Metrology for low cost CO₂ sensors applications: the case of Steady-State-Through-Flow (SS-TF) chamber for CO₂ fluxes observations

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Abstract. Soil CO₂ emissions are one of the largest contributions to the global carbon cycle, and a full understanding of processes generating them and how climate change may modify them is needed and still uncertain. Thus, a dense spatial and temporal network of CO₂ flux measurements from soil could help reduce uncertainty in the global carbon budgets.

In the present study the design, assembling and calibration of low cost Air Enquirer kits, including CO₂ and environmental parameters sensors, is presented. Different type of calibrations for the CO₂ sensors and their associated errors are calculated. In addition, for the first time this type of sensors have been applied to design, develop and test a new Steady-State-Through-Flow (SS-TF) chamber for simultaneous measurements of CO₂ fluxes in soil and CO₂ concentrations in air. Sensor's responses were corrected for temperature, relative humidity and pressure conditions in order to reduce the uncertainty of measured CO₂ values and of the following calculated CO₂ fluxes based on SS-TF. CO₂ soil fluxes measured by the proposed SS-TF and by a standard closed Non-Steady-State-Non-Through-Flow (NSS-NTF) chamber were shortly compared to ensure the reliability of the results.

The use of a multi-parametric fitting reduced the total uncertainty of CO₂ concentration measurements by 62% compared with the uncertainty if a simple CO₂ calibration was applied, and by a 90% when compared to the uncertainty declared by the manufacturer. The new SS-TF system allows continuous measurement of CO₂ fluxes and CO₂ ambient air with low cost (~1.2 k€), low energy demand (<5W) and low maintenance (twice per year due to sensor calibration requirements).

1 Introduction

Global soils store at least twice as much carbon as Earth’s atmosphere (Oertel et al., 2016; Scharlemann et al., 2014), and act as sources and/or sinks for greenhouse gases (GHGs) such as carbon dioxide (CO₂), methane (CH₄), and nitrous oxide (N₂O).

The total global emission of CO₂ from soils is recognized as one of the largest contributions in the global carbon cycle and is, among others, temperature dependent (Bond-Lamberty and Thomson, 2010a). However, soil respiration is probably the least
well constrained component of the terrestrial carbon cycle (Bond-Lamberty and Thomson, 2010b; Schlesinger and Andrews, 2000) and the degree to which climate change will stimulate soil-to-atmosphere CO2 flux remains highly uncertain (Pritchard, 2011). Continuous measurements of soil fluxes are therefore essential to understand changes in soil respiration of ecosystems in relation to climate variables such as atmospheric temperature. A high temporal and spatial resolution monitoring of CO2 fluxes at sensitive areas could offer useful data both for better understanding the processes at the sources and sinks and thus improving biogenic models (Agustí-Panareda et al., 2016; Randerson et al., 2009). In addition, a complete uncertainty budget of CO2 flux measurements will be essential for the evaluation and correction of global flux models and their associated uncertainties.

Gas interchange between the soil and the lower atmosphere is generally measured as the quantity of gas exhaled from the soil per unit of surface and time ($\mu$mol·m$^{-2}$·s$^{-1}$). It can be measured with different techniques, being the most common the Steady-State Through-Flow (SS-TF), also known as open dynamic chamber, and the Non-Steady-State Non-Through-Flow (NSS-NTF) or closed chamber (Pumpanen et al., 2004). In both cases, the CO2 fluxes are measured using a chamber installed on the soil surface. NSS-NTF measurements are based on the rate of CO2 concentration increase within the chamber, while in the SS-TF technique the CO2 efflux is continuously calculated as the difference between the CO2 concentration at the inlet and the outlet under determined hypothesis (Livingston and Hutchinson, 1995). In the case of NSS-NTF flux measurements, calibrated data is not strictly necessary as long as the sensor's calibration does not change during the measurement timespan because the flux is proportional to the slope of the CO2 concentration increase within the chamber. SS-TF based results need high accurate calibration sensors because the absolute value of the measured CO2 concentrations into the chamber are used. A literature survey suggests that generally NSS-NTF may underestimate CO2 fluxes by 4–14% probably due to: i) advective fluxes forced by small pressure gradients between the air into the chamber and outside it; ii) setting configurations, such as the installation depth of the chamber into the soil. No significant difference was observed when fluxes were measured using SS-TF chambers where no pressure gradients are created (Pumpanen et al., 2004; Rayment, 2000).

In recent years, Wireless Sensor Networks (WSN) are increasingly used for real time and high spatial resolution monitoring (Oliveira and Rodrigues, 2011). A WSN is composed of spatially distributed autonomous sensors to monitor physical, chemical or environmental conditions, and to cooperatively pass their data through the network to other locations. WSN can be used for local data recording for later analysis or for continuous transmission in real time to a remote laboratory for synchronous analysis.

So far low-cost sensors for CO2 atmospheric measurements have been largely used in industrial environments and for indoor air quality and ventilation rate studies (Fahlen et al., 1992; Mahyuddin and Awbi, 2012; Schell and Int-Hout, 2001). When low cost sensors are applied at high CO2 concentration areas and/or spots where air concentrations observed are in the order of thousands of parts per million (ppm), the total uncertainty of the measurement does not affect the quality of the study of the concentration variability under different conditions and sources/sinks. However, in the last decade, the improvement in precision and cost decrease of Non Dispersive InfraRed (NDIR) CO2 sensors have made them more useful for multiple purposes (Yasuda et al., 2012). Their low weight and dimensions allow their utilization in a wide variety of applications,
including Unmanned Aerial Vehicles (Kunz et al., 2018), CO₂ measurements network areas (Kim et al., 2018; Song et al., 2018) and for the study of the distribution of CO₂ in large regions, as in the case study of Switzerland (Müller et al., 2020). However, in order to be able to use these sensors in the outdoor atmosphere, a metrological effort is needed to: i) ensure a traceable and stable calibration; ii) evaluate and correct the influence of the environmental parameters, such as temperature, relative humidity and pressure, on the sensor response; iii) estimate the total uncertainty related with the sensors calibrations and corrections.

This work presents a low cost Air Enquirer Kit, including NDIR CO₂ and environmental parameters sensors and suggests new possible applications of it to reduce the cost and the maintenance of continuous CO₂ fluxes. The manuscript presents the results of the comparison of different calibration methodologies for NDIR CO₂ sensors. Furthermore, a new SS-TF system, based on 5 multi-sensors portable Air Enquirer Kits, is presented and shortly compared with a NSS-NTF system at a Spanish mountain site. The system has been designed and built to continuously monitor soil CO₂ fluxes with high temporal resolution, high accuracy and low cost and maintenance. This system also allows continuous measurements of ambient CO₂ concentration. The SS-TF is made by four Air Enquirer Kits fully characterized under laboratory conditions. The new prototype of the SS-FT chamber is also introduced after describing its theoretical basis as well as the NSS-NTF method. Finally, the results of the sensors calibrations and corrections and of the short NSS-NTF/SS-TF chambers comparison are presented and discussed together with further research steps.

2 Methods

2.1 Air Enquirer Kit

A multi-sensor portable kit, named Air Enquirer (Morguí et al., 2016), was designed and built in the mark of an EduCaixa project (www.educaixa.org). The kit consists of 5 low cost sensors controlled by an Arduino DUE Rev3 microcontroller board that measure: i) NDIR CO₂ concentration (in ppm); ii) relative humidity (%); iii) temperature (°C); iv) barometric pressure (hPa) and v) light intensity (lux). Data from sensors are automatically read and stored at a frequency of 0.2Hz in a microSD card. All sensors and the Arduino board controlling them are enclosed in a methacrylate box of 15x8x5 cm³ in size (Fig. 1). Table 1 shows the main features of each sensor, following specifications provided by their respective manufacturers. The total cost of each Air Enquirer (AE) kit is about 200€.

2.2 Calibrations and multi-parametric correction of the CO₂ sensors of the Air Enquirer kit

Low-cost CO₂ sensors are known to be temperature (T), humidity (H) and pressure (P) dependent (Arzoumanian et al., 2019; Martin et al., 2017). In this study, five AE kits were calibrated using different methodologies from the literature and their responses were corrected under different climate conditions. The simultaneous use of the CO₂ and the environmental parameters sensors allows a continuous correction of the response of the CO₂ sensor under different conditions of T, P and relative humidity (RH).
First of all, a theoretical correction of the CO₂ data was applied taking into account: i) the change from ppm of CO₂ in wet air to ppm of CO₂ in dry air following Wagner and Pruß, (2002); ii) the conversion from ppm of CO₂ measured under specific pressure to the declared using the ideal gas law equation.

The concentration of CO₂ in dry air ($C_{CO₂, dry}$) was calculated by Eq. (1):

$$C_{CO₂, dry} = \frac{C_{CO₂, wet} \cdot 1013}{V_{dry} \cdot p}$$  \hspace{1cm} (1)

being $V_{dry}$ the Volume of 1m$^3$ of dry air at 1013 hPa after removing the water volume. $V_{dry}$ can be calculated from Eq. (2):

$$V_{dry} = \frac{p-(p_{ws} \cdot \frac{RH}{100})}{p}$$  \hspace{1cm} (2)

being $P_{ws}$ the water vapour saturation directly calculated from Eq (3):

$$P_{ws} = A \cdot 10^{\left(\frac{m \cdot T_n}{T+T_n}\right)}$$  \hspace{1cm} (3)

$A$, $m$ and $T_n$ are constants with values 6.1164, 7.5914 and 240.73 respectively.

In a second step, an experimental multiparametric calibration of the CO₂ sensors was done using the data of the environmental sensors and a reference CO₂ instrument. A Picarro G2301 Cavity RingDown Spectroscopy Analyzer (CRDS) was used as a second reference standard. This CRDS has a precision better than 0.03 ppm for CO₂ (Crosson, 2008; Richardson et al., 2012).

The CRDS results were previously corrected for water vapour (Rella et al., 2013) and calibrated in the laboratory using six NOAA WMO-CO2-X2007 reference gases (primary standard) before and after each experiment following Tans et al. (2011). In order to calibrate the CO₂ sensors response for a wide range of temperature, pressure, humidity and CO₂ concentration, duplicate measurements were carried out using a temperature controlled box at two sites: i) at the Institut de Ciències del Clima laboratories (IC3), located at 20 meters above sea level (m.a.s.l.), in the city of Barcelona, Spain, and ii) at the Centre d’Recerca d’Alta Muntanya laboratories (CRAM, mountain town of Vielha, Spain, at 1582 m.a.s.l.). Each experiment lasted 7 days and was carried out using the scheme in Fig. 2. In order to remove high frequency variability, the sampled air was homogenised in a sealed pre-chamber prior to entering in the calibration chamber. Then, the air was pumped to the calibration box at a flow rate of 0.4 L·min$^{-1}$ and through the secondary standard reference instrument CRDS. Both experiments were performed in a temperature range between 20 ᵒC and 42 ᵒC and a relative humidity with diurnal cycles between 10% and 50%.

Temperature in the calibration box was set to be in increased in slopes of 10⁰C, although at low temperatures it fluctuated with room temperature. The pressure ranged between 1004 hPa and 1012 hPa in the calibration at IC3 and between 838 hPa and 850 hPa in the calibration at CRAM. The two calibration experiments at the CRAM and at IC3 stations were carried out with one month difference.

CO₂ concentration values measured by each NDIR CO₂ sensor and corrected for P and RH using Eq. (1) ($C_{CO₂, dry, kit}$), were calibrated by comparison with simultaneous CO₂ concentration measured by the CRDS ($C_{CO₂, CRDS}$) and considering the environmental conditions of T, absolute humidity (H) and P using Eq. (4):
\[ C_{O_2\, dry,\, kit} = \alpha + \beta C_{O_2\, CRDS} + \gamma T + \delta H + \varepsilon P \]  

(4)

A multiparametric fit of Eq. (4), yields the following calibrated/corrected CO\(_2\) values as reported in Eq. (5):

\[ C_{O_2\, corr} = \frac{-a}{\beta} + \frac{1}{\beta} C_{O_2\, dry,\, kit} - \frac{\gamma}{\beta} T - \frac{\delta}{\beta} H - \frac{\varepsilon}{\beta} P \]  

(5)

The \( C_{O_2\, corr} \) calibrated results were compared to those obtained with a simple bias correction using the averages of \( C_{O_2\, CRDS} \) and \( C_{O_2\, dry,\, kit} \) values and also to those obtained with a simple linear calibration of the \( C_{O_2\, dry,\, kit} \) values with the \( C_{O_2\, CRDS} \) values without taking in consideration the effect of T, P and H.

2.3 Steady-State Through-Flow chamber (SS-TF or Open Dynamic Chamber)

The prototype of the open SS-TF chamber consists of two methacrylate cells of 36 L, where two AE kits are installed in each of the chambers in order to continuously monitor the CO\(_2\) concentration and environmental variables. The duplicity of the AE kits is used to ensure the reliability of the measurements. The chamber dimensions were designed to avoid border effects and minimize measurement errors, as observed by Seneviratna et al. (2007). The first chamber is a hermetic closed chamber with a unique entry for ambient air (labelled here as Mixing chamber in Fig. 3). The second one (labelled here as Flux chamber), with an open base, was installed directly over the soil.

The Mixing chamber is used to mix the sampled air and to measure the CO\(_2\) concentration background of the atmospheric air \((C_{mix})\) before it enters into the Flux chamber. It contains two AE kits and a fan located at its top for mixing the sampled air. This chamber has only two openings for the inlet and outlet of atmospheric air at a flow of 6.5 L·min\(^{-1}\) (labelled 'q' in Fig. 3). Cable glands are used at the openings to prevent leakages. Using this configuration, high frequency variability of atmospheric air could be avoided and near steady-state conditions were reached.

The Flux chamber is bottomless and has to be positioned in the first 5 cm of the soil/vegetation layer where the soil fluxes are to be measured. Two AE kits and a vent fan were installed at the top of this chamber as well. A constant flow \( q \) between the two chambers was achieved with a membrane KNF pump and a flowmeter (labelled as FM in Fig. 3). Low flows, in comparison with the chamber volume, are needed to maintain near steady-state conditions during measurements.

Using the system depicted in Fig. 3, CO\(_2\) fluxes \((f_{CO_2} \, in \, \mu mol \cdot m^2 \cdot s^{-1})\) can be calculated for given time intervals within the Flux chamber using the mass balance in Eq. (6) (Gao and Yates, 1998), where, \( V \) and \( A \) are, respectively the volume of the Flux chamber and the emitted soil surface area, \( C_a(t) \) (\( \mu mol \cdot L^{-1} \)) is the spatially averaged concentration of target gas in the chamber headspace, \( C_{in}(t) \) (\( \mu mol \cdot L^{-1} \)) is the average CO\(_2\) concentration of inlet air in the flux chamber, \( C_{out}(t) \) (\( \mu mol \cdot L^{-1} \)) is the outflow CO\(_2\) concentration, \( J_g \) is the flux of the target gas at the enclosed soil surface and \( q_{in} \) and \( q_{out} \) are the inlet and outlet flow, respectively.

\[ dM(t) = VdC_a(t) = AJ_g(t)dt + q_{in}C_{in}(t)dt - q_{out}C_{out}(t)dt \]  

(6)
Assuming that for each measurement interval: i) the inflow and outflow rates are constant and equal (meaning no leakages present in the pneumatic circuit), thus $q_{in} = q_{out} = q$; ii) chamber reach a steady state condition, thus $C_{in}(t) = C_{in}$, $C_{out}(t) = C_{out}$ and $dM(t) = 0$, CO₂ flux can be calculated for each time interval from the simplified Eq. (7):

$$f_{CO₂} = J_g = \frac{q}{A} (C_{out} - C_{in})$$  \hspace{1cm} (7) 

Assuming that the fan completely mixes the air within the chamber and the CO₂ concentration at each of the boxes is homogeneous, outflow concentration is equal to Flux chamber concentration ($C_{out}(t) = C_{\alpha}(t)$), measured by the two AE kits within the flux chamber and inflow concentration is equal to the mixing concentration ($C_{in}(t) = C_{mix}(t)$), measured by the two AE kits within the Mixing chamber. The advantage of this system is that fluxes can be measured continuously with a very small energy requirement (<5 W) and, even using duplicate sensors, with a relative low cost (~1.2k€) in comparison with other automatic commercial flux chambers, priced at roughly 12 k€. The new system described here enables the feasibility of a network of continuous measurements and a replication of experiments to cope with soil flux variability.

2.4 Non-Steady-State Non-Through-Flow chamber (NSS-NTF)

CO₂ fluxes using the NSS-NTF chamber or closed static chamber are measured on the basis of the so-called linear accumulation method (Livingston and Hutchinson, 1995), which uses the initial rate of concentration increase in an isolated chamber that has been placed on the soil surface for a known period of time. Assuming ideal gas behaviour, the slope of the CO₂ concentration during the accumulation interval can be used to determine the CO₂ flux ($\mu$mol·m⁻²·s⁻¹) following Eq. (8):

$$f_{CO₂} = J_g = \frac{CO_2,\text{slope} \cdot P \cdot V}{A \cdot T \cdot R}$$  \hspace{1cm} (8) 

where $V$ (m³) and $A$ (m²) are the volume of the chamber and the enclosed soil surface area respectively, $CO_2,\text{slope}$ (ppm·s⁻¹) is the slope of the linear increment of the CO₂ concentration during the early accumulation time, $P$ and $T$ are the atmospheric pressure and the environmental temperature within the chamber, and $R$ (m³·Pa·K⁻¹·mol⁻¹) is the universal gas constant. It has been underlined that the linear approach of the accumulation method is only reliable for short time periods (Davidson et al., 2002; Grossi et al., 2012; Gutiérrez-Álvarez et al., 2020). Otherwise, gradients of environmental parameters between the inside and outside chamber could influence the measurement, probably yielding to leakages of unknown origin in the chamber. Luckily, high frequency measurements, as the ones performed by CO₂ sensors, allow to apply this method over a really short accumulation time ($T = 5$ min has been used in the present study), thus complying with the theoretical requirements. A necessary condition for the application of this method is that the initial CO₂ concentration within the chamber has to be equal to the atmospheric CO₂ concentration. Therefore, NSS-NTF chambers need to be ventilated after each measurement period (Davidson et al., 2002; Xu et al., 2006). This can be done manually or using automatic systems. In this study, a manual static chamber was used. A closed NSS-NTF chamber of methacrylate (25x25x25) cm³ was built at IC3 in order to perform a short campaign for the comparison of CO₂ fluxes measured by NSS-NTF and SS-TF systems. An AE (#03) and a fan were fastened at the top of the chamber. Both devices were run by a small external battery pack. An outer metallic sleeve was previously
fixed onto the soil to avoid leaks and other disturbances. However, the systemic comparison between these two systems is beyond the scope of this study.

3 Results and discussion

3.1 Comparison between different calibration/corrections approaches

The calibration and correction factors from Eq. (5) of the CO2 sensors installed in the five AE kits are shown in Table 2. The average bias (in ppm CO2) between the AE kit CO2 value after and before applying the theoretical corrections for P and dry air is also shown. The last five columns of Table 2 present, for the different methodologic approaches, the calculated Root Mean Square Error (RMSE) using Eq. (9):

$$\sqrt{\frac{\sum_{i=1}^{n}(x_i^p-x_i^k)^2}{n}}$$

where n is the number of values, $x_i^p$ are the CO2 values of the calibrated CRDS and $x_i^k$ are the CO2 values of the AE sensor for each case: kCO2: uncalibrated values; kCO2_dry: values corrected only for P and dry air; kCO2_dry-bias: values corrected for P and dry air and with the average bias from the CRDS data removed; kCO2_linear: values corrected for P and dry air and linearly calibrated with the CRDS data; kCO2-multi values corrected for P and dry air and calibrated with the CRDS data using a multiparametric correction with T, RH and P sensors data.

A single theoretical correction for P and RH is demonstrated that already reduces uncertainty by a factor of 5. However, this theoretical correction is not enough for applications where the absolute CO2 value is needed (e.g. for atmospheric composition or SS-TF measurements), as the bias value is extremely variable depending on the sensor unit and up to 50 ppm. When we remove the average bias between the sensor response corrected for P and RH and the CRDS CO2 reference value, the uncertainty is highly reduced and the RMSE of the corrected values ranges between 5.4 ppm and 10.8 ppm. This uncertainty, however, could still be too high for certain applications such as the measurements of small atmospheric variability or for small CO2 fluxes measurements both for the SS-TF and NSS-NTF chambers.

Calibrating these sensors through comparison with the CRDS secondary standard in the laboratory by linear fit allows reaching RMSE_simple values between 4.2 ppm and 10.9 ppm. However, when the influence of the environmental parameters in the response of the sensors is also taken into account, the RMSE_multi values range is shifted to the interval between 2.19 ppm and 5.92 ppm, the lowest ones. Figure 4 shows timeseries of the differences between the CO2 CRDS data and all CO2 sensors data after applying the simple calibration (CO2_linear) and the multiparametric regression (CO2_multi). Corresponding values of T and RH measured during the calibration experiments are also reported. Each CO2 sensor responses differently to the variations of T and RH, and so does the parametric coefficients. Therefore, a theoretical correction of the CO2 value for these variables won’t be applicable, and a specific multiparametric fitting is needed.

Figure 5 shows the relation between the reference CO2 values (CRDS) and the values measured by the CO2 sensors both for raw data than after the application of the different calibration methodologies. Four sensors show RMSE_multi values lower than
5 ppm, and just one of them (kit #04) greater than 5 ppm. However, this last sensor shows negative correlation with the ambient
temperature, unlike all the others where the CO2 values increased as temperature went up. Despite this kit was lately installed
within the CO2 fluxes chamber for the second part of the study, results from it were not used.

A variance and covariance analysis were also performed to check the influence of meteorological parameters on the CO2 sensor
response. A clear influence of temperature (T), absolute humidity (H) and pressure (P) was observed on the CO2 sensor's
response (p-value: < 10^{-6} for all variables). No cross-correlation was observed among variables. It is important to remark that
although the multiparametric calibration was done after applying the theoretical correction for P and RH, as explained
previously, pressure conditions have the highest influence on the sensor response. In fact, a reduction of 62% in the RMSE
was observed when pressure correction was applied. Moreover, parametric values for P diverge between sensors, so every
sensor seems to be differently influenced by atmospheric Pressure.

### 3.2 Comparison between the NSS-NTF and SS-TF systems

The new prototype of the SS-TF system, described in section 2.2, was shortly tested in a grassland area of the Pyrenees, near
CRAM, between the 1st and the 2nd of June of 2016 and compared with a manual NSS-NTF system. CO2 fluxes ($f_{CO2}$) were
 calculated for both SS-TF and NSS-NTF systems, using Eq. (7) and Eq. (8), respectively.

CO2 concentrations from each of the sensors installed in the SS-TF chamber (upper panel) and the corresponding calculated
$f_{CO2}$ time series (lower panel) are shown in Fig. 6. The differences between the ten minutes average of CO2 concentrations
measured by the two sensors within the Mixing chamber (AE Kits #1 and #2) were of 2.2 ± 5.3 ppm. This difference is coherent
with the RMSE$_{multi}$ of both sensors, and remains stable over time. The differences between the ten minutes average of CO2
concentrations measured by the two sensors within the Flux chamber (AE Kits #3 and #4) were greater (20 ± 8 ppm) and
temperature dependent with a significant correlation (p-value < 10^{-16} and $r^2$=0.95). As CO2 values of kit #4 were found to have
a different behaviour during the calibration events and the RMSE$_{multi}$ was greater than 5 ppm, values of this kit were discarded.

Each value of flux has been calculated using Eq. (7) and averaging the calibrated CO2 values of AE #1 and #2 for the mixing
chamber and using the calibrated CO2 values from AE #3 for the flux chamber. 10 min. averages were calculated from every
minute calculated flux data. The variability of the flux within the 10 minutes averages is represented in Fig. 6 as an associated
uncertainty of 2σ. The associated expanded uncertainty for each value has been calculated propagating the 2*RMSE$_{multi}$ of
the flux chamber CO2 sensor.

CO2 fluxes using the NSS-NTF chamber were calculated using the slope of the increase of the CO2 concentration within the
chamber and its associated uncertainty. Two examples of the CO2 concentrations measured by the CO2 sensor of kit #03 within
the NSS-NTF chamber (see section 2.3) are shown in Figure 7. Data of the first minute after manually closing the chamber
were discarded during the $f_{CO2}$ calculations in order to remove installation noise. Concentration gradients were linear over the
following 5 minutes, with a correlation coefficient $R^2$ >0.99 in all cases as calculated with Eq. (5). Positive fluxes were
measured during the afternoon and negative ones at morning as expected because of the photosynthesis phase of grassland plants.

The correlation between both NSS-NTF and SS-TF $f_{CO_2}$ results during the co-measurements carried out at CRAM grasslands during the 1st and the 2nd of June of 2016 is shown in Figure 8. CO$_2$ flux values change from close to zero up to 8 $\mu$mol·m$^{-2}$·s$^{-1}$. The obtained $f_{CO_2}$ values agree with CO$_2$ flux values observed in other studies in grasslands at a similar altitude, latitude and period of the year, where the range of night-time fluxes was reported to be between 2 and 4 $\mu$mol·m$^{-2}$·s$^{-1}$ (Bahn et al., 2008; Gilmanov et al., 2007). Although the short duration of this first comparison experiment, results help to strengthen the reliability of the new SS-TF chamber based on low cost sensors. However, the size of the comparison dataset does not allow a robust statistic and further long-term comparison should be carried out to fully characterize this new system. Indeed, the main goal of the present manuscript is not characterized the new SS-TF chamber but to offer a robust metrology for low cost CO$_2$ sensors and AE kits which can be easily applied for continuous CO$_2$ flux measurements with high precision, low cost and low maintenance.

CO$_2$ fluxes observations from NSS-NTF and SS-TF chambers agree for positive CO$_2$ fluxes while they do not for negative CO$_2$ fluxes. A plausible cause of this mismatch may be the different degree of opacity of the chambers which influence the sink effect of the soil during the sunlight hours. In fact, the NSS-NTF chamber was completely translucid while in the SS-TF chamber the top side was opaque.

### 3.3 Calibration and recalibration strategy

According to the RMSE results shown in Table 2, the multiparametric correction reduces the uncertainty of CO$_2$ measurements by a factor of 10 compared to those where only a theoretical correction for RH and P was applied and by a factor of 3 compared to a lineal calibration for CO$_2$. In the SS-TF, the flux calculation depends on the difference between the absolute concentrations values of different sensors in two chambers, and a bias between them of e.g. 10 ppm will cause, in this system, a fixed bias of 0.32 $\mu$mol·m$^{-2}$·s$^{-1}$ in the flux calculus. Therefore, the multiparametric correction of sensors for this application is strongly recommended, together with a periodical recalibration of the CO$_2$ sensors. Previous works with NDIR sensors have shown that at least every 6-months may be necessary to calibrate the sensors in order to take into account possible effects due to dust and soiling on their internal mirrors (Curcoll et al., 2019; Piedrahita et al., 2014) or the degradation of the IR light (CO2Meters, 2013). A mobile second reference standard could be displayed to perform in situ calibration of the low cost sensors. However, a periodical full calibration and calculation of correction factors for all environmental parameters could be difficult to carry out at field sites, and may even cause large errors if the range of temperature, humidity and pressure used is not large enough. For those cases where a full multiparametric recalibration couldn’t be performed each six months, a bias correction should be performed at least every six months. This could be done by placing CO$_2$ sensors in a mixing chamber at the same time and introducing air from a reference tank with known CO$_2$ concentration. Thus, taking in consideration the Eq. 4, this calibration will only adjust the $\alpha$ parameter, considering the effects of P, T and RH constant over the time.
For NSS-NTF applications, where only the slope of the CO₂ concentration is used, the bias has no effect on the calculus of the soil flux. Therefore, for this last case periodical corrections for the low cost sensors are not needed although they are advisable to improve the quality of the measurements. Finally, when no calibrations are possible, the recommendation is to calculate the CO₂ concentration in dry air and compensate for pressure. Actually, comparing NSS-NTF based flux data, only a difference of about 4% is observed when theoretical correction for P and RH or multiparametric calibration data are compared. However, when using the CO₂ AE kits values without any correction this difference rises up to a 23%.

4 Conclusions

Nowadays the improvement in precision and cost decrease of Non Dispersive InfraRed (NDIR) CO₂ sensors have made them more readily available for multiple purposes. However, in order to apply them for atmospheric measurements where low CO₂ concentrations or small CO₂ variability is observed a robust metrology is still needed to: i) ensure a traceable calibration; ii) evaluate and correct the influence of the environmental parameters on the sensor response; iii) estimate the total uncertainty related with the measurements.

In this study an analysis of different calibration methods is carried out for NDIR low cost CO₂ sensors using Air Enquirer kits, designed and built, including also environmental sensors. In addition, a new application of these sensors is presented to continuously measure CO₂ fluxes on soil with a dynamic chamber.

The lowest uncertainty for the CO₂ sensors was obtained by calibrating them using a secondary standard reference (CRDS monitor) and correcting the sensors response under different temperature, humidity and barometric pressure conditions. A multiparametric fitting was applied to calibrate and correct the sensor's responses, achieving a drastic reduction of 90% in the uncertainty of measured CO₂ concentrations. The multiparametric calibration will ensure the highest quality of the data and it will be advisable for SS-TF based CO₂ flux measurements or CO₂ atmospheric concentrations. For NSS-NTF based CO₂ flux measurements, a correction for P and RH of the CO₂ sensors will already give reliable results, although calibrating the sensors with a portable second reference standard is recommended.

The presented SS-TF chamber based on Air Enquirer kits allows continuous measurement of CO₂ fluxes from soil and continuous ambient air CO₂ concentration with low uncertainty, low cost (~1.2 k€), low energy demand and low maintenance. This system could be a good tool for creating CO₂ flux dense networks. In the present study it has only been shortly compared with a NSS-NTF chamber at Pyrenees area, showing CO₂ fluxes comparable between them and in agreement with the literature. However, a full characterization of this system needs to be carried out in the future by long-term comparison with commercial CO₂ flux systems.

Code availability

The software code for this paper is available from the corresponding author.
Data availability

The data for this paper are available from the corresponding author.

Author contributions

Josep Anton Morguí coordinated the design and manufacture of the AE kits, and promoted the building of the new low cost SS-TF chamber for CO₂ fluxes. Lidia Cañas collaborated in the mounting and tuning of the AE kits. Armand Karrang, during his bachelor degree project, participated in the laboratory and field campaigns. Roger Curcoll and Claudia Grossi, performed the laboratory and field experiments, analysed the data and coordinated the manuscript writing. Arturo Vargas participated in the development theoretical approach of the SS-TF methodology for gas fluxes. All authors participated in the data analysis, discussion of the results and writing of the manuscript.

Competing interests

The authors declare that they have no conflict of interest.

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References


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<table>
<thead>
<tr>
<th><strong>Measurement (Units)</strong></th>
<th><strong>Manufacturer</strong></th>
<th><strong>Accuracy</strong></th>
<th><strong>Range of measurement</strong></th>
<th><strong>Operating Temperature (°C)</strong></th>
<th><strong>Operating Relative Humidity (%)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂ (ppm)</td>
<td>CO₂ Engine K30 STA – Sense Air</td>
<td>±30 ppmCO₂</td>
<td>0 to 5000</td>
<td>0 to 50</td>
<td>0 to 95</td>
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<tr>
<td>Temperature (°C)</td>
<td>DS18B20 – Dallas</td>
<td>±0.5°C (within range -20 - +85°C)</td>
<td>-55 to +125</td>
<td>-55 to +125</td>
<td>-</td>
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<tr>
<td>Relative Humidity (%)</td>
<td>SparkFun HTU21D – Measurement Specialties</td>
<td>±2% (within range 20-80%)</td>
<td>0 to 100</td>
<td>-40 to +125</td>
<td>0 to 100</td>
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<td>Barometric pressure (hPa)</td>
<td>Adafruit BMP180 - Bosch</td>
<td>±1.0 hPa</td>
<td>300 to 1100</td>
<td>-40 to +85</td>
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<td>Light intensity (visible/IR)</td>
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<td>-</td>
<td>-30 to 70</td>
<td>0 to 60</td>
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</tbody>
</table>

**Table 1. Characteristics of the sensors included within the Air Enquirer kit.**

<table>
<thead>
<tr>
<th><strong>Kit</strong></th>
<th><strong>Intercept</strong></th>
<th>CO₂ Pred</th>
<th>T (°K)</th>
<th>H (hPa)</th>
<th>P (hPa)</th>
<th>Bias</th>
<th>Root Mean Square Error (ppm CO₂)</th>
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</thead>
<tbody>
<tr>
<td>(code)</td>
<td>α/β</td>
<td>1/β</td>
<td>γ/β</td>
<td>δ/β</td>
<td>ε/β</td>
<td>ppm CO₂</td>
<td>kCO₂</td>
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<tr>
<td>#01</td>
<td>59.15</td>
<td>1.1047</td>
<td>-0.395</td>
<td>-6.2·10⁴</td>
<td>-0.084</td>
<td>9.5</td>
<td>76.0</td>
</tr>
<tr>
<td>#02</td>
<td>52.53</td>
<td>1.0564</td>
<td>-1.594</td>
<td>-1.04·10⁴</td>
<td>-0.083</td>
<td>51.4</td>
<td>43.7</td>
</tr>
<tr>
<td>#03</td>
<td>93.22</td>
<td>1.1031</td>
<td>-1.150</td>
<td>-1.05·10⁴</td>
<td>-0.131</td>
<td>21.0</td>
<td>57.8</td>
</tr>
<tr>
<td>#04</td>
<td>49.26</td>
<td>1.0908</td>
<td>1.306</td>
<td>-5.5·10⁴</td>
<td>-0.139</td>
<td>1.8</td>
<td>68.6</td>
</tr>
<tr>
<td>#05</td>
<td>13.55</td>
<td>1.1030</td>
<td>-0.570</td>
<td>-1.17·10³</td>
<td>-0.048</td>
<td>14.9</td>
<td>58.0</td>
</tr>
</tbody>
</table>

**Table 2. Parametric fitting for calibration of CO₂ Air Enquirer sensors**
Figure 1. Air Enquirer kit, with sensors for measurements of temperature, humidity, barometric pressure, light intensity and CO$_2$ concentration in air.

Figure 2. System used at IC3 (Barcelona, Spain) and at the CRAM station (Vielha, Spain) for the calibration of CO$_2$ sensors mounted on the Air Enquirer kits.

Figure 3. Scheme of the Dynamic SS-TF Chamber designed and built at IC3 for continuous CO$_2$ flux measurements.
Figure 4. Timeseries of differences between CRDS CO₂ value and CO₂ AE kits value after simple calibration (grey) and after multiparametric fitting (black) for AE kit #1 (a), kit #2 (b), kit #3 (c) and kit #5 (d). Temperature values (red) and RH values (blue) are also plotted. Values before vertical green line correspond to the calibration at IC3, and after it to the calibration at CRAM.
Figure 5. CO$_2$ concentrations in air measured by each of the AE sensors during the experiment carried out at the CRAM and IC3 stations vs CRDS data using sensor with raw data (a), sensor data theoretically corrected by P and RH (b), sensor data corrected by P and RH and calibrated with the CRDS (c) and sensor data corrected by P and RH and calibrated using a multiparametric lineal model (d)
Figure 6. Time series of 10-min average CO₂ concentrations (upper panel) measured within the SS-TF chamber at the CRAM grassland between 1st and 2nd of June 2016, and calculated $f_{\text{CO}_2}$ (lower panel). The $2\sigma$ range for 10 minutes average variability and the extended error (adding 2 times the RSE of the multiparametric fit) are also plot.
Figure 7. Example of two cases where the linear accumulation method was applied within an NSS-NTF chamber to calculate positive (a) and negative (b) CO₂ fluxes with Kit #03.
Figure 8. Comparison of SS-TF and NSS-NTF CO₂ fluxes during a short campaign at the CRAM station between 1st and 2nd of June 2016.