High-precision quasi-continuous atmospheric greenhouse gas measurements at Trainou tower (Orléans forest, France)

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Abstract. Results from the Trainou tall tower measurement station installed in 2006 are presented for atmospheric measurements of CO2, CH4, N2O, SF6, CO, H2 mole fractions and radon-222 activity. Air is sampled from four sampling heights (180, 100, 50 and 5 m) of the Trainou 200 m television tower in the Orléans forest in France (47°57’53” N, 2°06’45” E, 131 m a.s.l.). The station is equipped with a custom-built CO2 analyser (CARIBOU), which is based on a commercial non-dispersive, infrared (NDIR) analyser (Licor 6252), and a coupled gas chromatography (GC) system equipped with an electron capture detector (ECD) and a flame ionization detector (FID) (HP6890N, Agilent) and a reduction gas detector (PP1, Peak Performer). Air intakes, pumping and air drying system are shared between the CARIBOU and the GC systems. The ultimately achieved short-term repeatability (1 sigma, over several days) for the GC system is 0.05 ppm for CO2, 1.4 ppb for CH4, 0.25 ppb for N2O, 0.08 ppb for SF6, 0.88 ppb for CO and 3.8 for H2. The repeatability of the CARIBOU CO2 analyser is 0.06 ppm. In addition to the in situ measurements, weekly flask sampling is performed, and flask air samples are analysed at the Laboratoire des Sciences du Climat et de l’Environnement (LSCE) central laboratory for the same species as well for stable isotopes of CO2. The comparison between in situ measurements and the flask sampling showed averaged differences of 0.08 ± 1.40 ppm for CO2, 0.7 ± 7.3 ppb for CH4, 0.6 ± 0.6 ppb for N2O, 0.01 ± 0.10 ppt for SF6, 1.5 ± 5.3 ppb for CO and 4.8 ± 6.9 ppb for H2 for the years 2008–2012.

At Trainou station, the mean annual increase rates from 2007 to 2011 at the 180 m sampling height were 2.2 ppm yr−1 for CO2, 4 ppb yr−1 for CH4, 0.78 ppb yr−1 for N2O and 0.29 ppt yr−1 for SF6. For all species, the 180 m sampling level showed the smallest diurnal variation. Mean diurnal gradients between the 50 m and the 180 m sampling level reached up to 30 ppm CO2, 15 ppb CH4 or 0.5 ppb N2O during nighttime whereas the mean gradients are smaller than 0.5 ppm for CO2 and 1.5 ppb for CH4 during afternoon.

1 Introduction

The atmospheric greenhouse gas (GHG) measurement group (RAMCES) at the Laboratoire des Sciences du Climat et de l’Environnement (LSCE) in Gif-sur-Yvette is currently running a global network of 8 in situ stations and 13 flask sampling sites at 12 fixed surface sites, and one on board a small aircraft. At LSCE the air samples are analysed for CO2 isotopes (δ13C and δ18O) and for CO2, CH4, N2O, SF6, CO and H2 mole fractions. The RAMCES CO2 and radon-222 monitoring program was initiated in 1980 at the Île Amsterdam observatory (Gaudry et al., 1992; Ramonet and Monfray, 1996) and was extended at Mace Head, Ireland in 1992 (Biraud et al., 2000, 2002) and at two further sites in France (Gif-sur-Yvette and Puy de Dome) in 2011. The three
western European sites reflect different environments from a marine site occasionally influenced by long range transport over Europe (Mace Head), to sites which are more influenced by rural (Puy de Dôme) and urban activities (Gif-sur-Yvette).

Europe has a long history of atmospheric CO$_2$ monitoring at continental stations, in which the LSCE network participates. To avoid the influence of local trace gas sources, mountain stations such as Schauinsland (Germany), Mt Cimone (Italy) or Puy de Dôme (France) were traditionally chosen, and a data selection is performed at these sites to obtain a greenhouse gas record reflecting regional scale continental sources and sinks (Schmidt et al., 2003; Cundari et al., 1990). However, such measurements are difficult to reproduce by atmospheric transport models due to the influence of topography on air mass transport and mixing (Geels et al., 2007). Consequently, other approaches needed to be developed to complement regional scale emission inventories, through airborne measurements or high towers that avoid being strongly influenced by topography. The frequency of the airborne measurements is still a limiting factor, as well as the bias towards good weather conditions (Stephens et al., 2007). A very promising approach is therefore to establish new stations on tall towers (> 100 m) which are used as television transmission towers (Bakwin et al., 1998; Haszpra et al., 2001). Gloor et al. (2001) showed that the concentration footprint of the Wisconsin 500 m tall tower is of the order of 10$^6$ km$^2$.

In the framework of the European projects CHIOTTO (Continuous HIgh-precisiOn Tall Tower Observations of greenhouse gases, http://www.chiotto.org) and CarboEurope IP (http://www.carboeurope.org), eight European tall towers have been equipped or upgraded with high precision measurement systems for continuous greenhouse gas monitoring (Vermeulen et al., 2007). Results and instrumental setups are published by Thompson et al. (2009), Popa et al. (2010), Vermeulen et al. (2011). Our group has equipped a new station within Orléans Forest, called Trainou tower, with instrumentation for atmospheric CO$_2$, CH$_4$, CO, N$_2$O, SF$_6$, H$_2$ and radon-222 measurements. The measurements at four levels of Trainou tower (5, 50, 100, 180 m) are completed by regular airborne measurements between 100 and 3000 m altitude close to the tower. In the framework of the TCCON (Total Carbon Column Observing) network, IUP Bremen installed a ground-based Fourier transform spectrometer (FTS) to analyze the column abundances of CO$_2$, CH$_4$ and N$_2$O (Messerschmidt et al., 2011).

In the following chapters of the paper, we present the station location, the instrumental setup, data transmission, quality control and the time series of atmospheric CO$_2$, N$_2$O, CH$_4$, SF$_6$, CO H$_2$, and $^{222}$Rn measurements.

2 Site description

The station (Trainou tower, TRN) for atmospheric greenhouse gas observation is located in the “Centre” region in France (47°57′53″ N, 2°06′45″ E, 131 m above sea level (a.s.l.) at a 200 m transmitter mast. We setup sampling at four levels on this tower at 5 m (TR0), 50 m (TR1), 100 m (TR2) and 180 m (TR3). Trainou station is about 15 km northeast of the city of Orléans (116,000 habitants) and about 100 km south of Paris. Five small villages (< 2000 habitants) are located nearby the station. In the surroundings of the station, the area is covered by forest (30%), and agriculture fields (50%) (INSEE 2008, National Institute of Statistics and Economic Studies, 2008, http://www.insee.fr).

Using the anthropogenic emissions estimated from bottom up studies for the Centre region for 2000 (CITEPA, 2005, http://www.citepa.org), we can identify the main local and regional sources by sector. For CO$_2$, the three main emission sectors are transport, agriculture and households. More than 50% of the CH$_4$ emissions in this region are released by livestock production (dairy cows) and about 25% from waste treatment (landfills). For N$_2$O, the agriculture sector (fertiliser application in croplands) contributes more than 90% of the anthropogenic emissions in this region.

Prior to installation, we studied the wind direction and velocity distribution using the Trajectory Viewer 1.1 developed at ECN (Energy Research Centre of the Netherlands, http://www.ecn.nl/) and the model output from the FLEXible PARTICle dispersion model (FLEXPART) (Stohl et al., 2005). We do not show the real wind measurements here because we have only one wind sensor at the tower and our measured wind direction and velocity are biased by the wind shadow of the tower. Figure 1 shows that the dominating wind direction at the 180 m level is southwest followed by northeast winds. The distribution of the wind velocity shows maximum value at 6 m s$^{-1}$.
3 Measurement systems

The atmospheric measurement station at Trainou tower consists of a combined air inlet system piloted by the custom-made CO$_2$ instrument (CARIBOU), a coupled GC analyser system for CO$_2$, CH$_4$, N$_2$O, SF$_6$, CO and H$_2$, a radon-222 analyser and a flask sampler unit. In the following chapters, these five parts are described in detail. Meteorological sensors (see Sect. 3.4) and seven independent inlet lines (DEKABON 1300, 1/2\textquoteright\textquoteright o.d.) are installed on the tower. Three inlet lines are placed at the 180 m level, one at the 100 m, two at 50 m level and one at the 5 m level. For radon-222 gas measurements, a 65 mm inner diameter rigid tubing is installed at the 180 m level. The inlet lines are transferred to the container, which houses the measurement systems.

3.1 Air inlet and drying system

The pumping and air drying system is controlled by the CARIBOU CO$_2$ analyser software (see below). It consists of three identical branches of pumps and cooling traps for the 50, 100 and 180 m levels, which allows dried air from each level to be delivered to the analysing instruments at any time (Fig. 2). Therefore we are able to analyse all three levels at least once every 30 min, without risk of incomplete flushing of air from the level which was passed before. The three air sampling lines are flushed constantly with 15 L of air min$^{-1}$ by the primary pumps (Neuberger N815KNE). To protect the pumps, two filters (40 and 7 µm, TF series from Swagelok) are installed. Depending on the meteorological conditions, the filters need to be changed at least every six months. From each inlet line, a secondary pump (Neuberger N86KTE) collects an air flow of 5 L min$^{-1}$, which is then dried in two steps. First, the air passes through a glass trap (decanting bowl 120 mL), which is hosted in a commercial refrigerator kept at 5°C in order to remove the main water content from air. To remove condensed water, the traps are evacuated automatically once a day. In a second step, air is further dried by passing through 335 mL glass traps cooled in an ethanol bath to $-60^\circ$C using a cryogenic cooler (Thermo Neslab CC-65). The cooling traps are filled with glass beads to increase the surface area for water vapour condensation. These cooling traps are changed once per week by a technician who maintains the station regularly. This setup allows us to provide dried air with a dew point of less than $-50^\circ$C to the analysing instruments.

3.2 In situ CO$_2$ analyser (CARIBOU)

LSCE and CEA/IRFU (Institute of Research into the Fundamental Laws of the Universe) developed new robust CO$_2$ continuous monitoring stations, named CARIBOU. Within the RAMCES network, four of these stations have been installed. The first one was installed at Biscarrosse, France (inlet at 116 m a.s.l.) in May 2005, the second one at Hanle, India (4517 m a.s.l.) in August 2006 and the third one at Trainou tower, France. Finally, in summer 2007, a fourth one was installed at Ivittuut, Greenland.
The CARIBOU systems are designed to make high precision measurements (short-term repeatability of the order of ±0.01 ppm), with high hardware reliability, and require low maintenance. The design also allows fully remote control, automated data retrieval and online display capability.

The CARIBOU system consists of two main subsystems: (a) an analysis unit which includes a commercial non-dispersive, infrared (NDIR) gas analyser (LI-6252, LI-COR, Lincoln, Nebraska, USA), pressure, flow and temperature regulators controlled by a programmable logic controller (PLC), and an industrial PC used to configure and control the equipment; (b) a pumping unit, which includes the pumps for three air inlets, and a refrigerator for preliminary drying of the air to be analysed (see Sect. 3.1).

The CARIBOU design is based on the CSIRO (Commonwealth Scientific and Industrial Research Organisation, Melbourne, Australia) Loflo CO\textsubscript{2} analyser (Da Costa and Steele, 1997). A schematic view of the system is shown in Fig. 3. The CARIBOU regulates the flow in the sample and reference cells \(20.00 ± 0.03\) mL min\(^{-1}\) and the pressure inside the cells at \(1080.00 ± 0.03\) mbar with a combination of four flow-controllers (Redwood Fluistor NC-1500 and Bronkhorst F201CHBD11V) piloted by fuzzy logic algorithms. The box containing the NDIR analyser and the hardware necessary to regulate the gas flow and the pressure is thermally regulated. Moreover, there is an additional temperature control of the NDIR cells themselves \(40.00 ± 0.05\) °C. The thermal and pressure control of the instrument minimize the drift of the NDIR analyser of 1 ppm per day specified by the manufacturer (Li-6252 Instruction manual) to approximately 0.07 ppm per day. In addition, we analyse the reference gas for 10 min per hour in analysis cell, and use these results to correct for the instrument drift. After drift correction we obtain a residual drift of the order of 1 ppb per day allowing a full-span calibrations to occur once per week. A low flow of \(20\) mL min\(^{-1}\) and the weekly calibration frequency allows the calibration gas tank to last more than 10 years. The system is implemented by a 16 position micro-electric valve (Valco Vici) to select the gas to be measured (ambient air, calibration gases, etc.). The parameters for temperature and pressure regulation are summarized in Table 1.

3.2.1 Online data transmission and remote controlling

The CARIBOU unit has its own control system with an internet link to the LSCE laboratory (Gif-sur-Yvette) which allows remote supervision, configuration and maintenance. The raw data are downloaded automatically every day (or more frequently if needed) by a central computer at LSCE, which allows for database operations. This computer is also

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**Table 1.** Parameters of pressure, temperature and gas flow regulation for the CARIBOU CO\textsubscript{2} analyser.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Licor Cell pressure regulation</td>
<td>(1080.00 ± 0.03) mbar</td>
</tr>
<tr>
<td>Thermal regulation</td>
<td>(40.00 ± 0.05) °C</td>
</tr>
<tr>
<td>Gas flow regulation</td>
<td>(20.00 ± 0.03) mL min(^{-1})</td>
</tr>
</tbody>
</table>
in charge of daily tasks such as checking station health. It triggers alarms or warnings in case any abnormal conditions are detected, such as poor pressure or temperature regulation. Remote controlling further allows the user to have full access to switching of pumps and valves, the regulation of temperature set points and flow rates and the duration of gas analysis. Measurement sequences are created via an built-in sequencer and most of the data (CO\textsubscript{2} mole fraction, pressure, temperature, valves state, etc.) can be plotted on screen for a rapid diagnostic.

### 3.2.2 Calibration and quality control

At Trainou station, the CARIBOU calibration of the full span of the instrument is carried out every eight days using six station standard gases with a concentration range of 340–450 ppm. These standard gases are produced by Deust Steininger (Mühlhausen, Germany) in a gas matrix of so-called natural air (mixture of nitrogen, oxygen and argon) and filled in 10 L aluminium cylinders (Luxfer). At Trainou station, we use two-stage nickel-plated brass pressure regulators (Model 14, Scott Speciality Gases, Breda, The Netherlands). The station standard gases have been calibrated against the WMO X2007 scale (Zhao and Tans, 2006) at the LSCE using the Loflo 2-D analyser in 2006 and 2011. The difference between the calibrations in 2006 and 2011 was for all six cylinders smaller than 0.03 ppm. Therefore we used the arithmetic mean of both calibrations. For each calibration sequence, we repeat a pyramid-like pattern eight times, injecting the calibration gases and the reference gas in ascending and descending order of concentrations; each standard gas analysis lasts 10 min. For the following calculations, a mean of the last 4 min of analysis is taken. To determine and correct the instrument drift, the reference gas is injected for 10 min in the sample cell once per hour. After drift correction using the hourly reference gas analysis, the calibration results showed that the residual drift is lower than 1 ppb per day, which allows the calibrations of the instrument to be done no more frequently than once every eight days to maintain precision at 0.01 ppm.

Between two full-span calibrations, the ambient air measurement cycle, which lasts 7 h, is repeated 24 times. The ambient air measurement cycle is a sequence composed of five 10 min passages of the air from 180, 100, 50, 180 and 50 m above ground level (AGL) which converts CO\textsubscript{2} to CO and H\textsubscript{2} (via nickel catalyst) and an electron capture detector (µECD) for N\textsubscript{2}O and SF\textsubscript{6} (Lopez et al., 2012). In October 2008, we coupled a second GC containing a reduction gas detector (PP1 analyser, Peak Laboratories) to the existing GC system to analyse simultaneously CO and H\textsubscript{2} (Yver et al., 2011).

Figure 4 shows a schematic of the gas flow though the GC system. The principal parameters are summarized in Table 2. A 16 position valve (Valco Vici, UWE, microelectronic actuator) #7 is used to switch between the four air inlets (180, 100, 50 and 5 m AGL), standard and target gases. The sample loops are flushed with dry ambient air or standard gas with a flow rate of 200 mL min\textsuperscript{-1}, controlled by the EPC (electronic pressure control, AUX 5). After 45 s, EPC AUX 5 is closed, allowing a pressure and temperature equilibration for 30 s. The three sample valves (#1a, #1b and #5) are then switched simultaneously in order to transfer the content of the sample loops to the separation columns. All valves are installed in a closed electrical rack system, to protect against short-term temperature variations.

For the CO\textsubscript{2}/CH\textsubscript{4} branch, nitrogen (5.0, 99.999 %) is used as carrier gas for the separation on a packed Haysep Q column. At a flow rate of 50 mL min\textsuperscript{-1} (EPC AUX 3), CH\textsubscript{4} elutes about 70 s after injection and CO\textsubscript{2} 120 s after injection. After the CH\textsubscript{4} peak is detected on the FID, valve #4 is switched from the bypass to the nickel catalyst at 390 °C, which converts CO\textsubscript{2} to CH\textsubscript{4} permitting the CO\textsubscript{2} analysis on the FID. The efficiency of the Ni catalyst for reduction of CO\textsubscript{2} to CH\textsubscript{4} is better than 96 %. To supply hydrogen to the FID and the Ni catalyst, we used hydrogen generators from Parker/Balston (model A9150 until summer 2008 and model 9200). In 2010, we replaced the hydrogen generator again with a NM-H\textsubscript{2} 250 (FDDBS). Synthetic air is supplied by a combination of a compressor (Jun Air) and a purifier (Parker, Chromgas 1000).

For the N\textsubscript{2}O/SF\textsubscript{6} branch, a 5 % CH\textsubscript{4} in Argon (ECD quality) is used as carrier gas. The separation of the peaks is performed on two packed Haysep columns. At 70 s after...
injection, when N₂O and SF₆ are on the analytical column, the pre-column is back flushed by turning valve #3. This prevents compounds with longer retention times from eluting with the next injection and therefore shortens the analysis time. N₂O and SF₆ are detected on the ECD 150 and 180 s respectively, after injections.

Electron capture detectors are known for their non-linear responses when measuring N₂O and the possible problem of co-elution of N₂O and CO₂, depending on the column setup. As described in more detail by Lopez et al. (2012) and Schmidt et al. (2001), we determine the non-linear response of the ECD using a reference sample diluted with N₂O-free and CH₄-free air. Methane is used to determine the dilution factor, as the response of the FID for CH₄ is linear within the chosen range. We found that the μECD at Trainou underestimates the N₂O concentration with increasing values, but a linear correction can be added to describe the response of the ECD using a reference sample diluted with N₂O from a reference sample. As described by Lopez et al. (2012), the influence of the CO₂ co-elution is negligible for the N₂O measurements at Trainou. They also reported a cross sensitivity of N₂O and SF₆, at Gif-sur-Yvette station, when SF₆ peaks exceeded 15 ppt. At Trainou station we never observed SF₆ mole fractions larger than 14 ppt, but were careful not to use calibration gases with elevated SF₆.

In order to add CO and H₂ measurements, we coupled a reduction gas detector (RGD) from Peak Performer, to the Agilent GC in 2008 (Yver et al., 2011). The sample loop of the RGD is connected in series with the sample loop of the FID and ECD. The analysis is performed by reduction of mercuric oxide and detection of mercury vapour by UV absorption. Synthetic air (Air Liquide, high purity) purified by an additional filter in the chromatograph is used as carrier gas. Aliquots of air are flushed through a 1 mL sampling loop. After equilibration, the sample is injected onto the columns. The pre-column (Unibeads 1S) is used to separate H₂ and CO from other components and the analytical one (Molecular Sieve 5Å) effectively separates H₂ and CO before being analysed by the mercury detector. The oven temperature is held at 105 °C and the catalytic chamber is heated to 265 °C.

The non-linearity of the RGD detector is regularly determined using a set of cylinders spanning H₂ mole fractions between 213 and 996 ppb and CO mole fractions between 157 and 523 ppb. The standard cylinders for H₂ were calibrated at the Max Planck Institute (MPI) Jena and for CO by NOAA/CMSL. During the test of the analyser in our laboratory, we performed a non-linearity test with 11 standard cylinders for H₂ and 5 for CO. During operation at Trainou station the number of standard cylinders had to be reduced to only four for both species due to logistical problems. In general, the standard cylinders are evaluated every 6–12 months. Over the described measurement period, the non-linearity of the RGD at Trainou did not change significantly for CO and H₂ in the range of observed mixing ratios. However, for H₂, a shift in the non-linearity was observed for mole fractions above 800 ppb. As the ambient air mole fractions are below 650 ppb, all data are corrected with the same third degree polynomial fit. Only the quality control gas (Target) with a H₂ mole fraction above 800 ppb is affected by this choice with an observed shift over time (see Sect. 3.3.2).

Table 2. Settings of the GC parameters.

<table>
<thead>
<tr>
<th></th>
<th>CO₂ and CH₄</th>
<th>N₂O and SF₆</th>
<th>CO and H₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oven temperature (column)</td>
<td>80 °C</td>
<td>80 °C</td>
<td>106 °C</td>
</tr>
<tr>
<td>Sample loops</td>
<td>15 mL</td>
<td>10 mL</td>
<td>1 mL</td>
</tr>
<tr>
<td>Flow through sample loops</td>
<td>200 mL min⁻¹</td>
<td>200 mL min⁻¹</td>
<td>200 mL min⁻¹</td>
</tr>
<tr>
<td>Carrier gas</td>
<td>Nitrogen 5.0</td>
<td>Air/CH₄ (5 %)</td>
<td>Synthetic air</td>
</tr>
<tr>
<td>Carrier gas (flow)</td>
<td>50 mL min⁻¹</td>
<td>40 mL min⁻¹</td>
<td>15 mL min⁻¹</td>
</tr>
<tr>
<td></td>
<td>(backflush: 55 mL min⁻¹)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Analytical columns</td>
<td>Hayesep-Q, 80–100 mesh (3.65 m × 4.7625 mm)</td>
<td>Hayesep-Q, 80–100 mesh (1.829 m × 4.7625 mm)</td>
<td>Molecular Sieve 5 Å 60–80 mesh (2.032 m × 3.175 mm)</td>
</tr>
<tr>
<td>Pre-column</td>
<td>Hayesep-Q, 80–100 mesh (1.219 m × 4.7625 mm)</td>
<td>Hayesep-Q, 80–100 mesh (1.219 m × 4.7625 mm)</td>
<td>Unibeads 1S column 60–80 mesh (0.762 m × 3.175 mm)</td>
</tr>
<tr>
<td>Detector</td>
<td>FID</td>
<td>μECD</td>
<td>RGD</td>
</tr>
<tr>
<td>Detector temperature</td>
<td>300 °C</td>
<td>395 °C</td>
<td>265 °C</td>
</tr>
<tr>
<td>H₂ Flow</td>
<td>50 mL min⁻¹</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Air Flow</td>
<td>400 mL min⁻¹</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methanizer</td>
<td>390 °C</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 4. Schematic of the two coupled GC systems. The FID and the ECD detector are housed in a GC 6890N from Agilent Technologies and the reduction gas detector (RGD) in a PP1 from Peak Laboratories. The three sample loops are flushed with the same sample and injected at the same time.

With this setup, we are able to achieve injection and detection of one sample within 5 min, allowing 12 ambient air or standard gas injections over 1 h.

3.3.1 Data acquisition and remote controlling

For the full control of the GC system, data acquisition and remote controlling, we use one PC, which is equipped with the chromatography software CHEMSTATION (A.09.03) from Agilent. To treat the signal of the PP1 GC in the same way and to synchronize the two GCs, we added an analog-to-digital converter (Agilent Technologies Inc 35900E). The CHEMSTATION software allows for scheduling the switching of all valves in order to control the temperature of the oven and the detectors and to regulate the flow of the carrier gases and the flushing of the sample loops. The detector signals of both GCs are recorded by the CHEMSTATION and at the end of each injection method, the integration of the peaks is performed.

Once a day, we automatically transfer the raw data (chromatograms and integrated peak areas and heights) via file transfer protocol to a central computer at LSCE. From there, peak areas and heights, sample information, etc., are read by the database and computed to concentration values. Chromatograms and other additional information are stored separately in the case of necessary data re-evaluation. In case modifications to the methods, sequences or integrations are needed, we can remotely access the CHEMSTATION at Trainou tower from our institute at any time.

3.3.2 Calibration and quality control

To correct for possible drift of the measurement system, we decided to perform the working standard analysis every 30 min. As the ECD and also the RGD show a non-linear behaviour, we inject a working standard low (WL) and a working standard high (WH) in order to apply a two-point calibration. These standards are produced by Deuste Steininger (Mühlhausen, Germany), and filled in 40L aluminium cylinders (Luxfer). The concentration range is chosen to bracket the atmospheric concentration range for all analysed gases at Trainou station. As the first working standards did not contain H₂, a working standard "special" (WS) is used to calculate the H₂ concentration. The non-linearity of the RGD is corrected using a third-degree polynomial fit. This fit is determined by measuring a range of four standard gases (Yver et al., 2011).

Like for CO₂ CARIBOU measurements, a target gas, with a known concentration value in between the WL and WH concentration, is analysed every 1–2 h. A typical 1 h sequence consist of WH, WL, WS, Target, TR3, TR2, TR1, WH, WL, TR3, TR2, TR1 and TR0. With this frequency of calibration, the theoretical lifetime of working standards is 1.5–2 years and for the target gas 3–4 years.

The target gas is used to calculate the short-term repeatability (on a daily basis) and the long-term reproducibility. Figure 5 shows the time series of the hourly target gas injection for CO₂, CH₄, N₂O, SF₆ and since July 2009, also for CO and H₂. The large gaps in winter 2006/2007 are due to electrical power supply problems. We then had to stop all measurement systems between October 2006 and February 2007. In November 2007, the ECD was broken, and could only be replaced in August 2008. For the whole period of analysis, we found a mean standard deviation of the hourly target gas injection of 0.14 ppm for CO₂, 3.2 ppb for CH₄, 0.7 ppb for N₂O, 0.08 for SF₆, 1.9 ppb for CO and 13 ppb for H₂. We can identify several periods where the GC was not working very well for different trace gas analyses, increasing the standard deviation of our target measurements. In 2009, we had repeated problems with hydrogen generators, which were not producing hydrogen that was sufficiently dry. After the replacement of the hydrogen generator in 2010, we achieved a CH₄ repeatability of 1.4 ppb. Similar problems, as mentioned above but caused by the ECD, are reported...
Figure 5. Long-term reproducibility of the target gases for CO₂ measured with the CARIBOU and for CO₂, CH₄, N₂O, SF₆, CO and H₂ measured with the coupled GC system. For all species the differences between measured data and their mean values are plotted against time. The drift of the target cylinder for H₂ can be explained by the elevated mole fraction of 841 ppb which is greatly above the typical high ambient air values of 650 ppb at Trainou. With our non-linearity test we were not able to correct such high values.

for the N₂O reproducibility. In 2009, the ECD worked without issues and the target gas showed a standard deviation of only 0.21 ppb. The hydrogen mole fraction of the target cylinder shows clearly a drift-like behaviour with decreasing values. As explained in Sect. 3.3, the non-linearity of the RGD-system changes for H₂ mole fraction above 800 ppb. The target cylinder has a H₂ concentration of 841 ppb which is much above typical high ambient air values of 650 ppb at Trainou. With our non-linearity test we were not able to correct such high values. Outside of the calibrated range has this target cylinder only a limited suitability for hydrogen and should be interpreted with care. To link the station to the international calibration scales, the working standards have been initially calibrated against gas tanks calibrated by the WMO calibration centres at NOAA (Zhao and Tans, 2006; Dlugokencky et al., 2005; Hall et al., 2007) or MPI Jena (Jordan and Steinberg, 2011). Before the end of the lifetime and in the case of drifts or jumps in the quality control gas, the working standards at the station are reanalysed in our laboratory in Gif-sur-Yvette. Details of the instrumentation used to calibrate our station standards in our laboratory and the WMO scales for different gases are summarized in Table 3.

3.4 Other instrumentation

3.4.1 Meteorological sensors

Meteorological sensors are installed at the 180 m (TR3) and at the surface level on the top of the container about 5 m above ground (TR0). To measure the wind speed and direction, we use a sensor from Young (Model 05103L) and for the temperature and humidity a sensor from Vaisala (HUMICAP HMP45/A). The data acquisition and transfer to the database is run by the CARIBOU software (see Sect. 3.2).

3.4.2 Flask sampling

Weekly flask sampling is performed from an independent sampling line at the 180 m level with a standard flask sampler used in the French GHG network. The sampling unit consist of a diaphragm pump (Model N86 KNDC, KNF Neuberger) which draws air through a chemical drying cartridge filled with anhydrous magnesium perchlorate [Mg(ClO₄)₂]. Air is collected in 1 L glass flasks sealed with PTFE (polytetrafluoroethylene) O-rings (Normag Labor- und Prozesstechnik, Ilmenau, Germany). Flasks are collected in pairs and pressurized to 2 bar absolute pressure. The sampling started with a low frequency in 2006. Since June 2007, a regular sampling has been performed during the weekly maintenance of the station. The flasks are transported to our central lab at Gif-sur-Yvette and analysed for mixing ratios of CO₂, CH₄, N₂O, SF₆, H₂, and CO by GC. The isotopic composition of CO₂ is determined using a Finnigan MAT252 isotope ratio mass spectrometer (IRMS), equipped with a trapping box for cryogenic separation of CO₂ from air.


www.atmos-meas-tech.net/7/2283/2014/
Table 3. Primary calibration scales at the LSCE central lab, which are used to calibrate the station standards. CO₂ calibration of the working standards is generally performed with the Loflo (NDIR) analyser. The other gases are calibrated by GC system.

<table>
<thead>
<tr>
<th>Component</th>
<th>Instrument</th>
<th>Range</th>
<th># of tanks</th>
<th>Scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂</td>
<td>Loflo</td>
<td>318–468 ppm</td>
<td>7</td>
<td>WMO X 2007</td>
</tr>
<tr>
<td>CO₂</td>
<td>GC (LSCE)</td>
<td>365–462 ppm</td>
<td>6</td>
<td>WMO X 2007</td>
</tr>
<tr>
<td>CH₄</td>
<td>GC (LSCE)</td>
<td>1634–2081 ppb</td>
<td>6</td>
<td>NOAA04 scale</td>
</tr>
<tr>
<td>N₂O</td>
<td>GC (LSCE)</td>
<td>302–340 ppb</td>
<td>6</td>
<td>NOAA 2006A</td>
</tr>
<tr>
<td>SF₆</td>
<td>GC (LSCE)</td>
<td>6–15 ppb</td>
<td>6</td>
<td>NOAA 2006</td>
</tr>
<tr>
<td>CO</td>
<td>GC (LSCE)</td>
<td>41–595 ppb</td>
<td>6</td>
<td>NOAA-GMD/WMO 2004</td>
</tr>
<tr>
<td>H₂</td>
<td>GC (LSCE)</td>
<td>430–810 ppb</td>
<td>3</td>
<td>MPI 2009</td>
</tr>
</tbody>
</table>

3.4.3 Radon-222 measurements

In June 2009, we installed a radon measurement instrument at Trainou, which was tested between January 2007 and May 2009 in Gif-sur-Yvette (Yver et al., 2009). The detection method based on a double filter was installed in cooperation with the Australian Nuclear Science and Technology Organisation (ANSTO). In the double filter method (Whittlestone and Zahorowski, 1998; Zahorowski et al., 2004), ambient air is continuously pumped through two 200 L cylinders and through a first filter to the analysis chamber. During this first step, radon-222 (²²²Rn) disintegrates as the time residence is ten times higher than its half-life and the filter retains all solid particles letting just ²²²Rn coming in the analysis chamber. In this chamber, ²²²Rn daughters are produced and collected on a second filter. The α-decay of these daughter elements is then counted. The instrument is regularly calibrated with known quantity of ²²²Rn emitted by a ²²⁶Ra source (type 2000A-20, Pylon). The inlet line is installed at the 180 m level.

3.5 Comparison of the different instruments and flask analysis

In 2007 and from 2009 to 2012, the GC system and the CARIBOU analysed CO₂ mole fractions at the 50, 100 and 180 m sampling level. The sampling of the different levels for both analysers is not synchronized, but both analysers sample each level between one and two times per hour. In the case of the GC, it is a single injection and in the case of the CARIBOU a 10 min interval with a frequency of 1 Hz. In order to compare both values, we used the 1 h mean values, which risks a maximum time shift of 50 min in the case of only one injection per analyser during the 1 h with a maximum time difference. The mean differences and the 1σ standard deviations are summarized in Table 4 for the different years and the different sampling level. The values correspond to the GC measurement minus CARIBOU measurement. Generally, the CO₂ difference between both analysers was of the order of 0.1 ppm or less, with the exception of the years 2007 and 2011, when the GC measured around 0.15 ppm higher values at all three sampling level. This indicates a possible leakages or a calibration problem of the GC, but afterwards we were not able to isolate it to one specific analyser.

In parallel to the GC measurements, weekly flask sampling at the 180 m level was carried out and analysed on the GC system in the LSCE central lab in Gif-sur-Yvette (see Sect. 3.4.2). During the period 2007–2012 we monitored a mean difference between GC Trainou and flasks of −0.08 ± 1.40 ppm CO₂, −0.7 ± 7.3 ppb CH₄, −0.64 ± 0.62 ppb N₂O, 0.01 ± 0.10 ppb SF₆, 1.5 ± 5.3 ppb CO and 4.85 ± 6.9 ppb H₂.

3.6 Additional instrumentation

In cooperation with the Institut für Umweltphysik (IUP) at the University of Heidelberg, we installed a sampler for ¹⁴CO₂ sampling. Integrated atmospheric ¹⁴CO₂ samples have been collected from the 180 m inlet line over a two week period in CO₂– free sodium hydroxide solution since January 2008. The flushing rate is adjusted in order to sample CO₂ from a total volume of 20–25 m³ of air. The sampling and analysis technique is described in detail by Levin et al. (1980).

Since December 2010, a vertically pointed aerosol lidar system (ALS300, Leosphere, Orsay, France) for monitoring atmospheric boundary layer (ABL) depths is routinely operated at Trainou tower. Full details of the instrumentation and the first time series of ABL depths can be found in Pal et al. (2012, 2014).

4 Measurement of ambient air

Figure 6 shows the hourly averaged time series of CO₂, CH₄, N₂O and SF₆, CO, H₂ mole fractions and ²²²Rn activity for the four sampling heights (180 m in blue, 100 m in green, 50 m in red and 5 m in black). The lowest sampling level (5 m) was only added in October 2010. As already described in Sect. 3.3.2, all measurement systems were stopped between October 2006 and February 2007. In November 2007, the ECD was broken, and could only be replaced in August 2008 leading to gaps in the time series of N₂O and SF₆.
Table 4. Mean annual CO$_2$ difference and 1$\sigma$ standard deviation for the three ambient air sampling lines at Trainou station (GC-CARIBOU). For the mean difference of the period 2007–2012 we applied a 2 sigma filter.

<table>
<thead>
<tr>
<th>Year</th>
<th>50 m sampling level $\Delta$ CO$_2$ (ppm)</th>
<th>100 m sampling level $\Delta$ CO$_2$ (ppm)</th>
<th>180 m sampling level $\Delta$ CO$_2$ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2007</td>
<td>0.12 ± 1.75</td>
<td>0.12 ± 1.0</td>
<td>0.15 ± 0.75</td>
</tr>
<tr>
<td>2009</td>
<td>0.11 ± 1.3</td>
<td>0.05 ± 1.01</td>
<td>−0.06 ± 1.0</td>
</tr>
<tr>
<td>2010</td>
<td>0.07 ± 1.05</td>
<td>0.03 ± 1.09</td>
<td>−0.02 ± 0.88</td>
</tr>
<tr>
<td>2011</td>
<td>0.13 ± 0.82</td>
<td>0.18 ± 1.1</td>
<td>0.21 ± 1.1</td>
</tr>
<tr>
<td>2012</td>
<td>−0.01 ± 1.26</td>
<td>0.07 ± 0.97</td>
<td>−0.07 ± 0.81</td>
</tr>
<tr>
<td>2007–2012</td>
<td>0.08 ± 1.00</td>
<td>0.09 ± 1.05</td>
<td>0.01 ± 0.93</td>
</tr>
</tbody>
</table>

4.1 Growth rates and seasonal variations

We determine the growth rate and the seasonal cycles for all trace gases at the highest sampling level at 180 m. We employ the curve fitting procedure described by Thoning et al. (1989) and Masarie and Tans (1995) to the daily mean values. The curve fit incorporates harmonic and quadratic functions, and an 80-day smoothing is applied to the residuals. A 3$\sigma$ filter is applied in the smoothing process to obtain the best fit curve, excluding statistical outliers.

The CO$_2$ time series shows a seasonal cycle with minimum value in summertime (August) and a wintertime maximum (January) as expected. Superimposed on this cycle are synoptic and diurnal peaks generally associated with regional pollution events and changes in the boundary layer height. Diurnal variations with CO$_2$ mole fractions of up to 550 ppm during summer have been monitored at the lowest sampling level, 5 m above ground, whereas at the 180 m level, elevated mole fractions of up to 450 ppm occurs from time to time in winter. The amplitude of the diurnal variations decreases with the increasing sampling height. We computed a CO$_2$ annual growth rate of 2.2 ppm per year for the 180 m sampling level between 2007 and 2012, considering all-day round measurements.

The CH$_4$ time series shows a mean seasonal cycle of approximately 70 ppb peak-to-peak amplitude with higher values during January and lower values at the end of July. High short-term (diurnal) variations occur during winter correlated with pollution events of other trace gases, when the nocturnal planetary boundary layer height is low. During these events the CH$_4$ mole fraction can increase to up to 2400 ppb at the lowest (5 m) and 2050 ppb at the 180 m sampling level. Averaged over the plotted measurement period, CH$_4$ increased 4.1 ppb per year at Trainou station (180 m sampling).

N$_2$O mole fractions at 180 m sampling level increase with an average annual growth rate of 0.78 ppb per year. Again the lower sampling levels show larger variability likely reflecting local emissions from soil and other sources. In the study of Lopez et al. (2012), N$_2$O emissions in the catchment area of Trainou station are discussed in detail.

The SF$_6$ increase rate at Trainou 180 m sampling level equals 0.29 ppt per year. Superimposed on this trend are a few events with elevated SF$_6$ mole fractions of nearly 15 ppt at lower sampling levels (50 m and 5 m) and approximately 10 ppt at the highest sampling level.
Amplitudes of the diurnal variation and also nighttime gradients between sampling levels are largest between May and October for CO₂ mole fractions. The largest diurnal variations are observed on the sampling level closest to the ground and therefore close to local sources, but we still observe small diurnal cycles of CO₂ mole fractions at the 180 m sampling height as the nocturnal boundary layer is sometimes above 180 m. Also, CO₂ accumulated during nighttime below the 180 m sampling height can be observed as a small peak in the late morning, when vertical mixing starts and increases. A maximum gradient between the 5 m level and the 180 m level (see Fig. 8) of 30 to 50 ppm CO₂ is reached at 05:00 GMT when the nocturnal boundary layer is below 180 m. The CO₂ gradients between 180 and 100 m, and 180 m 50 m sampling lines are approximately 5 and 10 ppm, respectively. In the afternoon, 12:00 to 16:00 GMT, the monthly mean CO₂ mole fractions of the sampling levels between 50 and 180 m agree within 0.5 ppm from March to November and within 1 ppm during the winter months (December to February), respectively.

During spring and summer at the Cabauw and Bialystok tall towers, Vermeulen et al. (2011) and Popa et al. (2010) observed lower daytime CO₂ mole fractions at the lower sampling heights compared to the highest sampling heights due to local CO₂ net uptake. Even though the Trainou tower is surrounded by forest, we have not observed lower daytime CO₂ mole fractions in the mean gradients for spring or summer at the lower levels.

Figure 7 summarises the mean diurnal variation of CO₂, CH₄, N₂O, SF₆, CO and H₂ mole fractions and ²²²Rn activity for the four sampling levels averaged by month and hour of day. The x axis of each month covers an interval between 00:00 and 24:00 (GMT), where midnight GMT corresponds to 01:00 local time at Trainou station. CO₂, CH₄ and N₂O mole fractions show larger increases during nighttime, with strong accumulations at ground level. During afternoon,

Yver et al. (2011) have described the first year of CO and H₂ time series. We confirmed their findings of seasonal cycles at Trainou station with minimum values for CO mole fractions in July and for H₂ in September. Superimposed on this signal are diurnal peaks and synoptic peaks. In winter, CO mole fractions of up to 700 ppb have been observed at the lowest sampling level, 5 m above ground, whereas at the 180 m level elevated mole fractions of up to approximately 500 ppb occurs occasionally. In summer, H₂ mole fractions can be smaller at the lower levels than at the 180 m level due to uptake of H₂ by the soil.

### 4.2 Mean diurnal variation and vertical gradient

Figure 7 summarises the mean diurnal variation of CO₂, CH₄, N₂O, SF₆, CO and H₂ mole fractions and ²²²Rn activity for the four sampling levels averaged by month and hour of day. The x axis of each month covers an interval between 00:00 and 24:00 (GMT), where midnight GMT corresponds to 01:00 local time at Trainou station. CO₂, CH₄ and N₂O mole fractions show larger increases during nighttime, with strong accumulations at ground level. During afternoon,

The mean diurnal cycles of CO₂ and CH₄ show a peak in the morning and a smaller peak in the late afternoon. The peak in the late afternoon is due to the soil uptake of CO₂ and the production of CH₄ by the soil.

Table 1 shows the mean diurnal cycles of CO₂, CH₄, N₂O, SF₆, CO and H₂ mole fractions and ²²²Rn activity for the four sampling levels averaged by month and hour of day. The x axis of each month covers an interval between 00:00 and 24:00 (GMT), where midnight GMT corresponds to 01:00 local time at Trainou station. CO₂, CH₄ and N₂O mole fractions show larger increases during nighttime, with strong accumulations at ground level. During afternoon,

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The mean diurnal cycles of CO₂ and CH₄ show a peak in the morning and a smaller peak in the late afternoon. The peak in the late afternoon is due to the soil uptake of CO₂ and the production of CH₄ by the soil.
The mean diurnal variation of the CH\textsubscript{4} mole fraction show similar patterns as for CO\textsubscript{2}, with a stratification of the different sampling levels during nighttime. However, the amplitude of the 5 m level is less distinctive, indicating weaker local emissions compared to CO\textsubscript{2}. The nighttime CH\textsubscript{4} gradients between 180 and 100 m, and 180 and 5 m sampling level are approximately 30 and 15 ppb respectively. The afternoon values of all sampling levels agree within 1–2 ppb.

At Trainou tower, the mean diurnal cycle amplitudes for N\textsubscript{2}O are approximately 0.5 ppb for the 50, 100 and 180 m sampling levels. During February, March, April, August and September, the 5 m sampling level clearly shows accumulation of N\textsubscript{2}O during nighttime, which corresponds to large N\textsubscript{2}O emissions from arable land shown by Lopez et al. (2012) for the catchment area of Trainou.

SF\textsubscript{6} shows only a weak diurnal cycle without accumulation during nighttime. No significant gradient built up between the 50, 100 and 180 m sampling levels. Peaks shown in Fig. 7 can be attributed to single SF\textsubscript{6} “events”.

CO shows a noticeable seasonal behaviour, with large diurnal cycles and vertical gradients between October and March and very small diurnal cycles between April and September with small vertical gradients, also during nighttime.

For H\textsubscript{2}, in winter, there is almost no difference between the three heights (50, 100, 180 m) whereas in summer, due to the H\textsubscript{2} uptake by soil, the gradient can reach up to 20 ppb between the 50 and the 180 m inlet and even up to 60 ppb between the 5 and the 180 m line. With the soil sink predominating during nighttime, when the boundary layer is low and the emissions reduced, the lowest mixing ratios are encountered at 5 m. A more detailed analysis of H\textsubscript{2} and CO is presented by Yver et al. (2011).

5 Conclusions

A new fully automated measurement system to analyse simultaneously atmospheric CO\textsubscript{2}, CH\textsubscript{4}, N\textsubscript{2}O, SF\textsubscript{6}, CO and H\textsubscript{2} was installed on a 200 m transmission tower at Trainou in the Orléans forest. The station was installed in summer 2006 starting with a CO\textsubscript{2} analyser and a GC system for CH\textsubscript{4}, N\textsubscript{2}O, SF\textsubscript{6}, CO and H\textsubscript{2} and consecutively upgraded with new instrumentation for CO and H\textsubscript{2}, \textsuperscript{222}Rn analysis and \textsuperscript{14}CO\textsubscript{2} sampling. After some logistical problems in the starting phase of the installation, the station has delivered high quality measurements fulfilling the requirements of the WMO recommendations. In June 2012, the electron capture detector developed a leak and therefore we had to close the station for some months and to take out of service the GC system. In June 2013, the station was upgraded with a CRDS analyser (G2401, Picarro Inc.) to measure CO\textsubscript{2}, CH\textsubscript{4} and CO, and in 2015 a second CRDS analyser for N\textsubscript{2}O will be added. The CARIBOU system will continue to be in service in order to have more than one year of overlap between the different analysers. The new instrumentation will have the advantage of requiring less maintenance than the GC system. In recent years the maintenance of the GC, operated remotely with 10 service visits per year, was quite time consuming taking into account the 180 km distance between Trainou and our laboratory in Gif-sur-Yvette.

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