Comparison of measured reactive trace gas profiles with a multi-layer canopy chemical exchange model in an Amazonian rainforest

WHAT ARE WE DOING IN THE DEEP RAINFOREST?



In 2010 and 2011 two 80 m high towers for atmospheric research were erected at the ATTO site (Amazon Tall Tower Observatory) (02°08'38 8"S 58°59'59 5"W) in the remote Amazonian rainforest In 2015 the construction of the 325 m high ATTO tower was completed The nearly pristine environment

allows biosphere-atmosphere studies within an ecosystem far away from large anthropogenic emission sources. Combined analyses

Fig. 1: Location of the ATTO site. The main map shows the access to the site via the road and riverboat connections (background man from Google Earth).

of measurements from all three Fig. 2: The 325 meter high towers promises new insights in ATTO tower several emission, deposition, chemical and transport processes

METEOROLOGY IN THE AMAZON

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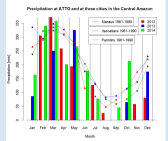
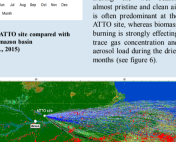


Fig. 8: Precipitation at the ATTO site compared with three different sites in the Amazon basin (adapted from Andreae et al., 2015)



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the South American trade wind zone, the predominant winds come from NE/ENE directions During the wet season almost pristine and clean air is often predominant at the ATTO site, whereas biomass burning is strongly effecting trace gas concentration and aerosol load during the drier

damage plant surfaces.

The climate in the Central

Amazon is characterized by

monthly average tempera-

tures of 26-28°C and strong

variations in precipitation

between the wet (Jul-Oct)

and the dryer season (Jan-

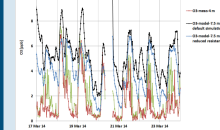
Apr) (see fig.5). As the

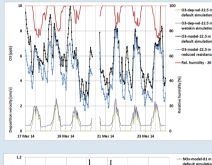
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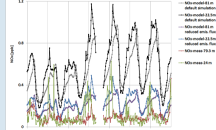
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SUMMARY AND OUTLOOK

Applying some sensitivity analyses within the MLC-CHEM, the effects of different deposition and emission scenarios on O₃ and NO₈ mixing ratios have been shown. The correlation between simulations and measuremnts could be improved with these analyses

The next planned steps to perform, regarding the application of MLC-CHEM on the measured profile data are:

- To compare different NO₂-soil emission flux-scenarios with NO₂-fluxes measured by different techniques.
- > To look, how nighttime O3-NO reactions and the "morning plume" are represented using different modifications in the simulations
- > To verify, how the O₂ deposition relates to the CO₂ exchange where these two processes share the stomatal uptake mechanism

MEASUREMENTS AND CHALLENGES







Fig 4: The reactive trace gas profile system inside the

Fig 5: The inlet tube at 54 m hight with 5um filter and

profiles of H₂O, CO₂ and O₃ were measured at 8 different heights between 2015 NO_x measurements are performed continuously.

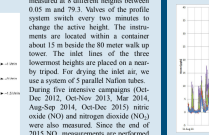


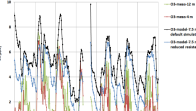
Fig. 6: Profile system with inlets and analysers [courtesy of A. Moravekl

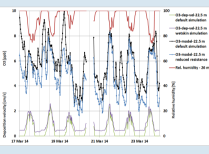
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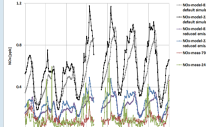
Fig. 8: Back trajectories for the ATTO site between 12 Aug 2014 and 28 Aug 2014 overlying a landscape map with fire sources. Dark blue lines represent conditions before 17 Aug, bright blue lines after 23 Aug, green line represent conditions between 17-23 Aug, black lines represent the peak time: 19 Aug 12:00 - 20 Aug 12:00 (Courtesy of C. Pöhlker, T. Könemann, D. Walter, Since April 2012 vertical mixing ratio J. Saturno, F. Ditas

effect of biomass burning events in August 2014 on the ozone concentrations The polluted air on 19/08 exhibited O3 mixing ratios which were about three times higher than those. measured during the days hefore In case of further and closer deforestation to the ATTO site we expect increasing O3 concentrations, which can effect and

Fig. 9: Ozone mixing ratios at the ATTO site between 11/08 and 29/18/2014 measured by a TEI 49i analyzer







NO_X analyze

Figure 7 shows the strong 19 Mar 14 21 Mar 14 23 Mar 16

fill gaps and to understand the controls on the exchange, the application of atmosphere-biosphere exchange models can help to complement the observations and support analysis of the observations (magnitude, temporal variability).

Through the combined use of detailed multi-compound long-term observations and the canopy exchange model "MLC-CHEM" we could improve our understanding of turbulent transport processes as well as emissions, deposition, and chemical processes occurring in the Amazon rainforest

Fig. 11: Comparison of the observed and simulated RWS-O

run for O, mixing ratios within the soil layer (measured, red

Fig. 12: Simulated O₃ mixing ratios for the default simulation

(green line) and the simulation with reduced wet skin uptake resistance for O₂ (black line) compared with changed denosition

velocities (green and purple lines) and relative humidity (red line)

Fig. 13: MLC-CHEM simulation using reduced NO emission fluxes

(purple and blue lines) compared with measurements (red and

green lines) for NO, mixing ratios within the crown layer and

Measurements of trace gases are generally limited spatially

(both horizontally and vertically). In order to generalize, to

and green, model, blue) for the wet season 2014

surface layer for the wet season

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The MLC-CHEM-model version we have used contains three different lavers

Soil layer (SOL) : 0 m - 15 m: Center at 7.5 m Crown layer (CRL): 15 m - 30 m. Center at 22.5 m Surface laver (SUL): 30 m - 110 m: Center at 70 m

which should correspond fairly well to the following measured heights: 4 m / 12 m. 24 m. 79.3 m.

The surface layer was constrained with measured O3 data from 79.3 m height. Simulated O3 values inside the canopy were some ppb higher than measured data. Sensitivity analyses with enhanced deposition velocities on wet surfaces led to better correlations

Yienger and Levy (1995) have made an estimation which leads to an NO emission flux of 2.6 ng m⁻²s⁻¹ for wet tropical soils. That parameter was proved for some tropical sites and matched fairly well with measured emissions. Within the MLC-CHEM-model that factor was used to constrain the NO-emissions. Simulated values were about twice as high as measured ones. Using a reduced NO emission flux by the factor of 2 within the model, NO

mixing ratios could be represented fairly well.