

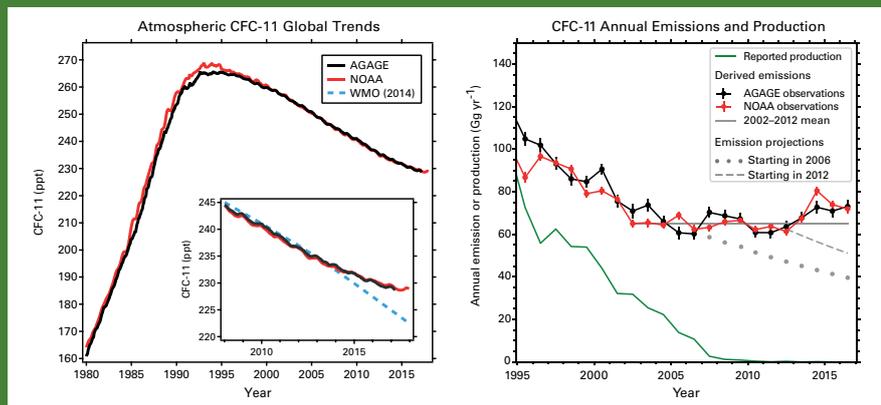


# WMO GREENHOUSE GAS BULLETIN

The State of Greenhouse Gases in the Atmosphere  
Based on Global Observations through 2017

No. 14 | 22 November 2018

## Unexpected Increases in Global Emissions of CFC-11



between 2002 and 2012 [1, 2]. These global trends are shown in the left graph of the figure for the Advanced Global Atmospheric Gases Experiment (AGAGE; shown in black) and the National Oceanic and Atmospheric Administration (NOAA; shown in red) measurement networks. Also shown in the inset to this graph is the trend that was predicted in 2014 by WMO (blue dashed) assuming adherence to the Montreal Protocol [3].

Modelling results lead to the robust conclusion that these changes are

**Measurements of the atmospheric abundance of the chlorofluorocarbon CFC-11, a potent greenhouse gas (GHG) and a stratospheric ozone-depleting substance (ODS) regulated under the Montreal Protocol on Substances that Deplete the Ozone Layer, show that since 2012 its rate of decline has slowed to roughly two thirds of its rate of decline during the preceding decade [1, 2]. The most likely cause of this slowing is increased emissions associated with production of CFC-11 in eastern Asia. This discovery illustrates the importance of long-term measurements of atmospheric composition, such as are carried out under the auspices of the Global Atmosphere Watch (GAW) Programme of WMO, in providing effective support and additional constraints for emissions-control legislation.**

The Montreal Protocol was designed to protect the stratospheric ozone layer by restricting the production of ODSs such as CFCs. As a consequence, CFC-11 (trichlorofluoromethane, or  $\text{CCl}_3\text{F}$ ) production reported under the Montreal Protocol declined to zero by 2010. As CFC-11 was phased out, its atmospheric abundance peaked in the early 1990s and then declined in a manner largely consistent with declining production combined with residual emissions of CFC-11 gradually escaping from stored “banks” in existing products and equipment.

Atmospheric measurements of CFC-11 made by independent global networks show that since 2012 the rate of decrease in atmospheric CFC-11 has slowed to roughly two thirds of the rate that was observed

predominately related to increased CFC-11 emissions rather than to other possible causes such as changing atmospheric transport. This conclusion is supported by recent increases in the northern to southern hemisphere difference in atmospheric concentration levels. Correlations between elevated abundances of CFC-11 and other measured gases further suggest that these increases originate from emissions in eastern Asia [1].

Separate CFC-11 emission trends resulting from model calculations taken from the 2018 WMO ozone assessment [2], based on data from each of the global measurement networks AGAGE (black) and NOAA (red), are shown in the graph on the right of the figure. They are contrasted to CFC-11 production as reported under the Montreal Protocol (green). These results show a levelling off of CFC-11 emissions around 2005, followed by an emission increase of about 15% after 2012. Emission scenario projections for the years 2006 and 2012 based on atmospheric data, reported production and releases from banks are shown as dots and dashes (grey), respectively.

This work demonstrates the importance of long-term measurements of atmospheric composition, such as are carried out under the auspices of the GAW Programme, in providing observation-based information to support national emission inventories, especially in the context of agreements to address anthropogenic climate change, as well as for the recovery of the stratospheric ozone layer.

## Executive summary

The latest analysis of observations from the WMO GAW Programme shows that globally averaged surface mole fractions<sup>(1)</sup> calculated from this in situ network for carbon dioxide (CO<sub>2</sub>), methane (CH<sub>4</sub>) and nitrous oxide (N<sub>2</sub>O) reached new highs in 2017, with CO<sub>2</sub> at 405.5 ± 0.1 ppm<sup>(2)</sup>, CH<sub>4</sub> at 1859 ± 2 ppb<sup>(3)</sup> and N<sub>2</sub>O at 329.9 ± 0.1 ppb. These values constitute, respectively, 146%, 257% and 122% of pre-industrial (before 1750) levels. The increase in CO<sub>2</sub> from 2016 to 2017 was smaller than that observed from 2015 to 2016 and practically equal to the average growth rate over the last decade. The influence of the El Niño event that peaked in 2015 and 2016 and contributed to the increased growth rate during that period sharply declined in 2017. For CH<sub>4</sub>, the increase from 2016 to 2017 was lower than that observed from 2015 to 2016 but practically equal to the average over the last decade. For N<sub>2</sub>O, the increase from 2016 to 2017 was higher than that observed from 2015 to 2016 and practically equal to the average growth rate over the past 10 years. The NOAA Annual Greenhouse Gas Index (AGGI) [4] shows that from 1990 to 2017 radiative forcing by long-lived GHGs (LLGHGs) increased by 41%, with CO<sub>2</sub> accounting for about 82% of this increase.

## Overview of the GAW in situ network observations for 2017

This fourteenth WMO Greenhouse Gas Bulletin reports atmospheric abundances and rates of change of the most important LLGHGs – CO<sub>2</sub>, CH<sub>4</sub> and N<sub>2</sub>O – and provides a summary of the contributions of the other gases. These three, together with CFC-12 and CFC-11, account for approximately 96%<sup>(4)</sup> of radiative forcing due to LLGHGs (Figure 1).

The GAW Programme (<http://www.wmo.int/gaw>) coordinates systematic observations and analysis of GHGs and other trace species. Sites where GHGs have

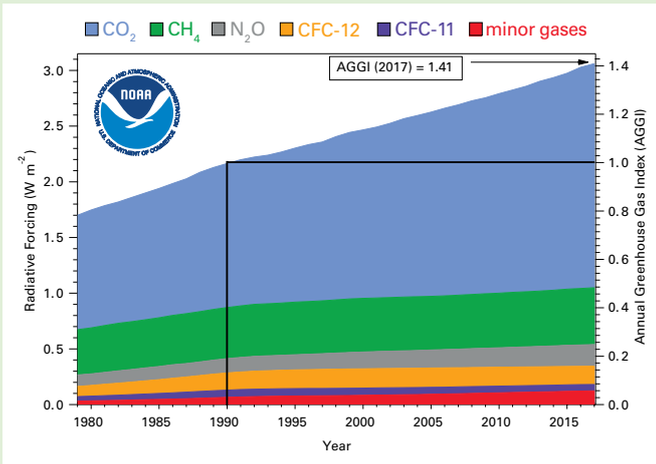


Figure 1. Atmospheric radiative forcing, relative to 1750, of LLGHGs and the 2017 update of the NOAA AGGI [4]

Table 1. Global annual surface mean abundances (2017) and trends of key GHGs from the WMO GAW global GHG observational network. Units are dry-air mole fractions, uncertainties are 68% confidence limits [5], and the averaging method is described in [7]. The numbers of stations used for the analyses are 129 for CO<sub>2</sub>, 126 for CH<sub>4</sub> and 96 for N<sub>2</sub>O.

	CO <sub>2</sub>	CH <sub>4</sub>	N <sub>2</sub> O
Global abundance in 2017	405.5±0.1 ppm	1859±2 ppb	329.9±0.1 ppb
2017 abundance relative to year 1750*	146%	257%	122%
2016–2017 absolute increase	2.2 ppm	7 ppb	0.9 ppb
2016–2017 relative increase	0.55%	0.38%	0.27%
Mean annual absolute increase of last 10 years	2.24 ppm yr <sup>-1</sup>	6.9 ppb yr <sup>-1</sup>	0.93 ppb yr <sup>-1</sup>

\* Assuming a pre-industrial mole fraction of 278 ppm for CO<sub>2</sub>, 722 ppb for CH<sub>4</sub> and 270 ppb for N<sub>2</sub>O.

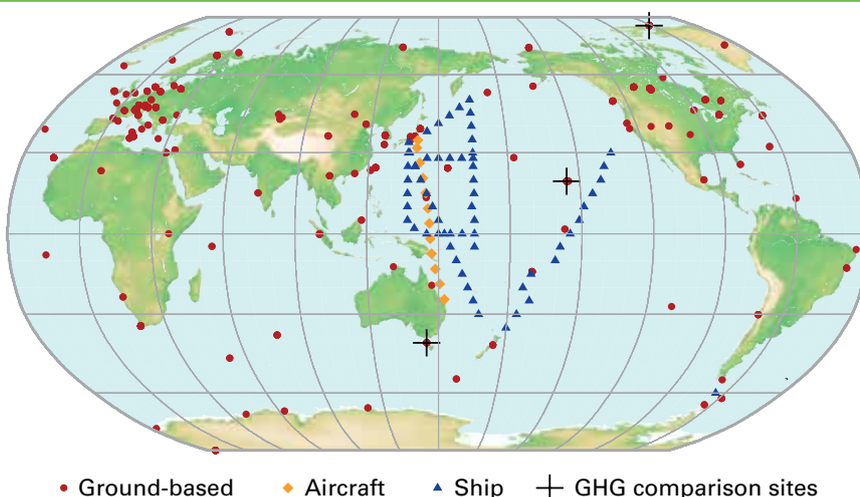
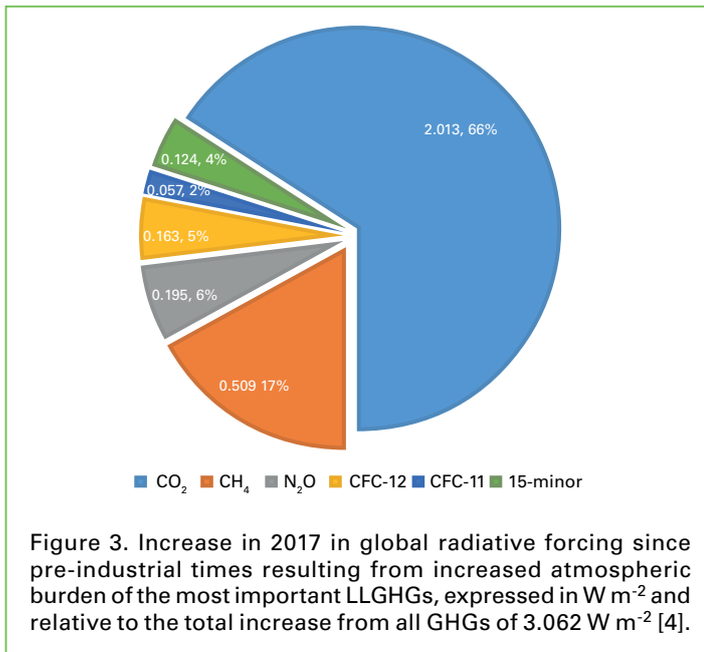


Figure 2. The GAW global network for CO<sub>2</sub> in the last decade. The network for CH<sub>4</sub> is similar.



been measured in the last decade are shown in Figure 2. Measurement data are reported by participating countries and archived and distributed by the WMO World Data Centre for Greenhouse Gases (WDCGG) at the Japan Meteorological Agency.

The results reported here by WDCGG for the global average and growth rate are slightly different from results reported by NOAA for the same years [6] due to differences in the stations used, differences in the averaging procedure and a slightly different time period for which the numbers are representative. WDCGG follows the procedure described in detail in [7].

Table 1 provides globally averaged atmospheric abundances of the three major LLGHGs in 2017 and changes in their abundances since 2016 and 1750. Data from mobile stations (blue triangles and orange diamonds in Figure 2), with the exception of NOAA sampling in the eastern Pacific, are not used for this global analysis.

The three GHGs shown in Table 1 are closely linked to anthropogenic activities and also interact strongly with the biosphere and the oceans. Predicting the evolution of the atmospheric content of GHGs requires quantitative understanding of their many sources, sinks and chemical transformations in the atmosphere. Observations from GAW provide invaluable constraints on the budgets of these and other LLGHGs, and they are used to support emission inventories preparation and evaluate satellite retrievals of LLGHG column averages. The Integrated Global Greenhouse Gas Information System (IG<sup>3</sup>IS), promoted by WMO, provides further insights on the sources of GHGs on the national and sub-national level. Some examples of the information that is delivered by the IG<sup>3</sup>IS projects can be found in the central insert of this Bulletin.

The NOAA AGGI [4] in 2017 was 1.41, representing a 41% increase in total radiative forcing<sup>(4)</sup> by all LLGHGs since 1990 and a 1.6% increase from 2016 to 2017 (Figure 1). The total radiative forcing by all LLGHGs in 2017 (3.062 W m<sup>-2</sup>) corresponds to an equivalent CO<sub>2</sub> mole fraction of 493 ppm [4]. Relative contributions of the other gases in the total radiative forcing since pre-industrial time are presented in Figure 3.

## Carbon dioxide

Carbon dioxide is the single most important anthropogenic GHG in the atmosphere, contributing approximately 66%<sup>(4)</sup> of the radiative forcing by LLGHGs. It is responsible for approximately 82%<sup>(4)</sup> of the increase in radiative forcing over the past decade and over the past five years. The pre-industrial level of 278 ppm represented a balance of fluxes among the atmosphere, the oceans and the land biosphere. Atmospheric CO<sub>2</sub> reached 146% of the pre-industrial level in 2017, primarily because of emissions from combustion of fossil fuels and cement production (the sum of CO<sub>2</sub> emissions was 9.9 ± 0.5 PgC<sup>(5)</sup> in 2016 [8]), deforestation and other land-use change (1.3 ± 0.7 PgC average for 2007–2016). Of the total emissions from human activities during the period 2007–2016, approximately 44% accumulated in the atmosphere, 22% in the ocean and 28% on land; the unattributed budget imbalance is 5% [8]. The portion of CO<sub>2</sub> emitted by fossil fuel combustion that remains in the atmosphere (airborne fraction), varies inter-annually due to the high natural variability of CO<sub>2</sub> sinks without a confirmed global trend.

The globally averaged CO<sub>2</sub> mole fraction in 2017 was 405.5 ± 0.1 ppm (Figure 4). The increase in annual means from 2016 to 2017, 2.2 ppm, is smaller than the increase from 2015 to 2016 (3.2 ppm) and practically equal to the average growth rate for the past decade (2.24 ppm yr<sup>-1</sup>). The higher growth rates in 2016 and 2015, in comparison with the years before 2016 and the increase from 2016 to 2017, are due in part to increased natural emissions of CO<sub>2</sub> related to the most recent El Niño event, as explained in the twelfth edition of this Bulletin.

## Methane

Methane contributes approximately 17%<sup>(4)</sup> of the radiative forcing by LLGHGs. Approximately 40% of methane is emitted into the atmosphere by natural sources (e.g., wetlands and termites), and about 60% comes from anthropogenic sources (e.g., ruminants, rice agriculture, fossil fuel exploitation, landfills and biomass burning). Atmospheric CH<sub>4</sub> reached 257% of the pre-industrial level (approximately 722 ppb) due to increased emissions from anthropogenic sources. Globally averaged CH<sub>4</sub> calculated from in situ observations reached a new high of 1859 ± 2 ppb in 2017, an increase of 7 ppb with respect to the

# ATMOSPHERIC OBSERVATIONS AND ANALYSIS IN SUPPORT OF GHG EMISSION MITIGATION – EXAMPLE PROJECTS OF THE GAW IG<sup>3</sup>IS PROGRAMME

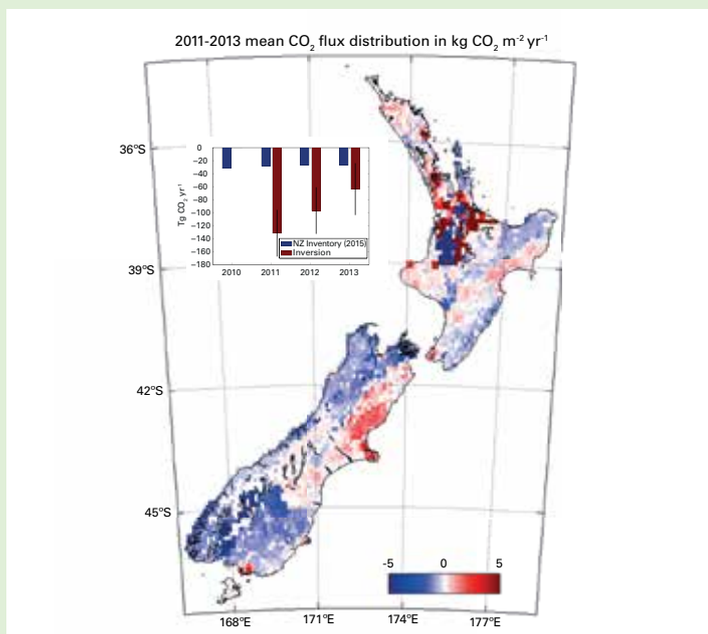


Figure 8. Geographic distribution of land-to-air CO<sub>2</sub> flux, averaged over 2011–2013 [10]. Blue and red regions indicate net carbon uptake and release, respectively. Per-area ocean fluxes are too small to show on this scale. Fossil fuel emissions are included and reach up to 20 kg CO<sub>2</sub> m<sup>-2</sup> yr<sup>-1</sup> in a few grid cells (Auckland area). The colour scale is capped to focus on natural fluxes. Inset: annual mean inverse model results [12] compared to the National Greenhouse Gas Inventory Report.

## 1. Atmospheric measurements reveal strong forest carbon sink in New Zealand

By Sara Mikaloff-Fletcher (National Institute of Water and Atmospheric Research Ltd, New Zealand) and Jocelyn Turnbull (GNS Science, New Zealand)

Net CO<sub>2</sub> uptake from land use, land-use change and forestry currently offsets approximately 30% of New Zealand’s GHG emissions [10]. These land carbon sinks played a key role in meeting New Zealand’s past GHG emission targets under the United Nations Framework Convention on Climate Change (UNFCCC), and they are expected to be a major component of the nation’s strategy for future GHG mitigation. New Zealand’s National Inventory Report (NIR) estimates forest carbon uptake based on tree diameter and height measurements at a national network of study sites, and allometric equations that infer carbon mass from these measurements. This approach, which is required by current Intergovernmental Panel on Climate Change (IPCC) guidelines [11], has substantial uncertainty.

Atmospheric CO<sub>2</sub> observations and inverse model simulations [12], illustrated in Figure 8, suggest that New Zealand’s forest carbon sink may far exceed estimates from

the NIR [10] and land process models [12]. Furthermore, the atmospheric observations reveal significant interannual variability that is not detected by the NIR methodology. This study combined in situ observations of atmospheric CO<sub>2</sub> at a network of sites with a high-resolution atmospheric model. The spatial pattern of the sink suggests that much of this missing carbon uptake occurs in Fiordland, a high rainfall region dominated by indigenous forests. The research team of New Zealand is launching a new research programme to further evaluate the processes that drive this sink. Through close engagement with users in the carbon accounting, land management and policy communities, this nationally funded programme will support the IG<sup>3</sup>IS mission to provide a bridge between science and policy for GHG monitoring and emission estimation.

## 2. Use of atmospheric observations of greenhouse gases to inform the United Kingdom national inventory

By Alistair Manning (UK Met Office)

To support the emission estimates that follow the IPCC protocol (“bottom-up”) [11] and are reported annually to UNFCCC, the United Kingdom uses a completely independent method (“top-down”) [13] for informing on its GHG emission estimates. The method uses a combination of atmospheric observations and modelling, and the results are also reported annually in the United Kingdom National Inventory Report to UNFCCC. Significant differences in the emissions estimated utilizing the two approaches are used by the United Kingdom Government Department of Business, Energy and Industrial Strategy (BEIS) to identify areas worthy of further investigation.

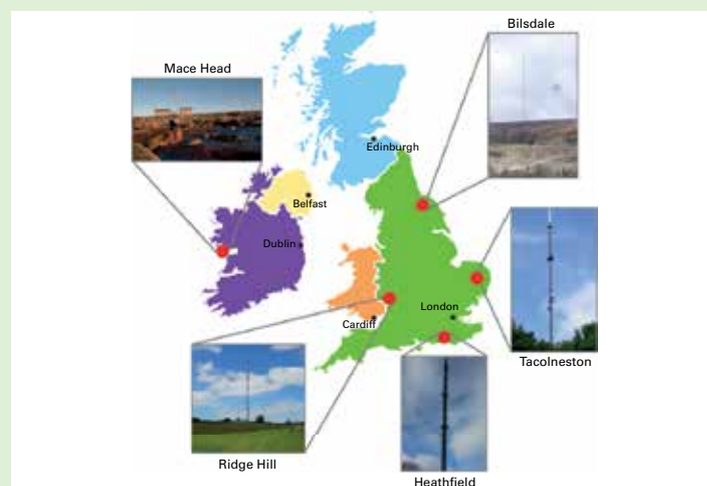


Figure 9. United Kingdom-funded DECC network of observation sites

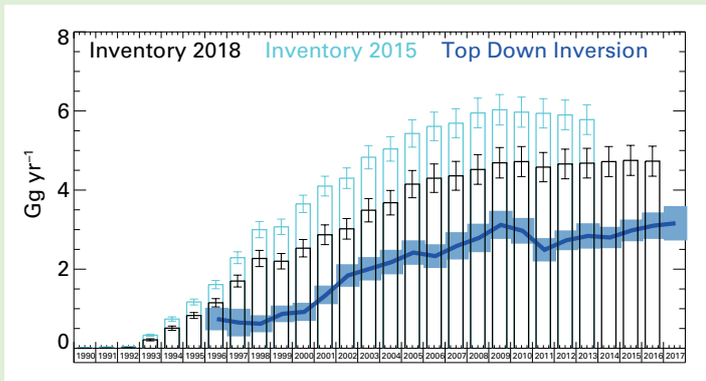


Figure 10. United Kingdom emission estimates of HFC-134a. Inventory estimates from two reporting years compared against top-down (InTEM) estimates.

In 2012, BEIS invested in a network of observation sites (Figure 9) called the UK Deriving Emissions related to Climate Change (DECC) network [14]. These are primarily tall-tower telecommunication masts equipped with state-of-the-art observation equipment measuring  $\text{CO}_2$ ,  $\text{CH}_4$ ,  $\text{N}_2\text{O}$ , HFCs, perfluorocarbons,  $\text{SF}_6$  and nitrogen trifluoride ( $\text{NF}_3$ ) to high precision and quality.

A recent example of how the top-down approach has been used to inform the bottom-up estimate is demonstrated in Figure 10. The country's 2015 bottom-up estimate for HFC-134a is shown in the figure as the light blue bars with an estimated uncertainty of 8%. The top-down estimate was consistently approximately 50% of this value throughout the time series from 1994, when the observations started, until 2013, the last year of that inventory. This result, and the subsequent work undertaken [15], motivated BEIS to investigate this further and an industry expert partly revised the United Kingdom HFC-134a inventory estimates.

The result of the revised bottom-up estimate is shown in Figure 10 as black bars – it has moved to be considerably nearer to the top-down estimates. The remaining discrepancy is believed to arise from the use of assumption on a refill rate.

### 3. Oil and gas methane emissions in Alberta, Canada: Collecting policy-relevant atmospheric data

By Daniel Zavala-Araiza (Environmental Defense Fund, United States of America)

The oil and gas sector in Canada accounts for roughly half of total  $\text{CH}_4$  emissions in the national inventory [16]. The federal government recently announced regulations that support the goal of a 40–45% reduction of  $\text{CH}_4$  emissions from the oil and gas system below 2012 levels by the year 2025 [17].

For emission reduction goals and policies to be realistically achieved, knowledge of the current emissions baseline as well as the characteristics of the major emitting sources is a necessary condition. Therefore, a multi-scale campaign, targeting oil and gas production regions in Alberta was conducted in the fall of 2016 [18–20] and the data were used to estimate the emissions (mass balance approach). These top-down estimates were then compared with spatially explicit, region-specific inventories and industry-reported emissions. In addition, ground-based mobile (downwind, site-wide characterization using dual tracer release and Gaussian dispersion modelling) measurements allowed the characterization of the emission distributions and major sources of emissions (see Figure 11a).

In the Lloydminster region of Alberta, the major source of emissions is related to direct venting of methane to the atmosphere from the production casing. The results based on atmospheric observations suggest that emissions are three to five times greater than inventories. This large discrepancy is particularly relevant in the context of proposed regulations and emission reduction policies in Canada. If these results are conservatively extrapolated to the larger population of similar sites in Alberta, actual methane emissions from oil and gas production in the province are likely to be 25–50% higher as illustrated in Figure 11b.

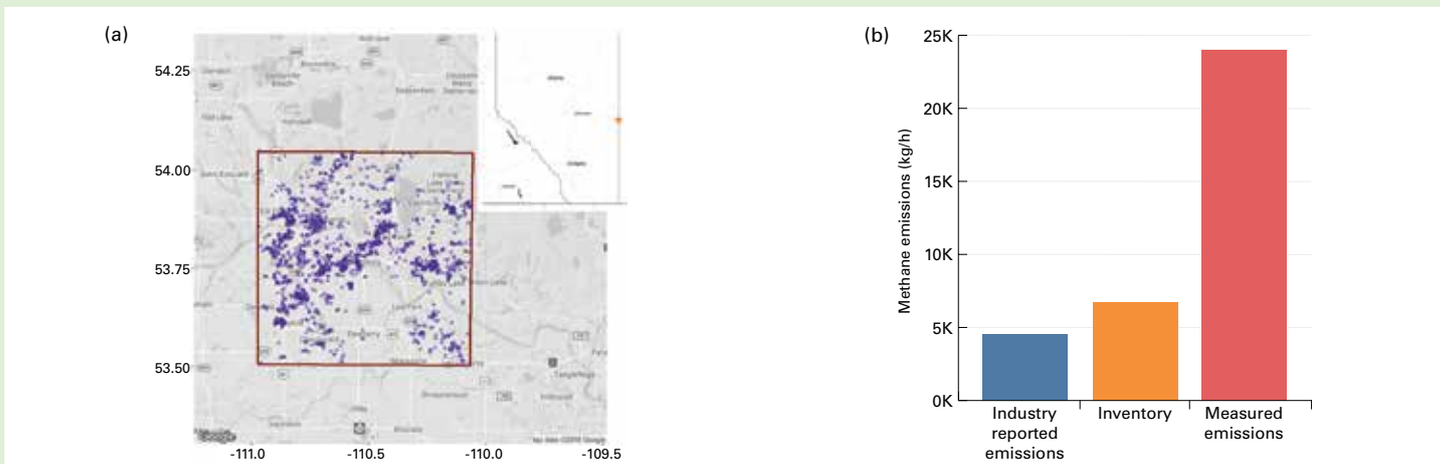


Figure 11. (a) Sampling region near Lloydminster, Alberta. The red box illustrates the source envelope where the aircraft took measurements. Purple dots inside the box represent active oil wells. (b) Comparison between measured  $\text{CH}_4$  emissions and “bottom-up” estimates based on inventory and industry reports.

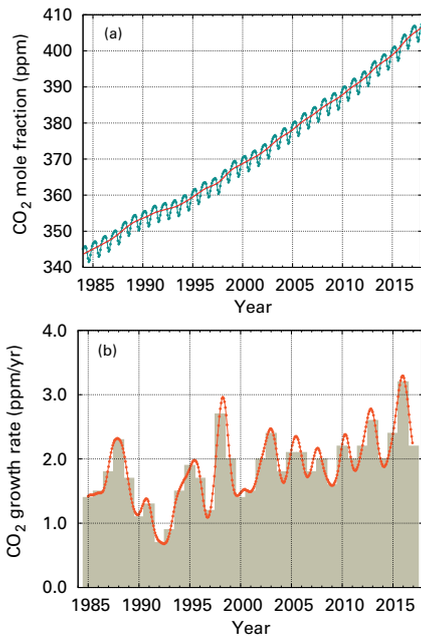


Figure 4. Globally averaged CO<sub>2</sub> mole fraction (a) and its growth rate (b) from 1984 to 2017. Increases in successive annual means are shown as the shaded columns in (b). The red line in (a) is the monthly mean with the seasonal variation removed; the blue dots and line depict the monthly averages. Observations from 129 stations have been used for this analysis.

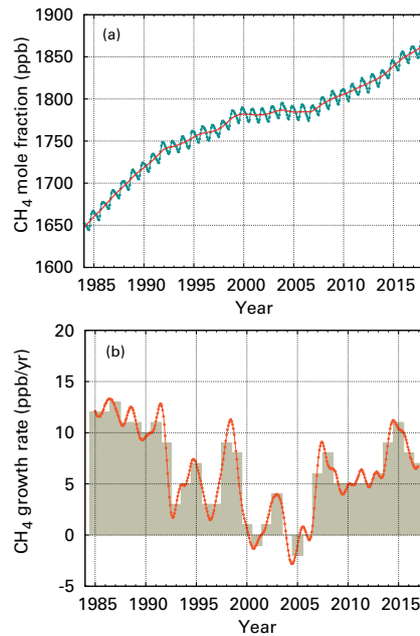


Figure 5. Globally averaged CH<sub>4</sub> mole fraction (a) and its growth rate (b) from 1984 to 2017. Increases in successive annual means are shown as the shaded columns in (b). The red line in (a) is the monthly mean with the seasonal variation removed; the blue dots and line depict the monthly averages. Observations from 126 stations have been used for this analysis.

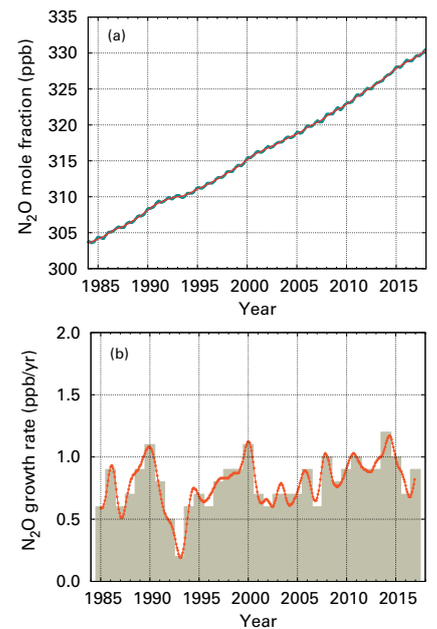


Figure 6. Globally averaged N<sub>2</sub>O mole fraction (a) and its growth rate (b) from 1984 to 2017. Increases in successive annual means are shown as the shaded columns in (b). The red line in (a) is the monthly mean with the seasonal variation removed; in this plot it is overlapping with the blue dots and line that depict the monthly averages. Observations from 96 stations have been used for this analysis.

previous year (Figure 5). This increase is lower than the increase from 2015 to 2016 but practically equal to the average annual increase over the past decade. The mean annual increase of CH<sub>4</sub> decreased from approximately 12 ppb yr<sup>-1</sup> during the late 1980s to near zero during 1999–2006. Since 2007, atmospheric CH<sub>4</sub> has been increasing again. Studies using GAW CH<sub>4</sub> measurements indicate that increased CH<sub>4</sub> emissions from wetlands in the tropics and from anthropogenic sources at mid-latitudes of the northern hemisphere are likely causes of this recent increase.

## Nitrous oxide

Nitrous oxide contributes approximately 6%<sup>(4)</sup> of the radiative forcing by LLGHGs. It is the third most important individual contributor to the combined forcing. N<sub>2</sub>O is emitted into the atmosphere from both natural (about 60%) and anthropogenic sources (approximately 40%), including oceans, soils, biomass burning, fertilizer use and various industrial processes. The globally averaged N<sub>2</sub>O mole fraction in 2017 reached 329.9 ± 0.1 ppb, which is 0.9 ppb above the previous year (Figure 6) and 122%

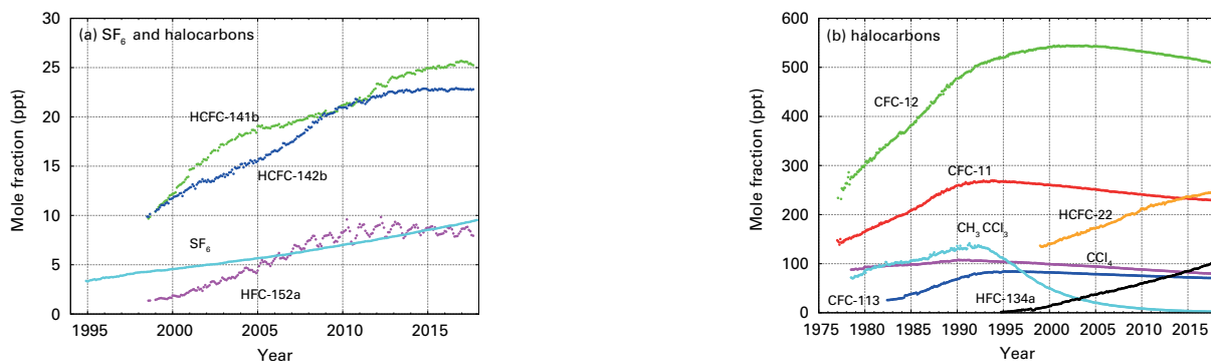


Figure 7. Monthly mean mole fractions of SF<sub>6</sub> and the most important halocarbons: (a) SF<sub>6</sub> and lower mole fractions of halocarbons and (b) higher halocarbon mole fractions. The numbers of stations used for the analyses are as follows: SF<sub>6</sub> (85), CFC-11 (23), CFC-12 (25), CFC-113 (21), CCl<sub>4</sub> (21), CH<sub>3</sub>CCl<sub>3</sub> (24), HCFC-141b (9), HCFC-142b (14), HCFC-22 (13), HFC-134a (10), HFC-152a (9).

of the pre-industrial level (270 ppb). The annual increase from 2016 to 2017 is higher than the increase from 2015 to 2016 and practically equal to the mean growth rate over the past 10 years (0.93 ppb yr<sup>-1</sup>). The likely causes of N<sub>2</sub>O increase in the atmosphere are an increased use of fertilizers in agriculture and increased release of N<sub>2</sub>O from soils due to an excess of atmospheric nitrogen deposition related to air pollution.

## Other greenhouse gases

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Sulphur hexafluoride (SF<sub>6</sub>) is a potent LLGHG. It is produced by the chemical industry, mainly as an electrical insulator in power distribution equipment. Its current mole fraction is more than twice the level observed in the mid-1990s (Figure 7a). The stratospheric ozone-depleting CFCs, together with minor halogenated gases, contribute approximately 11%<sup>(4)</sup> of the radiative forcing by LLGHGs. While CFCs and most halons are decreasing, some hydrochlorofluorocarbons (HCFCs) and hydrofluorocarbons (HFCs), which are also potent GHGs, are increasing at relatively rapid rates, although they are still low in abundance (at ppt<sup>(6)</sup> levels).

This Bulletin primarily addresses LLGHGs. Relatively short-lived tropospheric ozone [9] has a radiative forcing comparable to that of the halocarbons. Many other pollutants, such as carbon monoxide, nitrogen oxides and volatile organic compounds, although not referred to as GHGs, have small direct or indirect effects on radiative forcing. Aerosols (suspended particulate matter) are short-lived substances that alter the radiation budget. All gases mentioned herein, as well as aerosols, are monitored by the GAW Programme, with support from WMO Members and contributing networks.

## Acknowledgements and links

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Fifty-three WMO Members have contributed CO<sub>2</sub> and other GHG data to WDCGG. Approximately 41% of the measurement records submitted to WDCGG were obtained at sites of the NOAA Earth System Research Laboratory cooperative air-sampling network. For other networks and stations, see GAW Report No. 229. AGAGE also contributed observations to this Bulletin. Furthermore, the GAW observational stations that contributed data to this Bulletin, shown in Figure 2, are included in the list of contributors on the WDCGG web page (<https://gaw.kishou.go.jp/>). They are also described in the GAW Station Information System (<https://gawsis.meteoswiss.ch/>) supported by MeteoSwiss.

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## Contacts

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(1) Mole fraction = the preferred expression for abundance (concentration) of a mixture of gases or fluids. In atmospheric chemistry it is used to express the concentration as the number of moles of a compound per mole of dry air.

(2) ppm = number of molecules of the gas per million (10<sup>6</sup>) molecules of dry air.

(3) ppb = number of molecules of the gas per billion (10<sup>9</sup>) molecules of dry air.

(4) This percentage is calculated as the relative contribution of the mentioned gas(es) to the increase in global radiative forcing caused by all LLGHGs since 1750.

(5) 1 PgC = 1 petagram (10<sup>15</sup> gram) of carbon.

(6) ppt = number of molecules of the gas per trillion (10<sup>12</sup>) molecules of dry air.

## The Integrated Carbon Observation System (ICOS) atmosphere network



The ICOS atmosphere station network: yellow dots are combined atmosphere/ecosystem stations, red dots only observe the atmosphere. Not shown are the stations in French Guyana, La Reunion and Cabo Verde. Some example stations are: Pallas (Finland), Jungfrauoch (Switzerland), Svartberget (Sweden), Lampedusa (Italy).

Konsta Punkka and Alcide di Sarra

Since mid-2018, the European atmosphere station network of the ICOS Research Infrastructure (<https://www.icos-ri.eu>) is a GAW-contributing network consisting of 33 stations (of which 22 are tall towers). Many ICOS atmosphere stations have already been in operation a long time, but ICOS has now also been extended into new regions and with new sites. ICOS has developed community-defined standardized measurement designs and protocols that for atmospheric GHG observations build and extend upon the WMO recommendations with regards to compatibility, calibration to WMO mole fraction scales and transparency of the data lifecycle. All ICOS stations have to meet the agreed standards. All data are processed by the ICOS Atmosphere Thematic Centre and checked and annotated on a daily basis by the responsible station managers. The Central Analytical Laboratories perform analyses of flask samples, e.g. for  $^{14}\text{CO}_2$  radiocarbon detection of fossil fuel emissions, and provide all stations with WMO scale calibrated working standards. All fully quality-controlled ICOS atmosphere data are published as open data through the ICOS Carbon Portal (<https://data.icos-cp.eu/portal>) and are updated currently about twice per year. Near-real-time data, utilizing automatic quality control, are published with a maximum delay of one day from the time of the last final full quality-controlled release onwards. The atmospheric data will also be accessible through WDCGG, are part of the regular updates of the NOAA Obspack data products, and are delivered on a daily basis to the COPERNICUS services (<http://www.copernicus.eu/main/overview>).

### Research vessel I (RVI) *Investigator*, the first mobile station in the GAW network

The RV *Investigator* of the Australian Marine National Facility has two dedicated atmospheric sampling laboratories providing continuous, high-quality, in situ measurements of  $\text{CO}_2$ ,  $\text{CH}_4$  and  $\text{N}_2\text{O}$  along with other important trace gases such as carbon monoxide, tropospheric ozone and radon. A wide range of aerosol and meteorological parameters are also measured. In 2018, the *Investigator* became the first mobile station in the GAW network.

The *Investigator* sails 300 days a year in the waters around Australia, voyaging from the Equator to the Antarctic ice edge, perpetually collecting atmospheric composition data from highly under-sampled parts of the atmosphere. During its frequent voyages through the remote Southern Ocean, this new GAW station is providing insight into the closest analogue we have to a pristine or undisturbed atmosphere. This new understanding is invaluable for improving climate models.



The RV *Investigator* steams past a colossal iceberg in the Southern Ocean.

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