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Published in:
Atmospheric Measurement Techniques

DOI:
10.5194/amt-9-3687-2016

Publication date:
2016

Document Version
Publisher’s PDF, also known as Version of record

Citation for published version (APA):
A new set-up for simultaneous high-precision measurements of CO$_2$, $\delta^{13}$C-CO$_2$ and $\delta^{18}$O-CO$_2$ on small ice core samples

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Received: 13 November 2015 – Published in Atmos. Meas. Tech. Discuss.: 1 February 2016
Revised: 16 May 2016 – Accepted: 21 June 2016 – Published: 10 August 2016

Abstract. Palaeoatmospheric records of carbon dioxide and its stable carbon isotope composition ($\delta^{13}$C) obtained from polar ice cores provide important constraints on the natural variability of the carbon cycle. However, the measurements are both analytically challenging and time-consuming; thus only data exist from a limited number of sampling sites and time periods. Additional analytical resources with high analytical precision and throughput are thus desirable to extend the existing datasets. Moreover, consistent measurements derived by independent laboratories and a variety of analytical systems help to further increase confidence in the global CO$_2$ palaeo-reconstructions. Here, we describe our new set-up for simultaneous measurements of atmospheric CO$_2$ mixing ratios and atmospheric $\delta^{13}$C and $\delta^{18}$O-CO$_2$ in air extracted from ice core samples. The centrepiece of the system is a newly designed needle cracker for the mechanical release of air entrapped in ice core samples of 8–13 g operated at $-45$ °C. The small sample size allows for high resolution and replicate sampling schemes. In our method, CO$_2$ is cryogenically and chromatographically separated from the bulk air and its isotopic composition subsequently determined by continuous flow isotope ratio mass spectrometry (IRMS). In combination with thermal conductivity measurement of the bulk air, the CO$_2$ mixing ratio is calculated. The analytical precision determined from standard air sample measurements over ice is ±1.9 ppm for CO$_2$ and ±0.09 ‰ for $\delta^{13}$C. This number is good, though they are rather conservative estimates of the overall analytical precision achieved for single ice sample measurements. Facilitated by the small sample requirement, replicate measurements are feasible, allowing the method precision to be improved potentially. Further, new analytical approaches are introduced for the accurate correction of the procedural blank and for a consistent detection of measurement outliers, which is based on $\delta^{18}$O-CO$_2$ and the exchange of oxygen between CO$_2$ and the surrounding ice (H$_2$O).

1 Introduction

Polar ice cores are unique in providing direct information of the past atmospheric composition. Analysis of entrapped air allows the evolution of the atmospheric composition over the last 800 000 years to be reconstructed (e.g. Lüthi et al., 2008 and references therein; Bereiter et al., 2015). Knowledge of past natural CO$_2$ variations – only several ppm during the Holocene and up to about 100 ppm over glacial/interglacial changes – is crucial to improve predictions of future climate under continued anthropogenic CO$_2$ forcing. Changes in the global carbon cycle fluxes are imprinted in the stable carbon isotope signal of atmospheric CO$_2$ ($\delta^{13}$C, e.g. Köhler et al., 2006). However, carbon isotope measurements of ice core air samples are highly demanding and time-consuming. As a result, detailed measurements of $\delta^{13}$C are still limited to specific time periods (Francey et al., 1999; Indermühle et al.,
Alternatively, ice samples are ground in a ball mill when both the sample surface by a centrifugal ice microtome (CIM). Al- al., 1982). The system described by Bereiter et al. (2013) pul- arrived by sublimation systems.

and no difference has been observed compared to results de- Extraction efficiencies usually decrease by 10–20 %. For the transition zone, where air bub- be observed for \( \delta ^{13} C \) analysis of \( \text{CO}_2 \) (Schaefer et al., 2011 and references therein). In addition, for pure bubbly ice, the extraction efficiency is not a concern for \( \text{CO}_2 \) measurements and no difference has been observed compared to results derived by sublimation systems.

A variety of mechanical extraction systems are in use. In a needle cracker (NC), the ice is crushed to small pieces and air is released from the thereby opened bubbles (Zumbrunn et al., 1982). The system described by Bereiter et al. (2013) pulverizes ice samples by continuously shaving off thin layers of the sample surface by a centrifugal ice microtome (CIM). Alternatively, ice samples are ground in a ball mill when both the ice sample and stainless steel balls inside a small container are shaken (Barnola et al., 1995; Lourantou, 2009), or grated into small chips by shaking the ice in a vessel containing a perforated inner cylinder (“cheese grater”, Etheridge et al., 1996).

Only a few laboratories have the ability to do ice core analysis of both \( \text{CO}_2 \) concentrations and its stable isotopic composition. In the following, the published and recently operated analytical systems allowing measurements of both parameters on a single ice sample are summarized (see Table 1 for detailed system characteristics). All systems use isotopic ratio mass spectrometry (IRMS) to detect the different mass ratios between the stable \( \text{CO}_2 \) isotopologues (\( m/z \) 44, 45 and 46).

The Laboratory of Climate and Environmental Physics (KUP, Bern, Switzerland) operates two such systems. For the mechanical extraction system (NC) the released air is first expanded over a water trap into a small volume where the gas pressure is measured for evaluation of the \( \text{CO}_2 \) mixing ratio in combination with the IRMS signal. Using helium as a carrier gas, the gas sample is then flushed into a pre-concentration system (PreCon) to separate the main components of air. In order to avoid isobaric interference, \( \text{CO}_2 \) is separated from \( \text{N}_2 \text{O} \) and organic compounds (e.g. from drilling fluids) by gas chromatography (GC) before being injected into the IRMS via an open-split interface (Elsig et al., 2009). In the sublimation system, sublimated water is quantitatively removed before the liberated air is cryogenically collected. Then, the basic principle is similar to the system described before but extraction and GC–IRMS are decoupled (Schmitt et al., 2011). KUP additionally operates one system (CIM) solely dedicated to the analysis of the \( \text{CO}_2 \) mixing ratio (Bereiter et al., 2013), which replaces their initial NC system described by Zumbrunn et al. (1982) and modified by Lüthi (2009).

The Laboratoire de Glaciologie et Géophysique de l’ Environnement (LGGE, Grenoble, France) uses a ball mill for mechanical extraction before the air is directly released to the inlet system of a coupled GC–IRMS for \( \text{CO}_2 \) mixing ratio and \( \text{CO}_2 \) stable isotope analysis (Barnola et al., 1995; Lourantou, 2009).

The ice core and quaternary geochemistry lab at Oregon State University (OSU, USA) also uses a mechanical extraction system (cheese grater). A small aliquot of the extracted sample gas is isolated from the grater and finally trapped at \(-260^\circ \text{C}\) after water is removed at \(-100^\circ \text{C}\). The \( \text{CO}_2 \) mixing ratio is then determined by GC. The rest of the gas, again first passing a water trap at \(-100^\circ \text{C}\), is condensed in a second trap at \(-190^\circ \text{C}\) and finally analysed for \( \delta ^{13} \text{C} \) by IRMS dual-inlet measurement, applying a correction for the isobaric \( \text{N}_2 \text{O} \) interference. Interference from drilling fluid contamination can potentially be a problem for certain samples. The rather large sample size allows measurement of \( \text{N}_2 \text{O} \) in addition (Bauska et al., 2014). OSU also operates a NC system for the analysis of \( \text{CO}_2 \) only (Ahn et al., 2009).
CSIRO uses a cheese grater for mechanical extraction (Etheridge et al., 1996; MacFarling Meure et al., 2006; Rubino et al., 2013). The released air is cryogenically collected in an external trap (around $-260^\circ$C) after removing water at $-100^\circ$C. Subsequently, the sample is analysed using GC for determination of the CO$_2$ mixing ratio and by IRMS for $\delta^{13}$C without further GC separation and purification. A correction for the isobaric N$_2$O interference is applied. Interference from drilling fluid contamination can potentially be a problem for certain samples. The large sample size allows measurement of other trace gases from the same sample (CH$_4$, CO, and N$_2$O).

In this study, we present a new system built at the Centre for Ice and Climate (CIC, University of Copenhagen, Denmark) in the laboratory for atmospheric trace gas measurements in ice cores (Stowasser et al., 2012; Spelrich et al., 2013). The approach was to opt for small sample size, to allow simultaneous analysis of both CO$_2$ mixing ratios and its stable isotopic composition in the same sample, and to achieve high precision with reasonable throughput in order to pursue high resolution sampling schemes. We thereby followed the extraction principle of the NC using a modified design. Due to the intended small sample size, the extraction unit was coupled to a continuous flow GC–IRMS set-up, with the benefit of overcoming the problem of interference from isobaric N$_2$O and fragments of remaining contamination from drilling fluid.

## 2 Instrumental set-up and standards

### 2.1 Dry extraction unit

The dry extraction unit was designed based on the NC principle for small sample sizes of a few grams described by e.g. Lüthi (2009) and Ahn et al. (2009). However, some major modifications were implemented to achieve the following goals: (i) avoidance of mechanical friction within the system in order to reduce related contamination and adsorption/desorption effects (Zumbrunn et al., 1982; Stauffer et al., 1985; Lüthi, 2009); (ii) operation at very low temperatures to reduce the risk of CO$_2$ in situ production (within the extraction unit) from chemical reactions due to the presence of H$_2$O in the mobile phase; (iii) fast and simplified sample loading with minimal exposure of inner surfaces to ambient air in order to maximize sample throughput and to reduce artefacts from surface effects, respectively.

Our NC design for ice samples with maximum dimensions of $2.3 \times 2.5 \times 2.5$ cm$^3$ and a typical mass of 8–13 g is shown in Fig. 1. All inner parts are made from stainless steel (SS). Similar to Ahn et al. (2009), we use a compressible welded bellow (SS, Comvat, Germany, 5 in Fig. 1). This allows crushing of the ice by axial movement of the needles mounted with a hot/cold press fit (hardened SS, 1.5 mm OD, 30 mm length, Dema, Germany, 6 in Fig. 1). In comparison to the design described by Lüthi (2009), which requires a vacuum tight seal around a movable piston, the mechanical friction within our unit is thus strongly reduced. In addition, the bellow is mounted differently than in the design presented by Ahn et al. (2009), resulting in an inner volume of half the size ($\sim 110$ cm$^3$; $\sim 63$ cm$^3$ with the bellow compressed) and an inner surface area reduced by about two-thirds. A small volume is favourable in terms of evacuation speed and time required for transferring the gas out of the extraction unit for subsequent treatment. A small inner surface reduces the potential for surface effects (adsorption/desorption) which can bias the CO$_2$ stable isotope ratios due to isotopic fractionation processes. The extraction unit is connected to the rest of the set-up by 1/4 in. SS tubes welded to the NC and equipped with valves (SS–4H, Swagelok, USA, 11, 12 and 13).

<table>
<thead>
<tr>
<th>Laboratory</th>
<th>Extraction principle (design)</th>
<th>Operating temp. (°C)</th>
<th>Sample mass (g)</th>
<th>Extraction efficiency for bubbly (clathrate) ice</th>
<th>Daily sample throughput</th>
<th>Precision CO$_2$ (ppm)</th>
<th>Precision $\delta^{13}$C (‰)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KUP</td>
<td>mechanical (cheese grater)</td>
<td>$-20$ ($-35)^*$</td>
<td>5–6</td>
<td>$\sim 70$ ($\sim 50$) %</td>
<td>3–6</td>
<td>2.0</td>
<td>0.07</td>
</tr>
<tr>
<td>LGGE</td>
<td>sublimation</td>
<td>$-65$</td>
<td>40–50</td>
<td>$\sim 100$ ($\sim 100$) %</td>
<td>1–2</td>
<td>$\sim 2.0$</td>
<td>0.05</td>
</tr>
<tr>
<td>OSU</td>
<td>mechanical (ball mill)</td>
<td>$-60$</td>
<td>400–550</td>
<td>$\sim 60$ ($&lt;60$) %</td>
<td>1–2</td>
<td>1.9</td>
<td>0.02</td>
</tr>
<tr>
<td>CSIRO</td>
<td>mechanical (cheese grater)</td>
<td>$-20$</td>
<td>800–1000</td>
<td>$60–80$ (unknown) %</td>
<td>3–4</td>
<td>1.0</td>
<td>0.04</td>
</tr>
<tr>
<td>CIC (this study)</td>
<td>mechanical (NC)</td>
<td>$-45$</td>
<td>8–13</td>
<td>$70–80$ ($\sim 60$) %</td>
<td>3–4</td>
<td>2.0</td>
<td>0.11</td>
</tr>
</tbody>
</table>

* Initially reported at $-20^\circ$C, but lowered to $-35^\circ$C since (Leuenberger, 2009).

Table 1. Characteristics of published and recently operated analytical systems allowing measurement of CO$_2$ mixing ratios and the CO$_2$ stable isotopic composition on a single ice sample. Indicated precisions (1σ) are estimated from replicate analysis of natural ice samples. One should note that the thereby applied metric may not be entirely comparable (e.g. replicates either measured on different or on the same day).
A fixed soft copper seal connects the needle piston to the bellow base plate.

Both the crushing mechanism by axial compression of the bellow and the opening/closing mechanism are pneumatically actuated. To crush the ice sample, a pressure of 4.7 bar is applied to the upper cylinder (CP95SDB40–80, SMC, 1a in Fig. 1) and the needles are actuated via a 5-port solenoid valve (VQZ3120–5YZB–C10, SMC) controlled by an external logical device (homemade). The total number and frequency of strokes can be controlled and were typically set to 37 and ~3 Hz, respectively. Six bars of air pressure applied on the lower actuator (C95NDB80–250, SMC, 1b in Fig. 1) creates enough force for a vacuum tight sealing between the connection of the upper and lower part of the extraction unit using indium wire (1.5 mm OD, 99.99 %, Sigma-Aldrich, USA, 10 in Fig. 1). Although the wire needs to be replaced whenever a new sample is loaded, the indium can be reused when drawn into wire again. This sealing mechanism reduces the amount of time required to open and vacuum seal the vessel compared to systems using bolts and nuts. It takes less than 2 min, including the removal of previously crushed ice, cleaning and reloading of a new sample. To minimize contact of ambient air with the inner surfaces and avoid condensation/deposition of water vapour, both the lower and upper part of the device are flushed through the respective inlets with \( \text{N}_2 \) (99.999 %, Air Liquid, Denmark) whenever the system is opened.

To cool the well-insulated NC, we chose to use an air cooling set-up similar to that described by Schmitt (2006) instead of using a liquid cooling fluid. Pressurized air with an adjustable flow between 0 and ~60 L min\(^{-1}\) is dried in two sequential traps (filled with Molecular sieve 13X/4Å, Supelco, USA) and cooled in a copper heat exchanger mounted in a Dewar (D2). D2 is supplied with droplets of liquid nitrogen (LN) from a larger Dewar (D1) containing the LN reservoir. The droplets are pumped by applying 12.8 V to a heater (10Ω resistor) mounted in the widened inlet of an empty 1/4 in. tube submerged in the LN. Whenever heat is applied, LN evaporates around the heater and the evolving \( \text{N}_2 \) bubbles transport the above, still liquid nitrogen through the isolated tube to D2. This LN pump is regulated by the use of two coupled proportional–integral–derivative controllers (PID, iTRON 08, JUMO, UK). One PID is set to the desired final temperature measured in the NC (9 in Fig. 1), whereas the other is set to a minimum temperature of \(-180^\circ\text{C}\) in D2, preventing the system from eventual clogging by frozen remnant water in the air stream and from potential condensation of oxygen. The temperatures used for PID input and survey of the air stream are measured with platinum resistance thermometer PT100 elements (100Ω, −200 to 600 °C, Class 1/10, TC Direct, USA). By changing the settings for air flow and/or the set points for NC and D2, the air stream temperature is regulated. To cool the NC, the cold air stream is split in front of the unit and either directed through the cavities in the lower part of the massive steel unit (4b in Fig. 1) or the cooling jacket mounted around the compressible welded bellow (4a in Fig. 1). The minimum operating temperature of the NC is \(-55^\circ\text{C}\), whereas the standard operating temperature is set to \(-45^\circ\text{C}\) (stability ±1 °C) with significantly reduced build-up of ice on the vacuum sealing surfaces. While cooling down to \(-45^\circ\text{C}\), the air stream first regulates to about \(-80^\circ\text{C}\) before it stabilizes at \(-60^\circ\text{C}\). Cooling the NC to \(-45^\circ\text{C}\) takes around 70 min.

Compared to other systems allowing analysis of similarly small sample sizes (other NC designs, CIM) the operating temperature of our extraction chamber is significantly lower (\(-45^\circ\text{C}\) compared to around \(-35/-30^\circ\text{C}\)). This is beneficiary because the resulting lower water partial pressure in the extraction chamber (about 5-fold) reduces the risk of in situ \( \text{CO}_2 \) production by wet chemistry. This is supported by the findings of Bauska et al. (2014) which indicated that operating at low temperatures improves precision and decreases the blank of the method. For our design this is reflected in a reduced system offset compared to the KUP NC (Sect. 4.1.1), though most likely resulting from a combined positive effect of friction-reduced motion and lower operating temperature. The low water vapour partial pressure allowed a water trap to be omitted at the exit of the extraction unit.
2.2 Analytical system

Our analytical system allows simultaneous measurements of atmospheric CO₂ mixing ratios and its stable isotopic composition in the same sample. A schematic representation of the system is shown in Fig. 2. It can be divided into two main sections: section A (manually operated) for standard or sample gas loading, and section B (fully automated and in continuous flow mode) for separation, purification and final detection run in continuous flow mode. Highlighted in red and green are components for gas separation and detection, respectively. See main text for details (Sects. 2 and 3).

Section A consists of four main parts:

i. a vacuum line;

ii. a gas manifold for carrier-, protection- or standard-gas injection;

iii. a dry extraction unit (NC);

iv. a trap (T1) to quantitatively cryopump sampled gas out of the NC for subsequent and complete transfer from section A to B.

Section B consists of two main parts:

i. a gas separation part which allows initial trapping of the transferred sample (T2), separation of CO₂ (and N₂O) from the major air components (T3) and subsequent partitioning of the two fractions to individual lines for either final detection (main air fraction) or further purification by gas chromatography (GC–1 and GC–2);

ii. the detection systems including a thermal conductivity detector to quantify the amount of the main air fraction (TCD, VICI, USA, integral part of GC–1, TRACE GC Ultra, Thermo Scientific, USA), a pulsed discharge detector to survey CO₂ separation and purification (PDD, VICI, USA, integral part of GC–1) and an IRMS to quantify amount and isotopic ratios of the CO₂ fraction (Delta V Plus, Thermo Fisher, Germany).

All inner surfaces of the set-up are either made from SS or fused silica and the connections are either welded or sealed with metal or graphite/vespel ferrules to exclude artefacts due to outgassing (Sturm et al., 2004). Section A can either be evacuated or flushed with N₂ (inert gas also used for protection of inner surfaces when the extraction unit is opened) or He (99.9995 %, Air Liquide, Denmark) additionally purified by a getter (Gas Purifier, VICI, Valco Instruments Co. Inc, USA). Similar to the gas manifold, the vacuum line is made from 1/4 in. SS tubes and equipped with on/off valves (SS–4H, Swagelok, USA). It includes a low vacuum (LV) rotary vane pump (EDM2, Edwards, UK) and a turbo pump (TMU 071, Pfeiffer, Germany) to reach high vacuum (HV). The LV pump is used for the fast removal of large quantities of gas (e.g. after sample loading) and is protected from the analytical line by a liquid nitrogen trap (WT). All lines in section B are always above atmospheric pressure and continuously flushed with He used as a carrier gas for the entire system, allowing transfer of sample/standard gas from the extraction unit to the detection systems which are all run in continuous flow mode. Further details will be given in Sect. 3.
Table 2. CIC reference standards for CO₂ mixing and stable isotope ratios.

<table>
<thead>
<tr>
<th>Name</th>
<th>Reference Gas</th>
<th>CO₂ (ppm)</th>
<th>δ¹³C-CO₂ (% VPDB)</th>
<th>δ¹⁸O-CO₂ (% VPDB-CO₂)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Messer–649250¹</td>
<td>GS19/GS20 CO₂</td>
<td>−6.004 ± 0.008</td>
<td>−10.80 ± 0.13</td>
<td></td>
</tr>
<tr>
<td>CA08274²,³</td>
<td>NOAA/CIC Air</td>
<td>181.04 ± 0.06</td>
<td>(−35)²</td>
<td></td>
</tr>
<tr>
<td>CA08054²,⁴</td>
<td>NOAA Air</td>
<td>267.08 ± 0.01</td>
<td>−7.779 ± 0.002</td>
<td></td>
</tr>
<tr>
<td>CA08292²,³</td>
<td>NOAA/CIC Air</td>
<td>400.53 ± 0.02</td>
<td>(−35)³</td>
<td></td>
</tr>
<tr>
<td>AL-1³,⁵</td>
<td>CIC Air</td>
<td>215.8 ± 0.7</td>
<td>−9.26 ± 0.04</td>
<td></td>
</tr>
<tr>
<td>AL-2³,⁵</td>
<td>CIC Air</td>
<td>368.9 ± 0.5</td>
<td>−9.80 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>NEEM–2³,⁵</td>
<td>CIC Air</td>
<td>378.6 ± 0.5</td>
<td>−8.0 ± 0.1</td>
<td></td>
</tr>
</tbody>
</table>

¹ Stable isotopic composition calibrated at CIC by IRMS dual-inlet against GS19 and GS20 (Centre for Isotope Research, Groningen University, Netherlands). ² CO₂ mixing ratio calibrated and certified by NOAA ESRL/GMD (Boulder, Colorado, USA). ³ Stable isotopic composition calibrated at CIC against Messer–649250 and CA08054 with the setup described in this study but in “dry mode” (without ice). ⁴ Stable isotopic composition calibrated by the Stable Isotope Lab at INSTAAR (SIOL, University of Colorado, USA) in cooperation with NOAA. ⁵ CO₂ mixing ratio calibrated at CIC against the three NOAA standards CA08274, CA08054 and CA08292, with the setup described in this study and by WS-CRD spectroscopy. * Values outside of the reliable calibration range.

2.3 Standards

The reported CO₂ mixing ratios (also referred to as CO₂ concentrations in the literature) are defined as the dry air mole fraction expressed in parts per million by volume (ppm), and are linked to the World Meteorological Organization (WMO) mole fraction scale for CO₂ in air (Tans and Zhao, 2003; Zhao and Tans, 2006). Isotope ratios are reported relative to the international measurement standards (VPDB, VPDB-CO₂ and VSMOW for ¹³C-CO₂, ¹⁸O-CO₂ and ¹⁸O-H₂O, respectively) using the delta notation:

\[
\delta = \left( \frac{R_{\text{sample}}}{R_{\text{standard}}} \right) - 1,
\]

where \( R \) denotes the ratio of the heavy to light isotope in the sample and the standard, respectively. Our working standards were selected in order to cover the range of atmospheric CO₂ mixing and stable isotope ratios expected for glacial to interglacial conditions (Table 2). With these standards, system characterization, daily calibration and continuous quality control sample (QCS) measurements were performed (Sects. 3 and 4).

For referencing IRMS measurements, a working standard (WS) of pure CO₂ from a natural source (Messer–649250, Messer, Italy) is injected via an open-split interface (GP interface). Its stable isotopic composition was referenced at CIC against pure CO₂ reference gases GS19 and GS20 from the Centre for Isotope Research (Groningen University, Netherlands; Meijer, 1995) by IRMS dual-inlet measurement (Delta V Plus, Thermo Fisher, Germany). For air standards we used three synthetic air mixtures, in the following called CA08274, CA08054 and CA08292 provided by the Global Monitoring Division of the Earth System Research Laboratory at the National Oceanic and Atmospheric Administration (NOAA ESRL/GMD, Boulder, USA), two pressurized air tanks called AL-1 and AL-2 (Air Liquid, Denmark) and one of two atmospheric air tanks sampled in 2008 at a clean-air site of the NEEM deep ice core drilling camp called NEEM–2. The three NOAA tanks have been calibrated and certified by the NOAA ESRL/GMD Carbon Cycle Gases Group for CO₂ mixing ratios. The other tanks have then been calibrated against these three standards at CIC both with the setup described in this study and directly from the tanks by wavelength-scanned cavity ring-down spectroscopy (WS-CRDS; CFADS36 CO₂/CH₄/H₂O analyser, Picarro Inc., USA). The stable isotopic composition of CA08054 has been calibrated by the Stable Isotope Lab at INSTAAR (SIOL, University of Colorado) in cooperation with the NOAA Climate Monitoring and Diagnostics Division (CMDL). The stable isotopic composition of the two other NOAA cylinders and of AL-1, AL-2 and NEEM-2 has then been calibrated against these three standards at CIC both with the setup described in this study and directly from the tanks by wavelength-scanned cavity ring-down spectroscopy (WS-CRDS; CFADS36 CO₂/CH₄/H₂O analyser, Picarro Inc., USA). The stable isotopic composition of CA08054 has been calibrated by the Stable Isotope Lab at INSTAAR (SIOL, University of Colorado) in cooperation with the NOAA Climate Monitoring and Diagnostics Division (CMDL). The stable isotopic composition of the two other NOAA cylinders and of AL-1, AL-2 and NEEM-2 has then been calibrated against these three standards at CIC both with the setup described in this study and directly from the tanks by wavelength-scanned cavity ring-down spectroscopy (WS-CRDS; CFADS36 CO₂/CH₄/H₂O analyser, Picarro Inc., USA). The stable isotopic composition of CA08054 has been calibrated by the Stable Isotope Lab at INSTAAR (SIOL, University of Colorado) in cooperation with the NOAA Climate Monitoring and Diagnostics Division (CMDL). The stable isotopic composition of the two other NOAA cylinders and of AL-1, AL-2 and NEEM-2 has then been calibrated against these three standards at CIC both with the setup described in this study and directly from the tanks by wavelength-scanned cavity ring-down spectroscopy (WS-CRDS; CFADS36 CO₂/CH₄/H₂O analyser, Picarro Inc., USA). The stable isotopic composition of CA08054 has been calibrated by the Stable Isotope Lab at INSTAAR (SIOL, University of Colorado) in cooperation with the NOAA Climate Monitoring and Diagnostics Division (CMDL). The stable isotopic composition of the two other NOAA cylinders and of AL-1, AL-2 and NEEM-2 has then been calibrated against these three standards at CIC both with the setup described in this study and directly from the tanks by wavelength-scanned cavity ring-down spectroscopy (WS-CRDS; CFADS36 CO₂/CH₄/H₂O analyser, Picarro Inc., USA).

3 Measurement procedures and quality control

3.1 PreCon system

The pure CO₂-WS (Messer–649250) is injected via an open-split interface (GP interface) for referencing of IRMS measurements but can optionally also be passed through the automated section B of the system (Sect. 2.3, Fig. 2). This allows the assessment of potential fractionation effects from gas trapping and GC separation (Sect. 4.1.2). In the daily routine this is useful for an immediate control of conditions and stability of the PreCon, GC and detection systems. At the beginning of each measurement day, such injection runs, variable in the amount as well as blank runs for this part of the set-up, were therefore analysed (Table 3).
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Table 3. Example of a daily measurement sequence. See Table 2 for details on standard gases (WS, air standard) and the main text for analytical procedures. BFI denotes bubble-free ice.

<table>
<thead>
<tr>
<th>Daily run</th>
<th>Type of sample</th>
<th>Sample name</th>
<th>Comment</th>
<th>Set-up$^1$</th>
<th>Data processing$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>WS – 0.4 µL</td>
<td>Messer–649250</td>
<td>first run of the day</td>
<td>B (discard)</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Blank</td>
<td>Blank</td>
<td></td>
<td>B (discard)</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>WS – 0.4 µL</td>
<td>Messer–649250</td>
<td>first run after blank</td>
<td>B</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>WS – 0.2 µL</td>
<td>Messer–649250</td>
<td></td>
<td>B</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>WS – 0.4 µL</td>
<td>Messer–649250</td>
<td></td>
<td>B</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Air standard</td>
<td>CA08054</td>
<td>sample loaded – NC opened</td>
<td>A, B</td>
<td>(discard, NC surface effects)</td>
</tr>
<tr>
<td>7</td>
<td>Air standard</td>
<td>CA08054</td>
<td></td>
<td>A, B</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Air standard</td>
<td>CA08054</td>
<td></td>
<td>A, B</td>
<td>daily calibration &amp; drift correction</td>
</tr>
<tr>
<td>9</td>
<td>ICE</td>
<td>DE08–439</td>
<td>ice crushed</td>
<td>A, B</td>
<td>RESULT</td>
</tr>
<tr>
<td>10</td>
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<td>CA08054</td>
<td></td>
<td>A, B</td>
<td></td>
</tr>
<tr>
<td>11</td>
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<td>CA08054</td>
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<td>A, B</td>
<td>(discard, NC surface effects)</td>
</tr>
<tr>
<td>12</td>
<td>ICE</td>
<td>GRIP 250–12</td>
<td>ice crushed</td>
<td>A, B</td>
<td>RESULT</td>
</tr>
<tr>
<td>13</td>
<td>ICE</td>
<td>BFI (w or w/o CA08054)</td>
<td>BFI crushed (or not crushed)</td>
<td>A, B</td>
<td>procedural blank</td>
</tr>
<tr>
<td>14</td>
<td>Air standard</td>
<td>CA08054</td>
<td>sample loaded – NC opened</td>
<td>A, B</td>
<td>(discard, NC surface effects)</td>
</tr>
<tr>
<td>15</td>
<td>ICE</td>
<td>DE08–443</td>
<td>ice crushed</td>
<td>A, B</td>
<td>RESULT</td>
</tr>
<tr>
<td>16</td>
<td>Air standard</td>
<td>AL-2</td>
<td></td>
<td>A, B</td>
<td>daily calibration</td>
</tr>
<tr>
<td>17</td>
<td>Air standard</td>
<td>AL-1 (AL-2)$^3$</td>
<td></td>
<td>A, B</td>
<td>quality control sample (QCS)</td>
</tr>
<tr>
<td>18</td>
<td>Air standard</td>
<td>CA08054</td>
<td>sample loaded – NC opened</td>
<td>A, B</td>
<td>(discard, NC surface effects)</td>
</tr>
<tr>
<td>19</td>
<td>ICE</td>
<td>DE08–426</td>
<td>ice crushed</td>
<td>A, B</td>
<td>RESULT</td>
</tr>
<tr>
<td>20</td>
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<td>CA08054</td>
<td></td>
<td>A, B</td>
<td>daily calibration &amp; drift correction</td>
</tr>
<tr>
<td>21</td>
<td>Air standard</td>
<td>AL-1 (CA08292/CA08274)$^3$</td>
<td></td>
<td>A, B</td>
<td>daily calibration</td>
</tr>
</tbody>
</table>

$^1$ Section of system passed by the analysed gas sample (see Fig. 2). $^2$ See main text for details about data processing. $^3$ Alternative measurement option.

The following step by step description of the measurement procedure follows part numbering and abbreviations as indicated in Fig. 2. Connected in series, the cryogenic traps T2 (1/4 in. SS Swagelok tube, filled with around 5 cm of HayeSep D, 100/120 mesh, Sigma-Aldrich, Switzerland) and T3 (empty 1/16 in. SS Swagelok tube) are cooled by being immersed into liquid nitrogen. In order to prevent ambient air to be sucked into these traps while cooling down, the on/off valve (SS–4BK–TW–1C, Swagelok, USA) actuated, A210UWM, VICI, USA) switched compared to the sample flow (10 mL min$^{-1}$ for air standard and ice sample measurements – directing the sample gas flow (10 mL min$^{-1}$) set by a flow controller; VCD 1000, Porter, USA) through the TCD detector. After an idle time of 30 s allowing the flow to stabilize, the LN Dewar cooling T2 and T3 is automatically lowered by a double activated pneumatic cylinder (CD85N20–250B, SMC, Denmark) to a level at which T3 is still cooled. T2 is then heated by a rope heater (FGR–060, Omegalux, UK) to the set temperature of 150$^\circ$C regulated by a PID controller (iTRON 08, JUMO, UK). Thereby, the trapped gas is released and the amount of the main air components is detected by the TCD (no signal for pure CO$_2$ and for blanks) while both CO$_2$ and N$_2$O are trapped in T3 for later separation and detection in a second line (typical chromatograms for measurements of CO$_2$–WS, procedural blanks and ice core samples can be found in Supplement Figs. S1 and S2 and in Fig. 3, respectively). After 5 min, V2 is switched again, now redirecting the sample flow (again 20 mL min$^{-1}$) through this alternative detection line. After an idle time of 40 s allowing the flow to stabilize, the LN Dewar cooling T2 and T3 is further lowered and trap T3 quickly heated to 100$^\circ$C (resistance wire, 2.5Ω m$^{-1}$, 5 m, Conrad Electronics, Germany) regulated by a second PID controller (iTRON 32, JUMO, UK). Thereby, CO$_2$ and N$_2$O (not present in the pure CO$_2$–WS) are released and the different gases are subsequently separated by the gas chromatographic fused silica capillary column in the GC–1 (30$^\circ$C, CP–PoraBond–Q, 25 m × 0.53 mm ID, d$f = 10$ µm, Varian,
USA). The signal then being detected by the PDD is shown in Fig. S1 (Supplement). Shortly before the eluted CO₂ peak arrives at the PDD (discharge gas flow set to 30 mL min⁻¹), three parallel traps (fused silica capillaries, 250 µm ID, 1.8 m length, BGB, Germany) are immersed into LN to re-trap CO₂ and N₂O for cryogenic focus (CF) after splitting the sample stream at the PDD outlet valve V3 (GC built in 6-port valve, VICI, USA). To vent remaining H₂O and potential contaminants from drilling fluid (for ice samples), V3 is switched before the signal is detected in the PDD (~ 70 s after the maximum in the CO₂ peak). The GC column is then conditioned for the next sample by heating to 150°C. In the separated part of the line, the three CF capillaries are meanwhile lifted one after the other out of the LN, subsequently releasing the sample – now split in three aliquots – for further transport in a reduced He flow of 1.5 mL min⁻¹. To avoid isobaric interference, CO₂ and N₂O contained in these aliquots are again separated in a second GC column (GC–2; 35°C, CP–PoraBond–Q, 40 cm × 0.53 mm ID, df = 10 µm, Varian, USA) before the remnant water vapour is removed by a Nafion drying column (40 cm × 0.36 mm ID, Perma Pure Inc., USA). Finally, the sample gases are introduced to the IRMS via the open split of the GP interface. Before the three CO₂ sample peaks elute, the reference gas (CO₂-WS, here the same as the sample gas) is injected several times (reference port) in order to reach stable IRMS-source conditions. The peak amplitude is thereby adjusted to closely match the amplitude of the sample aliquots (not for blanks). For the same reason a constant CO background flow through the reference open-split port into the ion source was maintained as proposed in Elsig and Leuenberger (2010). The mean value of the peak before and after the sample is ultimately used for referencing. Splitting the sample in aliquots allows for three IRMS measurements on the same sample theoretically improving the analytical precision. In practice, the therefore required quantitative splitting in three evenly sized aliquots is difficult to achieve. For the calculated mean values (weighted by mass, i.e. size) no difference in final precision compared to a single measurement was observed. However, results from these multiple measurements could be statistically analysed and were useful, e.g., to evaluate the applied IRMS nonlinearity correction which reduces the standard deviation over the three replicates (Sect. 4.1.2).

The total measurement time from injection to final IRMS detection is 21 min. It is longer for air standard and ice sample measurements, as these include section A of the system (Sect. 3.2).

Figure 3. Chromatograms for the measurement of an ice sample as described in Sect. 3.2. Upper panel: IRMS signal intensity for m/z 44, 45 and 46. Injections of the WS via the open split are identifiable by the flat-topped peaks. Peaks 1–9 are used to reach stable source conditions while peaks 10 and 14 before and after the samples are used for referencing. The inset shows baseline details and N₂O separation in detail. Lower panel: PDD and TCD intensity signal for CO₂ and air, respectively. Stars indicate valve switching, resulting in small variations in the PDD signal due to changes in pressure and flow (see inset, not detected by the less sensitive TCD). Diamonds indicate immersion of the three capillary traps into liquid nitrogen for CO₂ cryofocusing and their subsequent one-by-one release resulting in the three peaks of the split sample shown in the upper panel (peaks 11–13). Over the time period, indicated by the grey bar, the GC–1 temperature is increased to 150°C in order to precondition the column for the next sample (release of water and remnants of drilling fluid contamination). The enlargement shows baseline details revealing the N₂O peak and small remains of N₂ and O₂ from the air sample, incompletely separated by the preceding cryogenic partition.

3.2 Standard air, ice samples and blanks

Air standards and blank measurements were performed regularly for calibration and system characterization, thereby following the exact procedure used to measure real (natural) ice samples, i.e. following the “identical treatment” principle (Werner and Brand, 2001). To simulate the entire measurement procedure as close as possible, artificial bubble-free ice (BFI) samples were used. BFI was produced from ultrapure water (MilliQ, 18.2MΩ cm at 25 °C) degassed for 60–90 min using a roughing LV pump (E2M0.7, Edwards, UK) and then slowly frozen from the bottom (20 cm in 48 h), thereby forcing the remaining gas out of the water. Results for calibrations and system characterization will be presented in Sect. 4.

Samples of air, either from tanks (e.g. atmospheric samples or standards) or extracted from ice core samples, pass all sections of the experimental set-up (A and B, Fig. 2). When the system is not in use (e.g. overnight), section B is constantly flushed with He while all lines in section A and the NC are pressurized slightly above atmospheric pressure with He and N2, respectively. Prior to analysis, ice core or BFI samples are cut to the required dimensions and the surfaces are decontaminated by removing the top layers with a scalpel.

A typical daily measurement sequence is listed in Table 3. The measurements can be separated into five main categories:

1. air standard measurements performed by injecting variable amounts of the standard gas over an ice sample of either natural or artificial origin (BFI) without crushing the ice; standards of different CO2 mixing ratios and isotopic composition were thereby used (Table 2);

2. system blank measurements performed by omitting addition of standard gas and hence resulting in sampling carrier gas only;

3. BFI measurements either performed with air standard added or omitted but with the ice being crushed;

4. measurements simulating the crushing procedure (i.e. needles moved) but using BFI which has been crushed in advance; in this case, artefacts from remnant gas potentially present in the initial, intact BFI sample can be excluded (Sect. 4); these measurements were also performed with and without the addition of air standard; the latter case will be further referred to as “procedural blank”;

5. measurements of air extracted from natural ice samples by crushing the ice.

The following step by step description of the measurement procedure follows part numbering and abbreviations as indicated in Fig. 2. Section A and the NC are prepared for measurements during the first five runs of the daily sequence (Table 3). The procedure for these runs is described in Sect. 3.1. After the air cooling system of the NC is started, the first ice sample is loaded once a temperature of −20 °C has been reached. While the unit is open, both the upper and lower parts are continuously flushed with N2 through the respective inlets (11 and 12) to prevent contamination from ambient air and condensation of water vapour on the cold inner surfaces. After the NC is closed and vacuum sealed again, inlets 11 and 12 are closed and the N2 flow is turned off.

Each time the NC is opened, the following sequence of steps is required for evacuation and reconditioning. The chamber is evacuated by the LV pump through inlet 11 for 1 min. Only port 11 is used in this evacuation step to prevent trapping of N2 on trap T1 (heated to 100 °C, 1/4 in. SS tube, filled with HayeSep D, 100/120 mesh, Sigma-Aldrich). This is essential because at this point the N2 abundance in the NC is much higher than in samples of recent or past atmosphere. When the system is evacuated, inlet 12 remains always closed. To avoid analytical artefacts, this inlet is only used with a one-directional flow into the NC, thus preventing water vapour from reaching the gas manifold and the vacuum lines used for the injection of standard air samples. When the pressure in the NC has dropped, trap T1 and the rest of the vacuum lines still filled with He or N2, respectively, are now also evacuated through the respective lines. After a few seconds the vacuum system is switched to the HV pump for another 5 min. To precondition the NC, the chamber is closed off from the vacuum (valves 11 and 13) and then filled with an aliquot of air standard through inlet 12. Since the standard added in this step is used only for conditioning, the injected amount is non-quantitative. After inlet 12 is closed again, T1 is cooled down by being immersed in LN before outlet 13 is opened to cryogenically pump the air aliquot onto T1. After a trapping time of 3 min, the line (vacuum line–NC–T1) is flushed with He from the inlet next to the gas manifold set to a flow of 80 mL min−1 (gas flow controller, Model 100, VICI, USA). Since this air aliquot is not to be analysed, the He gas flow is then directed through the 3-way valve (3-WV; SS–43GX54, Swagelok, USA) into the LV pump. To release the previously trapped air sample while flushing, T1 is heated by a rope heater (FGR–060, OmegaLux, UK) to the set temperature of 150 °C regulated by a PID controller (N2300, West Instruments, USA). After a flushing time of 3 min, outlet 13 and inlet 12 are closed and the temperature of T1 is reduced to 100 °C. The system is now in a similar state as prior to runs where the NC has not been opened beforehand (see Table 3).

To start an acquisition run, the NC, T1 and all vacuum lines are evacuated for ~10 min until reaching a pressure of 2 × 10−3 mbar monitored by P1 (Single Gauge Pirani Transmitter, Pfeiffer, Germany; in a dry system, a vacuum < 5 × 10−4 mbar, the lower detection limit of P1 is reached). The gas manifold allows selection between the different air standards (Air STD) and the amount to be injected. For this purpose, gas can be expanded into a defined volume adjustable in size by the valves included in each line. For each
expansion step, an equilibrium time of 2 min was applied. In every sequence, the first run which includes section A (run 6) is performed using an air standard of large size (aliquot expanded from the volume between the first and last valves of the gas manifold line). This allows for a first immediate control of system conditions and day by day stability after the idle time overnight. From the gas manifold the sample is expanded into a defined volume of ~11 cm$^3$ (DF), equipped with another pressure gauge, P2 (high-precision piezo pressure transmitter, PAA–35X, Keller AG, Switzerland). After DF has been closed off towards the manifold (valve 15) and the NC (inlet 12 has been closed initially), the pressure is recorded for later use (Sect. 4.1.1). At the same time, sample air contained in the rest of the lines is pumped off by the HV pump. After the pressure has been recorded and valves 11 and 13 have been closed, the gas is further expanded into the NC. The standard gas is kept in the NC for 2 min, simulating the procedure applied for natural ice samples which are crushed at this point, allowing the gas to subsequently expand from the ice. The valve to DF is thereby kept open, which allows us to record and survey the efficiency of gas extraction from the opened bubbles when measuring natural ice samples and subsequently the gas transfer out of the NC onto the pre-cooled trap T1 in the following 3 min cryogenic pumping steps. Afterwards, with the cryopump still active (i.e. T1 still cooled with LN), the NC and T1 are filled with He (80 mL min$^{-1}$) to 1100 mbar, which is slightly above the pressure in section B (~3 min). This prevents backflushing from section B to A when they are connected by switching the 3–WV. At this point, the air sample is released by heating T1 to the set temperature of 150°C and transferred to the pre-cooled trap T2 by the carrier gas (80 mL min$^{-1}$). This step takes another 3 min. The 3–WV is then closed, again separating section A and B. From here on, the sample is in a reduced carrier gas flow environment, and further sample treatment is handled by the automated part of the system as described in Sect. 3.1. Meanwhile section A can be prepared for the next sample. Therefore, the outlet valve of the NC is closed, the N$_2$ flush flow opened and the next ice cube loaded. If the next sample in the sequence (Table 3) is to be measured with the ice remaining in the NC, evacuation of the extraction unit and the vacuum lines is sufficient to prepare for the next run.

For the measurement of natural ice samples, the procedure is as described above but all steps related to the air standard addition can be skipped. Chromatograms of the TCD, PDD and IRMS for a typical ice sample measurement are shown in Fig. 3. Here, as opposed to measurements of pure CO$_2$-WS and blanks (Figs. S1 and S2, respectively), the peak from sample air shows up as a distinct TCD signal, and signals of H$_2$O from ice sample water vapour and remnants of drilling fluid contamination are detected by the PDD in addition.

Including section A of the system does not add much to the total measurement time given in Sect. 3.1 because most of the additional steps required can already be executed while the current measurement is running in the automated section B. Typically, the analysis takes around 30 min (run to run). However, it increases for runs requiring loading of a new ice sample (opening of the NC) and as a consequence the described evacuation and flushing steps (~50 min, e.g. runs 6, 11, 14 and 18 in Table 3).

4 Results and discussion

4.1 Evaluation and calibration

System-specific analytical bias for CO$_2$ concentration and CO$_2$ stable isotope measurements can result from fractionation in the analytical system related to adsorption and desorption processes, specifically in the extraction unit (Zumbrunn et al., 1982), non-quantitative trapping and releasing of gas and from GC separation. For stable isotope measurements, further bias arises from IRMS injection and source effects (Elsig and Leuenberger, 2010). The net effect of these processes was assessed and monitored on a regular basis to account for system drifts. Such drifts may occur when boundary conditions (e.g. room temperature) change and can appear within a measurement day, on a day to day basis and on longer timescales, here being either related to changes in the set-up (e.g. replacements of parts, change of carrier gas tanks or standards, adjustments in carrier gas flows) or small changes in the procedure. Following the identical treatment principle (Werner and Brand, 2001), we characterized all sections of our system for systematic effects using the various standards available (Table 2), ultimately allowing a reliable correction of the raw data. Runs of QCS to assess long-term stability of our final measurement results were carried out on a daily basis (results in Sect. 4.2.1).

4.1.1 CO$_2$

CO$_2$ mixing ratios can, in principle, be derived from the ratio between the IRMS CO$_2$ peak area (values given in the following always denote the mean for the three sample peaks) and the major air components detected as TCD peak area (Fig. 3). The TCD air peak signal was observed to be nonlinearly related to the amount of air with significant variability. This can be seen in the relation between the high-precision pressure readings of standard gas (described in Sect. 3.2) and the TCD air peak area investigated for different measurement periods (Fig. 4a). We also found variability in the relation between the IRMS peak area and the CO$_2$ amount; in this case, however, the relationship is linear. Reasons for the observed variability are manifold, e.g. small adjustments in flow or trapping procedures, or small changes in outer conditions such as room temperature (e.g. March and June 2012).

Calibration: To avoid nonlinearity in data processing, the TCD signal was converted to pressure using the second-order polynomial functions shown in Fig. 4a. These functions can be determined for any user-defined period of time with
the data obtained from daily performed measurements of air standards. In doing so, the large long-term variations in TCD sensitivity are accounted for and all TCD measurements are adjusted and referred to a common scale (pressure). The CO$_2$ mixing ratios known for the standard gases were then related to the derived ratio between the IRMS CO$_2$ peak area and the computed pressure. By including data of all available standards, the resulting linear calibration curve is well defined over a large concentration range (Fig. 4b). The data shown are based on repeated measurements series over the course of more than 1 year ($n = 4$). Each series was performed on a single day with the entire range of standards being measured at least three times. The ratios calculated for the individual series were matched for CA08054 in order to account for the long-term variation of the IRMS response, resulting in the average transfer function shown.

With the calibration curve covering the entire range of expected measurement results (Fig. 4b), daily calibration could be performed with a subset of standards only, thus significantly reducing the total sequence time. Summarized in Table 3, a typical daily measurement sequence includes the repeated measurement of standard CA08054 (run 8, 10, 13 and 20) to determine and subsequently adjust for (i) potential off-set in the calculated ratio compared to the calibration curve and (ii) system drift over the sequence measurement time. It further includes at least one additional standard gas measurement with a different CO$_2$ mixing ratio to adjust for potential variations in the slope of the linear calibration fit (runs 16, 21 and 17 if necessary). If not used in the daily calibration, run 17 is treated as a real sample in the post processing of the raw data, allowing us to assess the long-term consistency and precision of our measurements (QCS; see Sect. 4.2.1). In run 21, CA08054 is occasionally injected in variable amounts to ensure the independence of the final results from the sampled amount of gas.

**Procedural blank:** We find that independently of sample size, gas amount or CO$_2$ concentration, a constant amount of CO$_2$ is produced by the extraction itself. This amount was determined by measurements of BFI samples. To perfectly simulate measurements of natural ice samples according to the identical treatment principle (Werner and Brand, 2001), artificially produced bubble ice with entrapped standard gases would be required. Because this is technically not feasible, instead we added standard gas to a BFI sample loaded in the NC and crushed for the measurement. The CO$_2$ mixing ratios we observed were elevated by $4.6 \pm 2.6$ ppm on average ($n = 5$) compared to the expected value. However, for subsequent measurements with an identical procedure (including movement of the needle pins), but using the previously crushed BFI, the observed elevation was less. This indicates the presence of remnant gases in our BFI which is in agreement with independent results from another study ($\sim 3$ ppm for CIC BFI, Appendix A6 in Rubino et al., 2013). We thus considered the lower offset value of $2.3 \pm 2.0$ ppm (average, $n = 4$) to be representative of the purely system related blank.

This is a reduced offset compared to other systems, allowing the analysis of similarly small sample sizes (e.g. 4.9 ppm for the KUP NC; Bereiter et al., 2013) which most likely is a result of the friction-reduced motion and lower operating temperature in our NC design. Anyhow, this CO$_2$ enrichment – expressed in ppm before – is observed in the raw data as an elevated signal in the IRMS CO$_2$ peak area. The size of this extra signal can therefore be directly estimated from procedural blank measurements when using BFI which has been crushed in advance (Fig. S3). As an advantage, propagation of uncertainties associated with data post-processing can then be omitted. Combining the results from both approaches, the IRMS CO$_2$ peak area for the procedural blank was observed to be $0.5 \pm 0.2$ mVs on average ($n = 9$), i.e. elevated by $0.1 \pm 0.1$ mVs compared to the system blank (see

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**Figure 4.** CO$_2$ calibration. (a) Relationship between sample size and TCD peak area of air for different measurement periods. The lines are second-order polynomial fits through the data. Pressure in the injection volume is proportional to amount. The grey bar indicates the typical sample size range extracted from ice samples. (b) Calibration curve; known CO$_2$ mixing ratios of air standards vs. ratio of IRMS peak area for CO$_2$ (m/z 44) and TCD detected air amount transformed to pressure from the relationship shown in (a).

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![Graph showing relationship between sample size and TCD peak area of air for different measurement periods. The lines are second-order polynomial fits through the data.](https://www.atmos-meas-tech.net/9/3687/2016/)

![Calibration curve showing known CO$_2$ mixing ratios of air standards vs. ratio of IRMS peak area for CO$_2$ (m/z 44) and TCD detected air amount transformed to pressure.](https://www.atmos-meas-tech.net/9/3687/2016/)
Sect. 4.1.2 – Air amount dependence). In Sect. 4.1.3, the procedural blank correction applied to ice core samples will be discussed in detail.

4.1.2 $\delta^{13}$C-CO$_2$ and $\delta^{18}$O-CO$_2$

The individual sections of the set-up were characterized for systematic effects on $\delta^{13}$C and $\delta^{18}$O results. Calibrations, control of system stability and blanks, as well as QCS measurements to assure the long-term quality of our analysis (Sect. 4.2.1), were all performed on a daily basis (Table 3).

**IRMС nonlinearity:** We characterized the IRMS source effects – in the following referred to as IRMS nonlinearity – by measurements of reference gas (CO$_2$-WS) injected via the GP interface reference open split (Fig. 2). To reach stable source conditions, six injections were made prior to acquisition similar to the approach described in Elsig and Leuenberger (2010). Then, the injection amount was increased step-wise, reflected in rising amplitudes of the rectangular peaks. The signal amplitude thereby ranged between 400 and 8400 mV (m/z 44), with the reference peak always set to around 4000 mV (gain: $R = 10^8 \Omega$). This experiment was replicated on three different days distributed over a time period of 1 year. The resulting total number of acquisitions was 177. In Fig. 5a, the IRMS nonlinearity effect for $\delta^{13}$C is shown, measured as the deviation from the reference (~4000 mV). The same procedure was applied to investigate the $\delta^{18}$O nonlinearity (Fig. S3).

**PreCon–GC linearity:** A potential and possibly size-dependent effect on the stable isotope values of the sampled gas when passing the various traps and GC columns in the automated set-up section B (Fig. 2) was investigated. Therefore, various amounts of the CO$_2$-WS were injected onto trap T2 via the injV (Fig. 2) as described in Sect. 3.1. The relationship found will be denoted as PreCon–GC linearity in the following. In addition, the blank for this section was determined when injection of CO$_2$ was omitted, but otherwise the exact same procedure was followed (Sect. 3.1). To allow determination of this PreCon–GC-related effect independently from any other system-induced contribution, the obtained raw data were first corrected for the IRMS nonlinearity by applying the third-order polynomial fit shown in Fig. 5a. Subsequently, a blank correction using the values determined here was applied. Therefore, the following equation which closely approximates the isotopic composition in a pool ($\delta_\Sigma$) of isotopically different members ($m_i$ with corresponding $\delta_i$) was used:

$$\delta_\Sigma = \frac{\sum m_i \delta_i}{\sum m_i}.$$  

Equation (2) shows that the bigger the CO$_2$ blank-to-sample ratio and the difference between blank and sample isotopic composition, the bigger the blank correction will be. In other words, even when the blank is constant both in size and isotopic composition, the blank correction should vary depend-

Figure 5. Fractionation effects for $\delta^{13}$C: (a) IRMS nonlinearity; $\delta^{13}$C dependence on peak amplitude (top x-axis), $\Delta$-amplitude is the deviation in intensity (m/z 44) from the reference peak (ref, $\delta^{13}$C and $\Delta$-amplitude = 0). The data are obtained from a total of 177 measurements and mean values with the 1$\sigma$ standard deviation are shown. (b) PreCon–GC linearity (bottom x-axis); CO$_2$ sample size dependence for pure CO$_2$ working standard directly injected to section B. The data are obtained from a total of 318 measurements corrected for IRMS nonlinearity and blank; mean values with the 1$\sigma$ standard deviation are shown. (e) Air amount dependence (bottom x-axis); air sample size dependence for air standards/samples injected to section A. The data are obtained from a total of 46 measurements corrected for IRMS nonlinearity, PreCon–GC linearity and system blank. Mean values with the 1$\sigma$ standard deviation are shown. The grey bars indicate the typical procedural blank and sample size range of air extracted from ice samples, respectively.
Fig. S3 for $\delta^{18}$O were derived iteratively until changes in the converging values were well below the IRMS precision ($n = 5$). The final mean values for blanks ($n = 99$) which were reproducible for the acquired 2-year time period were $0.09 \pm 0.02$ mVs, $-24.2 \pm 1.9$ ‰ and $-41 \pm 5$ ‰ for CO$_2$ IRMS peak area, $\delta^{13}$C and $\delta^{18}$O, respectively. This demonstrates that the blank isotopic values are strongly depleted compared to atmospheric values. This is consistent with the assumption that the blank is related to tiny amounts of CO$_2$ constantly being adsorbed and desorbed from inner surfaces, trapping materials and GC columns, thereby undergoing heavy isotopic fractionation. To test if the measured values are reliable although measured on extremely small sample amounts, comparable amounts of CO$_2$-WS were directly injected via the sample open split, resulting in a similar peak shape ($n = 5$, IRMS peak area between 0.14 and 1.17 mVs). 

After correction for the IRMS nonlinearity effect, we obtained average values of $-6.5 \pm 0.6$ ‰ and $-11.7 \pm 1.6$ ‰ for $\delta^{13}$C and $\delta^{18}$O, respectively. These numbers are not significantly different from the expected values for the CO$_2$-WS ($-6.004 \pm 0.008$ ‰ and $-10.80 \pm 0.13$ ‰), which demonstrates the reliability of measurements for such small sample amounts and adds confidence to the determined blank values. 

**Air amount dependence:** To investigate the characteristics of the system section A, variable amounts of the CA08054 air standard were analysed over ice. Changing the gas matrix from pure CO$_2$ to trace amounts in air may cause alteration in the previously defined PreCon–GC linearity. This potential effect cannot be distinguished in this experiment and may contribute to the observations in the following being denoted as the "air amount dependence". In repeated series ($n = 10$), distributed over a time period of more than 1 year, 46 such measurements were made. In addition, 42 blanks (omitting sample injection but exactly following the procedure otherwise) were measured. Following the approach for determination of the PreCon–GC linearity, the blank contribution was separated from the effect investigated here. Identical to samples, blanks were corrected for IRMS nonlinearity, PreCon–GC linearity and the air amount dependence discussed here again in an iterative way. The blank value obtained, now representative of the entire set-up and accordingly denoted as "system blank", was $0.4 \pm 0.1$ mVs, $-27.6 \pm 1.2$ ‰ and $-30 \pm 3$ ‰ for CO$_2$ peak area, $\delta^{13}$C and $\delta^{18}$O, respectively. Because of the additional trap (T1) and large additional surface area from extra lines and the NC chamber, the bigger size of the system blank compared to the blank observed for the PreCon–GC section alone is expected. However, the obtained values for their isotopic composition are comparable. This indicates the responsible fractionation effects, likely related to adsorption and desorption processes, to be similar in the different sections of the set-up. Because the built-in parts are alike in surface and trapping material, this is not unexpected. The final relationship for the air amount dependence is presented in Fig. 5c. As described, it was obtained after correction for IRMS nonlinearity, PreCon–GC linearity and blank contribution (for $\delta^{18}$O see Fig. S3).

**Procedural blank:** The procedural blank determined in Sect. 4.1.1 was also analysed for its isotopic composition. The isotope ratios resulted in values of $-26.6 \pm 0.8$ ‰ for $\delta^{13}$C and $-29 \pm 3$ ‰ for $\delta^{18}$O ($n = 5$) when using BFI which has been crushed in advance to avoid artefacts from entrapped remnant gas. Because the available amount of gas for such measurements is very small ($0.5 \pm 0.2$ mVs), a second indirect approach based on larger sample sizes was applied. Thereby, the same standard was analysed twice, simulating the crushing procedure with BFI in the second measurement. Nine such sets were measured and in four of them, the BFI had already been crushed in advance. In addition to the already known amount of the CO$_2$ procedural blank contribution, for each set, the isotopic composition expected for the standard and the amount of the CO$_2$-total with its isotopic composition is defined by the first and second measurement, respectively. With these numbers as input values, the isotopic composition of the procedural blank could then be estimated using a reversed form of Eq. (2). The values derived in this way for the procedural blank were $-24 \pm 3$ ‰ and $-28 \pm 4$ ‰ for $\delta^{13}$C and $\delta^{18}$O, respectively ($n = 9$). The bias due to the small amounts of remnant gas in the BFI can be calculated (Eq. 2) to be of the order of $0.2$ ‰ and can thus be neglected considering the uncertainty of this approach. In any case, these results are in close agreement with those obtained by the direct measurement of the procedural blank and further verification of their strongly depleted isotopic composition. In Sect. 4.1.3, the procedural blank correction ultimately applied to ice samples will be discussed.

**Calibration:** All raw data were post-processed, correcting for the characterized effects in the given order: (1) IRMS nonlinearity, (2) PreCon–GC linearity, (3) air amount dependence and (4) blank contribution. The repeated measurements of air standards differing in their CO$_2$ mixing ratio and isotopic composition were then used for daily calibration to adjust for potential day by day offsets and daily drift (runs 8, 10, 13, 16, 20, 21 and 17 in some cases). If not used for calibration, run 17 was treated as a QCS (see Sect. 4.2.1). For run 21, CA08504 is occasionally injected in variable amounts to ensure the independence of final results from the sampled gas amounts. Results and achieved precision for the measurements of ice samples will be discussed.

4.1.3 Procedural blank correction

For any analytical application, a thorough assessment and quantification of the blank is crucial. Whereas the system blank usually is representative for the analysis of standards ultimately used for calibration of the results, the procedural blank – here related to the measurement of ice core samples – can include extra contribution from sample treatment (e.g. preparation and crushing).
CO$_2$: to account (i.e. correct) for the extra contribution of the procedural blank compared to the system blank, the common approach described in the literature for measurements of CO$_2$ and its stable isotopes in ice core samples is to ultimately subtract a constant offset (e.g. Elsig et al., 2009; Schmitt et al., 2011; Rubino et al., 2013). Here, for CO$_2$ results this offset would correspond to the 2.3 $\pm$ 2.0 ppm determined in Sect. 4.1.1. However, such an approach requires the assumption that the extra CO$_2$ contribution of the procedural blank is variable in terms of amount (i.e. moles) in such a way, that the offset results in a constant blank-to-sample ratio regardless of CO$_2$ concentration and amount of sample gas extracted from the ice. Only then will a constant offset in terms of ppm (parts per million by volume) result. As this is highly unlikely, a compressed scale for data covering a large range of CO$_2$ mixing ratios has to be expected. As an example, this would be the case for records of glacial to interglacial atmospheric conditions ranging from around 180 ppm to the current atmospheric level of around 400 ppm. Whereas for measurements with large sample sizes, i.e. low blank-to-sample ratios, such a bias might be negligible, a different, more accurate correction of the procedural blank should be applied particularly if using small samples. In agreement with our observations of the blank, we here assumed the extra CO$_2$ contribution to be constant in terms of the absolute amount (moles not mole fraction/i.e. ppm). Accordingly, for each ice sample we subtracted the signal determined for this additional CO$_2$ contribution (Sect. 4.1.1) from the measured IRMS CO$_2$ peak area prior to conversion of results into ppm. For the small sample sizes analysed here, the improvement of this approach is directly reflected in a reduced standard deviation for results from sets of replicate ice measurements (same site and sampling depth, variable in the sampled gas amount; see Sect. 4.2.2 for sample details). For these samples, the applied correction varied between 1.9 and 3.3 in terms of ppm (2.4 ppm on average, $n = 18$).

$\delta^{13}C$ and $\delta^{18}O$: for the procedural blank correction of isotopic values, the common approach of subtracting a constant offset is even more critical. As discussed in Sect. 4.1.2 (PreCon–GC linearity), even for blanks constant in CO$_2$ contribution and isotopic composition, the magnitude of the procedural blank correction should be dependent on the sample size as well as the sample CO$_2$ mixing ratio and isotopic composition. Obviously, the bigger the CO$_2$ blank-to-sample ratios and the difference between blank and sample isotopic composition, the bigger the correction will be. Considering the observed strongly depleted isotopic composition of the blank CO$_2$ (Sect. 4.1.2), the variation of the correction might be significant even for samples of larger size. For sample sizes 10 times bigger than the ones analysed in this study (i.e. $\sim$ 100 g ice), the scale compression bias resulting from the application of the conventional approach was calculated for an ice core record covering the Holocene (approximated range: 180 to 370 ppm in CO$_2$ and $-6.3$ to $-6.6\%e$ in $\delta^{13}C$). For the $\delta^{13}C$ of the procedural blank, the determined low value of $-26.6\%e$ was used (Sect. 4.1.2). The procedural blank for the conventional correction of CO$_2$ and $\delta^{13}C$ was assumed to be 1 ppm and 0.1%e, respectively. These are typical literature values and small compared to the numbers determined in this study. Employing Eq. (2) for calculation, a potential additional effect arising from variations in the ice sample size used to obtain the record is not considered. Nevertheless, the expected scale compression bias is calculated to be around 0.06%e for the commonly applied procedural blank correction. This demonstrates that even for larger sample sizes, this bias can be significant, considering the recent improvements in analytical precision. Obviously, it is of particular relevance for higher blank-to-sample ratios (e.g. small sample sizes). Here we thus applied a new, more accurate approach for the procedural blank correction. It is similar in principle to the description given in Sect. 4.1.2 for air standards and is based on Eq. (2). We used 0.5 $\pm$ 0.2 mVs, $-26.6 \pm -0.8\%e$ and $-29 \pm 3\%e$ for procedural blank size, $\delta^{13}C$ and $\delta^{18}O$, respectively (Sect. 4.1.2). The results and precision for measurements of natural ice samples will be discussed in Sect. 4.2.2.

4.2 System performance

4.2.1 Analytical precision for the measurement of air samples

The precision and long-term consistency of our measurements were assessed by repeated QCS measurements of the two air standards AL-1 and AL-2 injected over natural and artificial (BFI) ice both before and after crushing. These two standards are different in their CO$_2$ mixing ratios and isotopic compositions (Table 2) and were injected in variable amounts of gas. QCS measurements were treated similarly to real ice samples, both in the applied measurement procedure (except for the crushing step) and the post-processing of the acquired raw data. In Fig. 6, the resulting time series for CO$_2$, $\delta^{13}C$ and $\delta^{18}O$ covering a 2-year period are shown. To derive a completely independent assessment, QCS measurements used in the daily calibration routine were excluded for this analysis. For the time frame covered, no trend can be observed for either of the parameters analysed in the two standards. The determined and assigned values agree well within the uncertainties. However, a small systematic shift of unknown origin observed for $\delta^{18}O$ of AL-2 cannot be excluded. From the combined dataset of AL-1 and AL-2 shown in Fig. 6, our analytical precision for the measurement of air samples over ice was determined with 1.9 ppm, 0.09 and 0.16%e for CO$_2$, $\delta^{13}C$ and $\delta^{18}O$, respectively (standard deviations around the respective mean).
Figure 6. Repeated quality control sample (QCS) measurements of air standards AL-1 (in green) and AL-2 (in blue) over ice, covering a time period of 2 years. The grey bands indicate the assigned value of the standard with uncertainty as given in Table 2. Dashed lines indicate the 1σ standard deviation of the data points (see numerical values). (a) CO2 mixing ratios (ppm). (b) δ13C-CO2 (% VPDB). (c) δ18O-CO2 (% VPDB-CO2). (d) Injected amount of air (top, left axis) and related amount of CO2 (bottom, right axis). Fewer results are shown for the stable isotopes because standards used in the daily calibration routine (i.e. not post-processed similar to real samples) were excluded from this analysis.

4.2.2 Natural ice samples – laboratory comparison and reproducibility

From measurements of ice samples from various sites and depths (8–13 g), the extraction efficiency of our NC was determined by the amount of air liberated, divided by the expected air total in the sample. Typically the efficiency is around 70–80% for bubbly ice and around 60% for clathrate ice (with the gas release time after crushing extended by 4 min). This is in the similar range as reported for other NC designs (Ahn et al., 2009; Lüthi, 2009; see Table 1).

To demonstrate the system performance and reproducibility, we analysed six ice samples of the recent past (1851–1969 AD) from Law Dome, Antarctica (DE08, 66°43′S, 113°12′E). This was done in a comparison study with CSIRO. The range of sample ages allowed for a comparison across a relatively wide range of CO2 and δ13C values. These samples were also part of an independent study published earlier (Rubino et al., 2013). Note that DE08 was dry-drilled and therefore the difference in the measurement systems that addresses drilling fluid contamination (i.e. separation by GC in the CIC system) is not tested. We measured replicates (n = 2 to 4) on the egg-shaped pieces remaining after the samples have been processed at CSIRO using their cheese grater dry extraction system. To allow a realistic assessment of measurement reproducibility, the replicates were measured on different days. The pooled standard deviation, used as a measure to estimate the overall analytical precision of our system for single measurements, was 2.0 ppm and 0.11 ‰ for CO2 and δ13C, respectively (n = 18). Compared with the results from CSIRO, good agreement within the assigned 1σ uncertainties was found for both CO2 and δ13C (Fig. 7). While the agreement between δ13C results is high (average CIC – CSIRO = 0.02 ‰), a small systematic offset of +1.8 ppm seems to exist for CO2. An obvious explanation for this offset would be discrepancies in calibration between the two laboratories relying on independent standards.
This is however not supported by the good agreement for the comparison of air tank measurements with different CO₂ mixing ratios (−0.3 ppm on average, \( n = 3 \)). Alternatively, the observed offset could be a real signal, explained by the occlusion of ambient air in micro-cracks resulting from the fierce mechanical treatment in the CSIRO grater and the elevated temperatures recorded during ice transport preceding the CIC measurements (around −5°C). For the same reason, the deviation in results of replicate measurements may partly be influenced by the ice itself, i.e. causing analytically independent variability. We therefore consider the pooled standard deviation calculated for the Law Dome samples to be a rather conservative estimate of our overall analytical uncertainty for single ice sample measurements. The estimate is however in line with the uncertainty determined for the QCS measurements (Sect. 4.2.1) with a slightly larger uncertainty to be expected for the measurements of ice considering size and uncertainty of the procedural blank compared to the system blank correction. In any case, due to the benefit of the small sample size required by our method, execution of replicate measurements is feasible even though the estimated size and uncertainty of the procedural blank compared to the system blank correction is expected to be too high. The sample history discussed before (i.e. mechanical pre-treatment and transport) may be another contributing factor. Considering the differences in sample size and resolution used at CSIRO and CIC, this may be another contributing factor. Considering the differences in sample size and resolution used at CSIRO and CIC, this may be another contributing factor.

CO₂ exchange with the terrestrial biosphere dominates the signal of the \(^{18}\text{O}/^{16}\text{O}\) ratio in atmospheric CO₂. Therefore, atmospheric \(^{18}\text{O}-\text{CO}_2\) is a valuable proxy to constrain changes in terrestrial primary production and the hydrological cycle (e.g. Farquhar et al., 1993). However, \(^{18}\text{O}-\text{CO}_2\) measured in ice cores is affected by the exchange of oxygen with the ice matrix. This has been demonstrated by measurements of samples from different sites and time periods (Friedli et al., 1984; Siegenthaler et al., 1988; Bauska et al., 2014) as well as in firm gas (Assonov et al., 2005).

Here, we present a new approach for analytical quality control, based on \(^{18}\text{O}-\text{CO}_2\) measurements. It allows reliable and consistent rejection of results from samples which were affected by analytical problems or suffered contamination during e.g. storage or gas extraction. Different sections from GRIP and NGRIP with gas ages of 260 to 1770 years (bags GRIP-250 and GRIP-360, NGRIP-213 and NGRIP-696) and 25 000 years (bags GRIP-3628 and GRIP-3636) were analysed in high spatial and temporal resolution of 2.5 cm and < 1 year (ice age scale), respectively. Parallel samples (same depth) were measured for \(^{18}\text{O}-\text{H}_2\text{O}\) in similar resolution if no data existed already. Similar to the previ-
ous studies, high correlation between $\delta^{18}$O-CO$_2$ and $\delta^{18}$O-H$_2$O was observed (Fig. S5). With this dataset, the expected $\delta^{18}$O-CO$_2$ value in the following denoted as $\delta^{18}$O-(CO$_2$-ice)$_{eq}$ could be calculated for each sample from its corresponding $\delta^{18}$O-H$_2$O value, considering oxygen exchange in thermodynamic equilibrium between gaseous CO$_2$ and the surrounding ice matrix. We thereby followed the approach by Siegenthaler et al. (1988), combining the thermodynamic equilibrium oxygen isotope fractionation factors $\alpha$ for (CO$_2$-H$_2$O$_{ij}$), (H$_2$O$_{ij}$-H$_2$O$_{gj}$) and (H$_2$O$_{gj}$-H$_2$O$_{gi}$) given therein. It should be noted that this approach might not be valid particularly for ice from very cold sites because it assumes that those values for $\alpha$ which for obvious reasons could be determined experimentally only for temperatures above the freezing point of water can be linearly extrapolated to lower temperatures ($\leq$ 273.15 K). Using the same approach for a site with annual mean temperatures of $-$26°C (Berkner, Antarctica), the time to reach 50 % of the thermodynamic equilibrium ($T^{1/2}$) was empirically determined to be around 23 years (Assonov et al., 2005). The site temperatures for GRIP and NGRIP are comparable (around $-$30°C). It can therefore be assumed that even the youngest samples analysed in this study have reached complete equilibrium in the glacier. However, the GRIP and NGRIP ice cores were drilled in 1992 and from 1996 to 2004, respectively. From the time the cores have been recovered, a new thermodynamic equilibrium now driven by the storage temperature needs to be considered. For $T^{1/2}$ and storage durations of $\sim$ 15 and 20 years until the time of measurement, the degree of the new equilibrium reached is only 30 and 45 % for NGRIP and GRIP samples, respectively. To take the temperature exposure history of the ice into account, the equilibrium temperature was defined as the weighted mean of borehole temperature at sampling site depth (around $-$30°C; Johnsen et al., 1995; Dahl-Jensen et al., 2003) and freezer temperature ($-$23°C). Accordingly, the freezer temperature was weighted with 0.30 for NGRIP and 0.45 for GRIP. The uncertainty of the mean equilibrium temperature ($-$28°C) was estimated by propagating a 10 % uncertainty for the degree of the equilibrium and an error of 2°C each for borehole and storage temperatures. In Fig. 8, the correlation between the observed and the theoretical $\delta^{18}$O-CO$_2$ is shown. Overall, the agreement between measured and theoretical values is high ($R = 0.90$) with a small offset of 0.05‰ on average. This is a strong indication for the occurrence of oxygen exchange between CO$_2$ and H$_2$O within the ice archive and in agreement with previous observations (e.g. Siegenthaler et al., 1988). The 1σ standard deviation around the theoretical equilibrium line was 0.8‰, exceeding the estimated analytical precision by more than a factor of 2. Based on that, we defined an outlier identification criterion for samples where $\delta^{18}$O-CO$_2$ differed by more than 1.6‰ (2σ) from the theoretical value (Figs. 8 and S5). For these samples, the CO$_2$ and $\delta^{13}$C measured values were rejected from the analyses. In all cases the rejected samples showed suspicious values in both CO$_2$ and $\delta^{13}$C, but because no obvious reason such as issues with the analytical system was noticed (e.g. bad vacuum, trap temperatures being out of range), they could not have been consistently removed otherwise. Unnoticed micro-cracks in the ice might be the most likely explanation for these outliers.

5 Conclusions and outlook

This study describes a new analytical set-up for simultaneous measurements of atmospheric CO$_2$ mixing ratios and atmospheric $\delta^{13}$C and $\delta^{18}$O-CO$_2$ in air extracted from ice core samples. The centrepiece of the system is a newly designed needle cracker for mechanical dry extraction, operated at an extra low temperature of $-$45°C. With this set-up the throughput of three to four samples per day and the small amount of ice required (8–13 g) enables a high resolution.

We discussed analytical procedures, systematic linearity testing for the various system parts, daily calibration as well as data processing. Determined from repeated long-term quality control measurement of air samples over ice (natural and BFI), high analytical precision was achieved, being 1.9 ppm for CO$_2$ and 0.09‰ for $\delta^{13}$C. Law Dome ice core

![Diagram](https://www.atmos-meas-tech.net/9/3687/2016/fig8.png)

Figure 8. $\delta^{18}$O-CO$_2$ used as a quality control tool for ice core measurements. The correlation between measured (x axis) and expected (y axis) $\delta^{18}$O-CO$_2$ is shown for samples from GRIP and NGRIP measured in high spatial and temporal resolution for sections with gas ages of 260 to 1770 years and 25 000 years. The expected $\delta^{18}$O-CO$_2$ was derived considering thermodynamic equilibrium of gaseous CO$_2$ with the surrounding ice matrix and is accordingly denoted as $\delta^{18}$O-(CO$_2$-ice)$_{eq}$. The calculated theoretical equilibrium (black line) is shown with an uncertainty band (grey), accounting for the error associated with equilibrium temperature estimates. The dashed line indicates the 2σ standard deviation around the theoretical value considering all data points shown. For samples outside this range (red dots) CO$_2$ and $\delta^{13}$C-CO$_2$ results were rejected.

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samples were analysed in a laboratory intercomparision study with CSIRO, and good agreement between the two laboratories was found for CO$_2$ and $\delta^{13}$C. Replicate analysis of these samples was performed on different days and resulted in a pooled standard deviation of 2.0 ppm for CO$_2$ and 0.11 %o for $\delta^{13}$C. Due to the properties of the ice analysed, these numbers may be rather conservative estimates of the achieved overall analytical system precision for single measurements. The potential method precision is higher for results derived from replicate measurements, feasible because of the small sample requirement of the system. In conclusion, our system is well calibrated and the precision comparable to other systems using samples of similar small sizes.

Further, a new approach was proposed for the correction of the procedural blank, leading to more accurate results particularly for the measurements of small samples. Analysis of $\delta^{18}$O-CO$_2$ and $\delta^{18}$O-H$_2$O confirmed the previously observed exchange of oxygen between CO$_2$ and the surrounding ice matrix occurring within the archive. Based on this, we introduced a new approach for analytical outlier detection which allows reliable and consistent rejection of results from samples affected by analytical problems or some sort of contamination.

Methodological improvement could be achieved through higher extraction efficiency, further reduction in system blank size by optimizing dimensions of connecting lines and traps and by an increased system automation (gas manifold, vacuum lines).

**Acknowledgements.** This work was partly funded by the Centre for Ice and Climate through the Danish National Research Foundation (DNRF) and the European Union’s Seventh Framework programme (FP7/2007–2013) under agreement no. 243908, “Past4Future. Climate change – Learning from the past climate”. CSIRO’s contribution was supported in part by the Australian Climate Change Science Program, an Australian government initiative. Thanks are due to the teams involved in recovering and analysis of GRIP, NGRIP and Law Dome sample. We highly appreciate the excellent working environment, particularly to Dorthe Dahl-Jensen who made this all happen.

Edited by: R. Koppmann
Reviewed by: two anonymous referees

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