

Revised methodology for CO₂ and CH₄ measurements at remote sites using a working standard gas saving system

Motoki Sasakawa¹, Noritsugu Tsuda², Toshinobu Machida¹, Mikhail Arshinov³, Denis Davydov³, Aleksandr Fofonov³, Boris Belan³

¹Earth Science Division, Center for Global Environmental Research, National Institute for Environmental Studies, Tsukuba, 305-8506, Japan

²Global Environmental Forum, Tsukuba, 305-0061, Japan

³Institute of Atmospheric Optics, Russian Academy of Sciences, Siberian Branch, Tomsk, Russia

Correspondence to: Motoki Sasakawa (sasakawa.motoki@nies.go.jp)

Abstract. We have revised a calculation method of mole fractions and uncertainties for in-situ CO₂ and CH₄ measurements with a working standard gas saving system. It uses on-site compressed air to track the baseline drift of sensors. JR-STATION (Japan-Russia Siberian Tall Tower Inland Observation Network) was made up of this system, which was installed across nine different sites in Siberia. The system acquires semi-continuous data by alternating between sampling air from multiple altitudes through switched flow paths and recording several minutes of averaged data for each altitude. We estimated the measurement uncertainty (repeatability) in about one week based on the repeated measurement of on-site compressed air. The uncertainty for CO₂ and CH₄ mostly varied by less than 0.2 ppm and five ppb, respectively. Standard deviation (SD) of time-averaged data were sometimes higher than the measurement uncertainty for each period because the data include atmospheric variability during the measurement period of several minutes. Data users should consider the difference between the SD and uncertainty to select optimal data, depending on their focusing spatial scale. The CO₂ and CH₄ data measured with the NDIR and the tin dioxide sensor exhibited good agreement with those measured by the CRDS.

1 Introduction

It is known that accurate measurements of greenhouse gas mole fractions require the analyzers to be calibrated against a set of standard gas mixtures. At least one of them (target) should be used hourly to track an NDIR (Non-Dispersive Infrared) analyzer's baseline drift (Andrews et al., 2014). Delivering high-pressure cylinders to remote sites is a significant issue for long-term atmospheric monitoring. Thus, to reduce the consumption of gas, Watai et al. (2010) developed a system that utilizes on-site air as sub-working standard gas (SWS-gas) to track the baseline drift of the NDIR sensors. Watai et al. (2010) then installed this system at a remote tower site at Berezhnechka (56°08'45"N 84°19'55"E) in West Siberia in 2001 to measure CO₂ mole fractions semi-continuously. After this, in Central Siberia, Winderlich et al. (2010) developed a measuring system without dehumidification using a CRDS (Cavity Ring-Down Spectroscopy) analyzer to reduce the frequency of cylinder replacement.

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The CRDS is a more stable device, and a calibration frequency of every two weeks to every month is recommended (ICOS RI, 2020).

50 Concerning CH₄ measurement, a commonly used gas chromatograph with a flame ionization detector (GC/FID) requires hydrogen and carrier gases. It also needs significant power consumption. However, electric power is often restricted at remote sites. Thus, Suto and Inoue (2010) modified a tin dioxide sensor (TOS), which is widely used to detect natural gas leaks, to be able to measure CH₄ in the atmosphere. The developed TOS unit does not need hydrogen and carrier gases. The nominal power consumption for the unit, consisting of TOS, temperature-stabilizer mechanism, and electronic circuits for the sensor and
55 heater, is less than 10W.

We added the TOS unit to the system at the tower site in West Siberia, then expanded the tower observation network (Sasakawa et al., 2010; Sasakawa et al., 2012; Sasakawa et al., 2013). The tower network named JR-STATION (Japan-Russia Siberian Tall Tower Inland Observation Network) now consists of six tower sites in West Siberia. Recently, we added CRDS analyzers at Karasevov (58°14'44"N 82°25'28"E) in 2015 (Picarro G2401), and at Demyanskoe (59°47'29"N 70°52'16"E) and Noyabrsk
60 (63°25'45"N 75°46'48"E) in 2016 (Picarro G2301) to improve the robustness of the measurements.

The system follows the operational method conceived under the instrumentation constraints a quarter of a century ago (around 2000) and in remote areas with limited infrastructure (Watai et al., 2010). However, Watai et al. (2010) did not present an optimal calculation method for the measurement sequence of this system (nor did the TOS calculation method). Nor did they calculate uncertainties, especially as has been recommended in the GAW report (2020) in recent years.

65 Thus, we have updated the calculation method for calculating CO₂ and CH₄ mole fractions to derive their uncertainty for each data set simultaneously. Here, we describe the details of the modified measurement system and the calculation method. Furthermore, some sites have installed CRDS, which have been widely used for greenhouse gas observations (Kwok et al., 2015) and allow partial comparisons with conventional sensors, so the recalculated data were compared with the CRDS data to see how well they agree.

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70 2 Method

2.1 Measurement system

Ambient air was taken from air sample inlets at two different heights (four at Berezorechka) on television and radio-relay communication towers (Table 1). Each sample inlet was mounted several meters away from the tower at the end of an extension arm. The air from the inlets was pulled into the measurement system through the sampling lines (6-mm OD Decabon tube).

75 The measurement system was housed in a freight container insulated to reduce temperature variation. Two thermometers were mounted inside the container, one near the ceiling and the other near the floor. According to the upper thermometer, the room temperature in the container during the year was kept above 15°C and the temperature difference in the 12-hour calibration interval was kept below 3°C on average during the year. Since the introduction of the CRDS, a simple cooler was installed to prevent the temperature inside the container from rising too high during the summer months due to the heat generated by the

85 CRDS. A schematic diagram of the measurement system is shown in Fig. 1. The measurement system consists of a pump unit,
a selector unit, and an analyzer unit. The pump unit was located upstream of the selector and analyzer unit to keep the
downstream pressure higher than the ambient, which reduced the likelihood of bias in measurements due to any leak from
many connections in the system. Two diaphragm pumps (model N86KNE, KNF, Germany) delivered the sample air into the
system. The sampling lines were flushed continuously with a flow rate of about seven standard liters per minute, and excess
90 air was vented through the back-pressure valve ("BPV" in the pump unit). Then the air was dried by an adiabatic expansion in
a glass water trap ("WT" in the pump unit) that was purged every hour via an NC solenoid valve, which was opened twice for
10 seconds to remove the condensed water. The sample air was also dried using a semipermeable membrane dryer (PD-625-
24SS, Permapure, USA) ("Nafion" in the selector unit). The semipermeable membrane dryer removed water vapor from the
pressurized inner tube to an outer tube where the split gas flowed (split sample method). The air from the upper and lower-
95 level inlet, the three working standard gases (WS-gases), and the sub-working standard gas (SWS-gas) were selected through
a 6-port valve manifold. While the WS-gases or the SWS-gas flowed into the analyzer unit, the sample air was exhausted at
the 6-port valve. In the analyzer unit, the sampled air was extra dried with magnesium perchlorate, which was fed into a
stainless steel tube with a dimension of 2 cm in inner diameter and 10 cm in length ("Mg(ClO₄)₂" in the analyzer unit). There
were two tubes, and the flow path of the air switched from one to the other every month. The used magnesium perchlorate was
100 replaced before the next run. After being dried with the magnesium perchlorate, the air retained its dewpoint at around -50 °C
(39 ppm). The dehumidified air was then introduced into an NDIR analyzer (LI-820, LI-COR, USA; LI-7000 was used until
September 2008 at BRZ) at a constant flow rate of 35 standard cubic centimeters per minute (sccm) set by a mass flow
controller (SEC-E40, STEC, Japan). The CO₂ mole fraction was defined as the mole fraction in the dry air, and water vapor
correction was not adopted. After passing through the NDIR, the air flowed into the TOS unit. A chemical desiccant made of
105 P₂O₅ was installed in front of the TOS because it is necessary to keep water vapor below ten ppm in the sample air for this
type of sensor. The sensor was designed to operate in areas lacking the sufficient infrastructure to sustain a conventional
measurement system, such as a significant power source, carrier gas supply, and temperature-stabilized environment. The
sensor has been verified against a gas chromatograph equipped with a flame ionization detector (Suto and Inoue, 2010). We
additionally installed the CRDS (Picarro Inc.) analyzer at Karasevov (G2401) in 2015, and at Demyanskoe and Noyabrsk
110 (G2301) in 2016 to improve the system. The sampled air was split after leaving the 6-port valve, then fed into the CRDS at a
constant flow rate of 35 sccm set by a mass flow controller (SEC-E40, STEC, Japan) through a semipermeable membrane
dryer (model MD-050-72S-1, Permapure, USA). To protect the cavity of the CRDS from an inflow of the dissolved chemical
desiccant (Mg(ClO₄)₂ or P₂O₅) in the accidental case of a broken pump etc., we equipped the CRDS with two poppet check
valves ("PCV" in the analyzer unit). When the pumps in the pump unit stop and only the CRDS pump is running, the flow
115 stops at the PCV upstream of P₂O₅, and the increased suction pressure allows air in the container to enter from the PCV in
front of Nafion. The data from this process has been deleted.

Three WS-gases (STD1, STD2, STD3) were prepared from pure CO₂ and CH₄ (G1 grade, Japan Fine Products Corp. (JFP),
Japan) diluted with purified air (G1 grade, JFP), and their mole fractions were determined against the NIES 09 CO₂ scale

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(Machida et al., 2011) and NIES 94 CH₄ scale. Each scale was established by a series of standard gases prepared by the gravimetric method. Since the pure CO₂ gas is derived from burned petroleum, the isotopic CO₂ composition of the gases shows lighter than atmospheric CO₂. When the NDIR analyzer is calibrated against CO₂ standards with lighter-than-atmospheric CO₂ isotopic composition, the NDIR analyzer measures a lower CO₂ mole fraction in a sample air with known CO₂ mole fraction. The error in the apparent NDIR CO₂ mole fraction depends on its individual sensitivity to the optical filter property. Tohjima et al. (2009) reported that the errors for the three NDIR analyzers range from -0.04 to -0.08 ppm. The CO₂ values measured by the system may appear low in the range of up to 0.08 ppm. Compared to the WMO-CO₂-X2007 scale, the NIES 09 CO₂ scale is consistent within 0.1 ppm (Round Robin 5 and 6 Comparison Experiment). [Since there have yet to be published results with WMO-CO₂-X2019, scale conversion could be the linear function shown in Hall et al. \(2021\) \(X2019 = 1.00079 × NIES09 – 0.142 \(ppm\)\).](#) The NIES 94 CH₄ scale ranges from 3.0 to 5.5 ppb higher than the WMO-CH₄-X2004A scale (Round Robin 5 and 6 Comparison Experiment).

Table 1. Main features of the towers in the network of tall towers used for continuous long-term atmospheric CO₂ and CH₄ measurements over Siberia.

Identifying Code	Location	Latitude	Longitude	Air inlet heights (m)	Elevation at tower base (m a.s.l) ¹
BRZ	Berezorechka	56°08'45"	84°19'55"	5, 20, 40, 80	168
KRS	Karasevoe	58°14'44"	82°25'28"	35, 67	76
IGR	Igrim	63°11'30"	64°24'50"	24, 47	9
NOY	Noyabrsk	63°25'45"	75°46'48"	21, 43	108
DEM	Demyanskoe	59°47'29"	70°52'16"	45, 63	63
SVV	Savvushka	51°19'31"	82°07'42"	27, 52	495
AZV	Azovo	54°42'18"	73°01'45"	29, 50	110
VGN	Vaganovo	54°29'50"	62°19'29"	42, 85	192
YAK	Yakutsk	62°05'19"	129°21'21"	11, 77	264

¹Approximate estimates from Google earth.

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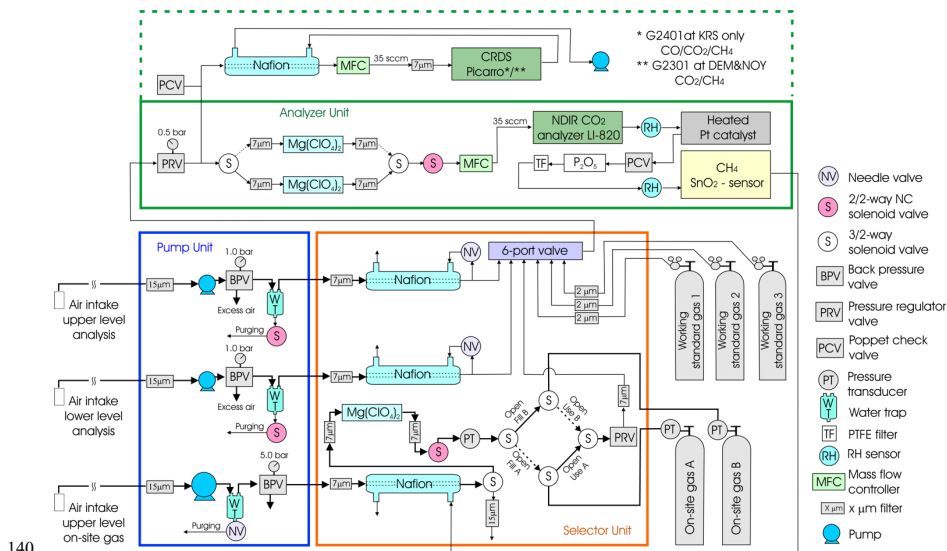


Figure 1. Schematic diagram of tower observation system.

A frequent calibration with WS-gases within 1-2 hours is necessary to conduct precise measurements of CO₂ and CH₄ because the output of the NDIR analyzer or the TOS could vary depending on the environment (atmospheric pressure etc.) in 1-2 hours. But if the calibration were done at this frequency, standard gases would be consumed in less than a year. Because delivering WS-gases to remote sites is a significant issue, we utilized on-site compressed air as SWS-gas to track the sensors' baseline drift, which reduced the consumption of the three standard gases. The on-site compressed air ("On-site gas A/B" in Fig. 1) was analyzed every hour, and the WS-gases were measured every 12 hours to calibrate the sensors (details of the sequence are shown below). The measurement protocol adhered to the procedure established by Watai et al. (2010) for NDIR analysis using this system. However, at sites where the CRDS was installed, the WS-gas measurement interval was extended to 48 hours to prolong the longevity of the WS-gas. An aluminum cylinder (0.048 m³) for SWS-gas was automatically exchanged when the inner pressure decreased below 0.1 MPa, then soon air from the highest inlet was compressed by a pump (LOA-P103-NO, GAST, USA) into the cylinder for about 5 hours to approximately 0.35 MPa, after having been passed through a similar triple dehumidification path as the sampled air (a stainless steel water trap, a semipermeable membrane dryer (SWF- M06-400, AGC, Japan), and magnesium perchlorate). It was preserved for approximately one week (three days with the CRDS) for usage until the inner pressure in one used for measurements decreased below 0.1 MPa (Table 2). Schibig et al.

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(2018) reported that the CO₂ mole fraction in a 29.5 L aluminum cylinder increases by $0.090 \pm 0.009 \mu\text{mol mol}^{-1}$ when dropping from 150 bar to 1 bar, but also note that this change is smaller if larger cylinders are used. Given the cylinder size and filling pressure in this system, mole fraction changes within the cylinder are considered negligible. The variations in SWS mole fraction with CRDS between WS-gases (48 hours) were in fact very stable regardless of the SWS mole fraction range (Fig. S1-2). Air temperature and relative humidity were measured at both heights on the tower using commercial sensors (HMP45D, Vaisala, Finland). A wind monitor (model 81000, R. M. Young, USA) determined wind direction and speed at the higher inlet. Solar radiation was measured by a pyranometer (CM3, Kipp & Zonen, Netherlands), and precipitation by a tipping bucket rain gauge (model 52202, R. M. Young, USA) on the top of the container laboratory.

The analysis operation and data logging were performed by a measurement and control system (CR10X datalogger, CAMPBELL, USA). Stored data were retrieved once a month when both a system check and replacement of consumables (e.g., chemical desiccants) took place.

Table 2. SWS-gas A/B measurement and filling sequence.

<u>Approximate elapsed time (h)</u>	<u>Trigger (inner pressure)</u>	<u>3-way solenoid valve in Figure 1</u>	<u>Cylinder A</u>	<u>Cylinder B</u>
<u>0</u>	<u><0.1 Mpa (B)</u>	<u>Solid line</u>	<u>starts flowing</u>	<u>starts being compressed</u>
<u>5</u>	<u>>0.35MPa (B)</u>	<u>-</u>	<u>-</u>	<u>stop compression</u>
<u>168</u>	<u><0.1 Mpa (A)</u>	<u>Dash line</u>	<u>starts being compressed</u>	<u>starts flowing</u>
<u>173</u>	<u>>0.35MPa (A)</u>	<u>-</u>	<u>stop compression</u>	<u>-</u>
<u>336</u>	<u><0.1 Mpa (B)</u>	<u>Solid line</u>	<u>starts flowing</u>	<u>starts being compressed</u>

2.2 Measurement sequence

To be able to measure air at two heights, the air-sampling flow path was rotated every 20 min with the 6-port valve in the selector unit; that is, the higher inlet was sampled from hh:00 to hh:20, the lower inlet from hh:20 to hh:40, and the SWS-gas from hh:40 to (hh+1):00. During the first 17 min of each 20-min sampling interval, the system is flushing to equilibrate the air sample after switching. The final three-minute readouts were averaged and reported as the representative output data for the applicable one-hour period. Measurement frequency was 3 sec; thus, only the average and standard deviation (SD) of 60 readouts in voltage were stored in the CR10X. This was to minimize the data size for the limited storage capacity. The timestamp was the end time of every 20-min measurement interval. The raw data collected with the CRDS analyzer were stored in the CRDS' hard disk and processed after downloading in our laboratory.

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Figure 2 shows the schematic measurement sequence for half a day. In the Fig. 2, we defined when the SWS-gas was measured just before an arbitrary series of WS-gas measurements as t_0 . Then we numbered the time of the following measurements in turn. We also defined the series of standard gas measurements at the beginning of the 12 hours as “B” and at the end as “E”.

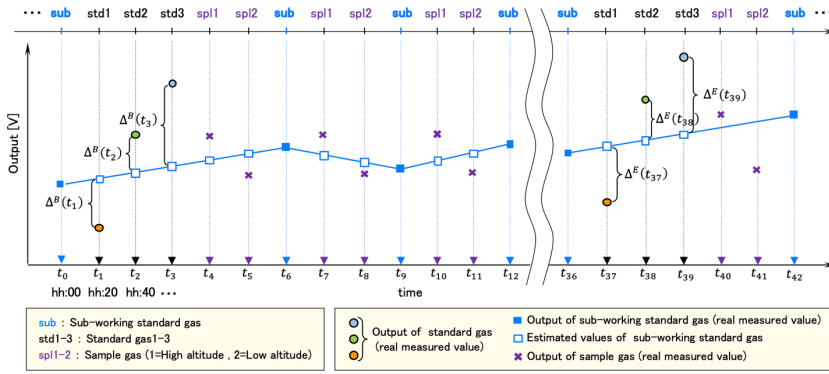


Figure 2. Measurement sequence for a half day between subsequent measurements of WS-gases.

2.3 Quality check of the standard gas measurements

First, we checked the relationship among three standard gas measurements. We calculated the differences ($\Delta^B(t_i)$, $\Delta^E(t_j)$) between the measured output voltages of the standard gases ($V_{std}(t_i)$, $V_{std}(t_j)$) and the estimated one of the SWS-gas at the time of the standard gas measurement (Fig. 2). Here $i = 1, 2, 3$, and $j = 37, 38, 39$. The output value of the SWS-gas was interpolated by time using the closest output of the SWS-gas before and after the series of standard gas measurements. Thus, these values and their variances are expressed as:

$$\Delta^B(t_i) = V_{std}(t_i) - \left(\frac{6-i}{6} \cdot V_{sub}(t_0) + \frac{i}{6} \cdot V_{sub}(t_6) \right) \quad (1)$$

$$(\sigma^B(t_i))^2 = (\sigma_{std}(t_i))^2 + \left(\frac{6-i}{6} \cdot \sigma_{sub}(t_0) \right)^2 + \left(\frac{i}{6} \cdot \sigma_{sub}(t_6) \right)^2 \quad (2)$$

$$\Delta^E(t_j) = V_{std}(t_j) - \left(\frac{42-j}{6} \cdot V_{sub}(t_{36}) + \frac{j-36}{6} \cdot V_{sub}(t_{42}) \right) \quad (3)$$

$$(\sigma^E(t_j))^2 = (\sigma_{std}(t_j))^2 + \left(\frac{42-j}{6} \cdot \sigma_{sub}(t_{36}) \right)^2 + \left(\frac{j-36}{6} \cdot \sigma_{sub}(t_{42}) \right)^2 \quad (4)$$

We estimated the output of STD1 at t_2 by adding $\Delta^B(t_i)$ to the estimated one of the SWS-gas at t_2 . We also evaluated the output of STD3 at t_2 by adding $\Delta^B(t_i)$ to the estimated one of the SWS-gas at t_2 . The same estimation was done at t_{38} . We then made

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a linear calibration line with the output of STD2 and the estimated outputs of STD1 and STD3. Only those sets of the three standard gas measurements whose coefficients of determination were higher than 0.999 for CO₂ and 0.99 for CH₄ were adopted for the following calculation.

The difference in output (voltage) between Δ^B and Δ^E for each standard gas was defined as follows:

$$\delta_{(1,37)} = \Delta^E(t_{37}) - \Delta^B(t_1) \quad (5)$$

$$\delta_{(2,38)} = \Delta^E(t_{38}) - \Delta^B(t_2) \quad (6)$$

$$\delta_{(3,39)} = \Delta^E(t_{39}) - \Delta^B(t_3) \quad (7)$$

The δ must be small unless the system is unstable, e.g., when the sensitivity of the sensors changes considerably for some reason. To exclude the data obtained during system malfunction, we determined a threshold for δ by converting it into **mole fraction** (<5.0 ppm for CO₂, <50 ppb for CH₄). Data showing values over the threshold were excluded from the calculation.

The difference in CO₂ **mole fraction** between sides *B* and *E* was calculated as follows:

$$\overline{\delta_{(i,j)}^B} = |\delta_{(i,j)} / S^B| \quad (8)$$

$$\overline{\delta_{(i,j)}^E} = |\delta_{(i,j)} / S^E| \quad (9)$$

where S^B and S^E are the slopes of the linear regression line at sides *B* and *E*. Because the x-axis of the calibration line for CH₄ is the logarithm of the **mole fraction**, the difference in CH₄ **mole fractions** was calculated as:

$$\overline{\delta_{(i,j)}^B} = C_i \cdot \left| e^{\frac{\delta_{(i,j)}}{S^B}} - 1 \right| \quad (10)$$

$$\overline{\delta_{(i,j)}^E} = C_i \cdot \left| e^{-\frac{\delta_{(i,j)}}{S^E}} - 1 \right| \quad (11)$$

where C_i is the **mole fraction** of the standard gas.

2.4 Calculation of the sample **mole fraction**, **SD**, and **system uncertainty**

The analysis precision for this system under laboratory conditions was uniformly estimated as 0.3 ppm for CO₂ (Watai et al., 2010). Concerning CH₄ precision, Sasakawa et al. (2010) estimated it as 3.0 ppb based on the result of Suto and Inoue (2010). However, the experiment condition by Suto and Inoue (2010) was different from the gas-saving system. Instead, they connected only the WS-gases to the TOS, then reported the **SD** of repeated measurements. The CH₄ analysis precision for this system thus could be more significant than 3.0 ppb. Furthermore, the sensitivity and stability of the sensor could differ depending on the individual sensor and the condition of the individual system. We thus have updated the method for calculating the CO₂ and CH₄ **mole fractions** to derive their **SD** for each data simultaneously.

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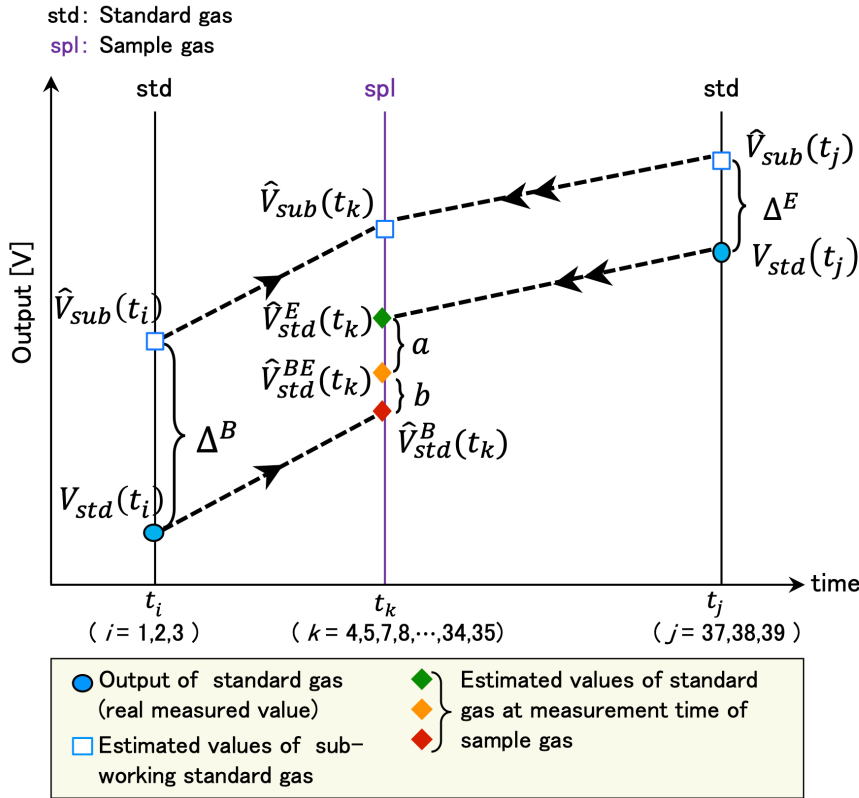
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2.4.1 Estimation of the output values of working standard gases and their **SD** at the time of the air sample measurements

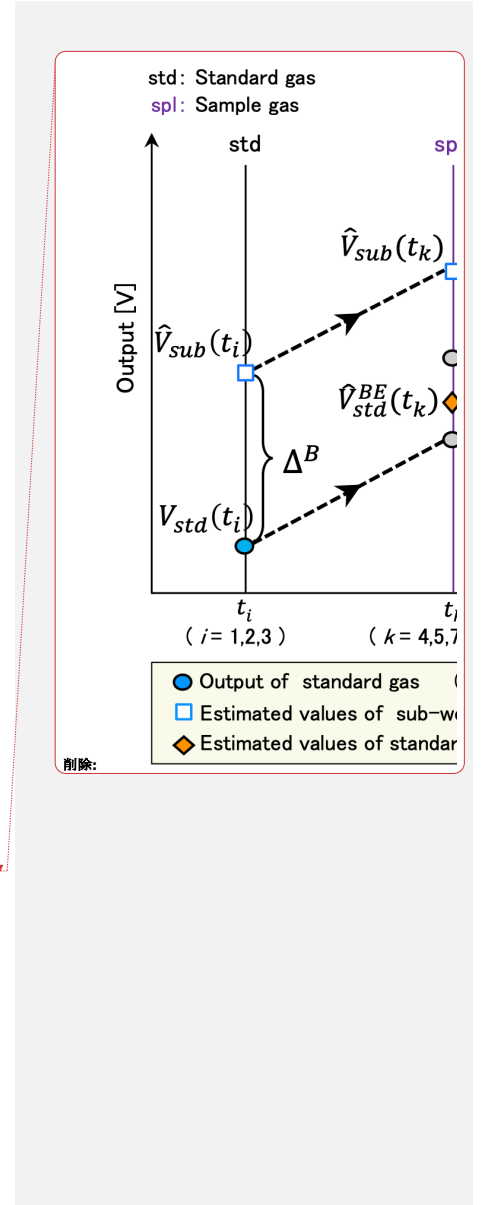
We estimated the outputs in voltage of three standard gases at each measurement time of the sample air by interpolating the outputs of the three WS-gases depending on the difference of the outputs in voltage of the SWS-gas only when both standard gas measurements satisfied the criteria described in section 2.3. Depending on the time difference between the targeted sample (t_k ; $k = 4, 5, 7, 8, \dots, 34, 35$) and standard gases at both sides B and E ((t_i, t_j) ; $(i, j) = (1, 37), (2, 38), (3, 39)$), the representative value ($V_{std(t_i, t_j)}^{BE}(t_k)$) and its variance ($\left(\sigma_{std(t_i, t_j)}^{BE}(t_k) \right)^2$) were estimated as follows (Fig. 3):

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255 Figure 3. Schematic diagram of the estimation method for the output of the standard gas at the time of sample air measurement.

$$\begin{aligned}
 V_{std(t_i, t_j)}^{BE}(t_k) &= \frac{a}{a+b} \cdot V_{std(t_i)}^B(t_k) + \frac{b}{a+b} \cdot V_{std(t_j)}^E(t_k) \\
 &= V_{sub}(t_k) + \frac{1}{a+b} \cdot \{a \cdot \Delta^B(t_i) + b \cdot \Delta^E(t_j)\} \\
 \left(\sigma_{std(t_i, t_j)}^{BE}(t_k) \right)^2 &= \left(\sigma_{sub}(t_k) \right)^2 + \left(\frac{a}{a+b} \cdot \sigma_{std(t_i)} \right)^2 + \left(\frac{a}{a+b} \cdot \frac{6-i}{6} \cdot \sigma_{sub}(t_0) \right)^2
 \end{aligned} \tag{12}$$



$$\begin{aligned}
& + \left(\frac{a}{a+b} \cdot \frac{i}{6} \cdot \sigma_{sub}(t_6) \right)^2 + \left(\frac{b}{a+b} \cdot \sigma_{std}(t_j) \right)^2 + \left(\frac{b}{a+b} \cdot \frac{42-j}{6} \cdot \sigma_{sub}(t_{36}) \right)^2 \\
& + \left(\frac{b}{a+b} \cdot \frac{j-36}{6} \cdot \sigma_{sub}(t_{42}) \right)^2
\end{aligned} \tag{13}$$

where $a : b = (t_j - t_k) : (t_k - t_i)$. Hat “^” means estimated value. $V_{sub}(t_k)$ was calculated by interpolating the output of the SWS-gas value nearest to the targeted sample as follows:

$$\begin{aligned}
265 \quad & \{spl1 \mid k = 4\} \\
& V_{sub}(t_4) = \frac{1}{3} \cdot V_{sub}(t_0) + \frac{2}{3} \cdot V_{sub}(t_6)
\end{aligned} \tag{14}$$

$$\begin{aligned}
& \{spl1 \mid k = 7, 10, 13, \dots, 34\} \\
& V_{sub}(t_k) = \frac{2}{3} \cdot V_{sub}(t_{k-1}) + \frac{1}{3} \cdot V_{sub}(t_{k+2})
\end{aligned} \tag{15}$$

$$\begin{aligned}
270 \quad & \{spl2 \mid k = 5\} \\
& V_{sub}(t_5) = \frac{1}{6} \cdot V_{sub}(t_0) + \frac{5}{6} \cdot V_{sub}(t_6)
\end{aligned} \tag{16}$$

$$\begin{aligned}
& \{spl2 \mid k = 8, 11, 14, \dots, 35\} \\
& V_{sub}(t_k) = \frac{1}{3} \cdot V_{sub}(t_{k-2}) + \frac{2}{3} \cdot V_{sub}(t_{k+1})
\end{aligned} \tag{17}$$

Here $V_{sub}(t_l) \{sub \mid l = 0, 6, 9, 12, \dots, 36\}$ is the measured value. In the following, a calculation example of $V_{std(t_i, t_j)}^{BE}(t_k)$ and the variance for the case $\{spl1 \mid k = 4, (i, j)\}$ are given without any estimated value.

$$\begin{aligned}
275 \quad & \{spl1 \mid k = 4, (i, j) = (1, 37), (2, 38), (3, 39)\} \\
& V_{std(t_i, t_j)}^{BE}(t_4) = \frac{1}{3} \cdot V_{sub}(t_0) + \frac{2}{3} \cdot V_{sub}(t_6) + \frac{a}{a+b} \cdot \left\{ V_{std}(t_i) - \left(\frac{6-i}{6} \cdot V_{sub}(t_0) + \frac{i}{6} \cdot V_{sub}(t_6) \right) \right\} \\
& + \frac{b}{a+b} \cdot \left\{ V_{std}(t_j) - \left(\frac{42-j}{6} \cdot V_{sub}(t_{36}) + \frac{j-36}{6} \cdot V_{sub}(t_{42}) \right) \right\}
\end{aligned} \tag{18}$$

$$\begin{aligned}
& \left(V_{std(t_i, t_j)}^{BE}(t_4) \right)^2 = \left(\frac{1}{3} \cdot \sigma_{sub}(t_0) \right)^2 + \left(\frac{2}{3} \cdot \sigma_{sub}(t_6) \right)^2 \\
& + \left(\frac{a}{a+b} \cdot \sigma_{std}(t_i) \right)^2 + \left(\frac{a}{a+b} \cdot \frac{6-i}{6} \cdot \sigma_{sub}(t_0) \right)^2 + \left(\frac{a}{a+b} \cdot \frac{i}{6} \cdot \sigma_{sub}(t_6) \right)^2 \\
280 \quad & + \left(\frac{b}{a+b} \cdot \sigma_{std}(t_j) \right)^2 + \left(\frac{b}{a+b} \cdot \frac{42-j}{6} \cdot \sigma_{sub}(t_{36}) \right)^2 + \left(\frac{b}{a+b} \cdot \frac{j-36}{6} \cdot \sigma_{sub}(t_{42}) \right)^2
\end{aligned} \tag{19}$$

2.4.2 Estimation of sample air μ mole fraction and its SD using a calibration line

We calculated a calibration line with the estimated outputs of standard gases ($\hat{V}_{std(t_i, t_j)}^{BE}(t_k)$) and their variances ($\sigma_{std(t_i, t_j)}^{BE}(t_k)$) at the time of the sample measurement obtained in the section 2.4.1. A linear line ($y = Sx + I$; y : output in voltage, x : μ mole fraction for CO₂ and log(μ mole fraction) for CH₄) was adopted for the calibration line (Fig. 4).

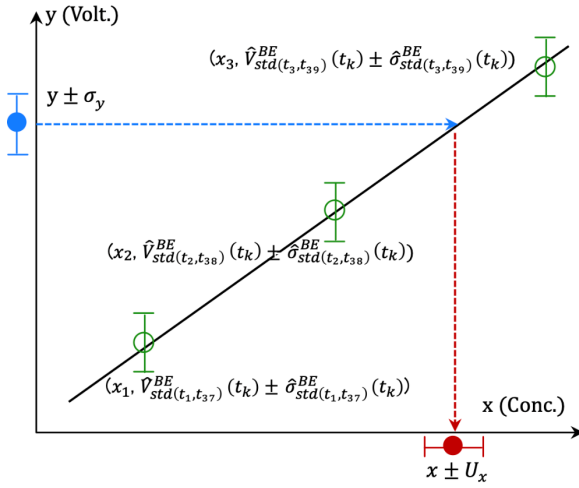


Figure 4. Schematic diagram for estimating the CO₂ and CH₄ μ mole fraction (x) of the gases and estimation $SD_x(U_x)$ from the output in voltage (y) with its $SD_y(\sigma_y)$. The gray line indicates the estimated linear calibration line ($y = Sx + I$).

290 Following the likelihood method, we identified the slope (S) and intercept (I) for every sample time (k) at the maximum of the

likelihood function (L). Solving the normal equation of $\begin{cases} \frac{\partial L}{\partial S} = 0 \\ \frac{\partial L}{\partial I} = 0 \end{cases}$, S and I were obtained as follows:

$$S(k) = \frac{(\sum w_{ijk})(\sum w_{ijk}x_i y_{ijk}) - (\sum w_{ijk}x_i)(\sum w_{ijk}y_{ijk})}{(\sum w_{ijk})(\sum w_{ijk}x_i^2) - (\sum w_{ijk}x_i)^2} \quad (20)$$

$$I(k) = \frac{(\sum w_{ijk}y_{ijk})(\sum w_{ijk}x_i^2) - (\sum w_{ijk}x_i y_{ijk})(\sum w_{ijk}x_i)}{(\sum w_{ijk})(\sum w_{ijk}x_i^2) - (\sum w_{ijk}x_i)^2} \quad (21)$$

where x_i is WS-gas **mole fraction** determined against the NIES scale. y_{ijk} is the estimated outputs of standard gas ($V_{std(t_i,t_j)}^{BE}(t_k)$)

and w_{ijk} is the reciprocal of the variance ($1/\left(\sigma_{std(t_i,t_j)}^{BE}(t_k)\right)^2$). Σ indicates the sum of i (three standard gases) and the same for the following discussion. As shown in Section 2.4.1, the combinations of (i, j) are (1, 37), (2, 38), and (3, 39). We omitted i, j , and k for the following expression. **The linearity of the calibration line was assessed using the correlation coefficient, and data were rejected when the linearity was deemed insufficient.**

The inverse function was used because we estimated the **mole fraction** from the output in voltage. Furthermore, because the calibration line passes through the weighted mean point $(x, y) = \left(\frac{\Sigma wx}{\Sigma w}, \frac{\Sigma wy}{\Sigma w}\right)$, we practically used the following line:

$$x = \frac{y - \bar{y}}{\bar{s}} + \bar{x} \quad (22)$$

The **variance** for the estimated **mole fraction** (x) was calculated with the following equation:

$$U_x^2 = \left(\frac{\partial x}{\partial y}\right)^2 U^2(y) + \left(\frac{\partial x}{\partial y}\right)^2 U^2(y) + \left(\frac{\partial x}{\partial s}\right)^2 U^2(s) + \left(\frac{\partial x}{\partial x}\right)^2 U^2(x) \quad (23-1)$$

where U_x^2 is **variance** for each component. The first term expresses the contribution from the variation in output of the measured air (σ_y) and 60 repeated measurements:

$$\left(\frac{\partial x}{\partial y}\right)^2 U^2(y) = \frac{\sigma_y^2}{s^2} \cdot \frac{1}{60}$$

315 The second term expresses the contribution from the variation in y :

$$\left(\frac{\partial x}{\partial y}\right)^2 U^2(y) = \frac{1}{s^2} \cdot \frac{\Sigma w^2 \sigma^2}{(\Sigma w)^2} = \frac{1}{s^2} \cdot \frac{1}{\Sigma w}$$

where σ^2 is the variance of the output for standard gases constituting the calibration line. The third term expresses the contribution from the variation in the slope of the calibration line (S):

$$\left(\frac{\partial x}{\partial s}\right)^2 U^2(s) = \frac{(y - \frac{\Sigma wy}{\Sigma w})^2}{s^4} \cdot \Sigma \sigma^2 \left(\frac{\partial s}{\partial y}\right)^2 = \frac{(y - \frac{\Sigma wy}{\Sigma w})^2}{s^4} \cdot \frac{\Sigma w}{(\Sigma w)(\Sigma wx^2) - (\Sigma wx)^2}$$

320 The fourth term expresses the contribution from the variation in x . The NIES09 scale is based on the gravimetric primary standard gases using a one-step dilution (Tohjima et al., 2006), and its overall uncertainty, including transfer, is estimated to be 0.043 ppm (Machida et al., 2011). **The production of standard gases is typically conducted concurrently using the same equipment (such as dilution vessels and pressure gauges). This process may introduce common biases, albeit to varying degrees, potentially resulting in correlated (non-independent) mole fractions. Consequently, the combined variance of the mean exhibits**

325 **a range contingent upon the strength of these correlations. In the case of complete independence, the value would be $0.043/\sqrt{3} = 0.025$, whereas if all correlation coefficients were unity, it would equal 0.043. Furthermore, as this value is common to all data points, it should be denoted separately. Summarizing all the terms, the SD for the estimated **mole fraction** (x) is as follows:**

$$U_x = \frac{1}{s} \cdot \sqrt{\frac{\sigma_y^2}{60} + \frac{1}{\Sigma w} + \frac{1}{s^2} \cdot \frac{((\Sigma w)y - \Sigma wy)^2}{(\Sigma w)^2 (\Sigma wx^2) - (\Sigma wx)^2}} \quad (23-2)$$

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340 Since the calculation was done for the logarithm of the **mole fraction** for CH₄, the **SD** for the estimated **mole fraction** was determined differently for the higher level ($U^+ = x(e^{U_x} - 1)$) and lower level ($U^- = x(1 - e^{-U_x})$). However, the average value is expressed as the **SD** since the difference is less than 0.1 ppb in real terms.

Figures S3 to S11 show the time series of **SD**(U_{x_sample}) for the ambient air CO₂ **mole fraction**. Most **SD**s were distributed at 0.05 ppm but can be higher than 0.3 ppm, especially during summer. Note that the **SD** of the output (σ_i) of the sample air could become significant due to large diurnal variation during summer since the output of the sample air could include a natural variation of the atmosphere during the measurement period of three minutes. Figures S12 to S17 show the time series of **SD**(U_{x_sample}) for the ambient air CH₄ **mole fraction**. Most are within five ppb but can be above ten ppb during summer. This is also due to the influence of natural variation and possibly heterogeneous CH₄ emissions from wetlands.

350 2.4.3 Measurement uncertainty (repeatability) with the SWS-gas measurement

We calculated a calibration line only when the SWS-gas measurements closest to both sides of the sample measurements were normal. The normality of the SWS-gas measurement was assessed as follows. The same on-site compressed air was measured several times (for about a week) since the air was used as SWS-gas until its pressure dropped below 0.1 MPa. The on-site compressed air output value would vary smoothly if the system were stable. When the system temporarily became unstable, the corresponding large changes of the analyzer output could not be corrected by the SWS to a sufficient degree. To identify such occasions, we first estimated the output of standard gases at the time of the target SWS-gas measurement by interpolating $\Delta^B(t_i)$ and $\Delta^E(t_j)$ based on the output value of the target SWS-gas itself (Section 2.4.1). Then, calculating a calibration line with the estimated output of standard gases, we obtained the **mole fraction** of the target SWS-gas. Second, we estimated the output value of the SWS-gas at the time of the target SWS-gas measurement by interpolating the outputs of the two adjacent SWS-gases. Then, we determined the **mole fraction** of the target SWS-gas in the same manner. If these estimated **mole fractions** differed from each other by more than one ppm for CO₂ and ten ppb for CH₄, we regarded the target SWS-gas data as abnormal. This assessment (referred to as “self-check-value (scv) for SWS-gases” in Figure 5) was done while the adjacent SWS-gas measurements were conducted for the same on-site compressed air.

We then checked the system's stability with the measurement of the SWS-gas. Interpolating the outputs of the SWS-gases adjacent to the standard gas measurements, we calculated the **mole fractions** of the SWS-gas with the calibration line at the time of STD2, which was used to assess the coefficient of determination in Section 2.3. We regarded the estimated **mole fractions** of the SWS-gas as independent; thus, we obtained 14 estimated **mole fractions** if the measurements for the same SWS-gas continued for a week. We determined a threshold for the **SD**(σ_{SWS} ; 1 ppm for CO₂, 10 ppb for CH₄) and the fluctuation range (3 ppm for CO₂ and 30 ppb for CH₄) obtained from the estimated independent data set. All the data that exceeded the threshold were deleted.

370 The uncertainty of this system should ideally be determined by conducting continuous measurements with a cylinder connected to the sample inlet. However, this approach has been impractical since the system's initial installation. Nevertheless, the

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aforementioned SWS-gas measurements involving continuous cylinder air sampling can be considered a reference value for estimating the measurement uncertainty (repeatability) for a given period (often a week). Regardless of the site or time of measurement, σ_{SWS} for CO₂ mostly was below 0.2 ppm, and for CH₄ was below five ppb (Supplement Figs. S3-S17). These values meet the extended network compatibility goal recommended in the GAW report (WMO, 2020). The extended network compatibility goals are provided as a guideline for the studies in which the smallest bias is not required, for example, a regionally focused study with large local fluxes or services related to urban air quality. After introducing the CRDS, the working standard gas measurement interval was changed to 48 hours. At the same time, since the consumption flow rate has doubled, the SWS-gas changeover time has decreased to approximately three days, which makes the number of SWS measurements with NDIR/TOS only three times at most. Although the σ_{SWS} values obtained from a few data are for reference only, they were distributed in almost the same range (red dots in Figures S3-S17). The CO₂ (CH₄) mole fractions observed in JR-STATION can fluctuate on the order of ppm (10 ppb), even during a few hours of daytime when the atmosphere is well mixed (Sawakawa et al., 2010, 2013). This indicates the observations were carried out in the vicinity of strong emission and absorption sources.

The SD obtained for each sample air mole fraction (U_{x_sample}) with equation 23-2 could be more considerable than the repeatability (σ_{SWS}). GAW report (WMO, 2020) mentioned that both values should be reported, and U_{x_sample} provides a quantitative indicator of the influence of nearby sources and can be used for data selection and weighting in applications such as inverse modeling. Following this guide, we showed U_{x_sample} and σ_{SWS} in our data set. A flow chart for the calculation method is shown in Fig. 5.

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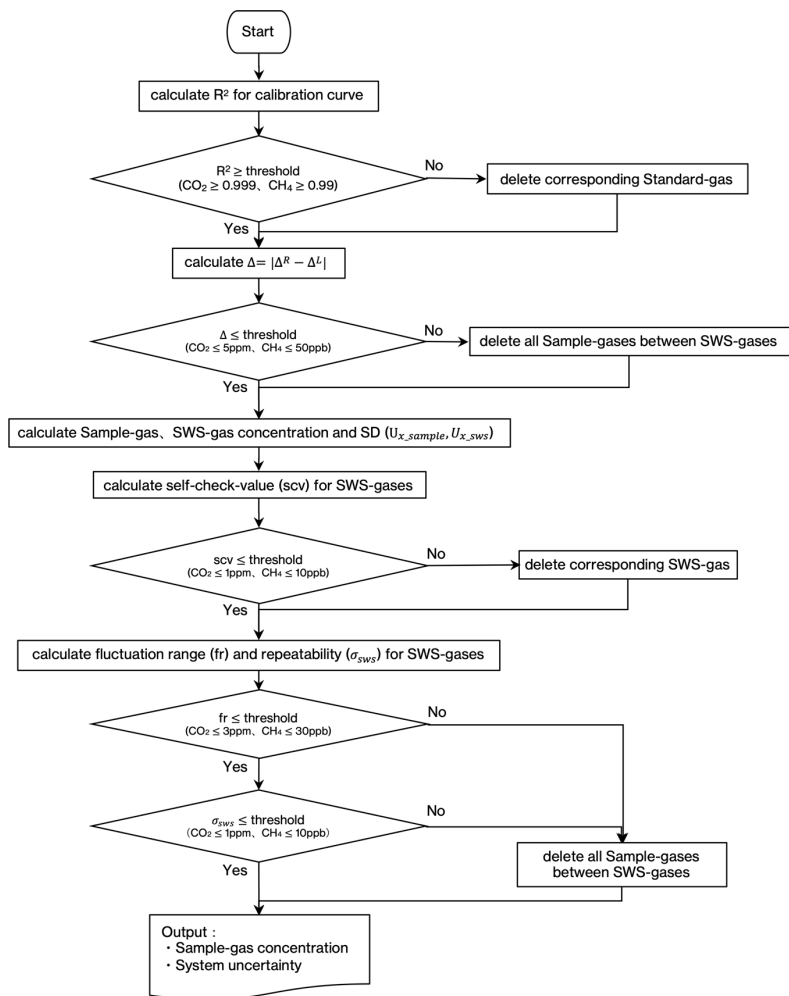
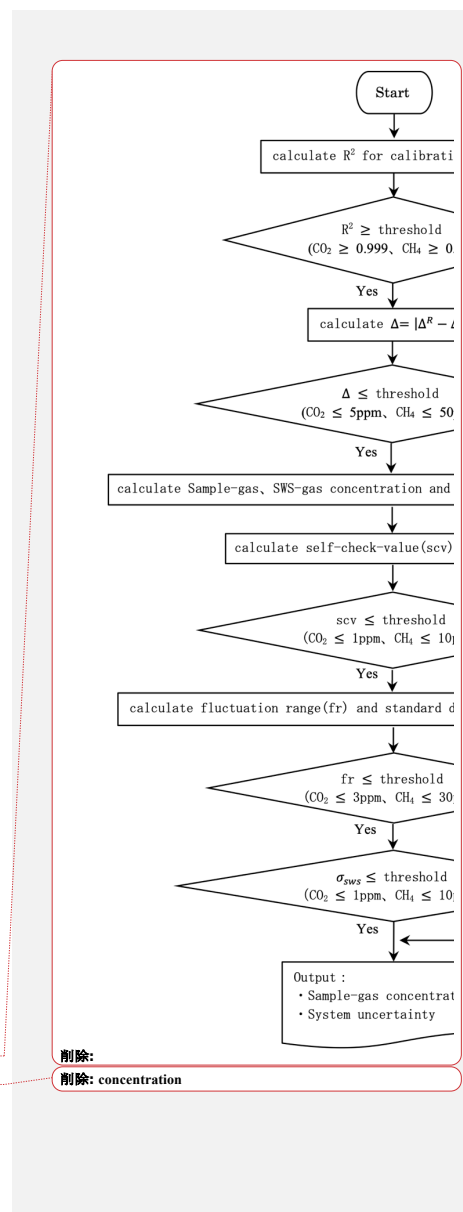


Figure 5. Flow chart of mole fraction calculation method.



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435 **2.5 Comparison between NDIR/TOS and CRDS**

The CRDS operated at KRS from July 2015, DEM from June 2016, and NOY from August 2016, albeit for a short period of time for JR-STATION. The CRDS is a highly stable analyzer used in greenhouse gas observations worldwide (Kwok et al., 2015). We compared the recalculated NDIR (and TOS) values with the CRDS values (Fig. 6, Fig. 7, Supplement Fig. S18-S21). The flow path branches off after the 6-port valve (Fig. 1), so the same air is analyzed. The CRDS operates independently of the existing system, so information on instrument error flags and valve switching timing is not linked to the measured data.

440 Therefore, by detecting the CH₄ mole fraction of the lowest standard gas, the timing of the standard gas measurement was captured. Only data from periods was extracted when the SWS results fulfilled the criteria outlined in the previous section. The CRDS output values were converted to the NIES scale based on the WS measurements and averaged over three minutes for comparison. The temperature in the warm box of CRDS (data column name is 'WarmBoxTemperature') was kept constant

445 (45.00°C). Still, it rarely changed by more than 0.03°C in 3 minutes, and the data was not used for comparison as the system was not considered stable during these periods. Since some observed values exceeded the highest mole fraction of the standard gas, only values within the mole fraction range of the standard gas were used to calculate the difference between the two. There was no significant difference in CO₂ mole fraction regardless of the inlet at both altitudes (Fig. 6, Supplement Figs. S18-S20).

450 As stated in Section 2.4.2, natural variability within the measurement time (3 minutes) caused a more significant error bar during summer (Supplement Fig. S20). No significant difference was found for CH₄ mole fraction either at KRS (Fig. 7). At NOY and DEM, it was discovered that the temperature controller of the catalytic unit was not functioning correctly. Since the TOS is sensitive to CO and H₂ in the air, it could produce unusually high values without a proper catalytic unit. For the period, only the data from the CRDS should be published. Since no catalytic unit errors were identified at the other sites, the ambient atmospheric values were detected, as is the case with KRS.

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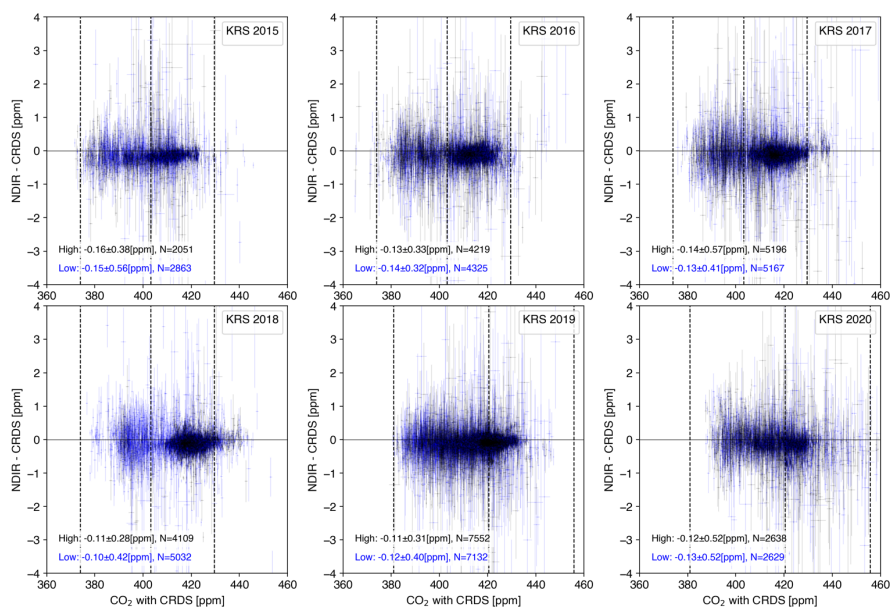


Figure 6. Relationship between the CO₂ mole fraction by the CRDS at KRS and the difference in respective CO₂ mole fractions measured by the NDIR and the CRDS (NDIR – CRDS): The CRDS values were averaged over the corresponding 3-minute period. Error bars on the vertical axis are the square root of the sum of the squares of U_{x_sample} and the SD of CRDS 3-minute measurements. Error bars on the horizontal axis are the SD of CRDS 3-minute measurements. The dotted lines indicate the mole fraction of standard gases. The average difference for each inlet is represented in the figure (mean ± SD). Only data that were within the standard gas mole fraction range were used.

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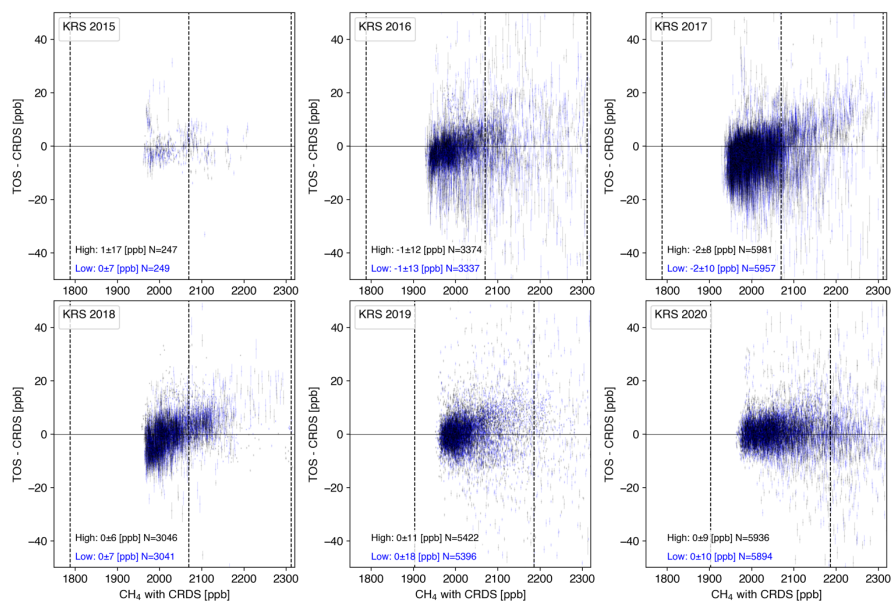


Figure 7. Relationship between the CH₄ mole fraction by the CRDS at KRS and the difference in respective CH₄ mole fractions measured by the TOS and the CRDS (TOS – CRDS): The CRDS values were averaged over the corresponding 3-minute period. Error bars on the vertical axis are the square root of the sum of the squares of U_{sample} and the SD of CRDS 3-minute measurements. Error bars on the horizontal axis are the SD of CRDS 3-minute measurements. The dotted lines indicate the mole fraction of standard gases. The average difference for each inlet is represented in the figure (mean ± SD). Only data that were within the standard gas mole fraction range were used.

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480 Data availability

The data are available from the Global Environmental Database, hosted by GED, CGER, NIES (<http://db.cger.nies.go.jp/portal/geds/index>).

Author contribution

MS and NT designed the study. MS wrote the manuscript. MA, DD, and AF conducted the measurements. All authors contributed to the discussion and preparation of the manuscript.

Competing interests

The authors declare that they have no conflict of interest.

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