively. These integrated fluxes are significantly larger than any single flux averaged over a large area. Basinwide wetland emissions determined from chamber measurements and remote sensing [Melack et al., 2004] of 29 Tg CH$_4$/yr distributed over 5 $\times$ 10$^6$ km$^2$ equate to 16 mg CH$_4$/m$^2$/day. Scaling direct plant emissions from laboratory chambers is uncertain [Ferretti et al., 2007; Houweling et al., 2006; Keppler et al., 2006; Kirschbaum et al., 2006], but here we use 4 mg CH$_4$/m$^2$/day (see auxiliary material); nighttime emissions of unknown origin measured at upland Amazonian sites [do Carmo et al., 2006] have a median of about 5 mg CH$_4$/m$^2$/day, but if they are plant emissions, they could be significantly larger in the presence of sunlight [Keppler et al., 2006]. Maximum fire emissions over a five year period from 65°W to the Atlantic coast between 5°S and the equator are estimated to be 5 $\times$ 10$^6$ g CH$_4$/month (G. van der Werf, personal communication, 2006). If we assume that fires occur one third of the days during the dry season, this equates to 5 mg CH$_4$/m$^2$/day for days when fires occur. Termitic sources ($0.5$ mg/m$^2$/day) [Martins et al., 1993], consumption by soils (<1 mg/m$^2$/day) [Keller et al., 2005], and consumption by OH (3 mg/m$^2$/day), also likely contribute. No individual process estimates can explain our mean values of 35 and 20 mg CH$_4$/m$^2$/day based on SAN and MAN observations, but all sources and sinks total 23 mg CH$_4$/m$^2$/day, closer to our observations. Wetland emissions are likely to be the most important source in the wet season, while during the dry season a combination of wetland, fire and other sources influence the observed CH$_4$ enhancements. We do not presently have a way of distinguishing direct-plant and wetland emissions, so at present we assume emissions from plants that are uniform throughout the year.

[13] One potential experimental bias is whether our air samples are representative of large areas or just the area near the sampling sites. There is no consistent bias between our two profiles taken 30 km apart (Figure S1), although there is significant spread at the lowest level suggesting some influence from local fluxes. Re-calculating fluxes using the non-tower profiles or by not using the lowest two (300 and 600 m) levels reduces the mean flux at the sites.
by only 0–5 mg CH$_4$/m$^2$/day, indicating that most of the integrated signal is not local ($\sim10^2$ km$^2$), but regional ($\sim10^5$ km$^2$).

14 Bias may also result from our measurements not exceeding about four km. Due to convection, some methane emissions will affect the profile above this height, and neglecting these altitudes will result in an underestimate of emissions. On the other hand, free troposphere FTL data show enhancements of about 15 ppb above two km (relative to the MBL); this translates to a positive bias of 6 mg CH$_4$/m$^2$/day (assuming two days travel time from FTL to SAN). Neglecting destruction of CH$_4$ introduces a negative bias of about 3 mg CH$_4$/m$^2$/day in a column from the surface to 4 km.

4. Conclusions

15 We interpret our measurements as a "climatological Lagrangian experiment," such that the mean fluxes are more reliable than any single flux determination. This is appropriate for our estimation strategy which intentionally avoids the use of detailed models of atmospheric transport, but instead relies upon the position of our measurement sites relative to the strong easterly trade winds. Furthermore, our use of SF$_6$ as tracer of Northern and Southern Hemisphere air allows us to improve the accuracy of our boundary condition calculation relative to using data from just a single site, like ASC. In the future, our observations can be used with detailed models of atmospheric transport to make more highly resolved estimates of surface CH$_4$ fluxes in eastern Amazônia.

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