The arctic seasonal cycle of total column CO2 and CH4 from ground-based solar and lunar FTIR absorption spectrometry

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Abstract. Solar absorption spectroscopy in the near infrared has been performed in Ny-Ålesund (78.9°N, 11.9°E) since 2002; however, due to the high latitude of the site, the sun is below the horizon from October to March (polar night) and no solar absorption measurements are possible. Here we present a novel method of retrieving the total column dry-air mole fractions (DMFs) of CO$_2$ and CH$_4$ using moonlight in winter. Measurements have been taken during the polar nights from 2012 to 2016 and are validated with TCCON (Total Carbon Column Observing Network) measurements by solar and lunar absorption measurements on consecutive days and nights during spring and autumn. The complete seasonal cycle of the DMFs of CO$_2$ and CH$_4$ is presented and a precision of up to 0.5% is achieved. A comparison of solar and lunar measurements on consecutive days during day and night in March 2013 yields non-significant biases of 0.66 ± 4.56 ppm for $x_{CO_2}$ and $-1.94 ± 20.63$ ppb for $x_{CH_4}$. Additionally a model comparison has been performed with data from various reanalysis models.

1 Introduction

Since 1992 a Fourier transform infrared (FTIR) spectrometer (FTS) in Ny-Ålesund (78.9°N, 11.9°E) has been used for the ground-based observation of total column trace gas abundances in the Arctic via solar absorption spectroscopy (Notholt and Schrems, 1994). The measurements are taken within the Infrared Working Group (IRWG) of the Network for the Detection of Atmospheric Composition Change (NDACC). Since 2002, measurements in the near-infrared (NIR) spectral region have been performed to retrieve the dry-air mole fractions (DMFs) of CO$_2$ and CH$_4$ (denoted here as $x_{CO_2}$ and $x_{CH_4}$) and other gases (Warneke et al., 2005, 2006). These are, since 2005, part of the Total Carbon Column Observing Network (TCCON). Today, these measurements are widely used as validation for satellite products, in model comparisons and studies of sources and sinks.

A large limitation of the availability of these measurements is the absence of sunlight in the polar winter. At Ny-Ålesund, between October and March, the sun is permanently below the horizon. However, during this period the moon is permanently above the horizon around full moon.

Moonlight has already successfully been used as a light source in retrievals of various trace gas concentrations via the FTS in Ny-Ålesund in the mid-infrared spectral region (Notholt et al., 1993, 1997; Notholt and Lehmann, 2003; Palm et al., 2010) and in Antarctica (Wood et al., 2004). Here the employment of liquid-nitrogen-cooled InSb and MCT detectors ensures low instrumental noise, even under low light conditions. In the NIR, i.e. $>4000\, \text{cm}^{-1}$, typically extended-range InGaAs diodes are used. Recently Fu et al. (2014) and Wong et al. (2015) showed the application of a thermoelectrically cooled InGaAs detector for the measurement of reflected sunlight spectra from the Los Angeles basin on a mountaintop site. The thermoelectrical cooling reduces the detector noise and allows for higher signal-to-noise ratios in the measured spectrum.

After initial tests at the Bremen TCCON site (Buschmann et al., 2015), a thermoelectrically cooled InGaAs diode detector was implemented in the Ny-Ålesund FTS and a time series of $x_{CO_2}$ and $x_{CH_4}$, the total column dry-air mole frac-
tion, was obtained from spectra measured during polar nights between 2012 and 2016. The resulting product is compared to TCCON solar measurements as well as model simulations from the MACC reanalysis model for CO\(_2\) (v14r2; MACCCO2, 2015) and for CH\(_4\) (v10; MACCCCH4, 2015), the Jena CO\(_2\) inversion CarboScope s04, v3.7 (JenaCO2, 2005) and the CarbonTracker 2015 model (CT2015, 2016). Together with the summer TCCON data from Ny-Ålesund, for the first time the whole seasonal cycle of xCO\(_2\) and xCH\(_4\) is presented.

In Sects. 2 and 3, this paper describes the measurement set-up and the methods used to retrieve the dry-air mole fractions. Section 4 describes the newly obtained time series and the comparison to TCCON. Finally we compare our results with model data in Sect. 5.

## 2 Set-up

### 2.1 Measurement site

The instrument, a Bruker IFS 120-5HR, is located at the AWIPEV research station in Ny-Ålesund (78.92° N, 11.92° E). Measurements are taken under cloud-free conditions for both the NDACC and TCCON networks during summer, and lunar absorption and atmospheric emission measurements are performed in winter. In 2014–2015 the measurement set-up was gradually changed to a semi-automated system. The new system is able to automatically start a set of measurements without the need of an operator, which considerably increased the number of measured spectra. The performance of the instrument is monitored by reference cell measurements on a monthly basis and it is ensured that the phase error is smaller than \(\pm 0.04\) rad and the modulation efficiency is \(\pm 2\%\) of 1.0 up to a maximum optical path difference of 180 cm. These values are indicative of a well-aligned instrument.

### 2.2 Thermoelectrically cooled InGaAs diode

The sensitivity of the extended InGaAs diode used as a detector in standard TCCON near-infrared measurements is too small to obtain a sufficient signal-to-noise ratio (S/N) from lunar irradiance. The introduction of a two-stage Peltier element cooling system attached to the back of the diode can reduce the dark current noise and thereby minimize overall detector noise. Generally the extension of the detectors spectral sensitivity range reduces the quantum efficiency. Therefore, a non-extended diode improves the signal-to-noise ratio; however, cooling the InGaAs diode affects its crystal structure and therefore widens the band gap, which leads to a shift of the diode’s sensitivity range. The commercially available diode used here has a cut-on frequency of about 5260 cm\(^{-1}\) in the uncooled and about 5450 cm\(^{-1}\) in the cooled state. The noise equivalent power of the cooled diode, i.e. the power of the incident light to achieve a signal-to-noise ratio of 1, is about 3 orders of magnitude smaller than that of a standard TCCON diode. The shift in sensitivity due to cooling, an example of an averaged lunar spectrum, and a picture of the diode are shown in Fig. 1.

### 2.3 Availability of moonlight

The total number of potential lunar measurement hours can be calculated by excluding all times where the lunar elevation is below the terrain height. Additionally, lunar phases with insufficient illumination (lunar phase < 85 %) and times where the solar zenith angle is smaller than 95° have to be excluded.

Depending on lunar orbital parameters, the maximum number of measurement hours ranges from about 886 h in 2012 to 634 h in 2016. This is much less than the potential yearly solar measurement time of 3883 h. The minimum lunar zenith angle is 57.13° (2012) and 60.84° (2016) compared to a minimum solar zenith angle of 55.47°.

The actual possible time available for near-infrared measurements, of course, further depends on clear-sky conditions and other scheduled FTS experiments. The number of measurements was increased by switching to a semi-automated measurement set-up that required less operator intervention in autumn 2015, as described above.

## 3 Method

### 3.1 Measurement set-up

The measurements follow the TCCON standard settings wherever possible. A solar (lunar) tracker is mounted on the roof of the AWIPEV observatory and the light is reflected into the laboratory underneath and into the FTS. Accurate tracking is ensured by usage of a four-quadrant diode with feedback to the solar tracker motor controller. The incident light is focused on an entrance aperture and afterwards parallelized to enter a Michelson interferometer arrangement of the Bruker IFS 120-5 HR. The movable retro-reflective mirror is mounted on a sledge on steel rods. Accurate tracking of the movable mirror’s position is provided by a stabilized internal HeNe laser reference. The light path arrives in the detector compartment of the instrument, where it is focused through a HeNe laser filter onto the InGaAs detector. The resulting signal is amplified and recorded together with the internal laser reference.

In a post processing step the spectra are calculated via a fast Fourier transform (FFT) routine by the instrument operating software OPUS (by Bruker). After changing the measurement routine in 2015 to a semi-automated set-up, less intervention from the operator is required. At the same time, the interferograms are read directly from the instrument, resulting in raw data slices that are processed to spectra via the i2s program shipped with the GGG2014 software suite used within TCCON.
All interferograms have been transformed using Boxcar apodization and the retrieval code adjusts for the resulting sinc-shaped distortion of the spectral lines. Using $i_2$s, the DC interferograms have been corrected for brightness fluctuations. However, the effect of the correction is expected to be minimal; because of the low resolution, thin cirrus clouds for example typically lead to brightness fluctuations between consecutive scans and to a lesser degree to fluctuations within one interferogram record.

The differences between the solar and lunar measurements include the detector, the spectral resolution, the integration time and the size of the entrance aperture. Decreasing the resolution leads to a shorter measurement time and therefore allows for integration of more interferograms in the same time frame. Increasing the entrance aperture allows for more incident light on the detector, which increases the signal-to-noise ratio. The impact of spectral resolution is further discussed in Sect. 3.4.

At full moon, the entrance aperture was set to 3.15 mm. Occasionally, a smaller entrance aperture is required, because if the moon is not full, its image on the aperture wheel requires a smaller aperture to still ensure that the aperture is uniformly lit. Additionally, the four-quadrant diode used in the tracking system sometimes has difficulty centring the non-full lunar image; using a smaller aperture in this case, again, ensures full illumination of the entrance aperture.

In the TCCON the small entrance aperture samples the centre of the solar disc and the corresponding solar lines are narrow. Sunlight reflected at the lunar surface will have a (solar) disc-averaged spectrum; i.e. the solar lines will be broadened as a result of the different Doppler shifted contributions from different parts of the solar disc. GFTI includes a setting that switches to a calculation of a disc-averaged spectrum when the moon is selected as the source. This approach leads to well-captured solar lines in the spectral fit residuum (see Fig. 2) and therefore indicates the absence of a bias from using different solar line shapes, as is to be expected. A potentially introduced bias would be within the limits of the total bias to the solar measurements (e.g. $0.66 \pm 4.56$ ppm for $xCO_2$ and $-1.94 \pm 20.63$ ppb for $xCH_4$) as discussed in Sect. 4.

### 3.2 Calculation of dry-air mole fractions

For this analysis the current TCCON standard processing code GGG2014 was used for both solar and lunar retrievals. The retrieval code returns vertical columns ($V_{C_{gas}}$), which have to be converted to dry-air mole fractions. There are two possibilities to do this. The standard TCCON processing uses the simultaneously retrieved vertical $O_2$ column to scale the target gas’ vertical column via

$$x_{Gas} = \frac{V_{C_{gas}}}{V_{O_2}}$$

The dry-air mole fraction of $O_2$ is well known and assumed constant; therefore systematic errors common to both vertical column retrievals cancel out using this approach.

However, for the retrieval of $O_2$ the spectral band at 1.27$\mu$m (7880 cm$^{-1}$) is used and the detector is much less sensitive in that region compared to the $CO_2$ and $CH_4$ windows between 5800 and 6400 cm$^{-1}$ (compare Fig. 1). This results in a noisier $O_2$ retrieval especially under low signal-to-noise conditions (see Fig. 2).

The second option to calculate the dry-air mole fraction involves the scaling to atmospheric surface pressure and a correction for the water contained in the column:

$$x_{Gas} = \frac{V_{C_{gas}}}{\frac{m_{H_2O}}{mol} \cdot \frac{m_{H_2O}}{mol}} - \frac{V_{CH_2O}}{\frac{m_{H_2O}}{mol}}$$

www.atmos-meas-tech.net/10/2397/2017/  
Here, xGas denotes the target species’ dry-air mole fraction, VC_gas the vertical column and p_0 the surface pressure. N_A is Avogadro’s number and the molecular masses of water, m_{H_2O} = 18.01534 g mol^{-1}, and dry air, m_{air} = 28.9644 g mol^{-1}, are given. \bar{g} denotes the column-averaged gravitational acceleration at the measurement site and is assumed to be \bar{g} = 9.81 m s^{-2}.

This approach requires accurate knowledge of the surface pressure p_0. Additionally systematic errors, e.g. pointing errors can affect the retrieval, as they are not cancelled out via ratio with O_2. The surface pressure measurement is performed at the Ny-Ålesund station of the Baseline Surface Radiation Network (BSRN), located adjacent to the AWIPEV observatory and thus the FTS. The raw pressure measurements are then scaled to compensate for the height difference to the FTS. The meteorological data are provided by AWIPEV and publicly available at https://doi.org/10.1594/PANGAEA.150000 for years until 2013, with corresponding updates for more recent years.

In the following, the approach described in Eq. (1) was used to retrieve xCO_2 and xCH_4. The second approach, in Eq. (2), was only used to derive xO_2 in Sect. 4, which covers the validation with solar measurements. The main retrieval windows and the fit residuals of an example spectrum are shown in Fig. 2. The vertical column of H_2O used for the water correction in Eq. (2) is retrieved simultaneously in several micro-windows in the same spectral region as the target species.

3.3 Atmospheric model

Information on the target gas is retrieved from the processed spectra by the least-square fitting algorithm GFIT (see Sect. 3.2). The software assumes an a priori profile of the target gas and calculates an artificial spectrum given additional information on the atmospheric profile. In TCCON the interpolation of the NCEP/NCAR reanalysis data (NCEPNCAR, 2016) to the sites latitude, longitude and local noon is used as an atmospheric model, resulting in one model profile per day. The NCEP/NCAR reanalysis data are publicly available and was provided via http://www.esrl.noaa.gov/psd/ (NCEP-NCAR, 2016). In case of lunar measurements, this presents a potential problem around midnight, as consecutive measurements would use different atmospheric models, i.e. the one interpolated to local noon.

Given that the reanalysis data are available in 6 h time intervals, we use the model profile interpolated to the site coordinates and the time of measurement, resulting in specific model profiles for each measurement. These profiles presumably better reflect the atmospheric conditions, especially at night. The increased computational effort for this per-spectrum-model approach is affordable for this comparatively small time series.

A comparison of the differences in retrieved xCO_2 and xCH_4 between the daily and spectrum-specific model profiles is shown in Fig. 3 for the lunar time series and for selected days in the TCCON time series in Fig. 4. The two retrievals show minimal differences at local noon (as they should), but differences of about ±0.5 ppm (CO_2) and about ±2 ppb (CH_4) can occur later in the day, under quickly varying atmospheric conditions distant in time from local noon. Note that the measurements showing potentially large deviations are typically filtered out within TCCON as they occur at high solar zenith angles.
3.4 Analysis of optimal resolution

The resolution used in the TCCON is better than 0.02 cm$^{-1}$, corresponding to a maximum optical path difference (OPD) of 45 cm. Initial tests showed that even with the cooled detector, the spectral signal-to-noise ratio did not allow for a robust retrieval unless a lot of spectra were averaged; however, the path of moonlight through the atmosphere changes rapidly with time. Although this is more prominent in lower latitudes, it still must be considered here, especially at large lunar zenith angles. To avoid bias from inaccurate knowledge of the viewing geometry, the integration time per measurement must be as small as possible.

One option to decrease the measurement time is to increase the velocity of the instrument’s scanning mirror; however, this has no effect on the spectral signal-to-noise ratio. The scanner velocity was therefore not changed and kept at 10 kHz to minimize potential differences from the solar absorption measurements. The second option is to decrease the spectral resolution, which increases the spectral signal-to-noise ratio. Additionally, it allows for shorter measurement times and thus for more spectra to be averaged within the same time, resulting again in an increased signal-to-noise ratio.

The influence of resolution on the retrieval can be analysed in further detail and to circumvent differences arising from a varying atmospheric state. Previously, Petri et al. (2012) investigated this for the TCCON standard retrieval windows. Here the analysis was repeated with emphasis on lower resolutions (down to 1.0 cm$^{-1}$) and additionally spectra with different signal-to-noise ratios were used.

A set of 60 consecutive solar spectra has been selected and the interferograms cropped at lengths corresponding to a range of maximum optical path differences between 45 cm (0.02 cm$^{-1}$) and 0.9 cm (1.0 cm$^{-1}$). The interferograms were...
In addition to this series of spectra, different magnitudes of white noise were added to the created spectra to simulate the effect of the lower signal-to-noise ratio expected in lunar spectra. The signal-to-noise ratios are calculated from the reprocessed spectra by dividing the maximum mean signal between absorption lines at about 6000 cm\(^{-1}\) by the root mean square of a blacked-out region of the spectrum. Figure 5 shows the results of the standard retrieval of \(xCO_2\) and \(xCH_4\) for the various combinations of resolution and signal-to-noise ratio of the series.

The decrease in resolution leads to an increase in S/N. Fig. 6 shows the increase in S/N measured as a function of spectral resolution. The improvement due to lower resolution (blue line) and averaging over larger number of spectra in the same time frame (red line) and the resulting relative S/N from both effects (black line). For better visibility, Fig. 7 shows a subset of the data from Fig. 5, showing the mean retrieved \(xCO_2\) and \(xCH_4\) DMFs at a given resolution. Two series have been selected, with high (red) and low (black) signal-to-noise ratios. The associated errors can be estimated by the standard deviation (1\(\sigma\)) of the arithmetic mean and do not change much with resolution for a given S/N. The mean errors and their standard deviation for \(xCO_2\) are 4.0±0.6 ppm for the low S/N case (black dots in Fig. 7) compared to 0.6±0.05 ppm for the high S/N case (red dots). Similarly the errors for \(CH_4\) are 18.5±3.2 ppb (low S/N, black dots) and 2.9±0.3 ppb (high S/N, red dots).

A distinct cut-off above 0.67 cm\(^{-1}\) can be identified in the \(xCO_2\). For higher resolutions, i.e. 0.02–0.67 cm\(^{-1}\), no significant difference is visible in high signal-to-noise conditions. In general, a lower signal-to-noise ratio of the spectra leads to increased scatter of the retrieved DMFs but to no significant bias. Table 1 shows the bias in the retrieved DMFs of high and low signal-to-noise ratio spectra for the two resolutions used in the measurement set-up later.

Gisi et al. (2012) showed that lower-resolution solar spectra can be used to retrieve DMFs with a low-resolution FTS (Bruker EM27/SUN). Recently Hedelius et al. (2016) investigated errors and biases from a 0.5 cm\(^{-1}\) FTS (Bruker EM27) for TCCON relevant species. The three studies (Petri et al., 2012; Gisi et al., 2012; Hedelius et al., 2016) report different biases in \(xCO_2\) when changing the resolu-
Figure 7. Mean of the retrieved xCO₂ and xCH₄ from cropped interferograms at different resolutions with low and high signal-to-noise ratio (S/N). Shown is the relative difference to the highest signal-to-noise ratio and highest resolution.

Table 1. Comparison of the biases, introduced by lower-resolution measurements and low signal-to-noise ratio (S/N). Subset of data points from Fig. 7.

<table>
<thead>
<tr>
<th>S/N</th>
<th>Resolution (cm⁻¹)</th>
<th>ΔxCO₂ (%)</th>
<th>ΔxCH₄ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt; 300</td>
<td>0.08</td>
<td>0.03 ± 0.57</td>
<td>0.28 ± 2.61</td>
</tr>
<tr>
<td></td>
<td>0.5</td>
<td>0.07 ± 0.65</td>
<td>0.76 ± 3.03</td>
</tr>
<tr>
<td>≈ 30</td>
<td>0.08</td>
<td>-0.13 ± 4.12</td>
<td>0.00 ± 15.03</td>
</tr>
<tr>
<td></td>
<td>0.5</td>
<td>-0.20 ± 4.50</td>
<td>0.79 ± 22.89</td>
</tr>
</tbody>
</table>

The effect of different resolutions on the retrieved columns can also be investigated by comparing different measurements taken consecutively with different resolutions. Figure 8 shows lunar absorption measurements of the target species on 7 October 2014. The first and third batch of measurements were taken with a resolution of 0.085 cm⁻¹ (OPD = 10.59 cm), while the second batch was measured with 0.5 cm⁻¹ (OPD = 1.8 cm) resolution. No significant bias is observed.

Decreasing the spectral resolution also changes the information content of the recorded spectral lines. This results in a change in shape of the measurements averaging kernels and is discussed below.

3.5 Averaging kernels

The sensitivity of the retrieved dry-air mole fraction of the target gas depends on the a priori information and the measurement’s altitude dependent sensitivity, i.e. the averaging kernels. The a priori profiles used are the default TCCON ones. The averaging kernel of a measurement strongly depends on the retrieval methodology and the information content of the corresponding spectrum. As such it depends on the viewing geometry as well as the resolution, the absorption strength and the signal-to-noise ratio. The weight different altitude levels have in the retrieval can be parameterized as a function of the zenith angle. As the instrument faces the light source at a certain zenith angle, the measurement samples different contributions from the various atmospheric layers. The pressure broadening of the absorption features shows a specific altitude dependent sensitivity and this information depends on the chosen resolution and the signal-to-noise ratio of the measurement.
The set-up of the lunar measurements is similar to that of TCCON measurements, and therefore the averaging kernels are quite similar, aside from effects of resolution and noise for a given zenith angle.

The top panel in Fig. 9 shows the averaging kernels for the lunar measurements. The middle panel shows the difference from the standard TCCON ones from Ny-Ålesund, interpolated to the corresponding zenith angles. The lines are colour gradient coded with their respective zenith angles and different colour schemes reflect different resolutions.

Pressure broadening leads to spectral lines originating from gases at low pressure being narrower than those at higher pressure. The narrow part of a spectral line sampled with fewer points therefore cannot give as much information as one with higher resolution. This leads to averaging kernels from low-resolution spectra being less sensitive to the stratosphere and more sensitive in the lower troposphere than their high-resolution counterparts. This can be seen in the lower panel of Fig. 9, where the difference between standard TCCON averaging kernels and their lower-resolution counterparts at the same zenith angle is shown. As expected, decreasing the spectral resolution leads to greater differences between the averaging kernels.

4 Validation with solar absorption spectroscopy

The validation of the measurements performed during the polar night is difficult. In the absence of other options, here we compare to solar absorption measurements taken within TCCON. In spring and autumn there are a few consecutive days around the full moon when solar absorption measurements during the day and lunar absorption measurements during the night are possible. Such comparison measurements were performed in March and September 2013. Here the DMFs of xCO₂ and xCH₄ for both solar and lunar measurements were retrieved using Eq. (1). For the comparison of xO₂ Eq. (2) was used.

Assuming the total column values do not change significantly during that time period, the means of the two retrievals can be compared directly. Figure 10 shows the comparison results and the calculated means for a comparison in September 2013. Table 2 shows the corresponding values of the arithmetic mean and its standard deviation as an indication of the error for both comparison campaigns in March and September 2013. The same analysis was performed on the available smoothed model output. The calculated standard deviations of the models of about 0.2 ppm and 0.3 ppm for CO₂ and 1.0 and 1.6 ppb for CH₄ for March and September, respectively, indicate that the assumption of stable DMFs for the observed time frame is reasonable.

The accuracy of the lunar measurements can be determined via the bias of the lunar compared to the solar measurements and can be deduced from Table 2 as well. In March 2013 the difference between solar and lunar measurements is 0.66 ± 4.56 ppm for xCO₂ and −1.94 ± 20.63 ppb for xCH₄. In the September 2013 campaign a bias of 1.01 ± 8.52 ppm for xCO₂ and −3.36 ± 41.13 ppb for xCH₄ can be observed. The diurnal variability of the lunar measurements is used to define the precision. As the later measurements have a higher precision, a typical value achieved in the 2014–2015 winter is used. Here the standard deviations of the daily mean of 2 ppm for xCO₂ and 10 ppb for (xCH₄), corresponding to 0.5 % in both cases.

The target accuracy can be estimated via the detrended year-to-year wintertime variability. Here model output can be used as a proxy. In the smoothed, detrended MACC CO₂ and CH₄ model (see Sect. 5.1) the arithmetic mean of the first week of January differs by 0.55 ppm in xCO₂ and 9.84 ppb in xCH₄ between 2012 and 2014. At the same time, the stan-

Figure 8. Comparison of retrieved xCO₂ and xCH₄ for different resolutions from low (OPD = 1.8 cm⁻¹, black) and higher (OPD = 10.59 cm⁻¹, red) resolution measurements on 7 October 2014.
The standard deviation of all values for the first week of January between 2012 and 2014 is about 1.8 ppm for xCO₂ and 18.8 ppb for xCH₄. However, these estimates are potentially subject to unknown biases in the models, i.e. the model could be biased similarly every year. Additionally, the seasonal variability surely is an upper limit for the target precision. Here the seasonal cycle amplitude measured by solar FTS is about 15 ppm for xCO₂ and about 40 ppb for xCH₄.

As described in Sect. 3.2 (see Eq. 1), the dry-air column is calculated using the vertical column of O₂, retrieved from the 7885 cm⁻¹ spectral region. Here airglow emissions in the high atmosphere could potentially disturb the O₂ spectra.
Figure 10. Comparison of the solar and lunar measurements of xCO₂ and xCH₄ in September 2013 (dots) and the corresponding arithmetic means (lines). Values are given in Table 2.

Table 2. Comparison of the retrieved solar, lunar and model DMFs for the two comparison time periods. Note that xO₂ was calculated using the surface pressure and the offset to the true atmospheric value of 20.95 % is caused by spectroscopic errors.

<table>
<thead>
<tr>
<th></th>
<th>xCO₂ (ppm)</th>
<th>xCH₄ (ppb)</th>
<th>xO₂ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>March 2013</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solar</td>
<td>397.47 ± 0.67</td>
<td>1773.78 ± 2.99</td>
<td>21.33 ± 0.08</td>
</tr>
<tr>
<td>Lunar</td>
<td>396.81 ± 3.89</td>
<td>1775.72 ± 17.64</td>
<td>21.34 ± 0.36</td>
</tr>
<tr>
<td>Jena CO₂</td>
<td>398.01 ± 0.13</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>CT15 CO₂</td>
<td>396.89 ± 0.22</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MACC CO₂</td>
<td>397.16 ± 0.18</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MACC CH₄</td>
<td>–</td>
<td>1784.09 ± 1.06</td>
<td>–</td>
</tr>
<tr>
<td>September 2013</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solar</td>
<td>393.16 ± 0.49</td>
<td>1810.26 ± 3.11</td>
<td>21.38 ± 0.06</td>
</tr>
<tr>
<td>Lunar</td>
<td>392.15 ± 8.03</td>
<td>1813.62 ± 38.02</td>
<td>21.40 ± 0.60</td>
</tr>
<tr>
<td>Jena CO₂</td>
<td>391.56 ± 0.26</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>CT15 CO₂</td>
<td>391.29 ± 0.24</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MACC CO₂</td>
<td>392.07 ± 0.39</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MACC CH₄</td>
<td>–</td>
<td>1800.79 ± 1.58</td>
<td>–</td>
</tr>
</tbody>
</table>

This can typically be ignored in solar absorption spectra, as the magnitude of the emissions is negligible, when viewing directly into the sun. In case of lunar spectra, however, air-glow emissions could potentially fill in the spectral lines and influence the measurements. To test this, xO₂ was retrieved using the surface pressure to calculate the dry-air column as described in Eq. (2).

In both comparison periods, no significant difference between the solar and lunar retrievals of xO₂ can be observed. Note that xO₂ retrieved via surface pressure shows an offset of 0.4 % in both cases (lunar and solar). This offset originates in the line parameters used for the O₂ retrieval and is compensated in the xCO₂ and xCH₄ retrieval with the TC-CON in situ correction. Washenfelder et al. (2006) reported values that are 2.27 ± 0.25 % larger if the surface pressure retrieved dry column was used. Here we find a mean difference of 1.96 ± 0.14 % when calculating the mean and standard deviation of the solar and lunar mean xO₂ values shown in the sidebars in Fig. 11. Note that these retrievals were performed with updated spectroscopy available within GGG2014 compared to that used by Washenfelder et al. (2006).

5 Seasonal cycle and model comparison

5.1 Method – model comparison

The rigorous comparison of ground-based column measurements of a trace gas to model simulations requires resampling the model profile as if it was measured by the instrument.

The smoothed column dry-air mole fraction \(\hat{c}\) can be calculated following Rodgers and Connor (2003), Connor et al. (2008) and Wunch et al. (2010) by adding the column-integrated a priori profile \(c_a\) to the difference between the
model (x) and the dry TCCON a priori profile (x_a) weighted with the averaging kernel (a):
\[
\hat{c} = c_a + h^T a^T (x - x_a).
\]  
(3)

Here, h represents the pressure weighting function (see Connor et al., 2008).

Given a vertical model profile, the measurement’s averaging kernel and the vertical columns of water vapour and the a priori profile of the target gas, the smoothed dry-air mole fraction of the model output can be calculated. Due to the high random error of the lunar FTS measurements, daily means have been calculated for both the measurements and the model data, after the smoothing was applied.

5.2 Results – time series

In this section the FTIR time series is compared to CO₂ model results from three different models: the MACC CO₂ model version 14r2 (MACCCO2, 2015), the CarbonTracker 2015 (CT2015, 2016) model and the Jena CO₂ inversion version s04_v3.7 (JenaCO2, 2005). In case of the CH₄ time series, the MACC CH₄ v10 (MACCCH4, 2015) is used. As described in Sect. 5.1 the model’s DMF profile has been smoothed with the corresponding a priori and averaging kernel of the lunar and solar measurement, respectively. For times when there are no FTS measurements available, an averaging kernel was calculated using the solar zenith angle of the corresponding time. In winter the lunar zenith angle was used instead. For times where no FTS measurements were possible at all, e.g. sun and moon are below the horizon, a mean zenith angle of 65° was assumed.

The resulting model time series can now be compared directly to the FTS measurements. Figure 12 shows the comparison of the FTS and the smoothed model time series for CO₂. The CH₄ comparison is shown in Fig. 13.

5.3 Results – seasonal cycle

The detrended seasonal cycles of both target species are similar from year to year. In the following, the detrended seasonal cycles are compared to the models already discussed in Sect. 5.2.

Figure 14 shows the seasonal cycle of xCO₂ as observed with the Ny-Ålesund FTS between 2012 and 2016, detrended with a linear increase of 2.6 ppm yr⁻¹, an offset of 380.0 ppm on 1 January 2012 and condensed to 1 year. The seasonal cycle of xCO₂ shows little difference between the three models, and therefore the comparison can be performed with an model average. The shaded area in Fig. 14 shows the 3σ standard deviation around the daily mean of the combined model data points of all three models (MACC, CarbonTracker and Jena). The weighted average of all FTS measurements during one full moon period is shown (green dots) with error bars corresponding to the standard error (σ/√N) of the daily mean calculated from N measurements. The weights are chosen to be the inverse squared residual of the spectral fits.

The difference between the models and the TCCON measurements in summer is quite small, except for a phase shift in the onset of the downward slope at the beginning of the growing season decline. In winter the models agree well with the FTIR lunar absorption measurements, within the given error margin.

In the case of CH₄ a similar comparison has been performed and the results can be seen in Fig. 15. Here the xCH₄
time series have been linearly detrended with an annual increase of 10.6 ppb yr$^{-1}$ and an offset of 1760.0 ppb on 1 January 2012. Figure 15 shows the 3σ standard deviation around the daily means of the MACC CH$_4$ model (shaded area) compared to the FTS measurements (red and blue dots) averaged over one full moon measurement cycle. The error bars correspond to the 1σ standard deviation of the mean.

In spring/summer the FTS measurements show generally smaller values than the model and a larger spread. From late summer throughout the winter the measurements are in better agreement with the model. At specific events in spring, the FTS measurements show sudden decreases of xCH$_4$ (compare Figs. 13 and 15). This could be due to the model not being able to capture vertical transport very well, which has

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**Figure 12.** Comparison of the daily means of lunar (blue) and solar (red) xCO$_2$ FTIR measurements to the AK-smoothed MACC CO$_2$ model v14r2 (top panel, grey). Error bars show the standard error ($\sigma/\sqrt{N}$, with $N$ number of measurements). The lower panels show the model–measurement difference for all models.

**Figure 13.** Comparison of the daily means of lunar (blue) and solar (red) xCH$_4$ FTIR measurements to the AK-smoothed MACC CH$_4$ model v10 (grey). Error bars show the standard error ($\sigma/\sqrt{N}$, with $N$ number of measurements). The lower panel shows the model–measurement difference.
Figure 14. Comparison of solar (red) and lunar (blue) xCO₂ FTIR measurements. Error bars show 1σ standard deviation of the daily mean. The lunar data points have been averaged over one full moon period each. The shaded grey area shows the 1σ standard deviation of the three model daily means (MACC, CarbonTracker and Jena) as shown in Fig. 12.

Figure 15. Comparison of solar (red) and lunar (blue) xCH₄ FTIR measurements. Error bars show 1σ standard deviation of the daily mean. The lunar data points have been averaged over one full moon period each. The shaded grey area shows the 1σ standard deviation of the MACC CH₄ model daily means as shown in Fig. 13.

been shown previously by Ostler et al. (2016). Here, stratospheric intrusions during the breakdown of the polar vortex in spring can lead to large, short-term decreases in xCH₄. This is currently being investigated by using a stratospheric species as a tracer to separate the xCH₄ column in a tropospheric and stratospheric part and exceeds the scope of this paper.

6 Conclusions

Measurements of the column-averaged dry-air mole fractions of CO₂ and CH₄ have been performed in the polar night from 2012 to 2016 to complement the established solar absorption measurements within the TCCON. The newly employed thermoelectrically cooled InGaAs detector allows the usage of reflected sunlight on the full lunar disc to serve as a light source above the atmosphere to perform lunar absorption spectroscopy in the near-infrared spectral region.

Aircraft or AirCore profiles are not yet available for Ny-Ålesund. The lunar absorption measurements have therefore been validated with standard TCCON measurements in spring and autumn 2013 and the comparison shows no significant biases. The decrease of spectral resolution allows for an increase of the spectral signal-to-noise ratio, which in turn decreases the random error significantly. Under optimal conditions, lunar measurements with standard deviation of the daily mean (1σ) of about 2 ppm for xCO₂ and about 10 ppb
for xCH4 can be achieved using this approach. This corresponds to a precision of about 0.5 % for each gas.

The newly created time series has been compared to different model simulations. All three CO2 models (MACC CO2 model v. 14r2, CarbonTracker 2015, Jena CO2 inversion s04_v3.7) are generally in good agreement with the FTIR measurements. The xCH4 time series shows large deviations in spring/summer and an overall good agreement in autumn/winter.

Data availability. The TCCON data are publicly available from the TCCON archive at http://tccon.ucar.edu/. The solar and lunar measurement data used in this study have been uploaded to the Pangaea database and are available at https://doi.pangaea.de/10.1594/PANGAEA.872007 (Buschmann et al., 2017). The MACC model data can be accessed via http://www.gmes-atmosphere.eu/. The Jena CO2 inversion is provided via http://www.bgc-jena.mpg.de/CarboScope and the CarbonTracker data can be found at http://carbontracker.noaa.gov. The NCEP/NCAR reanalysis is provided by NOAA via http://www.esrl.noaa.gov/psd/. The surface meteorology data from the Ny-Ålesund BSRN station used here are available from https://doi.org/10.1594/PANGAEA.150000.

Author contributions. The measurements where taken by MB, the co-authors and the AWIPEV station staff. MB performed the analysis and prepared the manuscript with contributions from all co-authors.

Competing interests. The authors declare that they have no conflict of interest.

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