Emission ratio and isotopic signatures of molecular hydrogen emissions from tropical biomass burning

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Abstract. In this study, we identify a biomass-burning signal in molecular hydrogen (H2) over the Amazonian tropical rainforest. To quantify this signal, we measure the mixing ratios of H2 and several other species as well as the H2 isotopic composition in air samples that were collected in the BARCA (Balanço Atmosférico Regional de Carbono na Amazônia) aircraft campaign during the dry season. We derive a relative H2 emission ratio with respect to carbon monoxide (CO) of 0.31 ± 0.04 ppb ppb⁻¹ and an isotopic source signature of −280 ± 41 ‰ in the air masses influenced by tropical biomass burning. In order to retrieve a clear source signal that is not influenced by the soil uptake of H2, we exclude samples from the atmospheric boundary layer. This procedure is supported by data from a global chemistry transport model. The ΔH2/ΔCO emission ratio is significantly lower than some earlier estimates for the tropical rainforest. In addition, our results confirm the lower values of the previously conflicting estimates of the H2 isotopic source signature from biomass burning. These values for the emission ratio and isotopic source signatures of H2 from tropical biomass burning can be used in future bottom-up and top-down approaches aiming to constrain the strength of the biomass-burning source for H2. Hitherto, these two quantities relied only on combustion experiments or on statistical relations, since no direct signal had been obtained from in-situ observations.

1 Introduction

Molecular hydrogen (H2) is the second most prominent reduced trace gas in the atmosphere with an average abundance of about 530 parts per billion (ppb) (Ehhalt, 1999; Novelli et al., 1999). Through its removal by reaction with the hydroxyl radical (OH), H2 is indirectly linked to many other compounds, for example the important greenhouse gases methane (CH4), tropospheric ozone, or carbon monoxide (CO). This is because the abundance of OH determines the removal of many pollutants and greenhouse gases, such as CH4, for which the Amazon region is a large source (Miller et al., 2007; Beck et al., 2012). Consequently, the oxidation of H2 implies an indirect greenhouse effect (Schultz et al., 2003; Derwent et al., 2006). This active role of H2 in tropospheric chemical cycles elicits a detailed assessment of the impacts of H2 emissions and an investigation and quantification of the individual terms in the global H2 budget.

Table 1 summarises the most important global tropospheric sources and sinks of H2 and their relative importance according to two recent studies (Ehhalt and Rohrer, 2009; Pieterse et al., 2013). The oxidation of methane and volatile organic compounds is the major source, followed by biomass burning, fossil-fuel combustion and nitrogen fixation over land and ocean with respective decreasing strength of the source. The major sink of atmospheric H2 is the uptake by the soil followed by the oxidation by OH.

Crutzen et al. (1979) point out the importance of natural and anthropogenic vegetation fires as a source of many
atmospheric trace gases, especially from tropical regions. Table 1 shows that about 20 % of the total H\(_2\) source is ascribed to biomass burning. The major sources of H\(_2\) from biomass burning are vegetation fires in the tropical rainforests, biofuel burning, and fires in the savanna forests and grasslands (Andreae and Merlet, 2001). Minor contributors are the fires in extratropical forests and burning of agricultural waste (Andreae and Merlet, 2001). Table 2 lists three estimates of the H\(_2\) tropical biomass-burning source strength. Accordingly, about 3.9 % to 8.4 % of the total H\(_2\) emissions (Table 1) can be attributed to this source. Thus, human induced and natural biomass burning from tropical rainforests is a considerable source of H\(_2\) on the global scale, but an even more important and strong local source during the wildfire season. The model study by Pieterse et al. (2011), for example, shows a large plume of H\(_2\) over the Amazonian rainforest and other southern tropical areas in the austral spring and summer (cf. their Fig. 6). This illustrates that the local H\(_2\) emissions from biomass burning in the tropical rainforest can be a major seasonal contributor to the total source and might locally affect atmospheric levels of trace gas concentrations.

Global H\(_2\) emissions are often derived in a bottom-up approach using emission factors and the total biomass that is burned (Andreae and Merlet, 2001; Ehnhalt and Rohrer, 2009) or by linking H\(_2\) emissions to the better constrained CO emissions via the H\(_2\)/CO emission ratio (Price et al., 2007; Yver et al., 2011). Crutzen et al. (1979) and Novelli et al. (1999) use the latter approach with the H\(_2\)/CO ratio to carbon dioxide (CO\(_2\)) emission ratio. Emission ratios vary according to the type of biomass burned and the efficiency of the combustion process (Andreae and Merlet, 2001; Röckmann et al., 2010a). Consequently, estimates for the tropical rainforest differ from those for a boreal forest, for example. Bottom-up approaches to approximate the H\(_2\) source strength bear uncertainties arising from the emission ratio or factor, on the one hand, and the estimates of the amount of biomass burned or the CO and CO\(_2\) fluxes, on the other hand.

Top-down approaches were applied by Rhee et al. (2006) and Xiao et al. (2007). Rhee et al. (2006) used the stable isotopic composition together with a simple mass balance model. Xiao et al. (2007), on the other hand, did not use the isotopic composition but inverse modelling techniques. These methods are completely independent of emission ratios and may contribute to reducing the uncertainties. However, knowledge about the isotopic signatures of different sources needs to be improved to constrain the global budgets obtained from such an approach (Pieterse et al., 2011). So far, the only estimates of the biomass-burning source signature are obtained in laboratory experiments (Gerst and Quay, 2001; Röckmann et al., 2010a) or indirectly from in-situ observations using a statistical approach (Rhee et al., 2006). Röckmann et al. (2010a) discuss the differences between the contradicting values reported in these studies.

So far, no signal from biomass burning in the tropical rainforest has been obtained from in-situ measurements. In this study, we retrieve this signal in air samples collected during the BARCA (Balanço Atmosférico Regional de Carbono na Amazônia) flight campaign over the Amazonian rainforest. We derive the H\(_2\)/CO emission ratio directly from measurements of the mixing ratio and compare them to former estimates. Further, we characterise the isotopic signature of the biomass-burning source. These two estimates support both bottom-up and top-down approaches to quantify the contribution of tropical biomass burning to the global and local H\(_2\) budget.

After a short description of the measurement campaign and the air sample collection, we describe the measurement techniques to derive the chemical compounds and the isotopic composition in Sect. 2. The H\(_2\)/CO emission ratio is derived and discussed in Sect. 3, and the isotopic composition of H\(_2\) from biomass burning is studied in Sect. 4. In Sect. 5, we compare our results to output from a global chemistry transport model, TM5 (Krol et al., 2005; Pieterse et al., 2013).

### Table 1. Global tropospheric sources and sinks of H\(_2\) in Tg yr\(^{-1}\) and approximate percentage of the total source or sink in brackets.

<table>
<thead>
<tr>
<th>Source/Mechanism</th>
<th>Ehnhalt and Rohrer (2009)</th>
<th>Pieterse et al. (2013)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Photochem. production (by CH(_4) &amp; VOC)</td>
<td>41 ± 11 (54 %)</td>
<td>37 ± 4 (48 %)</td>
</tr>
<tr>
<td>Biomass burning</td>
<td>15 ± 6 (20 %)</td>
<td>15 ± 4 (19 %)</td>
</tr>
<tr>
<td>Fossil fuel combustion</td>
<td>11 ± 4 (14 %)</td>
<td>17 ± 4 (22 %)</td>
</tr>
<tr>
<td>N(_2)O fixation, ocean</td>
<td>6 ± 3 (8 %)</td>
<td>5 ± 2 (6 %)</td>
</tr>
<tr>
<td>N(_2)O fixation, land</td>
<td>3 ± 2 (4 %)</td>
<td>3 ± 2 (4 %)</td>
</tr>
<tr>
<td>Total</td>
<td>76 ± 14</td>
<td>77 ± 16</td>
</tr>
<tr>
<td>Soil deposition</td>
<td>60 ± 120 (76 %)</td>
<td>53 ± 4 (70 %)</td>
</tr>
<tr>
<td>Oxidation by OH</td>
<td>19 ± 5 (24 %)</td>
<td>23 ± 2 (30 %)</td>
</tr>
<tr>
<td>Total</td>
<td>79 ± 30</td>
<td>76 ± 6</td>
</tr>
</tbody>
</table>

* Their scenario S3c.

### 2 Data and methodology

The samples that were collected over the Amazonian tropical rainforest have been analysed in two different laboratories for mixing ratios of several trace gases (Max Planck Institute for Biogeochemistry, Jena; hereafter MPI-BGC) and the isotopic composition of H\(_2\) (Institute for Marine and Atmospheric Research Utrecht, Utrecht; hereafter IMAU). In this section, we provide a brief description on how the datasets were obtained. The CO and H\(_2\) mixing ratios as well as the H\(_2\) isotope data used for our analysis in this study are available in the Supplement.
Table 2. Estimates of the tropical biomass-burning $H_2$ source strength, emission ratio, and isotopic signature.

<table>
<thead>
<tr>
<th>Study</th>
<th>$H_2$ Flux [Tg yr$^{-1}$ (%)]</th>
<th>$H_2$/CO by mass [g mol$^{-1}$/g mol$^{-1}$]</th>
<th>$\Delta H_2/\Delta CO$ [ppb/ppb$^{-1}$]</th>
<th>$\delta$D$^a_b$ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laursen et al. (1992)</td>
<td>3.0 (3.9 $^a$)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Andreae and Merlet (2001)</td>
<td>5.1 (6.7 $^a$)</td>
<td>0.037 ± 0.005$^c$</td>
<td>0.52 ± 0.07$^c$</td>
<td>–</td>
</tr>
<tr>
<td>Gerst and Quay (2001)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>(−290 ± 60$^e$)</td>
</tr>
<tr>
<td>Rhee et al. (2006)</td>
<td>–</td>
<td>0.036 ± 0.007$^e$</td>
<td>0.5 ± 0.1</td>
<td>(−290 ± 20)</td>
</tr>
<tr>
<td>Röckmann et al. (2010a)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>(−262 ± 27$^f$)</td>
</tr>
<tr>
<td>Yver et al. (2011)</td>
<td>6.4 (8.4 $^a$)$^d$</td>
<td>(0.02$^f$g$^e$)</td>
<td>(0.28$^c$ f$^e$)</td>
<td>–</td>
</tr>
<tr>
<td>Vollmer et al. (2012)</td>
<td>–</td>
<td>(0.018 ± 0.004$^e$)</td>
<td>(0.25 ± 0.05$^e$)</td>
<td>–</td>
</tr>
<tr>
<td><strong>This study</strong></td>
<td>–</td>
<td>0.022 ± 0.002</td>
<td>0.31 ± 0.04</td>
<td>−280 ± 41</td>
</tr>
</tbody>
</table>

$^a$ With respect to total source in Table 1,  
$^b$ 1975–1980,  
$^c$ late 1990s,  
$^d$ mid-2006–mid-2009,  
$^e$ calculated from the referenced values,  
$^f$ from Hauglustaine and Ehhalt (2002),  
$^g$ global estimate.

2.1 Sample acquisition

Air samples were collected during the first of the two aircraft measurement campaigns of the BARCA project as part of the Large Scale Biosphere–Atmosphere (LBA) program. In general, the BARCA-LBA project aimed at providing in-situ measurements of mainly carbon fluxes from the biosphere of the tropical rainforest to the atmosphere on a scale between global and local.

The BARCA-A campaign was performed towards the end of the biomass-burning season, in November and December 2008. The second part of the program, BARCA-B, was carried out at the end of the wet season, in May 2009. As we are interested especially in the biomass-burning signal, we only analysed samples from the BARCA-A flights for this study. Figure 1 shows an overview of the biomass burning conditions during the BARCA-A period. The fire radiative power from wildfires over South America (Fig. 1a) can be considered as proportional to the biomass burned. The data is a satellite product from the MODIS instrument and obtained from the Global Fire Assimilation System (GFASv1.0) by Kaiser et al. (2012). It shows extensive fires in the eastern part of the Amazon Basin close to the coast. Further, the European Centre of Medium Range Weather Forecast (ECMWF) ERA-Interim reanalysis wind vectors on the 800 hPa level (Dee et al., 2011). (b) Enlargement of the box in (a). Mean GFASv1.0 scaled $H_2$ emissions from biomass burning (Kaiser et al., 2012) and BARCA-A sample sites (circles) used in this study (black: above 1100 m a.s.l.; white: below 1100 m a.s.l.; and green (highlighted by the arrow): highly contaminated sample in Figs. 2 and 3).

Several flight tracks span more than 2000 km of the Amazon Basin that were influenced by air masses from the regions of extensive biomass burning. Samples were taken in vertical profiles along the tracks from near-surface up to more than 4000 m a.s.l. in the atmospheric boundary layer and the lower free troposphere above. The samples that we analyse in this study are indicated by circles in Fig. 1b. Black circles indicate high-altitude samples (above 1100 m a.s.l.) and white circles low-altitude samples (below 1100 m a.s.l.). They were all either collected directly in the wildfire region visible in Fig. 1 or downwind of this region. The area where the samples were collected and the region to the east are mostly covered by dense forest (Saatchi et al., 2007;
Simon et al., 2009). Thus, we expect a biomass-burning influence mainly from C3 plants. In the south-eastern corner of the enlarged box in Fig. 1b, however, the dominant vegetation cover is savanna grassland (Saatchi et al., 2007; Simon et al., 2009). This could have a small influence on some of the samples. Andreae et al. (2012) and Beck et al. (2012) give more details on the sampling and a description of the flight tracks. Some of the samples were collected directly inside biomass-burning plumes along the tracks which gives us the possibility to distinguish samples with high contamination.

Next to the air sampling, numerous chemical species were analysed continuously on the aircraft (Chen et al., 2010). In this project, we have only used data from the air samples analysed in the laboratories. A more in depth analysis of CO and aerosols as well as CH4 from both flight campaigns is provided by Andreae et al. (2012) and Beck et al. (2012), respectively.

2.2 Mixing ratio measurements

The 174 flask samples taken during the BARCA-A campaign have been measured for the mixing ratio of several gases (among others CO, CO2, CH4, and H2) at the MPI-BGC Gaslab. Measurements in the laboratory were conducted 120 to 300 days after the sampling (Andreae et al., 2012). Between the time of sampling and the measurements the samples were kept in 11 borosilicate 3.3 glass flasks which were sealed with PCTFE valves (Batenburg et al., 2011; Andreae et al., 2012). In order to avoid any photochemical production of H2, the flasks were enclosed by black flexible tubes and stored in closed metal boxes. The stabilities of several species were tested at the MPI-BGC Gaslab. Most of the species showed no significant drift except for CO which increased by 2.6 to 6.5 ppb during the storage period (Andreae et al., 2012). Flask analysis results were combined with the continuous data (tracers, meteorological parameters, position) using weighting functions that mimic the temporal averaging during the flask sampling process (Chen et al., 2012). For a number of tracers measured continuously as well as in the flask samples comparisons resulted in good agreement (cf. Andreae et al., 2012; Beck et al., 2012).

In this study, we use H2 and CO mixing ratios of 72 samples for which data was available. These were determined using a Trace Analytical Reduction Gas Analyser (RGA), which uses gas chromatographic separation with subsequent HgO reduction and photometric UV detection of Hg. Each sample analysis was bracketed by a working tank analysis. The response of the RGA was determined on a daily basis using multiple standards bracketing the range of sample mixing ratios. CO values are reported on the WMO 2004 CO mole fraction scale (Novelli et al., 2003). Note that CO values from flask sample analysis were not corrected for the offset of 6.3 ± 2.1 ppb compared to the continuous measurements (Andreae et al., 2012). This offset is likely due to a temporal increase between sample collection and analysis. However, it has no significant impact on the results for emission ratios. In contrast to the CO values, H2 values are reported on the CSIRO94 calibration scale, which is also used by the Advanced Global Atmospheric Gases Experiment (AGAGE) network (Simmonds et al., 2000). Note that this is different to the new WMO H2 scale maintained at MPI-BGC (Jordan and Steinberg, 2011). The range of the H2 and CO mixing ratios in the samples that we analyse in this study is within the calibrated range of the RGA and the non-linearity was explicitly corrected for by a suite of reference gases.

2.3 Hydrogen stable isotopic composition

The fractionation of the two isotopes is coupled to the process in which H2 is involved. Reciprocally, it is often possible to identify a process from an air sample once the isotopic characteristics are known. The method of detecting sources and sinks by retrieving their isotopic hydrogen signature was successfully applied in many studies before (e.g. Gerst and Quay, 2001; Röckmann et al., 2010a; Batenburg et al., 2011, 2012).

We analysed 39 samples from the BARCA-A campaign for the H2 isotopic composition in the isotope laboratory at the IMAU. The measurement of the H2 isotopic composition with a gas-chromatography isotope-ratio mass spectrometer system (GC-IRMS) followed the procedure developed by Rhee et al. (2004) and updated by Röckmann et al. (2010b) and Batenburg et al. (2011, 2012): at first H2 was purified in three steps. A sample volume of about 750 ml was filled with the air sample to a pressure of about 600 hPa. Subsequently, the sample was exposed to a “cold head” (of about 40 K) where all gases except for the most volatile ones condensed. The remaining gases were flushed by a helium (He) carrier gas into a 5 Å molecular-sieve pre-concentration trap that was immersed in liquid nitrogen (cooled down to the N2 triple point by pumping on the gas phase) to collect the H2. The gases were then flushed to a cryo-focus trap that was immersed in liquid nitrogen (77 K) and afterwards to a GC column at 323 K. This procedure separated H2 from any other gas. Between the sample measurements the cold head, the pre-concentration trap, the cryo-focus trap, and the GC column were heated in order to avoid any contamination from the previous sample.

The isotopic composition was determined online from the purified H2 by an IRMS. Here, it is reported in the common δ-notation, defined as

$$\delta D = \left( \frac{R_{SA}}{R_{ST}} - 1 \right) \times 1000 \text{‰},$$

where R denotes the ratio of the “heavy” isotope, deuterium (D), to the “light” isotope (H) and the subscripts SA and ST abbreviate the sample and the standard, respectively. The international standard D/H ratio is the one of Vienna Standard Mean Ocean Water (VSMOW) with a value of
155.76 ± 0.08 ppm (de Wit et al., 1980; Gonfiantini et al., 1993). The H$_2$ sample peak was preceded by seven injections of our H$_2$ laboratory working gas and followed by two more injections of the reference gas into the IRMS from a custom made open split system (Röckmann et al., 2003).

In total, the samples were measured on ten different days. On each of these days, we determined the H$_2$ isotopic composition of two different laboratory reference air cylinders with known values of $\Delta$D (198.2 ± 0.5‰ and 207.0 ± 0.3‰). Additionally, a blank measurement was performed every day. These measurements were used to monitor the performance of the system and calibrate the results. The mean standard deviation obtained from the measurements of the two laboratory reference air cylinders was 6.7‰, which was slightly higher than the 4.5‰ reported by Batenburg et al. (2011). This larger scatter might have been caused by small fluctuations in the system performance or technical replacements in the set up. The area of the blank peak in the chromatogram was on average 1.7% of the sample peak area, so less than the 4% in the experiments by Batenburg et al. (2011), indicating an even smaller systematic error.

The $\Delta$D values retrieved for the samples were calibrated using the two measured $\Delta$D values from the laboratory reference-air flasks. We smoothed these reference values with a five-day moving average (cf. Batenburg et al., 2011). However, we did not apply the empirical bias correction (9.5‰) and scale contraction (1.0613) factors determined by Batenburg et al. (2011). An analysis of another reference gas that was measured as well on a daily basis revealed that its measured $\Delta$D value without any correction applied was very close (the difference is 4‰) to the certified value by the manufacturer (Messer Griesheim) and within the provided range of uncertainty. However, if we had applied the scale contraction factor, there would have been a large deviation from the certified value (outside the uncertainty range). The empirical bias correction factor had a smaller effect, but the deviation from the certified value was still larger than without this correction. Not using this bias correction factor was further motivated by the facts that no cause of the sudden increase in the values measured by Batenburg et al. (2011) had been determined, and that further technical changes were made on the system set up since this value had been determined. We tested the results presented in this study by applying all corrections as well and found that they are still within the range of uncertainty that we provide in Sects. 3 and 4.

As a consequence of the above discussion, the sample measurements were calibrated as follows:

$$\delta D_{c_{1,2}} = \delta D_s + (\delta D_R_{1,2} - \delta D_{1,2}),$$  \hspace{1cm} (2)

where $\delta D_s$ and $\delta D_{1,2}$ are the measured values from the sample and the two laboratory reference gas flasks, and $\delta D_R_{1,2}$ are the known reference values that were smoothed with the five-day moving average. The final calibrated $\Delta$D value is, then, the arithmetic mean of the two corrected values obtained from Eq. (2) ($\delta D_{c_{1,2}}$).

### 2.4 Model simulation

In order to support our analysis applied in Sects. 3 and 4, we performed a simulation with the global chemistry transport model TM5 (Tracer Model, version 5) (Krol et al., 2005; Huijnen et al., 2010). We used the two-way nested set up developed by Krol et al. (2005) for a high-resolution zoom over the Amazon Basin. Resolution increases from 6° × 4° (longitude × latitude), through 3° × 2° to 1° × 1°. In the vertical the model consists of 25 hybrid levels.

This full chemistry model was forced with scaled surface emissions of the year 2003 from Schultz and Stein (2006) as described by Pieterse et al. (2011). These emissions were fixed throughout the simulation and did not vary from one year to the other. Note that in this model configuration the surface emission data from Schultz and Stein (2006) are different from those shown in Fig. 1 for illustration. The meteorological forcing was the ECMWF ERA-Interim reanalysis data (Dee et al., 2011). For a detailed description of parameterisations applied, the H$_2$ chemistry in the model, and a model evaluation, we refer the reader to Pieterse et al. (2013).

In this study, we investigate the mean model output for the same season as the BARCA-A campaign (November and December) in order to resemble the meteorological conditions. We only use data from the fifth year of model integration. Even though surface fluxes are not from the year 2008, the model provides sufficient information on the relative strength of the source to analyse the general biomass-burning signal in this region. We will use vertical profiles of H$_2$ and $\Delta$D from the model simulation to compare the effect of vertical transport by turbulent mixing and convection to the observations.

### 3 Hydrogen to carbon monoxide ratio from biomass burning

Distinct emission ratios between several atmospheric trace gases are related to different combustion processes. These relations can be used to identify samples that are strongly contaminated by a certain process or to estimate the strength of the source of one species if the other one is known.

Comparing the H$_2$ and CO mixing ratios of the 72 BARCA-A samples available, we obtain a significant Pearson correlation coefficient of about 0.64 (at a 99% confidence level using a t test). Hence, an analysis of the apparent pyrogenic signal seems promising. No correlation between H$_2$ and CO$_2$ or between H$_2$ and CH$_4$ is present. Therefore, we derive the emission ratio between H$_2$ and CO. Values are commonly reported as elevations over a typical background mixing ratio, thus as $\Delta$H$_2$ = H$_2$ – H$_2$$_{bg}$ and $\Delta$CO = CO – CO$_{bg}$. In this study, however, we do not use the background mixing ratio to estimate the $\Delta$H$_2$/$\Delta$CO ratio but the slope of the regression line through the measured mixing ratios of H$_2$ and CO. Therefore, we need to assume that any deviations from the background mixing ratio of the
considered species almost solely result from biomass burning and all other source and sink terms are minor. We will later discuss to what extent this assumption is valid for our set of samples and how we can reduce potential interferences. Moreover, the assumption of a single source in the samples translates to a constant background value for both species. During the BARCA-A campaign the background mixing ratio of CO is about 79 ± 7 ppb (cf. Andreae et al., 2012). There is no estimate for the background mixing ratio of H2 for the BARCA-A period. Here, we will thus use the CO background mixing ratio and our value for the ΔH2/ΔCO ratio to estimate the H2 background concentration as well.

Figure 2 shows the relation between the H2 and CO mixing ratios determined from the BARCA-A samples. In order to retrieve a ΔH2/ΔCO ratio from all available samples (coloured and grey), the Williamson–York bivariate total least squares linear regression algorithm suggested by Cantrell (2008) has been applied. The iterative algorithm takes the statistical errors of both the H2 and CO mixing ratios into account. The resulting ΔH2/ΔCO ratio is 0.29 ± 0.05. Despite the significant correlation, there is a large scatter between these samples, especially for lower H2 mixing ratios. This probably results from other sinks or sources that influence the H2 or CO mixing ratios.

Next to the biomass-burning source, the uptake of H2 by the soil has the largest influence on the mixing ratio in the tropics (cf. Fig. 8 in Pieterse et al., 2011). Presumably, the effect of this sink is particularly strong in the lower atmospheric boundary layer by persistent mixing of air masses to the surface. Consequently, here, we expect the strongest signals of these sinks. In contrast, the source signal from the vegetation fires can reach high up in the atmosphere due to convection. This high uplift of air masses influenced by the biomass-burning source is caused either by the heat produced by these extensive fires themselves, by natural convection occurring at this time of the year, or by a combination of these two effects. During November and December – at the end of the dry season – the atmospheric stratification over the Amazon Basin is mostly unstable and frequent convection is very likely to occur (Fu et al., 1999; Fu and Li, 2004). Also the circulation pattern shown in Fig. 1a, that is dominantly easterly, suggests a transition period to the wet season and favourable conditions for convection.

In order to be consistent throughout this study, including the derivation of the isotopic signature, we will proceed to only analyse those 39 samples for which the isotopic composition was measured as well. Figure 2 shows these rather arbitrarily selected samples (colour) spread along the regression line of all collected samples. The selected samples cover the full horizontal and vertical space of the sampling region (cf. Fig. 1) and we selected them to be representative in terms of their range in the H2–CO space.

At this point, the fact that the sampling during the flight campaign was carried out in vertical profiles is very useful. A separation of the samples into two groups, higher and lower than 1100 m a.s.l. (red and blue in Fig. 2, respectively), visualises the effect of altitude in the measurements. We have chosen this altitude as it provides the clearest separation between the samples and the correlation coefficient for each group of samples is optimal. The value of about 1000 m above ground level is consistent with the approximate height of a daytime convective mixed boundary layer over land surfaces (cf. e.g. Stull, 1988). Samples closer to the ground have much lower H2 mixing ratios and carry interfering signals of the biomass-burning source and the soil uptake.

The highest 20 samples (red in Fig. 2) are very likely almost solely influenced by the biomass-burning source, reflected by a very high correlation coefficient of about 0.95 (significant at a 99 % confidence level). However, one should notice that the sample with very high mixing ratios, visible in Fig. 2, strongly increases the correlation. Removing this sample from the selection yields still a significant correlation coefficient of 0.85. Beck et al. (2012) also report a very high contamination by the biomass-burning source in this specific sample. Considering its location highlighted in Fig. 1, we are confident that this sample was taken in a large biomass-burning plume. Thus, we incorporate this sample in the further analysis.

The slope of the regression line through the samples above 1100 m a.s.l. (red in Fig. 2) with a value of 0.31 ± 0.04 is our best estimate of the ΔH2/ΔCO ratio of biomass burning over tropical rainforest. Excluding the discussed high mixing ratio sample results in a ratio that is still within the given range of uncertainty. The H2 background mixing ratio using the regression line through these high altitude samples at the intercept with the estimate of 79 ± 7 ppb (cf. Andreae et al., 2012) for the CO background mixing ratio is about 521 ± 7 ppb.

The correlation between H2 and CO is most significant for the higher altitude samples. However, also the 14 samples below 1100 m a.s.l. – for which an interference of the soil-sink signal is likely – carry a biomass-burning signature (blue in Fig. 2). The correlation coefficient for these samples has a significant value of 0.86 (at a 99 % confidence level). The resulting regression line has a slope of 0.21 ± 0.04. Closer to the ground H2 is taken up by the soil lowering the H2 values in these samples and, thus, altering their ΔH2/ΔCO ratio.

A group of five samples with very low H2 mixing ratios (below 510 ppb) in the lower boundary layer (below 300 m a.s.l.) is highlighted in green in Fig. 2. The low values are likely the result of a weak biomass-burning source signature and a very strong influence of the soil sink. This could be caused by little advection of contaminated air into this region and/or a strong mixing in the lower boundary layer that enhances the influence of the soil uptake. Our interpretation is supported by the fact that these samples are among those collected very close to ground in the atmospheric surface layer (about 60 to 190 m above ground level). Consequently, we excluded the five samples marked (green) in Fig. 2 from the
analysis (they are only considered for the black regression line in Fig. 2).

Considering only the selected samples that show a clear biomass-burning signal at higher altitudes, the relations between H$_2$ and CH$_4$ and between H$_2$ and CO$_2$ are still not significant. This makes it impossible to derive reliable estimates of the respective emission ratios. Probably, there are other sources and sinks of CH$_4$ and CO$_2$ that are rather dominant locally and mask the biomass-burning signal in some samples.

Beck et al. (2012) report a significant relation between CO and CH$_4$ for all BARCA-A samples analysed in the laboratory of the MPI-BGC. They found that several samples have a $\Delta$CO to $\Delta$CH$_4$ ratio that is close to that of biomass burning. Among these samples is the mentioned single outlier in our Fig. 2 that has been shown to be strongly contaminated by a biomass-burning plume (Beck et al., 2012). Its location very close to the regression line of the samples in the upper convective atmospheric layer in Fig. 2 confirms our approach to separate the samples.

Vollmer et al. (2012) summarise different $\Delta$H$_2$/$\Delta$CO and $\Delta$H$_2$/$\Delta$CH$_4$ ratios from biomass burning that can be found in literature. They derive an average $\Delta$H$_2$/$\Delta$CO ratio 0.25±0.05 and an average $\Delta$H$_2$/$\Delta$CH$_4$ ratio of 3.6. Their $\Delta$H$_2$/$\Delta$CO ratio is close to the value derived by Hauglustaine and Ehhalt (2002) that was adopted by Yver et al. (2011) for an estimation of the tropical H$_2$ source strength (cf. Table 2).

Most estimates of the emission ratios, including those by Vollmer et al. (2012) and Hauglustaine and Ehhalt (2002), represent global values. Biomass-burning characteristics in the tropical rainforest might, however, differ considerably from those in other regions. Andreae and Merlet (2001) address these deviations and list estimated emission factors of each species (compared to the burned dry matter) from tropical rainforests. We have converted these estimates to the relative $\Delta$H$_2$/$\Delta$CO emission ratio of 0.52±0.07 in Table 2. This result is much higher than those by Vollmer et al. (2012) and Hauglustaine and Ehhalt (2002) on a global scale, and also the $\Delta$H$_2$/$\Delta$CH$_4$ ratio derived from Andreae and Merlet (2001) yields a higher value (9.6±2.3) than the global scale estimate.

Higher emission ratios from burning of tropical plants mainly result from a comparably higher H$_2$ emission factor. A factor that could possibly alter the $\Delta$H$_2$/$\Delta$CO ratio regionally is the combustion efficiency. However, Röckmann et al. (2010a) did not find any significant relation. In contrast, Popa et al. (2011) find that the combustion efficiency might well influence the H$_2$/CO ratio of fossil fuel combustion considerably. To what degree it might alter the biomass-burning signal is not well understood.

Rhee et al. (2006) report a biomass-burning signal that is detected through an indirect analysis of air samples collected close to the tropical tropopause. Even though no significant direct signal of a biomass-burning source was visible in the H$_2$ mixing ratios of these samples, both the $\Delta$H$_2$/$\Delta$CO ratio and the isotopic signature were retrieved using the signal that could be seen in peaks of the CO mixing ratios. They interpreted the missing higher values of H$_2$ as an overlapping effect from the biomass-burning source and the soil sink. The resulting estimate of the $\Delta$H$_2$/$\Delta$CO ratio from biomass burning over the tropics ranges between 0.4 and 0.6 in their Fig. 4a. Additionally, the findings of Rhee et al. (2006) imply lower $\Delta$H$_2$/$\Delta$CO ratios for savanna and extratropical forests than for the tropical forests.

Our $\Delta$H$_2$/$\Delta$CO ratio with a value of 0.31±0.04 is significantly lower than those derived from the analysis by Rhee et al. (2006) and Andreae and Merlet (2001) for tropical rainforests (cf. Table 2). However, it is more in line with –
though slightly higher than – the upper-end global estimates summarised by Vollmer et al. (2012) and the estimate by Hauglustaine and Ehhalt (2002). Our direct measurements from a comparatively large set of samples thus support the lower $\Delta H_2 / \Delta CO$ estimates.

4 Hydrogen isotopic signature from biomass burning

It is known that $H_2$ from biomass burning is depleted in $\delta D$ relative to the global average isotopic composition of about $130\%e$ (Gerst and Quay, 2001; Röckmann et al., 2010a). However, its precise isotopic signature from biomass burning is still under discussion.

In this study, we derive the $\delta D_{bb}$ for tropical rainforest fires applying the selection procedure presented in Sect. 3 to the isotopic composition. The clearest biomass-burning signature should, consequently, be visible in the higher altitude samples. Figure 3 shows a Keeling plot of the inverse $H_2$ mixing ratios and the $\delta D$ values measured at the IMAU laboratory for all three groups of samples. As in Fig. 2, they are clearly separated:

1. The high-altitude samples bear a biomass-burning signature, thus they show a more depleted $\delta D$ for higher $H_2$ mixing ratios (red).
2. The samples in the boundary layer carry a similar signature but are overall shifted towards lower $H_2$ mixing ratios (blue).
3. As we expected, all samples that are probably strongly influenced by the soil uptake and are collected within the surface layer have high $\delta D$ values (green).

These observations confirm our approach and interpretation in Sect. 3.

Pearson’s correlation coefficient between the inverse $H_2$ mixing ratio and the $\delta D$ values for all samples at higher altitudes (red in Fig. 3) is $-0.89$ at a 99 % confidence level. Omitting the highly depleted sample yields a lower but still significant correlation of $-0.82$. Using the same bivariate total least squares linear regression algorithm as in Sect. 3 (Cantrell, 2008), we are able to estimate the $y$ axis intercept of these samples. The resulting $\delta D_{bb}$ signature from the tropical rainforest is $-280 \pm 41 \%e$.

The highly depleted sample in the lower left corner of Fig. 3 is the same sample as the one in the upper right corner of Fig. 2. This confirms the strong biomass-burning signature in this sample detected in Sect. 3 and by Beck et al. (2012). The signature without this sample would result in a signature of $-289 \%e$ and, thus, has only a minor effect on the result. Moreover, this sample has the largest offset to the regression line. The offset might not only be due to the uncertainty of the measurement but could also include the effect of the combustion efficiency (see below) on the $\delta D$ value.

So far, three independent estimates of $\delta D_{bb}$ exist (cf. Table 2). Gerst and Quay (2001) estimate the global average $\delta D_{bb}$ with $-290 \pm 60 \%e$ from experimental fire measurements by burning pine needles and branches that were, subsequently, extrapolated to a global scale. A similar overall value of $-262 \pm 27 \%e$ is given by Röckmann et al. (2010a) who performed combustion experiments under varying conditions with many wood samples from different locations. Even though this result has not been extrapolated to a global scale, it confirms the estimate by Gerst and Quay (2001). In contrast, the global mean $\delta D_{bb}$ derived by Rhee et al. (2006) is, with a value of $-90 \pm 40 \%e$, much higher. Röckmann et al. (2010a) do not find a relation between $\delta D_{bb}$ and the $\Delta H_2 / \Delta CO$ ratio that was used by Rhee et al. (2006) to derive their estimate of the global mean signature.

The regional $\delta D_{bb}$ depends mainly on the latitude due the variations of $\delta D$ in the precipitation, on the combustion efficiency, and the plant type. The precipitation effect leads to an enriched $\delta D_{bb}$ in the tropics compared to the global
mean value (Röckmann et al., 2010a). The combustion efficiency is extremely variable between different ecosystems, seasons, primary and secondary forests, and natural and anthropogenic fires (Seiler and Crutzen, 1980); it is determined (among other factors) by the availability of oxygen and the temperature of the combustion process (Röckmann et al., 2010a). In terms of plant type, C3 plants are depleted in δD compared to C4 plants (Leaney et al., 1985; Vigano et al., 2009, 2010).

The isotopic biomass-burning signature from the selected BARCA-A samples of $-280 \pm 41 \, \text{‰}$ is in line with the estimates given by Gerst and Quay (2001) and Röckmann et al. (2010a) that were retrieved from combustion experiments (cf. Table 2). The comparatively high δD values of the precipitation at lower latitudes would result in a slightly higher estimate from our samples than the global mean value. Figure 6 of the study by Röckmann et al. (2010a) suggests that this latitudinal effect in the biomass results in about 10 to 20 ‰ higher δD values than the global mean. However, this effect lies within the range of uncertainty that we provide.

We conclude that the isotopic biomass-burning signature provided by Rhee et al. (2006) for tropical rainforest is too high and values are rather within the suggested range by Gerst and Quay (2001) and Röckmann et al. (2010a).

5 Vertical structure of the biomass-burning signature in the model simulation

By analysing the model output from the TM5 simulation for the Amazon Basin as described in Sect. 2.4, we aim to verify our approach to separate the samples by altitude. We expect to obtain a similar signal as in the observations in a region of biomass burning, thus a mixture of the soil uptake and biomass-burning source signals close to the ground and a signal dominated by biomass burning above the atmospheric boundary layer. We do not attempt to provide a model–observations intercomparison since we use surface emissions from the year 2003. Here, we rather aim to support our analysis by studying how turbulent mixing in the...
boundary layer and vertical transport by convection change the vertical profiles of $H_2$ and $\delta D$ and distort the $\delta D_{bb}$ signal.

Figure 4a shows the TM5 zoom region over the Amazon Basin and the mean $H_2$ mixing ratio in the atmospheric surface layer for the months November and December after five years of integration. One should keep in mind that due to the different surface emissions, biomass-burning regions will not coincide with those displayed in Fig. 1. However, meteorological conditions should be representative for the vertical transport and boundary layer mixing during that time of the year. In the centre of the domain there is an area of highly elevated $H_2$ mixing ratios that are clearly the result of biomass burning (indicated by the green box). Furthermore, it should be noticed that this region is at a different location to the sampling region (cf. Fig. 1b) with, probably, a very different vegetation cover (mixed grassland, secondary forest, and rainforest; Saatchi et al., 2007; Simon et al., 2009). However, this will not affect our results since the model does not distinguish C3 and C4 plants in terms of their $\delta D$ signature, which has a fixed input value for biomass burning of $-262 \‰$.

The sixth model level at about 870 hPa (about 1300 m a.s.l.) in this region is just above the altitude of 1100 m a.s.l. that we determined to be optimal for the separation of the samples in Sect. 3. At this altitude the mean $H_2$ mixing ratio is almost everywhere fairly constant with values between 540 and 560 ppb (Fig. 4b). A mixing ratio of 560 ppb is only exceeded in the mentioned biomass-burning region and in another region close to the coast towards the north-east of the domain that is also influenced by biomass burning. While the biomass-burning signal is clearly present, we expect that the direct impact of surface deposition vanishes at an altitude of 1300 m a.s.l. The same effect is visible when considering the $H_2$ isotopic composition shown in Fig. 4c and d for the two model levels with strongly depleted values in the indicated region.

The average vertical profiles of $H_2$ and $\delta D$ for the region within the green box (cf. Fig. 4e) show elevated $H_2$ mixing ratios and lower $\delta D$ values up to a height of about 700 hPa (or about 3000 m a.s.l.), indicating a high reaching effect of the biomass burning. Within the surface layer, below 1000 hPa, the $H_2$ mixing ratio is strongly depleted by the soil uptake of $H_2$ leaving a peak mixing ratio at 1000 hPa. At higher altitudes the biomass-burning signal dominates. We propose that deep and shallow convection over the tropical South American continent carries the biomass-burning signal to an altitude of 1000 m a.s.l. and higher. In the TM5 model, we emit 40% of the biomass burning emissions between 1000 and 2000 m a.s.l. to account for plume rise above the most intensive fires. The 60% emitted at lower altitudes, however, stay within the atmospheric boundary layer. Turbulence repeatedly mixes the air to the surface and, thus, allows the soil uptake to alter the biomass-burning signal.

The influence of the soil uptake on the vertical $H_2$ mixing ratio profile apparent in the simulation likely also influenced the BARCA-A samples. It is very likely the reason for the lower $\Delta H_2 / \Delta CO$ ratio in the lower altitude samples (cf. Sect. 3). A vertical discrepancy in the $\Delta H_2 / \Delta CO$ ratio is also reported by Popa et al. (2011) who find an influence of altitude on the ratio from fossil-fuel-burning source and conclude that atmospheric mixing and the soil uptake close to the ground might be responsible.

We further test our hypothesis by comparing the known model input value for $\delta D_{bb}$ of $-262 \‰$ to a Keeling plot analysis of samples from each model level (Fig. 4f). The regression analysis is performed through all grid points within the green box. In the surface layer (blue line in Fig. 4f), the data points seem to carry a biomass-burning signature. However, the Keeling plot intercept is with $-290 \‰$ somewhat lower than the original source value of $-262 \‰$ that was applied in the model. This discrepancy may be caused by interfering effects from the soil sink near the surface, for example. If we perform the Keeling plot analysis at higher model levels, the intercept increases towards the input value. At the sixth model level, the value ($-265 \‰$) is the closest to the input value ($-262 \‰$) and the correlation coefficient between $\delta D_{bb}$ and the inverse $H_2$ mixing ratio is highest (cf. red line in Fig. 4f). This confirms that at this level the distortion of the biomass-burning signal by the soil uptake has nearly vanished completely and that the fixed input value for the biomass-burning emissions at the surface can be recovered at this altitude.

Comparing these observations to our selection procedure of Sects. 3 and 4 suggests that the actual separation level of 1100 m a.s.l. is reasonable in terms of vertical transport. The regression line from the sixth level of the model region is also added in Fig. 3. The offset between the lines of best fit (purple and red) results from the slightly different input signature in the model of $-262 \‰$ compared to the value of $-280 \pm 41 \‰$ that we derive for the tropical biomass burning from the measurements. Furthermore, the slope of the regression line is almost identical and within the range of uncertainty. Because of these two latter findings and the fact that the model derived biomass-burning signature is very close to the model forcing value, we are confident that the signal retrieved from the selected BARCA-A samples is a pure biomass-burning signature and that our approach is justified.

From both Figs. 3 and 4f, we conclude that closer to the ground the isotopic biomass-burning signature would be biased by the soil uptake towards lower $\delta D_{bb}$ values. This is the result of a shift in the $H_2$ mixing ratio towards lower values (to the right along the x axis in the Keeling plot) when $H_2$ is continuously deposited at the surface. As a consequence, the y axis intercept is shifted towards lower values resulting in an apparent lower isotopic biomass-burning signature.

6 Summary and conclusions

During the BARCA-LBA project air samples were collected over the Amazonian tropical rainforest during the dry season
in regions of extensive biomass burning or downwind of these regions at different altitudes. We have measured the mixing ratios of several species and the isotopic composition of \( \text{H}_2 \) of 39 air samples. From these measurements, we have derived a signal of the biomass-burning source that can be used to estimate \( \text{H}_2 \) emissions from tropical forest fires and better constrain global chemistry models. We have determined the \( \Delta \text{H}_2 / \Delta \text{CO} \) ratio, on the one hand, and the isotopic signature of \( \text{H}_2 \), on the other hand, to characterise the biomass-burning source signal.

For the analysis, the samples have been separated into groups of higher and lower altitude in order to avoid an interfering signal from the soil uptake of \( \text{H}_2 \). Mixing in the lower boundary layer leads to lower \( \text{H}_2 \) mixing ratios in those samples due to soil uptake. In contrast, the strong convection induced by the heat from the fires and the naturally unstable boundary layer results in high-reaching biomass-burning plumes. This enabled us to derive the signal from the samples collected at higher altitudes. We have obtained best results when using an altitude of 1100 m a.s.l. for the separation. A group of samples with very low \( \text{H}_2 \) mixing ratios has been separated from the analysis as they are probably strongly influenced by the soil uptake.

The selection of the samples from higher altitude yielded a high and significant correlation coefficient of 0.95 between the \( \text{CO} \) and \( \text{H}_2 \) mixing ratio and thus provides a clear biomass-burning signal. From these samples, we have derived a \( \Delta \text{H}_2 / \Delta \text{CO} \) emission ratio of 0.31 ± 0.04. The correlation coefficient from the lower altitude samples is lower. This implies an influence of the \( \text{H}_2 \) uptake process by the soil.

We have studied the isotopic composition in the same selection of samples from the top of the boundary layer. A Keeling plot of these samples shows a signal from biomass burning in the Amazonian tropical rainforest with an isotopic source signature \( \delta D_{\text{bb}} = -280 ± 41 \pm 6 \). The \( \delta D \) values and the inverse of the \( \text{H}_2 \) mixing ratio correlate significantly with a coefficient of −0.89. The biomass-burning signal identified by Beck et al. (2012) in the methane isotopic composition of a sample that was collected while the airplane was flying through a dense plume is also visible in our results.

From simulations with a chemistry transport model we have obtained a comparable vertical structure of the biomass-burning signal with an interfering signal from the soil uptake of \( \text{H}_2 \) closer to the ground likely due to vertical mixing to the surface. At an altitude of about 1300 m a.s.l. \( \delta D_{\text{bb}} \) converges to the imposed biomass-burning signature and to the highest correlation between \( \delta D \) and the inverse \( \text{H}_2 \) mixing ratio. The air masses at these high altitudes are only influenced by the biomass-burning source since presumably convection transported the signal to the free troposphere and the soil-sink process has vanished. The model results confirm our approach to separate the samples and also the selected altitude.

Our results suggest that earlier estimates of the \( \Delta \text{H}_2 / \Delta \text{CO} \) ratio from biomass burning over the tropical rainforest (Andreae and Merlet, 2001; Rhee et al., 2006) are probably too high. This should be further examined with additional measurements. Moreover, the \( \delta D_{\text{bb}} \) signature from the tropical rainforest is much lower than the estimate by Rhee et al. (2006) and confirms the range of values suggested by Gerst and Quay (2001) and Röckmann et al. (2010a).

Supplementary material related to this article is available online at http://www.atmos-chem-phys.net/13/9401/2013/acp-13-9401-2013-supplement.zip.

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