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CONTENTS

SUMMARY

This *Data Summary* reports the results of basic analyses of greenhouse and some reactive gas data submitted to the WMO World Data Centre for Greenhouse Gases (WDCGG). This issue covers observations from 1968 through 2010, based on data reported to the WDCGG by October 2011. The *Data Summary* includes analyses of global, hemispheric and latitudinal monthly mean mole fractions of greenhouse and some reactive gases, and provides information on the current state of mole fractions of these gases at the stations.

Although only monthly mean mole fractions were used for the analyses, the WDCGG greatly appreciates those stations that submit daily and hourly mean mole fractions, which are important for analysing variations on shorter time scales. All data submitted to the WDCGG are available on its web site, http://ds.data.jma.go.jp/gmd/wdcgg/.

To represent dry mole fractions, this *Data Summary* uses the units ppm, ppb and ppt, which correspond to the SI units μ mol/mol, nmol/mol and pmol/mol, respectively.

Variations in the mole fractions of some gases are presented as combinations of seasonal cycles and deseasonalized long-term trends. Growth rates are presented as time derivatives of the long-term trends. The analytical results are summarized below for each greenhouse and reactive gas.

Carbon Dioxide (CO₂)

The level of carbon dioxide (CO₂), which contributes the most to increases in anthropogenic induced radiative forcing, has been increasing since the beginning of the industrial era. The global average mole fraction of CO₂ reached a new high of 389.0 ppm in 2010, 139% of that in the pre-industrial period (1750). The annually averaged increase of 2.3 ppm from 2009 to 2010 was larger than the average growth rate for the 1990s (about 1.5 ppm/year) and for the past decade (about 2.0 ppm/year).

The global growth rate of CO_2 has a significant interannual variation driven by natural processes. Growth rates higher than 2 ppm/year in 1987/1988, 1997/1998, 2002/2003 and 2009/2010 resulted from warmer conditions caused by El Niño-Southern Oscillation (ENSO) events. The anomalously strong El Niño event in 1997/1998 resulted in greater annual increases in CO_2 worldwide in 1998 than during any other one-year period. The high growth rates in 2006 may have been related to the global high temperature during the same year. The exceptionally low growth rates in 1992, including negative values in northern high latitudes, may have been due to low global temperatures following the eruption of Mount Pinatubo in 1991. Variations in CO_2 mole fraction can be seen both on seasonal and long-term scales. The seasonal amplitudes are large in northern high and mid-latitudes and small in the Southern Hemisphere. In southern low latitudes, there is no clear annual cycle, but a semiannual cycle can be determined.

Methane (CH₄)

Methane (CH₄) is the second most significant greenhouse gas which is largely influenced by anthropogenic activity and whose level has been increasing since the beginning of the industrial era. The annual average mole fraction was 1808 ppb in 2010, an increase of 5 ppb since 2009. The mole fraction is now 258% of that in the pre-industrial period. This is the fourth year of strong methane increases after the levelling-off in the beginning of this century.

The latitudinal gradient of CH_4 mole fraction is large from the northern mid-latitudes to the tropics, suggesting that the major sources of CH_4 are located in the Northern Hemisphere.

CH₄ growth rates decreased significantly in some years, including 1992, when negative values were recorded in northern high and mid-latitudes. However, both hemispheres experienced high growth rates in 1998, caused by an exceptionally high global mean temperature. The global growth rates were generally low from 1999 to 2006, except during the El Niño event of 2002/2003. The global growth rate averaged over the period 1984–1990 was 11.5 ppb/year, but decreased markedly in the 1990s. The average global growth rate for the period 2000–2010 was 2.6 ppb/year, but in the last four years through 2010, the global mole fraction increased by a total of 24 ppb.

 CH_4 mole fractions vary seasonally, being relatively high in winter and low in summer. Unlike CO_2 , the seasonal amplitudes of CH_4 are large, not only in the Northern Hemisphere but also in southern high and mid-latitudes as those are connected with a chemical sink of methane. In southern low latitudes, a distinct secondary maximum in boreal winter overlies the annual cycle.

Nitrous Oxide (N₂O)

Nitrous oxide (N₂O) is an important greenhouse gas whose level is increasing globally. N₂O data submitted to the WDCGG show that mole fractions are increasing in both hemispheres. The global mean mole fraction reached a new high of 323.2 ppb in 2010, 0.8 ppb higher than that in the previous year. This mole fraction corresponds to 120% of that in the pre-industrial period. The mean growth rate of the global mean mole fraction over the period 2000–2010 was 0.75 ppb/year and the inter-hemispheric gradient in N_2O is 1.1 ppb (averaged over the years 1980 to 2010), indicating that majority of N_2O sources are situated in the Northern hemisphere.

Halocarbons and Other Halogenated Species

Halocarbons, most of which are anthropogenic and generated from 20^{th} Century, are potent greenhouse gases, with some also acting as ozone-depleting compounds. Levels of some halocarbons (e.g. CFCs) increased in the 1970s and 1980s, but this increase has almost ceased by now, due to the production and comsumption of halocarbons under the Montreal Protocol on Substances that Deplete the Ozone Layer and its subsequent Adjustments and Amendments. However, some substances targeted by the Kyoto Protocol but not regulated by the Montreal Protocol, such as HFCs and SF₆, are increasing.

The mole fraction of CFC-11 peaked around 1992 and then started decreasing. The growth rate of CFC-12 increased until around 2005 and then started decreasing gradually. The mole fraction of CFC-113 stopped increasing in the 1990s, followed by a slight decrease over the last decade. The mole fractions of HCFCs, used mainly as substitutes for CFCs, have increased significantly during the last decade, but the growth of HCFC-141b decelerated rapidly in the second half of the decade. The mole fraction of Halon-1211 has not increased since 2005, whereas the mole fraction of Halon-1301 is increasing. The mole fraction of CCl₄ was maximal around 1991 and has since decreased slowly. The mole fraction of CH₃CCl₃ peaked around 1992 and decreased thereafter. The mole fractions of HFC-134a, HFC-152a and SF₆ are increasing.

Surface Ozone (O₃)

Ozone (O_3) plays important roles in the atmospheric environment through radiative and chemical processes. It absorbs solar UV radiation in the stratosphere, influencing the vertical temperature profile as well as terrestrial IR radiation, and contributing to the greenhouse effect as a greenhouse gas. Ozone is also involved in the chemical transformations of the primary air pollutants, as its mole fraction in the boundary layer serves as an indicator of air quality.

The mole fraction of O_3 near the surface, so-called surface ozone, reflects various processes. While some of the O_3 in the troposphere comes from the stratosphere, the rest is chemically produced in the troposphere through oxidation of CO or hydrocarbons in the presence of NO_x .

The mole fraction of surface ozone is measured at many locations in various environments. Due to its uneven geographic distribution, however, it is difficult to identify a global long-term trend of surface O_3 (WMO, 2011b).

Carbon Monoxide (CO)

Carbon monoxide (CO) is not a greenhouse gas itself but influences the mole fractions of greenhouse gases by affecting hydroxyl radicals (OH). Its mole fractions in northern high latitudes have been increasing since the mid-19th century. In 2010, the global mean mole fraction of CO was about 93 ppb. The mole fraction is high in the Northern Hemisphere and low in the Southern Hemisphere, suggesting substantial anthropogenic emissions in the Northern Hemisphere.

There was a large fluctuation in growth rate of CO. High positive rates followed by high negative rates in northern latitudes and southern low latitudes from 1997 to 1999. The growth rates in the Northern Hemisphere increased again in 2002. Both growth rate increases are attributed to large scale biomass burnings related with El Niño events.

The monthly mean mole fractions show seasonal variations, with large amplitudes in the Northern Hemisphere and small amplitudes in the Southern Hemisphere with opposite phase.

Nitrogen Monoxide (NO) and Nitrogen Dioxide (NO₂)

Nitrogen oxides (NO_x, i.e., NO and NO₂) are not greenhouse gases are involved in the photochemical production of ozone in the troposphere. In the presence of NO_x, CO and hydrocarbons are oxidized to produce ozone (O₃), which affects the Earth's radiative balance as a greenhouse gas and the oxidization capacity of the atmosphere by reproducing OH.

Most of the stations that have so far reported NO_x data to the WDCGG are located in Europe. NO_x has a large temporal and geographic variability, and it is difficult to identify its long-term global trend based on a spatially limited dataset.

Sulphur Dioxide (SO₂)

Sulphur dioxide (SO_2) is not a greenhouse gas but a precursor of atmospheric sulphate aerosols. Sulphate aerosols are produced by SO_2 oxidation through photochemical gas-to-particle conversion. SO_2 has also been a major source of acid rain and deposition throughout the industrial era.

Most of the stations reporting SO_2 data to the WDCGG are located in Europe. Global analysis can not be performed on this data set.

1. INTRODUCTION

Human activities have had major impacts on the global environment. Since the beginning of the industrial era, mankind has increasingly made use of land, water, minerals and other natural resources, and continuous growth of the world human population and economies may further increase our impact on the environment. As the climate, biogeochemical processes and natural ecosystems are closely interlinked, changes in any one of these may affect the others and be detrimental to humans and other organisms. Emissions of man-made gaseous species and particulate matter alter the energy balance of the atmosphere, which in turn has implications for the interactions in the system atmosphere - hydrosphere -Complex interactions in this system, biosphere. including numerous feedbacks, are not completely understood in particular due to the lack of observations.

The World Meteorological Organization (WMO) established the Global Atmosphere Watch (GAW) Programme in 1989 to promote systematic and reliable observations of the global environment, including but not limited to, greenhouse gases (e.g., CO2, CH4, CFCs, and N_2O) and some reactive gases (e.g., O_3 , CO, NO_x , and SO₂) in the atmosphere. In October 1990, WMO designated the Japan Meteorological Agency (JMA) in Tokyo to serve as the World Data Centre for Greenhouse Gases (WDCGG). The WDCGG is responsible for collecting, archiving and providing data on greenhouse and reactive gases in the atmosphere and oceans from a number of observation sites throughout the world that participate in GAW and other scientific monitoring programmes (Appendix: LIST OF OBSERVING STATIONS). In August 2002, the WDCGG took over the role of the World Data Centre for Surface Ozone from the Norwegian Institute for Air Research (NILU).

With regard to the issue of climate change the Kyoto Protocol to the United Nations Framework Convention on Climate Change entered into force in February 2005. In March 2006, WMO commenced annual publication of the WMO Greenhouse Gas Bulletin, which summarizes the state of greenhouse gases in the atmosphere. The seventh issue of the Bulletin was published in November 2011. The WDCGG contributes to the production of the bulletin through timely and adequate collection and analysis of data in cooperation with the contributors of the data.

Since its establishment, the WDCGG has provided its users with data and other information through its regular publications, including the *Data Summary* and *DVD* (Appendix: LIST OF WMO WDCGG PUBLICATIONS). In accordance with the GAW Strategic Plan: (2008–2015) and its Addendum, all data and information have been made available on the WDCGG web site, improving the accessibility of data, information and products (WMO, 2007a; WMO, 2011a). The WDCGG published the Data Submission and Dissemination Guide in 2007 (WMO, 2007b), which, with its revision in 2009 (WMO, 2009b), is designed to facilitate submission of observational data and access to archived data in the WDCGG.

The GAW Strategic Plan requests that World Data Centres assist data users by providing the data and analysis related to atmospheric observations. To this end, the WDCGG provides global and integrated diagnostics on the state of greenhouse and some reactive gases as analytical information in the *Data Summary*. The WDCGG global analysis methods have been described in a GAW technical report (WMO, 2009a). The content of the *Data Summary* is revised and improved based on comments from data contributors and scientists. We hope the diagnostic information presented here will promote the use of data on greenhouse and reactive gases and will enhance appreciation of the value of the GAW Programme.

All users are required to accept the following statement endorsed by the Commission for Atmospheric Sciences (CAS) at its thirteenth session: "For scientific purposes, access to these data is unlimited and provided without charge. By their use you accept that an offer of co-authorship will be made through personal contact with the data providers or owners whenever substantial use is made of their data. In all cases, an acknowledgement must be made to the data providers or owners and to the data centre when these data are used within a publication." The WDCGG requests data users to make appropriate acknowledgements. The principal investigators and other contacts can be obtained from the WDCGG website, as well as from the GAW Station Information System (GAWSIS) website. http://gaw.empa.ch/gawsis/. Information on these websites is updated in cooperation with the data contributors and the WMO Secretariat.

Finally, the WDCGG would like to thank all data contributors worldwide, including those involved in on-site measurements, for their efforts in maintaining the observational programme and for continuous data provision.

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2. ANALYSIS

The WDCGG collects, archives and provides observational data on the dry mole fractions of greenhouse and some reactive gases, and publishes diagnostic information on these gases based on the reported data.

The long-term trends and seasonal variations in the mole fractions of CO_2 , CH_4 and CO are calculated for the whole globe (global means) and for latitudinal belts (zonal means). Only global and hemispheric long-term trends are calculated for N₂O. Global long-term trends in the surface O_3 , are not analysed due to its substantial spatial gradients, and its uneven geographic distribution which is poorly covered by observation sites. For halocarbons, NO_x and SO₂, only monthly mean mole fractions over time are presented without global, hemispheric or zonal averaging, due to insufficient number of reporting sites for each compound.

The units used in this analyses are ppm, ppb, and ppt, rather than the SI units of μ mol/mol, nmol/mol, and pmol/mol, respectively.

The method of analysis for CO₂, CH₄, CO and N₂O are summarized below. The details of the global analysis methods for CO₂, CH₄, and N₂O are provided in the *Technical Report of Global Analysis Method for Major Greenhouse Gases by the World Data Centre for Greenhouse Gases*, published as a GAW technical report (WMO, 2009a). Additional uncertainty can be expected in the result of CO global analysis due to diversity of scales. When assessing long-term trends for CO₂, CH₄ and N₂O, the growth rates at both ends of the period were assumed to be simple linear extensions of the adjacent year, thus avoiding end effects. For simplicity, the rates for the rest of the period were approximated by linear expressions.

(1) Site selection

For CO₂, CH₄ and N₂O, the diagnostic analyses, including global, hemispheric and zonal means, were based on data from sites that have adopted a standard scale traceable to the Primary Standard designated under GAW. These analyses also utilize data on other standard scales that are convertible to the WMO/GAW scale through a proven equation. The letters encouraging data submitters on the most recent WMO scales are sent out regularly since 2010.

Selection of observation sites have also been based on whether they provide data representing a reasonably large geographical area, considering the fact that some sites may be susceptible to local sources and sinks. Sites are selected objectively using data submitted to the WDCGG. Only those sites for CO₂, CH₄ and CO that provide annual mean mole fractions falling within a range of $\pm 3\sigma$ from a curve fitted to the LOESS model curve (Cleveland *et al.*, 1988) have been selected, with outliers rejected in an iterative manner. This procedure does not affect the datasets present in the WDCGG, and these data may be useful for purposes other than global analyis, such as sources and sinks identification.

The sites selected according to the above criteria are marked with asterisks in Plate 3.1 for CO_2 , Plate 4.1 for CH_4 , Plate 5.1 for N_2O and Plate 8.1 for CO, which represent 128, 125, 53 and 53 of the submitted datasets respectively.

(2) Analysis of long-term trends

The mole fractions of greenhouse and reactive gases over time, measured under unpolluted conditions, exhibit variations on different time scales. The two major components are seasonal variations and long-term trend. Various attempts have been made to divide measured data into these two components, including objective curve fitting (Keeling *et al.*, 1989), digital filtering (Thoning *et al.*, 1989; Nakazawa *et al.*, 1991), or both (Conway *et al.*, 1994; Dlugokencky *et al.*, 1994).

In this report, seasonal variations derived from components of Fourier harmonics and long-term trends are extracted by low-pass filtering with a cut-off frequency of 0.48 year⁻¹ for each selected site. Details are described in WDCGG Data Summary No. 22 (WMO, 2000).

(3) Estimation for missing periods and gaps

The number and distribution of sites used to assess trends during the analysis period should be kept as constant as possible to avoid the effects of changes in the availability of data over time. However, only a small number of sites provided data throughout the entire analysis period; others may have covered shorter periods or had gaps in measurements due to different reasons. To use as many sites as possible, missing values were estimated for the calculation of zonal means, using interpolation and extrapolation as described below.

For the former, gaps were interpolated linearly based on the data, by subtracting the seasonal variation calculated from the longest consecutive period of data with Lanczos filters (Duchon, 1979). The subtracted variation was added back to the data to obtain estimated mole fractions in a single sequence.

For the latter, long-term trends calculated from the interpolated series of data were extrapolated based on zonal mean growth rates calculated from other sites in the same latitudinal zone. The seasonal variation was added to the extrapolated long-term trend to obtain estimated mole fractions for the entire period of analysis. Using these statistical procedures, the future addition of new stations should not affect the consistency in global estimates over time.

(4) Calculation of global, hemispheric and zonal means

Zonal means were calculated by determining the arithmetic average of the mole fractions in each latitudinal zone, based on consistent datasets derived as above.

Global and hemispheric means were calculated as the weighted averages of the zonal means taking account of the area of each latitudinal zone.

Deseasonalized long-term trends and growth rates for the globe, each hemisphere and each latitudinal zone were calculated from the global, hemispheric and zonal means, respectively, using the low-pass filter mentioned above and the time derivatives after filtering.

CARBON DIOXIDE (CO₂)



This map shows locations of the stations that have submitted data for monthly mean mole fraction.



Plate 3.1 Monthly mean CO_2 mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south. In the case where data are reported for two or three different altitudes, only the data at the highest altitudes are illustrated. In the case where monthly means are not reported, the WDCGG calculates them from hourly or other mole fractions reported to the WDCGG by simple arithmetic mean. The data from the sites with an asterisk at the end of the station index are used for the analysis shown in Plate 3.2. (see Chapter 2)









Plate 3.2 Variation of zonally averaged monthly mean CO_2 mole fractions (top), deseasonalized long-term trends (middle), and growth rates (bottom). The zonally averaged mole fractions are calculated for each 20° zone. The deseasonalized trends and growth rates are derived as described in Chapter 2.

3. CARBON DIOXIDE (CO₂)

Basic information on CO₂ with regard to environmental issues

Carbon dioxide (CO₂) has strong absorption bands in the infrared region and is the biggest anthropogenic contributor to the greenhouse effect. CO₂ accounted for about 64% of total increase in the radiative forcing due to long-lived greenhouse gases in the atmosphere from 1750 to 2010 (WMO, 2011b).

The balance between its emission and absorption over the continents and oceans determines the mole fraction of CO_2 in the atmosphere. About 762 gigatonnes of carbon are present in the atmosphere as CO_2 (IPCC, 2007) and annual anthropogenic emissions mainly due to fossil fuel combustion gigatonnes reached 8.4±0.5 2009 in (http://www.globalcarbonproject.org). Carbon in the atmosphere is exchanged with two other large reservoirs, the terrestrial biosphere and the oceans. CO₂ exchanges between the atmosphere and terrestrial biosphere occur mainly through absorption by photosynthesis and emission from the respiration of plants and the decomposition of organic soils. These biogenic activities vary seasonally, resulting in large seasonal variations in the level of CO_2 . CO_2 exchange between the atmosphere and oceans occurs in direction determined by the gradient of CO₂ mole fraction, and varies in time and space.

The current mole fractions of atmospheric CO_2 far exceed historic records, dating back 650,000 years (Solomon et al., 2007). Based on the results of ice core studies, the mole fraction of atmospheric CO_2 in pre-industrial times was about 280 ppm (IPCC, 2007). The emission of CO_2 due to human activities has increased dramatically since the beginning of the industrial era, impacting CO2 exchange rates between different reservoirs and CO₂ levels not only in the atmosphere but in the oceans and terrestrial biosphere. The global carbon cycle, which is comprised mainly of CO₂, is not fully understood. About half of anthropogenic CO₂ emissions have remained in the atmosphere, with the remainder removed by sinks, including the terrestrial biosphere and oceans, and non-anthropogenic sources. However, the amount of CO₂ removed from the atmosphere varies significantly over time (Figure 3.1).

Carbon isotopic studies have shown the importance of the terrestrial biosphere and oceans as sources and sinks of CO_2 (Francey *et al.*, 1995; Keeling *et al.*, 1995; and Nakazawa *et al.*, 1993, 1997). In contrast, the atmospheric content of O_2 depends primarily on its removal by the burning of fossil fuels and on its release from the terrestrial biosphere. Therefore, the uptake of carbon by the terrestrial biosphere and oceans can be estimated from the combination of measurements of O_2 (O_2/N_2) and CO_2 (Manning and Keeling, 2006). CO2 equilibrates largely between the atmosphere and the ocean and, depending on associated increases in acidity and in ocean warming, typically about 20% of added tons of CO₂ remain in the atmosphere while about 80% are mixed into the ocean. A quasi-equilibrium amount of CO₂ is expected to be retained in the atmosphere by the end of the millennium that is surprisingly large: typically 40% of the peak concentration enhancement over preindustrial values (280 ppmv) (Solomon *et al.*,2009).

Large amounts of CO_2 are exchanged among these reservoirs, and the global carbon cycle is coupled with the climate system on seasonal, yearly and decadal time scales. Accurate understanding of the global carbon cycle is essential for estimating future CO_2 mole fractions in the atmosphere.



Fig. 3.1. Annual mean growth rates of CO₂ in the atmosphere, calculated from observational data (red columns) and from data for anthropogenic emissions (green curve). The estimated growth rates were calculated from CO₂ emissions (from CDIAC, Boden et al., 2011) .The values from 2009 to 2010 is quick estimation of Carbon Dioxide Information Analysis Center (CDIAC), expressed as moles divided by the total mass of gas in the atmosphere (5.2 petatonnes) converted to moles based on the mean molar weight of air (about 29). The observed growth rates were calculated by the WDCGG. The observational CO2 abundance is expressed as mole fraction with respect to dry air, CO_2 while the amount calculated from anthropogenic emissions is based on the atmosphere, including water vapor, usually in a mole fraction less than 1%.

Mole fractions of CO_2 can be analyzed utilizing data submitted to the WDCGG from fixed stations and some ships. The observation sites from which data were used for the analysis are shown on the map at the beginning of this chapter. They include fixed stations performing continuous measurements as well as flask-sampling stations, including those in the NOAA/ESRL cooperative air sampling network. In addition, mobile stations on ships and aircraft and other stations observing on an event basis report their data to the WDCGG (see Appendix: LIST OF OBSERVING STATIONS), which are not used for global analysis.

Annual variations of CO₂ mole fraction in the atmosphere

The monthly mean mole fractions of CO₂ used in the analysis are shown in Plate 3.1, with mole fraction levels illustrated in different colours. Global. hemispheric and zonal mean mole fractions were analysed based on data from selected stations under unpolluted conditions (see the caption to Plate 3.1). Latitudinally averaged mole fractions of atmospheric CO₂, together with their deseasonalized components and growth rates, are shown as three-dimensional representations in Plate 3.2. These plots show that the seasonal variations in mole fraction are large in northern high and mid-latitudes, but are indistinct in the Southern Hemisphere. The increases in the Northern Hemisphere precede those in the Southern Hemisphere by one or two years, and the interannual



Fig. 3.2 Global monthly mean mole fraction of CO_2 from 1983 to 2010, including deseasonalized long-term trend shown as a red line (top) and annual growth rate (bottom).

variation in growth rate is larger in the Northern Hemisphere.

Figure 3.2 shows global monthly mean CO_2 mole fractions and their growth rates from 1983 to 2010. The global average mole fraction reached a new high of 389.0 ppm, in 2010 which is 138% of the pre-industrial level of 280 ppm. The annually averaged increase from 2009 to 2010 was 2.3 ppm, larger than the average of annual increases for the 1990s (about 1.5 ppm/year) and that of the past decade (about 2.0 ppm/year).

The global growth rate shows large interannual variations, with an instantaneous maximum of about 3 ppm/year in 1998 and a minimum below 1 ppm/year in 1992. There were short periods of high rates in 1987/1988, 1997/1998, 2002/2003, 2005/2006, 2007 and 2009/2010.

Figure 3.3 shows monthly mean mole fractions and long-term trends from 1983 to 2010 for each 30° latitudinal zone, indicating that there were clear long-term increases in both hemispheres and seasonal variations in the Northern Hemisphere.



Fig. 3.3 Monthly mean mole fractions of CO_2 from 1983 to 2010 for each 30° latitudinal zone (dots) and their deseasonalized long-term trends (red lines).

As shown in Figure 3.4, the growth rates for each 30° latitude zone fluctuated between -0.3 and 3.6 ppm/year, with the largest interannual variability in northern high latitudes. High growth rates for all 30° latitude zones were observed in 1987/1988, 1997/1998, 2002/2003, 2005, 2007, with negative rates recorded in northern high latitudes in 1992.

Changes in growth rate are partly associated with El Niño-Southern Oscillation (ENSO). The El Niño events in 1982/1983, 1986–1988, 1991/1992, 1997/1998 and 2002/2003coincided with high growth rates of CO₂, with an exception in 1992. The growth rates of CO₂ observed by aircraft at high altitudes (8–13 km) over the Pacific Ocean were also associated with ENSO (Matsueda *et al.*, 2002). The latest El Niño event has occurred in 2009/2010.

During El Niño events, the up-welling of CO₂-rich ocean water in the eastern equatorial Pacific is suppressed, resulting in reduced CO₂ emissions from this area. In contrast, El Niño events induce high temperature anomalies in many areas, particularly in the tropics, resulting in increased CO₂ emissions from the terrestrial biosphere due to the enhanced respiration of plants and activated decomposition of organic matter in soil (Keeling et al., 1995). This effect is enhanced by the suppression of plant photosynthesis in areas of anomalously low precipitation, particularly in the tropics. These oceanic and terrestrial processes during El Niño events have opposing effects, but Heimann and Reichstein (2008) suggested that the latter was the main cause of the variation in the CO₂ growth rate.



Fig. 3.4 Long-term trends in the mole fraction of CO_2 for each 30° latitudinal zone (top) and their growth rates (bottom).

However, an exceptionally low CO₂ growth rate occurred during the El Niño event in 1991/1992. The injection of 14-20 Mt of SO2 aerosols into the stratosphere by the Mount Pinatubo eruption in June 1991 affected the radiation budget and atmospheric circulation (Hansen et al., 1992; Stenchikov et al., 2002), resulting in a drop in global temperature. Angert *et al.* (2004) suggested that the low CO_2 growth rate observed during this El Niño event was due to reduced CO₂ emissions caused by consequent changes in the respiration of terrestrial vegetation and the decomposition of organic matter (Conway et al., 1994; Lambert et al., 1995; Rayner et al., 1999), and by enhanced CO₂ absorption due to intensive photosynthesis caused by an increase in diffuse radiation (Gu et al., 2003).

Seasonal cycle of CO₂ mole fraction in the atmosphere

Figure 3.5 shows average seasonal cycles in the mole fraction of CO_2 for each 30° latitudinal zone. The seasonal cycles are clearly large in amplitude in northern high and mid-latitudes and small in the Southern Hemisphere. The seasonal cycle in the Northern Hemisphere is mainly dominated by the land biosphere (Nevison et al., 2008), and it is characterized by rapid decreases from June to August and large returns from September to December.



Fig. 3.5 Average seasonal cycles in the mole fraction of CO_2 for each 30° latitudinal zone obtained by subtracting long-term trends from the zonal mean time series. Error bar expresses standard deviation of each month.

The mole fractions of CO_2 in northern low latitudes lagged behind that in high latitudes by one or two months. Minimum values appeared in August in northern high and mid-latitudes and in September in northern low latitudes.

In the Southern Hemisphere, seasonal variations showed small amplitudes with a half-year delay due to small amounts of net emission and absorption by the terrestrial biosphere. Seasonal variations in both northern and southern mid-latitudes were apparently superimposed in southern low latitudes $(0-30^{\circ}S)$. The direct influence of sources and sinks in the Southern Hemisphere may be partially cancelled by the propagation of an antiphase variation from the Northern Hemisphere.

Figure 3.6 shows latitudinal distributions of the mole fractions of CO_2 in January, April, July and October 2010, from sites marked with an asterisk in Plate 3.1. In latitudes north of 30°N, the mole fractions increased towards higher latitudes in January and April, and decreased towards higher latitudes in July, corresponding to the large seasonal variations in northern high and mid-latitudes, variations associated with activities of the terrestrial biosphere.



Fig. 3.6 Latitudinal distributions of the monthly mean mole fractions of CO_2 in January, April, July and October 2010.

METHANE (CH₄)

- : CONTINUOUS STATION
- △ : FLASK STATION
- □ : FLASK MOBILE (SHIP)
- ▼ : REMOTE SENSING STATION



This map shows locations of the stations that have submitted data for monthly mean mole fraction.





Plate 4.1 Monthly mean CH4 mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south. In the case where data are reported for two or three different altitudes, only the data at the highest altitudes are illustrated. In the case where monthly means are not reported, the WDCGG calculates them from hourly or other mole fractions reported to the WDCGG by simple arithmetic mean. The data from the sites with an asterisk at the end of the station index are used for the analysis shown in Plate 4.2. (see Chapter 2)









Plate 4.2 Variation of zonally averaged monthly mean CH_4 mole fractions (top), deseasonalized long-term trends (middle), and growth rates (bottom). The zonally averaged mole fractions are calculated for each 20° zone. The deseasonalized trends and growth rates are derived as described in Chapter 2.

4. METHANE (CH₄)

Basic information on CH₄ with regard to environmental issues

Methane (CH₄) is the second most important anthropogenic greenhouse gas, with an estimated global warming potential per molecule 25 times greater over a 100 year horizon and 72 times greater over a 20 years horizon than CO₂. Between 1750 and 2010, CH₄ accounted for about 18% of total increase in radiative forcing due to long-lived greenhouse gases in the atmosphere (WMO, 2011c).

Analyses of air trapped in ice cores from Antarctica and the Arctic revealed that the current atmospheric CH₄ mole fraction is the highest it has been over the last 650,000 years (Solomon et al., 2007). The mole fraction of CH₄ remained steady at 700 ppb from 1000 A.D. until the start of the industrial era (Etheridge et al., 1998), after which it began to increase. Measurements in ice cores have shown that the interpolar differences in CH₄ mole fractions between Greenland and Antarctica ranged from 24 to 58 ppb between 1000 and 1800 A.D. (Etheridge et al., 1998). This difference reached a value of 136 ppb as follows from the comparison of northern high and southern high latitude averaged over the years 1984 to 2010, Fig. 4.3. This fact reflects emissions from the Northern Hemisphere, where major anthropogenic and natural sources are situated.

CH₄ is emitted by both natural and anthropogenic sources, including natural wetlands, oceans, landfills, rice paddies, enteric fermentation, fossil fuel production and consumption and biomass burning. Denman et al. (2007) estimated the global emission of CH_4 is 582 teragrams (Tg) CH_4 per year, with more than 60% related to anthropogenic activities. CH₄ removed from the atmosphere by reaction with hydroxyl radicals (OH) in both the troposphere and stratosphere, and by reaction with chlorine atoms and $O(^{1}D)$, an excited state of oxygen, in the stratosphere. CH₄ is one of the most important sources of water vapour in the stratosphere and has an atmospheric lifetime of about 10 years. More information regarding sources and sinks of CH₄ must be collected to better understand the budget of atmospheric CH₄.

Mole fractions of CH_4 are analyzed by using data submitted to the WDCGG from fixed stations and some ships. These observational sites are shown on the map at the beginning of this chapter.

Annual variation of CH₄ mole fraction in the atmosphere

The monthly mean mole fractions of CH_4 used in this analysis are shown in Plate 4.1, with the mole fraction levels illustrated in different colours. Global, hemispheric and zonal mean mole fractions have been

calculated based on data from selected stations under unpolluted conditions (see the caption for Plate 4.1). Latitudinally averaged atmospheric CH₄ mole fractions, together with their deseasonalized components and growth rates, are shown as three-dimensional representations in Plate 4.2. These plots show that the seasonal variations in CH₄ mole fractions are larger in the Northern than in the Southern Hemisphere and that the increase in the Northern Hemisphere propagates to the Southern Hemisphere. The growth rates vary on a global scale. These features are similar to those for CO_2 (see Section 3). There is a large latitudinal gradient in CH₄ mole fraction from the northern mid-latitudes to the tropics, suggesting major sinks in the tropics, where the mole fraction of OH radicals is higher.



Fig. 4.1 Global monthly mean mole fraction of CH_4 from 1984 to 2010, including deseasonalized long-term trend in red line (top) and annual growth rate (bottom).

Figure 4.1 shows global monthly mean mole fractions and the global growth rates for CH_4 from 1984 to 2010. The mean mole fraction was 1808 ppb in 2010, an increase of 5 ppb since 2009. Mole fraction did not change much between 1999 and 2006. The average growth rate over the period 2000–2010 was

2.6 ppb/year. The current mole fraction is 258% of its pre-industrial level, 700 ppb.

Figure 4.2 shows monthly mean mole fractions from 1984 to 2010 for each 30° latitudinal zone. The smallest magnitude of the seasonal variations is registered in the latitudinal zone between the equator and 30° S.



Fig. 4.2 Monthly mean mole fractions of CH_4 from 1984 to 2010 for each 30° latitudinal zone (dots) and their deseasonalized long-term trends (red lines).

Figure 4.3 summarizes deseasonalized long-term trends for each 30° latitudinal zone and their growth As it is the most distinctly seen in the rates. deseasonalized long-term trends, a latitudinal gradient between the northern high and mid-latitudes with higher mole fractions and the southern latitudes with lower mole fractions is clearly pronounced, while mole fractions in most latitudinal belts have similar tendency. In the 1990s, the growth rates clearly decreased in all latitudinal zones, while remaining positive nevertheless. The declined growth rate was especially evident during the second half of 1992 in 1996, and almost even in 1999 and in 2004/2005, when growth rates were less than 5 ppb/year in all latitudes. In 1998, the global growth rate increased to about 11 ppb/year (Fig. 4.1). Maximum increases occurred in northern high and mid-latitudes, where the growth rates were over 15

ppb/year. In 2000 and 2001, the global growth rate decreased to around -1 ppb/year. Around 2002/2003, the growth rates increased in the Northern Hemisphere, especially in northern high and mid-latitudes where they reached about 10 ppb/year. The global growth rate was -3 ppb/year in 2004 and 1 ppb/year in 2005. Despite the large growth rates in 1998 and 2002/2003, during El Niño events, the global mean mole fraction was relatively stable between 1999 and 2006. However, the global mean mole fraction increased again by a total of 24 ppb in the four years starting in 2007.



Fig. 4.3 Long-term trends in the mole fraction of CH_4 for each 30° latitudinal zone (top) and their growth rates (bottom).

The large increase in CH_4 growth rate in 1991 may have been caused by decreased levels of OH radicals in the atmosphere due to reduced UV radiation resulting from the eruption of Mount Pinatubo in 1991 (Dlugokencky *et al.*, 1996), and the subsequent decrease in 1992 may have been due to an increase in OH radicals resulting from the depletion of stratospheric ozone following this eruption (Bekki *et al.*, 1994).

In 1998, the growth rates were high in all latitudes, which may have been due to increased emissions in northern high latitudes and tropical wetlands caused by high temperatures and increased precipitation, as well as by biomass burning in boreal forests, mainly in Siberia (Dlugokencky *et al.*, 2001). In contrast, Morimoto *et al.* (2006) estimated from isotope observations that the contribution of biomass burning to the increase in 1998 was about half that of wetlands. The growth rates were low from 1999 to 2006, with an exception during the El Niño event of 2002/2003. The causes of these decreases in CH₄ growth rates remain

unclear.

Since 2007, atmospheric CH_4 has increased significantly throughout the entire monitoring network (Rigby *et al.*, 2008; Dlugokencky *et al.*, 2009). Although these increases may have been caused by emissions from natural sources in northern latitudes and the tropics, the reasons for renewed methane growth are not fully understood (WMO, 2011b).

The WMO/GAW observational network includes the observation of carbon stable isotopes in methane, with 19 datasets submitted to the WDCGG Such observations can be useful for the identification of primary methane sources.

Seasonal cycle of CH₄ mole fraction in the atmosphere

Figure 4.4 shows seasonal cycles in the mole fraction of CH₄ for each 30° latitudinal zone. The seasonal cycles are driven mainly by reaction with OH radicals, a major CH₄ sink in the atmosphere. Seasonal cycles are also affected by the magnitude and timing of CH₄ emissions from sources such as wetlands and biomass burning as well as by its atmospheric transport. The seasonal cycles are large in amplitude in the Northern Hemisphere. Unlike CO2, amplitudes were also large in southern high and mid-latitudes. Seasonally, the Northern Hemisphere shows minima in summer and maxima in winter, while the Southern Hemisphere shows a seasonal cycle lagging two-thirds to three-quarter years behind. The seasonal variations in the mole fraction of CH4 were almost consistent with those of the OH radical that reacts with CH₄. Southern low latitudes have a distinct antiphase annual component with that of the seasonal cycle arising from southern mid-latitudes. The maximum in the former component occurs in boreal winter due to the interhemisphere transportation of CH4 from the Northern Hemisphere.



Fig. 4.4 Average seasonal cycles in the mole fraction of CH_4 for each 30° latitudinal zone obtained by subtracting long-term trends from the zonal mean time series. The standard deviation is indicated by vertical error bars.

NITROUS OXIDE (N₂O)

• : CONTINUOUS STATION

△ : FLASK STATION



This map shows locations of the stations that have submitted data for monthly mean mole fraction.



Plate 5.1 Monthly mean N_2O mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south. The data from the sites with an asterisk at the end of the station index are used for the analysis shown in Fig.5.1. (see Chapter 2)









Plate 5.2 Variation of zonally averaged monthly mean N_2O mole fractions (top), deseasonalized long-term trends (middle), and growth rates (bottom). The zonally averaged mole fractions are calculated for each 30° zone. The deseasonalized trends and growth rates are derived as described in Chapter 2.

5. NITROUS OXIDE (N_2O)

Basic information on N₂O with regard to environmental issues

Nitrous oxide (N_2O) is a relatively stable greenhouse gas in the troposphere with an "adjustment-time" of 114 years. Between 1750 and 2010, N2O accounted for about 6 % of total increase in radiative forcing due to long-lived greenhouse gases (WMO, 2011c). N₂O is the third most important anthropogenic greenhouse gas in the atmosphere. The mole fraction of N₂O in the atmosphere has increased steadily from about 270 ppb in pre-industrial times to its current value, which is 20% higher. N_2O is emitted into the atmosphere from natural and anthropogenic sources, including the oceans, soil, combustion of fuels, biomass burning, use of fertiliser and various industrial processes. The amount of N₂O emitted into the atmosphere by human activities is approximately equal to that derived from natural systems (oceans, chemical oxidation of ammonia in the atmosphere, and soils). Most of the anthropogenic N₂O found in the atmosphere comes from the transformation of fertilizer nitrogen into N₂O and its subsequent emission from agricultural soils. N₂O breaks down mainly by photo-dissociation in the stratosphere, forming nitrogen oxides that trigger ozone depleting reactions, so it plays an important role in ozone depletion (Ravishankara et al., 2009). However, the cycling of N₂O as a part of global nitrogen cycle is not well understood yet.



Fig. 5.1 Global monthly mean mole fraction of N_2O from 1980 to 2010, including deseasonalized long-term trend shown as a red line (top) and annual growth rate (bottom).

Annual variation of N₂O mole fraction in the atmosphere

Mole fractions of N₂O are analysed by using the data submitted to the WDCGG from fixed stations and some ships. The observational sites that supplied data used for this analysis are shown on the map at the beginning of this chapter. The monthly mean mole fractions of N₂O used in the global analysis are shown in Plate 5.1, with the various mole fraction levels illustrated in different colours. The data submitted to the WDCGG show that N₂O mole fractions have increased at almost all stations. Latitudinally averaged atmospheric N₂O mole fractions, together with their deseasonalized components and growth rates, are shown as three-dimensional representations from 1980 to 2010 in Plate 5.2. Figure 5.1 shows global monthly mean N₂O mole fraction from 1980 to 2010 and its long-term trend. The global mean mole fraction reached a new high of 323.2 ppb in 2010, an increase of 0.8 ppb over the previous year. The mean growth rate of the global mean mole fraction during the period 2000 to 2010 was 0.75 ppb/year. Atmospheric growth rate showed substantial variability (from 0.6 to 1.0 ppb/year) from the beginning of observations. There was an inter-hemispheric gradient in the mole fraction of N₂O, of 1.1ppb averaged over the years 1980 to (Figure 5.2 upper panel) indicating that 2010 majority of the N₂O sources is situated in the Northern Hemisphere (mostly agriculture).



Fig. 5.2 Monthly mean mole fractions of N_2O from 1980 to 2010 (top) and annual growth rates (bottom), averaged over the Northern and Southern Hemispheres.

HALOCARBONS AND OTHER HALOGENATED SPECIES

• : CONTINUOUS STATION

△ : FLASK STATION



This map shows locations of the stations that have submitted data for monthly mean mole fraction.



Plate 6.1 Monthly mean (a) CFC-11, (b) CFC-12, (c) CFC-113, (d) Halon-1211, (e) Halon-1301, (f) HCFC-22, (g) HCFC-141b, (h) HCFC-142b mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south.



Plate 6.2 Monthly mean (a) CCl_4 , (b) CH_3CCl_3 , (c) HFC134a, (d) HFC152a, (e) CH_3Cl , (f) SF_6 mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south.

6. HALOCARBONS AND OTHER HALOGENATED SPECIES

Basic information on halocarbons with regard to environmental issues

Halocarbons are carbon compounds containing one or more halogens, i.e., fluorine, chlorine, bromine or with most being industrial products. iodine, Halocarbons are classified into chlorofluorocarbons (CFCs), which contain fluorine and chlorine; the hydrochlorofluorocarbons (HCFCs), which contain hydrogen in addition to fluorine and chlorine; and the halons, which contain bromine and other halogens. Perfluorocarbons (PFCs) are carbon compounds in which all hydrogen atoms are replaced by fluorine atoms, and hydrofluorocarbons (HFCs) are halocarbons that contain hydrogen and fluorine but no chlorine. Sulphur hexafluoride (SF₆), although not a halocarbon, behaves similarly to halocarbons and it is a potent long-lived greenhouse gas. Carbon tetrachloride (CCl₄) and methyl chloroform (CH₃CCl₃) are produced industrially, whereas methyl chloride (CH₃Cl) has natural sources. Although the mole fractions of the halocarbons are relatively low in the atmosphere, they have high global warming potentials. The halocarbons have been shown to account for about 12% of the total increase in radiative forcing due to long-lived greenhouse gases from 1750 to 2010 (WMO, 2011c).

The halocarbons are colourless, odourless and innocuous substances that can be readily gasified and liquefied and have low surface tension. Thus, they were commonly used as refrigerants, propellants and detergents for semiconductors, resulting in a rapid increase in their mole fractions in the atmosphere until the mid-1980s. Halocarbons containing chlorine and bromine led to the depletion of the ozone layer. Since the mid-1990s, the Montreal Protocol on Substances that Deplete the Ozone Layer and its subsequent Adjustments and Amendments have progressively increased the regulation of the production, consumption and trade of ozone-depleting compounds.

The CFCs are destroyed mainly by ultraviolet radiation in the stratosphere, and their lifetimes are generally long (*e.g.*, about 50 years for CFC-11). However, the HCFCs and CH₃CCl₃, which contain hydrogen, react with hydroxyl radicals (OH) in the troposphere and have relatively short lifetimes (*e.g.*, about 5 years for CH₃CCl₃). As the reaction with OH in the troposphere is a major sink for CH₃CCl₃, global measurements of CH₃CCl₃ provide an accurate estimate of the global mole fraction of OH (Prinn *et al.*, 2001). Due to substantial decrease of CH₃CCl₃ in the atmosphere the other compounds are considered now to trace OH mole fraction changes.

The Kyoto Protocol to the United Nations Framework Convention on Climate Change (UNFCCC), which entered into force on 16 February 2005, specifies HFCs, PFCs and SF_6 as targets for quantified emission limitation and reduction commitments.



Fig. 6.1 Time series of the monthly mean mole fractions of CFC-11, CFC-12 and CFC-113. Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.

Annual changes in the levels of halocarbons in the atmosphere

The map at the beginning of this chapter shows observational sites that have submitted data on halocarbons and other halogenated species to the WDCGG. Plates 6.1 and 6.2 show all the monthly mean mole fractions of these gases submitted to the WDCGG. The figures (6.1 - 6.7) in this chapter plot the monthly mean data reported to the WDCGG without spatial averaging. Some discrepancies in the absolute mole fractions were observed for several stations, suggesting that these stations may have

adopted different standard scales. Observational data based on identical standard scales revealed that the differences in the mole fractions between the two hemispheres were large in the 1980s for CFCs, CCl₄ and CH₃CCl₃ but have since narrowed as the emissions have been suppressed and the existing constituents have been mixed across the hemispheres.

Figure 6.1 shows monthly mean mole fractions of CFC-11 (CCl₃F), CFC-12 (CCl₂F₂) and CFC-113 (CCl₂FCClF₂) over time. The mole fractions of CFC-11 were maximal around 1992 in the Northern Hemisphere, followed by a maximum about one year later in the Southern Hemisphere. The mole fractions of CFC-113 were maximal around 1992 in the Northern Hemisphere and around 1997 in the Southern Hemisphere. The mole fractions of these gases have since been decreasing slowly in both hemispheres. The mole fraction of CFC-12 increased until around 2005 and then started decreasing gradually.

Figure 6.2 shows time series of the monthly mean mole fractions of Halon-1211 (CBrClF₂) and Halon-1301 (CBrF₃). The mole fraction of Halon-1211 has not increased since 2005, whereas the mole fraction of Halon-1301 is increasing.



Fig. 6.2 Time series of the monthly mean mole fractions of Halon-1211 and Halon-1301. Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.

Figure 6.3 shows time series of the mole fractions of HCFC-22 (CHClF₂), HCFC-141b (CH₃CCl₂F) and HCFC-142b (CH₃CClF₂). The mole fractions of these gases increased significantly during the last

decade as a result of their continued use as substitutes for CFCs. However, the growth of HCFC-141b decelerated rapidly in the second half of the decade.



Fig. 6.3 Time series of the monthly mean mole fractions of HCFC-22, HCFC-141b, and HCFC-142b. Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.

Figure 6.4 shows time series of the mole fractions of CCl_4 and CH_3CCl_3 . The mole fractions of CCl_4 in both hemispheres were at a maximum around 1991. The mole fractions of CH_3CCl_3 were at a maximum around 1992 in the Northern Hemisphere and around 1993 in the Southern Hemisphere. The mole fractions of these gases have since been decreasing.

Figure 6.5 shows time series of the monthly mean mole fractions of HFC-134a (CH_2FCF_3) and HFC-152a (CH_3CHF_2). The mole fractions of HFC-134a and HFC-152a have increased by 4 to 5-fold over the last 10 years. These increases have been larger in the Northern than in the Southern Hemisphere, suggesting that their predominant sources are located in the Northern Hemisphere.

Figure 6.6 shows a time series of the monthly mean mole fractions of methyl chloride (CH_3Cl). The mole fraction of CH_3Cl has remained steady although the signs of the seasonal variation can be noticed in the dataset.

Figure 6.7 shows a time series of the monthly mean mole fractions of SF_6 . The mole fraction of SF_6 in 2010 was double that in the mid-1990s increasing nearly linearly with a rate of 0.24 ppt/year (WMO, 2011c).



Fig. 6.4 Time series of the monthly mean mole fractions of CCl_4 and CH_3CCl_3 . Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.



Fig. 6.5 Time series of the monthly mean mole fractions of HFC-134a and HFC-152a. Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.



Fig. 6.6 Time series of the monthly mean mole fractions of CH_3Cl . Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.



Fig. 6.7 Time series of the monthly mean mole fractions of SF_6 . Solid circles show mole fractions measured in the Northern Hemisphere and open circles show mole fractions in the Southern Hemisphere.
SURFACE OZONE (O_3)



This map shows locations of the stations that have submitted data for monthly mean mole fraction.



Plate 7.1 Monthly mean O_3 mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south.

7. SURFACE OZONE (O_3)

Basic information on surface ozone (O_3) with regard to environmental issues

Ozone (O_3) in the atmosphere exists mostly in the stratosphere, with less than 10% in the troposphere. However, O₃ in the troposphere plays an important role in the atmospheric environment through radiative and chemical processes. O₃ absorbs UV radiation in the stratosphere, thus influencing the vertical profile of temperature and circulation in the stratosphere. Moreover, as a greenhouse gas in the troposphere, O_3 absorbs IR radiation. The latter effect is more significant in the upper troposphere, and tropospheric O₃ is the third most important anthropogenic greenhouse gas after CO₂ and CH₄ (Denman *et al.*, 2007; IPCC, 2007). Tropospheric O₃ in the northern extratropics was the greatest contributor to global warming during the 20th century, and increases in tropospheric O₃ from industrialization in developing countries was found to contribute to accelerated warming in the tropics during the latter half of the century (Shindell et al., 2006). Furthermore, by reacting with water vapour in the presence of UV radiation, O3 produces OH radicals, which control atmospheric mole fractions of many greenhouse gases, such as CH₄, through chemical reactions.

The observational results at high altitudes around 1990, compared with those from the end of the 19th century to the first half of 20th century, show increases in tropospheric O_3 in urban areas (Staehelin *et al.*, 1994). However, ozonesonde measurements in the troposphere show stable or decreasing trends in northern mid-latitudes (Oltmans et al., 2006). There is as yet no consensus on the global trend of tropospheric O_3 . Recently an attempt has been done to make a systematic review of the observed trends. It has shown that In most regions of the world — the noteworthy exception being East Asia - surface and free tropospheric ozone concentrations have not risen significantly after the year 2000. Prior to the 1990s almost all records indicate a strong rise, while during the 1990s the picture is very diverse (http://igac.jisao.washington.edu/Newsletter/ IGAC Newsletter Oct11.pdf).

Tropospheric O_3 originates from flux/mixing from the stratosphere and in-situ photochemical production. O_3 is destroyed in various processes, including chemical reactions with NO, the hydroperoxyl radical (HO₂) and OH, and deposition at the Earth's surface. The lifetime of tropospheric ozone varies from one or a few days in the boundary layer to a few tens of days or even a few months in the free troposphere.

In the troposphere, the mole fractions of O_3 are high in high and mid-latitudes in both hemispheres, and low in the Tropics over the Atlantic (Marenco and Said, 1989) and the Pacific (Tsutsumi *et al.*, 2003) Oceans. The localised sources of ozone precursors and generally short lifetime of surface O_3 make its distribution spatially non-uniform and time-variant.

Annual variation of surface O₃ mole fraction

The observational sites that have submitted data for surface O_3 to the WDCGG are shown on the map at the beginning of this chapter. The monthly mean mole fractions of O_3 that have been reported from these observational sites are shown in Plate 7.1, with different mole fraction levels illustrated in different colours. Data for the mole fractions of surface O_3 are reported in two different units, *i.e.*, mole fraction (ppb) and weight per volume ($\mu g/m^3$) at 25°C. The latter is converted to the former using the formula:

 $X_{p}[ppb] = (R \times T / M / P_{0}) \times 10 \times X_{g}[\mu g/m^{3}]$

where R is the molar gas constant (8.31451 [J/K/mol]), T is the absolute temperature reported from each station, M is the molecular weight of O₃ (47.9982), and P₀ is the standard pressure (1013.25 [hPa]).

If temperature is not reported in the data the units are converted using 25 °C.

The mole fraction of surface O_3 was found to vary from station to station, though many of these stations are located in Europe. Moreover, the seasonal and interannual variations were found to be relatively large at most stations, making it difficult to identify a global long-term trend in the mole fraction of surface O_3 .

The seasonal cycles of monthly mean mole fraction of surface O_3 averaged for each 30° latitudinal zone are shown in Figure 7.1. The latitudinal mean mole fractions were found to be elevated in spring in most latitudinal zones. However, several patterns of seasonal-diurnal cycles were observed at different locations, including a pronounced spring maximum, a spring maximum at night and a summer maximum during the day, a wide spring-summer maximum, and a pronounced winter maximum (Tarasova *et al.*, 2007).



Fig. 7.1 Average seasonal cycles in the mole fraction of O_3 for each 30° latitudinal zone obtained from the seasonal cycle of each station. The standard deviation from the difference of the seasonal cycle for each station within the latitudinal zone is indicated by vertical error bars.

CARBON MONOXIDE (CO)

• : CONTINUOUS STATION

- △ : FLASK STATION
- □ : FLASK MOBILE (SHIP)



This map shows locations of the stations that have submitted data for monthly mean mole fraction.

CO Monthly Data

20 40 60 80 100 120 140 160 180 200 220 ppb



Plate 8.1 Monthly mean CO mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south. The data from the sites with an asterisk at the end of the station index are used for the analysis shown in Plate 8.2. (see Chapter 2)









Plate 8.2 Variation of zonally averaged monthly mean CO mole fractions (top), deseasonalized long-term trends (middle), and growth rates (bottom). The zonally averaged mole fractions are calculated for each 20° zone. The deseasonalized trends and growth rates are derived as described in Chapter 2.

8. CARBON MONOXIDE (CO)

Basic information on CO with regard to environmental issues

Carbon monoxide (CO) is not a greenhouse gas; it absorbs hardly any infrared radiation from the Earth. However, CO influences the oxidation capacity of the atmosphere through its interaction with hydroxyl radicals (OH), which react with methane, halocarbons and tropospheric ozone. CO has been monitored due to its indirect influence on greenhouse gases through such reactions.

Sources of atmospheric CO include fossil fuel combustion and biomass burning, along with the oxidation of methane and non-methane hydrocarbons (NMHC). Major sinks include reactions with OH and surface deposition; the reaction of CO and OH accounts for all of the chemical loss of CO in the troposphere (Seinfeld and Pandis, 1998). CO has a relatively long atmospheric lifetime, ranging from 10 days in summer in the tropics to more than a year over the polar regions in winter. Thus, unlike CO₂, anthropogenic CO emissions do not lead to CO accumulation in the atmosphere. Furthermore, the uneven distribution of sources causes large spatial and temporal variations in the CO mole fraction.

Measurements of trapped air in ice cores have shown that the pre-industrial CO mole fraction over central Antarctica during the last two millennia was about 50 ppb and the CO level increased to 110 ppb by 1950 in Greenland (Haan and Raynaud, 1998). Beginning in 1950, the CO mole fraction increased at a rate of 1% per year but started to decrease in the late 1980s (WMO, 1999). Between 1991 and 2001, the global average mole fraction of CO decreased at an annual rate of about 0.5 ppb, excluding temporal enhancements from large biomass burning events (Novelli *et al.*, 2003).

Annual variation of CO mole fraction in the atmosphere

The monthly mean mole fractions of CO that have been reported from fixed stations and some ships to the WDCGG are shown in Plate 8.1, in which different mole fraction levels are plotted in different colours. The observational sites that supplied data for this analysis are shown on the map at the beginning of this chapter.

Latitudinally averaged mole fractions of CO in the atmosphere, together with their deseasonalized mole fractions and growth rates, are shown in Plate 8.2 as three-dimensional representations.

Data for the mole fractions of CO are reported in various units, *i.e.*, ppb, $\mu g/m^3$ -25°C, $\mu g/m^3$ -20°C and mg/m³-25°C. Units other than ppb were converted to ppb using the formulas:

 $\begin{array}{l} X_p \left[ppb \right] = \left(R \times T \, / \, M \, / \, P_0 \right) \times 10 \times X_g \left[\mu g / m^3 \right] \\ X_p \left[ppb \right] = \left(R \times T \, / \, M \, / \, P_0 \right) \times 10^4 \times X_g \left[mg / m^3 \right] \\ \end{array}$

where R is the molar gas constant (8.31451 [J/K/mol]), T is the absolute temperature reported from each station, M is the molecular weight of CO (28.0101), and P₀ is the standard pressure (1013.25 [hPa]).

If temperature is not reported in the data the units are converted using 25 °C.

Plate 8.2 shows that the seasonal variations of CO were larger in the Northern Hemisphere and smaller in the Southern Hemisphere, and that the deseasonalized mole fractions were the highest in northern mid-latitudes and the lowest in the Southern Hemisphere, having a large latitudinal gradient from northern mid- to southern low latitudes. This is likely due to the presence of numerous CO anthropogenic sources in the northern mid-latitudes, combined with the destruction of CO in the tropics, where OH radicals are abundant.



Fig. 8.1 Global monthly mean mole fraction of CO from 1992 to 2010, including deseasonalized long-term trend in red line (top) and annual growth rate (bottom).

Figure 8.1 shows global monthly mean CO mole fractions and their growth rates. Growth rates were high in 1993/1994, 1997/1998 and 2002, and low in 1992 and 1998/1999. The global annual mean mole fraction was about 93 ppb in 2010, which was calculated irrespective of the difference in observation scales.

Figure 8.2 shows monthly mean mole fractions of CO for each 30° latitudinal zone. Seasonal variations were observed in both hemispheres, with mole fractions being higher in winter. Amplitudes of the seasonal cycle were larger in the Northern Hemisphere than in the Southern Hemisphere.



Fig. 8.2 Monthly mean mole fractions of CO from 1992 to 2010 for each 30° latitudinal zone (dots) and their deseasonalized long-term trends (red lines).

Figure 8.3 summarizes deseasonalized long-term trends for each 30° latitudinal zone and their growth rates. The CO mole fractions were highest in northern mid-latitudes. There was a decline in CO mole fractions around 1992, almost coinciding with the decrease in the growth rate of CH₄ mole fractions, most likely due to variations in their common sinks (OH). The enhanced stratospheric ozone depletion due to increased volcanic aerosols following the eruption of Mount Pinatubo in 1991 may have increased atmospheric OH radicals, which react with both CO and CH₄ (Dlugokencky *et al.*, 1996).

Increases in CO mole fractions were observed from 1997 to 1998 in northern latitudes and in southern low latitudes. These increases were attributed to large biomass burning events in Indonesia in late 1997 and in Siberia in the summer and autumn of 1998 (Novelli *et al.*, 1998).



Fig. 8.3 Deseasonalized long-term trends of CO for each 30° latitudinal zone (top) and their growth rates (bottom).

The CO mole fractions returned to normal after 1999, but the growth rates in the Northern Hemisphere increased substantially again in 2002. The latter may have been due to large biomass burning. Large-scale boreal forest fires occurred in Siberia and North America from 2002 to 2003. Strong forest fires also occurred in Russia in summer 2010 which can be seen in the bottom panel of Figure 8.3.



Fig. 8.4 Average seasonal cycles in the mole fraction of CO for each 30° latitudinal zone obtained by subtracting long-term trends from the zonal mean time series. The standard deviation is indicated by vertical error bars.

Seasonal cycle of CO mole fraction in the atmosphere

Figure 8.4 shows average seasonal cycles in the mole fraction of CO for each 30° latitudinal zone. The seasonal cycle was driven mainly by seasonal variations in OH abundance as a CO sink. Additional factors include emission and oxidation from CO sources and large-scale transport of CO, despite a relatively weak seasonality in emission and oxidation compared with OH abundance. This seasonality and a short lifetime of about a few months resulted in a sharp decrease in early summer followed by a relatively slow increase in autumn. The levelling-off in the beginning of the year seen in the southern low latitudes may be attributed to the transport of CO from the Northern Hemisphere.

NITROGEN MONOXIDE (NO) AND NITROGEN DIOXIDE (NO₂)



This map shows locations of the stations that have submitted data for monthly mean mole fraction.



Plate 9.1 Monthly mean (a) NO and (b) NO_2 mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south.

9. NITROGEN MONOXIDE (NO) AND NITROGEN DIOXIDE (NO₂)

Basic information on NO and NO₂ with regard to environmental issues

Nitrogen oxides (NO_x, *i.e.*, NO and NO₂) are not greenhouse gases. Nevertheless, these compounds have a central regulatory role in the free radical and oxidising photochemistry of the troposphere. This photochemistry regulates the lifetime of methane and the production of tropospheric O_3 and secondary aerosols, all of which have important roles in the natural and anthropogenic greenhouse effect. The O_3 produced in the atmosphere as a result of the nitrogen oxides presence can affect vegetation growth and human health.

Sources of NO_x include energy production, transport, lightning, soils and biomass burning (Reis et al., 2009). They constitute major causes of acid rain and deposition. The dominant sink of NO_x in the atmosphere is its conversion into nitric acid (HNO₃) and peroxyacetylnitrate (PAN), which are eventually removed by dry or wet deposition. In some cases, NO_x is removed from the atmosphere directly by dry deposition. NO_x abundance varies in both space and time because of their short lifetimes and uneven source distribution. Some regional assessments are done based on satellite information to clarify such variations and trends.

Annual variation of NO and NO₂ mole fractions in the atmosphere

The observation stations that have submitted data for NO and NO_2 to the WDCGG are shown on the map at the beginning of this chapter. Most of these stations are located in Europe.

The monthly mean mole fractions of NO and NO₂ reported to the WDCGG are shown in Plate 9.1, in which different mole fraction levels are plotted in different colours. Data for NO_x are reported in various units, *i.e.*, ppb, $\mu g/m^3$ -25°C, $\mu g/m^3$ -20°C, $\mu gN/m^3$ -25°C and mg/m^3 -25°C. Units other than ppb were converted to ppb using the formulas:

- $$\begin{split} X_p \left[ppb \right] &= (R \times T / M / P_0) \times 10 \times X_g \left[\mu g / m^3 \right] \\ X_p \left[ppb \right] &= (R \times T / M / P_0) \times 10^4 \times X_g \left[mg / m^3 \right] \\ X_p \left[ppb \right] &= (R \times T / M_N / P_0) \times 10 \times X_g \left[\mu g N / m^3 \right] \end{split}$$
- where R is the molar gas constant (8.31451 [J/K/mol]),

T is the absolute temperature reported from each station,

M is the molecular weight of NO (30.00614) or NO₂ (46.00554),

 $M_{\rm N}$ is the atomic weight of N (14.00674), and

P₀ is the standard pressure (1013.25 [hPa]).

If temperature is not reported in the data the units are converted using 25 °C.

The distributions of NO and NO₂ are spatially non-uniform and variable over time. Due to the high temporal variability in the mole fraction of NO₂ at each observation site, it was difficult to identify a long-term trend. A number of stations located in southern Europe showed higher mole fractions, and some stations reported increased NO₂ in winter.

As there are few observational sites for NO, it was difficult to identify whether the global average NO mole fraction increases or decreases.

10. SULPHUR DIOXIDE (SO₂)



This map shows locations of the stations that have submitted data for monthly mean mole fraction.



Plate 10.1 Monthly mean SO_2 mole fractions that have been reported to the WDCGG. The mole fractions are illustrated in different colors. The sites are listed in order from north to south.

10. SULPHUR DIOXIDE (SO₂)

Basic information on SO₂ with regard to environmental issues

Sulphur dioxide (SO_2) is not a greenhouse gas, but it is a precursor of atmospheric sulphuric acid (H_2SO_4) and sulphate aerosol. SO_2 is oxidised by hydroxyl radicals (OH) to form sulphuric acid, which then becomes aerosols through photochemical gas-to-particle conversion. While SO_2 reacts much more slowly with OH than does NO_2 , SO_2 dissolves readily in suspended liquid droplets in the atmosphere. The global sulphur cycle affects atmospheric chemistry, including tropospheric ozone (Berglen *et al.*, 2004).

Sources of SO_2 include fossil fuel combustion by industry, biomass burning, volcanic release and the oxidation of dimethylsulphide (DMS) from the oceans (IPCC, 2007). Major SO_2 sinks are oxidation by OH and deposition onto wet surfaces. Anthropogenic SO_2 has caused acid rain and deposition throughout the industrial era. The mole fractions of SO_2 have shown large variations in both space and time because of the short lifetime and uneven anthropogenic source distribution of SO_2 .

Annual variation of SO₂ mole fraction in the atmosphere

The observational sites that have submitted data for SO_2 to the WDCGG are shown on the map at the beginning of this chapter. Most of these stations are located in Europe.

The monthly mean mole fractions of SO₂ that have been reported to the WDCGG are shown in Plate 10.1, with different mole fraction levels illustrated in different colours. Data for SO₂ are reported in various units, *i.e.*, ppb, μ g/m³, mg/m³ and μ gS/m³. Units other than ppb were converted to ppb using the formulas:

$$\begin{split} X_p \left[ppb \right] &= (R \times T \, / \, M \, / \, P_0) \times 10 \times X_g \left[\mu g / m^3 \right] \\ X_p \left[ppb \right] &= (R \times T \, / \, M \, / \, P_0) \times 10^4 \times X_g \left[mg / m^3 \right] \\ X_p \left[ppb \right] &= (R \times T \, / \, M_S \, / \, P_0) \times 10 \times X_g \left[\mu g S / m^3 \right] \end{split}$$

where R is the molar gas constant (8.31451 [J/K/mol]),

T is the absolute temperature reported from each station,

M is the molecular weight of SO_2 (64.0648), M_S is the atomic weight of S (32.066), and P_0 is the standard pressure (1013.25 [hPa]).

Although some stations in southern Europe have reported higher mole fractions, it has been difficult to identify an increasing or decreasing trend for SO_2 .

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APPENDICES

CALIBRATION AND STANDARD SCALES

1. Calibration System in the GAW programme

Under the Global Atmosphere Watch (GAW) programme, the World Calibration Centres (WCCs) are responsible for maintaining calibration standards for certain compounds, establishing instrument calibrations and providing training to the stations. A Reference Standard is designated for each variable to be used for all GAW measurements of that variable at the Central Calibration Laboratories. Table 1 lists the organizations that serve as WCCs and CCLs for GAW [*WMO*, 2007]. For CFCs and SO₂, no central facilities or quality control systems have so far been established within the GAW program, while central facilities for NOx have only recently been formulated.

Table 1. Overview of the GAW Central Calibration Laboratories (GAW-CCL, Reference Standard) and World Calibration Centres for greenhouse and other related gases. The World Calibration Centres have assumed global responsibilities, except where indicated (Am, Americas; E/A, Europe and Africa; A/O, Asia and the South-West Pacific)

Compounds	Central Calibration Laboratory (Host of Primary Standard)	World Calibration Centre
Carbon Dioxide (CO ₂)	NOAA/ESRL	NOAA/ESRL (round robin) Empa (audits)
carbon isotopes	MPI-BGC	
Methane (CH ₄)	NOAA/ESRL	Empa (Am, E/A) JMA (A/O)
Nitrous Oxide (N ₂ O)	NOAA/ESRL	IMK-IFU
Chlorofluorocarbons (CFCs)		
Sulfur Hexafluoride (SF ₆)	ESRL	KRISS&KMA
Molecular Hydrogen (H ₂)	MPI-BGC	
Surface Ozone (O ₃)	NIST	Empa
Carbon Monoxide (CO)	NOAA/ESRL	Empa
Volatile Organic Compounds (VOCs)	NPL (8 components)	IMK-IFU
Sulphur Dioxide (SO ₂)		
Nitrogen Oxides (NO _x)		IEK-8

2. Carbon Dioxide (CO₂)

In 1995, the National Oceanic and Atmospheric Administration's Earth System Research Laboratory (NOAA/ESRL, formerly CMDL; Climate Monitoring and Diagnostics Laboratory) in Boulder, Colorado, USA, took over the role of the Central Calibration Laboratory (CCL) from the Scripps Institution of Oceanography (SIO) in San Diego, California, USA. Since then, NOAA/ESRL has served as the CCL responsible for the maintenance of the GAW Primary Standard for CO₂. As the World Calibration Centre (WCC) for CO₂, NOAA/ESRL maintains a high-precision manometric system for absolute calibration of CO₂ as the reference for GAW measurements throughout the world [*Zhao et al.*, 1997]. It has been recommended that the standards of the GAW measurement laboratories be calibrated every two years at the CCL (WMO, 2003).

Under the WMO calibration system, there have been several calibration scales for CO₂, *e.g.*, SIO-based X74, X85, X87, X93 and X2002 scales and the

NOAA/ESRL-based WMO Mole Fraction Scale partially based on previous SIO scales. The NOAA/ESRL and SIO are working to resolve the possible small differences between their scales. The CCL adopted the WMO X2005 scale, reflecting historical manometric calibrations of the CCL's set of cylinders and the possible small differences between SIO and NOAA/ESRL calibrations. The most current WMO Mole Fraction Scale is the WMO-X2007 scale.

To assess the differences in standard scales among measuring laboratories, NOAA/ESRL organizes intercomparisons or Round Robin experiments endorsed by WMO every few years. Many laboratories participated in the experiments organized in 1991–1992, 1995–1997, 1999–2000, and 2002–2006. Table 2 shows the results of the experiments performed in 2002–2006, in which the mole fractions measured by various laboratories are compared with the mole fractions measured by NOAA/ESRL [*Zhou et al.*, 2009]. In addition, many laboratories compare their standards bilaterally or multilaterally.

Table 3 lists laboratories and sites used in the present issue of the *Data Summary* with standard scales of reported data and history of participation in WMO intercomparison experiments.

Table 2. Round robin results for the mole fraction of carbon dioxide. Differences between the mole fractions measured by various laboratories and the mole fractions measured by NOAA (Laboratory minus NOAA, ppm).

Laboratory	Analysis Date	Mole fraction Difference (ppm)		
		Low	Medium	High
		340–350 ppm	350–360 ppm	370–380 ppm
Tohoku Univ.	Jan-03	-0.11	-0.19	-0.29
NIES	Apr-03	-0.10	-0.15	-0.14
MRI	Jul-03	-0.16	-0.16	-0.08
AIST	Sep/Dec-03	-0.11	-0.22	-0.29
JMA	Jan-04	0.13	0.00	-0.02
KMA	Mar/Jun-04	-0.44	-0.12	-0.08
CMA (WLG)	Jul-04	-0.05	-0.19	-0.10
CMA (BJ)	Aug-04	-0.03	-0.20	0.02
Scripps (CMM)	Jun-05	0.23	0.17	0.20
Scripps (ECM II)		0.10	0.02	0.02
LSCE	Oct/Nov-05	-0.05	-0.11	-0.09
Monte Cimone	Oct-02	0.08	0.02	-0.03
Lampedusa	Nov-02	0.05	-0.15	-0.26
Plateau Rosa	Dec-02	-0.02	0.00	-0.05
HMS	Feb-03	0.06	-0.21	-0.06
EC	May-05	0.06	-0.05	-0.06
Penn State Univ.	Sep-05	0.09	-0.07	-0.05
Univ. Heidelberg	Sep/Oct-02	-0.01	-0.06	-0.06
UBA	Oct-02	0.05	-0.11	-0.21
LSCE	Nov/Dec-02	0.10	0.03	0.05
FMI	Jan-03	-0.02	-0.04	-0.14
Univ. Groningen	Oct/Nov-03	0.01	0.02	0.04
MPI-BGC	Nov/Dec-03	0.04	0.02	-0.02
HMS	Mar-04	-0.19	-0.36	-0.59
NIWA	May-05	-0.08	-0.08	-0.09
CSIRO	Sep/Oct-05	-0.01	-0.03	-0.08
Cape Point	Dec-05	-0.02	-0.09	-0.18
NCAR	May/Jun-06	0.07	-0.04	-0.04

Laboratory	WDCGG Filename Code	Calibration Scale	WMO Inter- comparison
AEMET	IZO128N0000	WMO	91/92, 96/97, 99/00
Aichi	MKW234N0000	WMO	
AIST	TKY236N0000	AIST	96/97, 99/00, 02/06
BoM & CSIRO	CGO540S0000,CGO540S0010	WMO	
СМА	WLG236N0000	WMO	96/97, 99/00, 02/06
CNR-ICES & DNA-IAA	JBN762S0000	WMO	
CSIRO	ALT482N0003, CFA519S0003, CGO540S0003, CRI215N0000, CYA766S0001, ESP449N0003, MAA767S0003, MLO519N0003, MQA554S0003, SIS660N0003, SPO789S0003	WMO	91/92, 96/97, 99/00, 02/06
EC	ALT482N0000, ALT482N0005, CDL453N0000, CHM449N0000, CSJ451N0000, EGB444N0100 , ESP449N0000, ETL454N0000 , FSD449N0000, LLB454N0100, WSA443N0000, WSA443N0001	WMO	91/92, 96/97, 99/00, 02/06
EMA	CAI130N0000		
Empa	JFJ646N0000	WMO	
ENEA	LMP635N0001	WMO	91/92, 96/97, 99/00, 02/06
FMI	PAL667N0000	WMO	02/06
НКО	HKG222N0001, HKO222N0001	WMO	
ПКО	HKO222N0000	NIST	
HMS	HUN646N0000, KPS646N0000	WMO	91/92, 96/97, 99/00, 02/06
IAFMS	CMN644N0000	WMO	91/92, 96/97, 02/06
IGP	HUA312S0000	WMO	
IMK-IFU	WNK647N0000, ZUG647N0014	WMO	99/00
INRNE	BEO642N0000	WMO	
IOEP	DIG654N0000		
ITM	ZEP678N0000	WMO	96/97, 99/00
JMA	MNM224N0000, RYO239N0000, YON224N0000	WMO	91/92, 96/97, 99/00, 02/06
KMA	AMY236N0000	KRISS	02/06
KSNU	ISK242N0000		
KUP	JFJ646N0003	WMO	
LSCE	AMS137S0000, BGU641N0000, FIK635N0000, LPO648N0000, MHD653N0002, PDM642N0000, PUY645N0000	WMO	91/92, 96/97, 99/00, 02/06
MGO	BER255N0001, KOT276N0001, KYZ240N0001, STC652N0001, TER669N0001	WMO	
MMD	DMV504N0000	WMO	
MRI	TKB236N0002		91/92, 96/97, 99/00, 02/06

Table 3. Status of standard scales and calibration/intercomparison for CO₂ at laboratories.

NIER	GSN233N0103	WMO	
NIES	COI243N0000, HAT224N0000	NIES	96/97, 99/00, 02/06
NIMR	GSN233N0001	WMO	96/97
NIPR & Tohoku Univ.	SYO769S0000		Tohoku Univ.:91/92, 96/97, 99/00, 02/06
NIWA	BHD541S0000	WMO	91/92, 96/97, 99/00, 02/06
NMA	FDT645N0002		
NOAA/ESRL	BRW471N0000, MLO519N0000, SMO514S0000, SPO789S0000, NOAA/ESRL flask network [*]	WMO	91/92, 96/97, 99/00, 02/06
Osaka Univ.	SUI234N0000		
RIVM	KMW653N0000	NIST	
RSE	PRS645N0000	WMO	99/00, 02/06
Saitama	DDR236N0000, KIS236N0000, URW235N0000	WMO	
SAWS	CPT134S0000	WMO	99/00, 02/06
Shizuoka Univ.	HMM234N0000		
UBA	BRT648N0000, DEU649N0000, LGB652N0000, NGL653N0000 SNB647N0000, SSL647N0000, SSL647N0002, WES654N0000, ZGT654N0000, ZSF647N0010, ZUG647N0000	WMO	91/92, 96/97, 99/00, 02/06

NOAA/ESRL flask network:

ABP312S0001,ALT482N0001,AMS137S0001,ASC107S0001,ASK123N0001,AVI417N0001,AZR638N0001,BAL655N0001,BHD541S0001, BKT500S0001,BME432N0001,BMW432N0001,BRW471N0001,BSC644N0001,CBA455N0001,CGO540S0001,CHR501N0001,CMO445N0001, CRZ146S0001,EIC327S0001,GMI513N0001,GOZ636N0001,HBA775S0001,HPB647N0003,HUN646N0001,ICE663N0001,ITN435N0001, IZO128N0001,KCO204N0001,KEY425N0001,KUM519N0001,KZD244N0001,KZM243N0001,LEF445N0001,LLB454N0001,LLN223N0001, LMP635N0003,MBC476N0001,MEX419N0001,MHD653N0001,MID528N0001,MKN100S0001,MLO519N0001,NMB123S0001,NWR440N0101, OPW448N0001,PAL667N0001,POC900N0001,POC905S0001,POC910N0001,POC910S0001,POC915N0001,POC915S0001, POC920N0001,POC920S0001,POC925S0001,POC925S0001,POC930S0001,POC935S0001,PSA764S0001,PTA438N0001, RPB413N0001,SCS903N0001,SCS906N0001,SCS912N0001,SCS915N0001,SCS918N0001,SCS921N0001,SCS921N0001,SCY104S0001, TDF354S0001,THD441N0001,UTA439N0001,UUM244N0001,WIS631N0001,WLG236N0001,ZEP678N0001

3. Methane (CH₄)

The GAW programmes have established two WCCs for CH₄, the Swiss Federal Laboratory for Materials Testing and Research (Empa), Dübendorf, Switzerland; and the Japan Meteorological Agency (JMA), Tokyo, Japan [*WMO*, 2007]. In addition, the Central Calibration Laboratory for CH₄ has been established at NOAA/ESRL [*Dlugokencky, et. al.*, 2005; *WMO*, 2007].

The NOAA04 scale has been designated as the Primary Standard of the GAW programme. This scale results in CH_4 mole fractions that are a factor of 1.0124 higher than the previous NOAA scale [*Dlugokencky et al.*, 2005].

Table 4 summarises the methane standard scales used by laboratories contributing to the WDCGG and lists tentative multiplying conversion factors applied for analysis in this issue of the *Data Summary*. The standard is the NOAA04 scale, and conversion factors were calculated from the results of comparisons with other laboratories performed bilaterally or multilaterally before the establishment of the GAW Standard.

The former CMDL scale is lower than an absolute gravimetric scale [*Aoki et al.*, 1992] by ~1.5% [*Dlugokencky et al.*, 1994] and lower than the AES (Atmospheric Environment Service, currently EC) scale by a factor of 1.0151 [Worthy et al., 1998]. The former CMDL scale can be converted to the Tohoku University standard by multiplying by 1.0121 [*Dlugokencky et al.*, 2005]. The conversion factors 1.0124 / 1.0151 = 0.9973 and 1.0124 / 1.0121 = 1.0003 have been adopted for comparisons with the NOAA04 scale.

Laboratory	aboratory WDCGG Filename Code		Conversion Factor	
AEMET	IZO128N0000	NOAA04	1	
AGAGE	CGO540S0011, CGO540S0013, CMO445N0011, MHD653N0011, MHD653N0013, RPB413N0000, RPB413N0011, SMO514S0014, SMO514S0016, THD441N0000	Tohoku Univ.	1.0003	
CHMI	KOS649N0000	CHMI	0.9973	
CMA	WLG236N0000	NOAA04	1	
CSIRO	ALT482N0003, CFA519S0003, CGO540S0003, CRI215N0000, CYA766S0001, ESP449N0003, MAA767S0003, MLO519N0003, MQA554S0003, SIS660N0003,SPO789S0003	NOAA04	1	
EC	ALT482N0000, CDL453N0000, CHM449N0000, EGB444N0100, ESP449N0000, ETL454N0000, FSD449N0000, LLB454N0100, WSA443N0000	NOAA04	1	
Empa	JFJ646N0000	NOAA04	1	
ENEA	LMP635N0001	NOAA04	1	
FMI	PAL667N0000	NOAA04		
ISAC	CMN644N0000	NOAA04	1	
JMA	MNM224N0000, RYO239N0000, YON224N0000	NOAA04	1	
KMA	AMY236N0000	KRISS		
KSNU	ISK242N0000			
LSCE	AMS137S0002, BGU641N0000, LPO648N0000, PDM642N0000, PUY645N0001	NOAA83	1.0124	
	FIK635N0000, MHD653N0007			
MGO	TER669N0001	NOAA04	1	
MRI	TKB236N0000		0.9973	
NIER	GSN233N0103	NOAA04	1	
NIES	COI243N0000, HAT224N0000	NIES	0.9973	
NIMR	GSN233N0001	SIO X97		
	BRW471N0000, MLO519N0000, NOAA/ESRL flask network*	NOAA04	1	
NOAA/ESRL	KPA431N0001, LEF445N0001, MCM777S0001, NZL543S0001, POC935S0001, SGI354S0001, SIO432N0001	NOAA/CMDL	1.0124	
RIVM	KMW653N0000	NIST	0.9973	
RSE	PRS645N0000	NOAA04	1	
SAWS	CPT134S0000	NOAA04	1	
UBA	DEU649N0000, NGL653N0000, SSL647N0000, ZGT654N0000, ZSF647N0010, ZUG647N0000	NOAA04	1	

Table 4. Status of the standard scales of CH₄ at laboratories with conversion factors.

^{*}NOAA/ESRL flask network:

ABP312S0001, ALT482N0001, AMS137S0001, ASC107S0001, ASK123N0001, AVI417N0001, AZR638N0001, BAL655N0001, BKT500S0001, BME432N0001, BMW432N0001, BRW471N0001, BSC644N0001, CBA455N0001, CGO540S0001, CHR501N0001, CMO445N0001, CRZ146S0001, EIC327S0001, GMI513N0001, GOZ636N0001, HBA775S0001, HPB647N0003, HUN646N0001, ICE663N0001, ITN435N0001, IZO128N0001, KEY425N0001, KUM519N0001, KZD244N0001, KZM243N0001, LLB454N0001, LLN223N0001, LMP635N0003, MBC476N0001, MEX419N0001, MHD653N0001, MID528N0001, MKN100S0001, MLO519N0001, NMB123S0001, NWR440N0101, OPW448N0001, OXK650N0001, PAL667N0001, POC900N0001, POC905N0001, POC905S0001, POC910S0001, POC915N0001, POC915S0001, POC920N0001, POC920S0001, POC925N0001, POC925S0001, POC930N0001, POC930S0001, PSA764S0001, PTA438N0001, RPB413N0001, SCS903N0001, SCS906N0001, SCS909N0001, SCS912N0001, SCS915N0001, SCS918N0001, SCS921N0001, SEY104S0001, SGP436N0001, SHM452N0001, SMO514S0001, SPO789S0001, STM666N0001, SUM672N0001, ZSP678N0001

4. Nitrous Oxide (N₂O)

The Halocarbons and other Atmospheric Trace Species (HATS) Group of NOAA/ESRL maintains a set of standards for N₂O [*Hall et al.*, 2001]. The NOAA-2006 N₂O scale [*Hall et al.*, 2007] has been designated as the Primary Standard of the GAW programme. This group analyses the standards of laboratories, including those of Environment Canada (EC) and the Australian Commonwealth Scientific and Industrial Research Organisation (CSIRO). The Fraunhofer Institut für Atmosphärische Umweltforschung (IFU) in Garmisch-Partenkirchen, Germany, serves as the GAW WCC.

The SIO-98 scale is essentially equivalent to the NOAA-2006 scale, with an average difference of 0.01% over the range of 299–319 ppb; the NOAA-2000 scale can be converted to the 2006 scale by using the factor 0.99402 [*Hall et al.*, 2007]. A constant ratio of 1.0017 between CSIRO and AGAGE data was used by Huang *et al.* (2008), and a factor of 1 / 1.0017 = 0.9983 has been used in this report to convert CSIRO scale to the NOAA-2006 scale.

Table 5. Status of the standard scales of N_2O at laboratories.

Laboratory	WDCGG Filename Code	Calibration Scale	Conversion Factor
AEMET	IZO128N0000	NOAA-2006	1
AGAGE	ADR651N0010, CGO540S0011, CGO540S0012, CGO540S0013, CMO445N0010, CMO445N0011, MHD653N0011, MHD653N0013, RPB413N0000, RPB413N0010, RPB413N0011, SMO514S0014, SMO514S0015, SMO514S0016, THD441N0000	SIO 1998	1
CSIRO	ALT482N0003, CFA519S0003, CGO540S0003, CRI215N0000, CYA766S0001, ESP449N0003, MAA767S0003, MLO519N0003, MQA554S0003, SIS660N0003, SPO789S0003	NOAA-2006	1
Empa	JFJ646N0000	SIO 1998	1
ENEA	LMP635N0001	NOAA-2006	1
ISAC	CMN644N0000	NOAA-2006	1
JMA	RYO239N0000	NOAA-2006	1
KMA	AMY236N0000	KRISS	
MRI	MMB243N0000		
Nagoya Univ.	NGY235N0000		
NIER	GSN233N0103	NOAA-2006	1
NIES	HAT224N0000		
NILU	ZEP678N0000		
NIMR	GSN233N0001	WMO X97	
NOAA/ESRL	ALT482N0001, BRW471N0001, BRW471N0011, CGO540S0001, KUM519N0001, MLO519N0001, MLO519N0011, NWR440N0001, NWR440N0011, SMO514S0001, SMO514S0011, SPO789S0001, SPO789S0011	NOAA/CMDL	0.999402

NOAA/ESRL	ALT482N0004, ALT482N0006, BRW471N0003, BRW471N0005, BRW471N0010, CGO540S0009, CGO540S0014, KUM519N0002, MHD653N0008, MLO519N0005, MLO519N0006, MLO519N0010, NWR440N0003, NWR440N0004,NWR440N0010, PSA764S0000, SMO514S0008, SMO514S0009, SMO514S0010, SPO789S0005, SPO789S0006, SPO789S0010, SUM672N0000, SUM672N0002, THD441N0002	NOAA-2006	1
SAWS	CPT134S0000	NOAA/CMDL	0.999402
UBA	SSL647N0000, ZSF647N0010	SIO 1998	1

5. Surface Ozone (O₃)

The National Institute of Standards and Technology (NIST) has developed and deployed Standard Reference Photometers (SRPs) in the USA and other countries. The GAW has designated SRP #2 maintained by NIST as the Primary Standard for the GAW programme, making NIST the CCL for O_3 . The Swiss Federal Laboratory for Materials Testing and Research (Empa) maintains NIST SRP #15 as the reference and is the GAW WCC for surface ozone

[*Hoferetal.*, 1998]. The traceability and uncertainty of O_3 within the GAW network were reported by Klausen *et al.*, (2003). Regional Calibration Centres have been established at the Solar and Ozone Observatory, Hradec Kralove, Czech Republic, and Observatorio Central Buenos Aires, Argentina [*WMO*, 2007]. The former maintains SRP #17 directly purchased from NIST.

Table 6. Status	of surface ozone s	tandard scales a	t laboratories

Laboratory	WDCGG Filename Code	Calibration Scale	Audit Empa-WCC
	IZO128N0000	WMO (NIST & Empa)	96, 98, 00, 04
AEMET	DON637N0000,MHN639N0000,NIA642N0000, ROQ640N0000,SPM639N0000	NPL (U. K.)	
AQRB	ALG447N0000, BRA450N0000, CHA446N0000, EGB444N0000, ELA449N0000, EST451N0000, KEJ444N0000, SAT448N0000		
AWI	NMY770S0000		
BMKG & Empa	BKT500S0000	WMO (NIST & Empa)	99,01,04,07,08
BoM & CSIRO	CGO540S0000	WMO (NIST & Empa)	02
CHMI	KOS649N0000	WMO (NIST & Empa)	
DEFRA	EDM655N0000		
DWD	HPB647N0000	WMO (NIST & Empa)	97,06
EARS	IRB645N0000, KVK646N0000, KVV646N0000, ZRN646N0000	WMO (NIST & Empa)	
EMA	CAI130N0000		

Empa	JFJ646N0000, PAY646N0000, RIG646N0000	WMO (NIST & Empa)	Jungfraujoch: 99, 06
Empa & KMD	MKN100S0000	WMO (NIST & Empa)	
FMI	AHT662N0000, OUL666N0000, PAL667N0000, UTO659N0000, VIR660N0000		Pallas-Sammaltunturi: 97, 03, 07
HMS	KPS646N0000	WMO (NIST & Empa)	
IM	ANG638N0000, BEJ638N0000, CAS639N0000, FUN132N0000, LIS638N0000, MVH638N0000, PEN640N0000		
INRNE	BEO642N0000	WMO (NIST & Empa)	
IOEP	DIG654N0000	WMO (NIST & Empa)	
ISAC	CMN644N0000, PYR227N0000	WMO (NIST & Empa)	
IVL	VDL664N0000	Stockholm Univ. (Sweden)	
JMA	MNM224N0000, RYO239N0000, SYO769S0002, TKB236N1004, YON224N0000	WMO (NIST & Empa)	Ryori: 05
KSNU	ISK242N0000		
LEGMA	DBL656N0000, RCV656N0000, ZSN657N0000	WMO (NIST & Empa)	
MMD	DMV504N0000, TAR504N0000		
NILU	ZEP678N0000	WMO (NIST & Empa)	97, 01, 05
NIWA	BHD541S0000	WMO (NIST & Empa)	
NMA	FDT645N0002		
NOAA/ESRL	ARH777S0000, BMW432N0004, BRW471N0004, ICE663N0004, LAU545S0004, MCM777S0004, MLO519N0004, NWR440N0002, NWR440N0204, RPB413N0004, SMO514S0004, SPO789S0004, SUM672N0004, THD441N0004	WMO (NIST & Empa)	Mauna Loa: 03 Barrow: 08
NUI	MHD653N0000	WMO (NIST & Empa)	96, 98, 02, 05
ONM	ASK123N0000	WMO (NIST & Empa)	03, 07
PolyU	HKG222N0000		
RIVM	KMW653N0001,KMW653N0002		
Roshydromet	DAK654N0000, SHP659N0000		
RSE	PRS645N0000		
SAWS	CPT134S0000	WMO (NIST & Empa)	97, 98, 02, 06

SMN	USH354S0000,USH354S0001	WMO (NIST & Empa)	98, 03
UBA	BRT648N0000, DEU649N0000, LGB652N0000, NGL653N0000, SNB647N0000, SSL647N0000, WES654N0000, ZGT654N0000, ZSF647N0010, ZUG647N0000	WMO (NIST & Empa)	Zugspitze: 96, 97, 01 Sonnblick: 98 Zugspitze/Schneefern erhaus: 06
Univ. Malta	GLH636N0000	UMEG	
Univ. York	CVO116N0001	NPL	

6. Carbon Monoxide (CO)

The Swiss Federal Laboratory for Materials Testing and Research (Empa) serves as the WCC under GAW based on its secondary standards calibrated against the

standard at NOAA/ESRL designated as the Primary Standard for GAW.

Laboratory	WDCGG Filename Code	Calibration Scale	Audit Empa-WCC
AEMET	IZO128N0000	WMO 2004 (NOAA/ESRL & Empa)	
AGAGE	CGO540S0011, MHD653N0011	CSIRO	
BMKG & Empa	BKT500S0000	WMO 2000 (NOAA/ESRL & Empa)	
CHMI	KOS649N0000	CHMI	
CSIRO	ALT482N0003, CFA519S0003, CGO540S0003, CRI215N0000, CYA766S0001, ESP449N0003, MAA767S0003, MLO519N0003, MQA554S0003, SIS660N0003, SPO789S0003	CSIRO	Cape Grim: 02
DWD	HPB647N0000	WMO (NOAA/ESRL & Empa)	06
EARS	KVV646N0000	CHMI	
EC	ALT482N0000,CDL453N0000, CHM449N0000, EGB444N0100, ESP449N0000, ETL454N0000, FSD449N0000, LLB454N0100, WSA443N0000	WMO (NOAA/ESRL & Empa)	
Empa	JFJ646N0000, PAY646N0000, RIG646N0000	WMO 2000 (NOAA/ESRL & Empa)	Jungfraujoch: 99,06
Empa &KMD	MKN100S0000	WMO 2000 (NOAA/ESRL & Empa)	
INRNE	BEO642N0000	WMO (NOAA/ESRL & Empa)	
ISAC	CMN644N0000, CMN644N0001	WMO 2004 (NOAA/ESRL & Empa)	
	MNM224N0001,RYO239N0001, YON224N0001	JMA	
JMA	MNM224N0002, RYO239N0002, YON224N0002	WMO 2000 (NOAA/ESRL)	Ryori:05
LSCE	AMS137S0000	WMO 2004 (NOAA/ESRL & Empa)	

Table 7. Status of carbon monoxide standard scales at laboratories

NOAA/ESRL	NOAA/ESRL flask network*	WMO (NOAA/ESRL & Empa)	Mauna Loa: 03 Barrow: 08
PolyU	HKG222N0000		
RIVM	KMW653N0000, KTB653N0000		
SAWS	CPT134S0000	WMO (NOAA/CMDL)	98, 02, 06
SMN	USH354S0000, USH354S0001	WMO (NOAA/ESRL & Empa)	98, 03
UBA	NGL653N0000, SNB647N0000, SSL647N0000, ZUG647N0000	WMO (NOAA/CMDL)	Zugspitze: 01 Sonnblick: 98
Univ. Malta	GLH636N0000		
Univ. York	CVO116N0001	WMO 2000 (NOAA/ESRL & Empa)	

NOAA/ESRL flask network:

ALT482N0001,ASC107S0001,ASK123N0001,AZR638N0001,BAL655N0001,BHD541S0001,BKT500S0001,BME432N0001,BMW432N0001, BRW471N0001,BSC644N0001,CBA455N0001,CGO540S0001,CHR501N0001,CMO445N0001,CRZ146S0001,EIC327S0001,GMI513N0001, GOZ636N0001,HBA775S0001,HPB647N0003,HUN646N0001,ICE663N0001,ITN435N0001,IZO128N0001,KEY425N0001,KUM519N0001, KZD244N0001,KZM243N0001,LEF445N0001,LLN223N0001,LMP635N0003,MBC476N0001,MHD653N0001,MID528N0001,MLO519N0001, NMB123S0001,NWR440N0101,OXK650N0001,PAL667N0001,POC900N0001,POC905N0001,POC905S0001,POC915N0001,POC915S0001,POC920S0001,POC925S0001,POC925S0001,POC935S0001,POC935S0001,POC935S0001,POC920S0001,POC925S0001,SCS903N0001,SCS905N0001,SCS905N0001,SCS912N0001,SCS912N0001,SCS915N0001, SCS918N0001,SCS921N0001,SEY104S0001,SIM452N0001,SIM452N0001,SW6514S0001,SPO789S0001,TME666N0001,SY0769S0001, TAP236N0001,TDF354S0001,THD441N0001,UTA439N0001,UUM244N0001,WIS631N0001,WLG236N0001,ZEP678N0001

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LIST OF ABBREVIATIONS IN THE CALIBLATION AND STANDARD SCALES

AEMET	Agancia Estatal da Matacrología (Spain)		
AGAGE	Agencia Estatal de Meteorología (Spain) Advanced Global Atmospheric Gases Experiment		
Aichi	Advanced Global Atmospheric Gases Experiment		
AIST	Aichi Prefecture (Japan) National Institute of Advanced Industrial Science and Technology (Japan)		
AQRB	Air Quality Research Branch, Meteorological Service of Canada (Canada)		
AWI			
BMKG	Alfred Wegener Institute for Polar and Marine Research (Germany)		
BoM	Agency for Meteorology, Climatology and Geophysics (Indonesia)		
СНМІ	Commonwealth Bureau of Meteorology (Australia)		
	Czech Hydrometeorological Institute (Czech Republic)		
CMA CND LCES	China Meteorological Administration (China)		
CNR-ICES	International Center for Earth Sciences, Consiglio Nazionale delle Ricerche (Italy)		
CSIRO DEEDA	Commonwealth Scientific and Industrial Research Organisation (Australia)		
DEFRA	Department for Environment, Food and Rural Affairs (United Kingdom)		
DNA-IAA	Direccion Nacional del Antartico-Instituto Antartico Argentino (Argentina)		
DWD	Deutscher Wetterdienst (German Meteorological Service, Germany)		
EARS	Environmental Agency of the Republic of Slovenia		
EC EMA	Environment Canada (Canada)		
	Egyptian Meteorological Authority (Egypt)		
Empa ENFA	Swiss Federal Laboratories for Material Testing and Research (Switzerland)		
ENEA	Italian National Agency for New Technology, Energy and the Environment		
FMI	(Italy) Finnish Mataorological Institute		
GAGE	Finnish Meteorological Institute		
GAGE GAW	Global Atmospheric Gases Experiment		
HATS	Global Atmosphere Watch (WMO)		
НКО	Halocarbons and other Atmospheric Trace Species Group, NOAA/ESRL		
HMS	Hong Kong Observatory (Hong Kong, China)		
IAFMS	Hungarian Meteorological Service (Hungary)		
IEK-8	Italian Air Force Meteorological Service (Italy) Institute for Energy and Climate Research: Troposphere (IEK-8), Research		
11/18-0	Center Juelich GmbH (Germany)		
IGP	Instituto Geofísico del Perú (Peru)		
IM	Instituto Geonsico del Peru (Peru) Instituto de Meteorologia (Portugal)		
IMI IMK-IFU	Instituto de Meteorologia (Fortugar) Institut für Meteorologie und Klimatologie, Atmosphärische		
	Umweltforschung, Forschungszentrum Karlsruhe (Germany)		
INMH	National Meteorological Administration (Romania)		
INRNE	Institute for Nuclear Research and Nuclear Energy (Bulgaria)		
IOEP	Institute of Environmental Protection (Poland)		
ISAC	Istituto di Scienze dell'Atmosfera e del Clima, Consiglio Nazionale delle		
10/10	Ricerche (Italy)		
ITM	Department of Applied Environmental Science, Stockholm University,		
	(Sweden)		
IVL	Sweden) Swedish Environmental Research Institute, Göteborg (Sweden)		
JMA	Japan Meteorological Agency (Japan)		
KMA	Korea Meteorological Administration (Republic of Korea)		
KMD	Kenya Meteorological Department (Kenya)		
KRISS	Korea Research Institute of Standards and Science (Republic of Korea)		
KSNU	Kyrgyz State National University (Kyrgyzstan)		
KUP	Physics Institute, Climate and Environmental Physics, University of Bern		

	(Switzerland)					
LEGMA	Latvian Environment, Geology and Meteorology Agency (Latvia)					
LSCE	Laboratoire des Sciences du Climat et de l'Environnement (France)					
MGO	Main Geophysical Observatory, Roshydromet (Russian Federation)					
MPI-BGC	Max-Planck Institute (MPI) for Biogeochemistry in Jena (Germany)					
MMD	Malaysian Meteorological Department					
MRI	Meteorological Research Institute, JMA (Japan)					
Nagoya Univ.	Nagoya University (Japan)					
NIER	National Institute of Environmental Resaearch (Republic of Korea)					
NIES	National Institute for Environmental Studies (Japan)					
NILU	Norwegian Institute for Air Research (Norway)					
NIMR	National Institute of Meteorologocal Reserch, KMA (Republic of Korea)					
NIPR	National Institute of Polar Research (Japan)					
NIST	National Institute of Standards and Technology (USA)					
NIWA	National Institute of Water & Atmospheric Research (New Zealand)					
NMA	National Meteorological Administration (Romania)					
NOAA/ESRL	Earth System Research Laboratory, NOAA (USA)					
NPL	National Physical Laboratory (United Kingdom)					
NUI	National University of Ireland, Galway (Ireland)					
ONM	Office National de la Météorologie (Algeria)					
Osaka Univ.	Osaka University (Japan)					
PolyU	Hong Kong Polytechnic University (Hong Kong, China)					
RIVM	National Institute for Health and Environment (Netherlands)					
Roshydromet	Federal Service for Hydrometeorology and Environmental Monitoring					
	(Russian Federation)					
RSE	Ricerca sul Sistema Elettrico (Italy)					
Saitama	Saitama Prefecture (Japan)					
SAWS	South African Weather Service (South Africa)					
	Shizuoka University (Japan)					
SMN	Servicio Meteorológico Nacional (Argentina)					
Tohoku Univ.	Tohoku University (Japan)					
UBA	Umweltbundesamt (Germany)					
Univ. Malta	University of Malta (Malta)					
Univ. York	University of York (United Kingdom of Great Britain and Northern Ireland)					
WDCGG	World Data Centre for Greenhouse Gases, operated by JMA, Japan (WMO)					
WMO	World Meteorological Organization					
Station	Country/Territory	Index Number	Latitude (° ')	Location Longitude (° ')	Altitude (m)	Parameter
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REGION I (Africa)						
Amsterdam Island	France	AMS137S00	37 47 S	77 31 E		CH4, CO, CO2, VOCs
Amsterdam Island	France	AMS137S00	37 47 S	77 31 E		CH4, CO ₂
Ascension Island	United Kingdom of Great Britain and Northern Ireland	ASC107S00	7 55 S	14 25 W		¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Assekrem	Algeria	ASK123N00	23 16 N	5 37 E	2710	O3
Assekrem	Algeria	ASK123N00	23 16 N	5 37 E		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Cairo	Egypt	CAI130N00	30 04 N	31 16 E	35	CO ₂ , O ₃
Cape Point	South Africa	CPT134S00	34 21 S	18 28 E		CH4, CO, CO2, N2O, O3
Cape Point	South Africa	CPT134S00	34 21 S	18 28 E		CH4, CO ₂
Cape Verde Observatory	Cape Verde	CVO116N00	16 50 N	24 52 W		CO, NOx, O ₃ , VOCs
Crozet	France	CRZ146S00	46 27 S	51 51 E		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Funchal	Portugal	FUN132N00	32 38 N	16 52 W	58	
Gobabeb	Namibia	NMB123S00	23 34 S	15 01 E		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂
Izaña (Tenerife)	Spain	IZO128N00	28 18 N	16 30 W		CH4, CO, CO2, N2O, O3, SF6
Izaña (Tenerife)	Spain	IZO128N00	28 18 N	16 30 W		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Mahe Island	Seychelles	SEY104S00	4 40 S	55 10 E		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Mt. Kenya	Kenya	MKN100S00	0 03 S	37 17 E		CO, O_3
Mt. Kenya	Kenya	MKN100S00	0 03 S	37 17 E	3678	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO ₂ , VOCs
REGION II (Asia)						
Anmyeon-do	Republic of Korea	AMY236N00	36 31 N	126 19 E		CFCs, CH4, CO2, N2O
Bering Island	Russian Federation	BER255N00	55 12 N	165 58 E		CO ₂
Cape Ochi-ishi	Japan	COI243N00	43 08 N	145 30 E		CH4, CO2
Cape Rama	India	CRI215N00	15 04 N			¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Everest - Pyramid	Nepal	PYR227N00	27 57 N	86 48 E	5079	
Gosan	Republic of Korea	GSN233N00	33 16 N	126 10 E		CFCs, CH4, CO ₂ , N ₂ O
Gosan	Republic of Korea	GSN233N01		126 05 E		CFCs, CH ₄ , CO ₂ , N ₂ O
Hamamatsu	Japan	HMM234N00		137 43 E		CO ₂
Hateruma	Japan	HAT224N00	24 03 N	123 47 E		CH_4 , CO_2 , N_2O
Hok Tsui	Hong Kong, China	HKG222N00	22 12 N	114 15 E		CO ₂
Hok Tsui	Hong Kong, China	HKG222N00	22 12 N	114 15 E		CO, O_3
Issyk-Kul	Kyrgyzstan	ISK242N00	42 37 N	76 58 E		CH_4, CO_2, O_3
Kaashidhoo Kingla Dark	Maldives	KCO204N00	4 58 N	73 28 E		¹³ CO ₂ , CH ₄ , CO ₂ CO ₂
King's Park Kisai	Hong Kong, China	HKO222N00 KIS236N00	22 18 N 36 04 N	114 10 E 139 33 E		CO_2 CO_2
Kotelny Island	Japan Russian Federation	K0T276N00	30 04 N 76 00 N	139 33 E 137 52 E		CO_2 CO_2
Kyzylcha	Uzbekistan	KYZ240N00	40 52 N	66 09 E	340	
Lulin	China	LLN223N00	40 32 N 23 28 N	120 52 E		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂
Memanbetsu	Japan	MMB243N00	43 55 N		32.9	
Mikawa-Ichinomiya	Japan	MKW234N00	34 51 N	137 25 E		CO ₂
Minamitorishima	Japan	MNM224N00	24 16 N	157 25 E 153 58 E		CH ₄ , CO, CO ₂ , O ₃
Mt. Dodaira	Japan	DDR236N00	36 00 N	139 10 E	840	
Mt. Waliguan	China	WLG236N00	36 16 N	100 54 E		CH ₄ , CO ₂
Mt. Waliguan	China	WLG236N00	36 16 N	100 54 E	3810	¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂

LIST OF OBSERVING STATIONS

Nagoya Japan NGY235N00 35 08 N 136 58 E 35 NgO Plateau Assy Kazzdkhstan RZM243N00 43 15 N 77 52 E 2519 ¹⁴ CO ₂ , C ¹⁴ O ₂ , C ¹⁴ a, CO, CO ₂ , H ₂ Ryori Japan RYO239N00 39 01 N 141 49 E 260 CCl4, CPC3, CH4, CO, CO ₂ , H ₂ Shangdianzi China SD2240N00 44 27 N 75 34 F 41 C CO ₂ , C ¹⁴ O ₂ , C ¹⁴ , CO, CO ₂ , H ₂ Ship between Isigaki Japan SH2240N00 44 27 N 75 34 F 41 C, CO ₂ , C ¹⁴ O ₂ , C ¹⁴ O ₂ , C ¹⁴ , CO, CO ₂ , H ₂ South China Sea (03N) N/A SCS990N00 3 00 N 105 00 E 15 ¹¹ CO ₂ , C ¹⁴ O ₂ , C ¹⁴ , CO, CO ₂ , H ₂ South China Sea (03N) N/A SCS991N00 10 00 P E 15 ¹¹ CO ₂ , C ¹⁴ O ₂ , C ¹⁴ , CO, CO ₂ , H ₂ South China Sea (03N) N/A SCS991N00 10 00 P E 15 ¹¹ CO ₂ , C ¹⁴ O ₂ , C ¹⁴ , CO, CO ₂ , H ₂ South China Sea (13N) N/A SCS91N00 13 00 N 113 00 E 15 ¹¹ CO ₂ , C ¹⁴ O ₂ , C ¹⁴ , CO, CO ₂ , H ₂ South China Sea (13N) N/A SCS91N00					Location		
Platear Assy Kazakhstan KZM243N00 43 15 N 77 52 E 2519 ¹³ CO2, C ¹⁰ O, CH4, CO, CO2, H2 Ryori Japan RYO239N00 39 01 N 141 49 E 260 CCl4, CFCs, CH3, CCl3, CH4, CO, CO2, H2 Sary Taukum Kazakhstan KZD244N00 44 27 N 75 34 E 412 ¹² CO2, C ¹⁰ O, CH4, CO, CO2, H2 Ship between Ishigaki Japan SIH224N00 24 07 N 123 49 E 5 CO2 South China Sea (05N) N/A SCS903N00 30 0 N 107 00 E 15< <td>¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (05N) N/A SCS903N00 90 0N 109 00 E 15¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS903N00 100 N 113 00 E 15¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 18 00 N 113 00 E 15¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 36 0N 137 05 E 140 07 E 26 South China Sea (12N) N/A SCS913N00</td>	¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (05N) N/A SCS903N00 90 0N 109 00 E 15 ¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS903N00 100 N 113 00 E 15 ¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 18 00 N 113 00 E 15 ¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 36 0N 137 05 E 140 07 E 26 South China Sea (12N) N/A SCS913N00	Station	Country/Territory	Index Number		-	
Platear Assy Kazakhstan KZM243N00 43 15 N 77 52 E 2519 ¹³ CO2, C ¹⁰ O, CH4, CO, CO2, H2 Ryori Japan RYO239N00 39 01 N 141 49 E 260 CCl4, CFCs, CH3, CCl3, CH4, CO, CO2, H2 Sary Taukum Kazakhstan KZD244N00 44 27 N 75 34 E 412 ¹² CO2, C ¹⁰ O, CH4, CO, CO2, H2 Ship between Ishigaki Japan SIH224N00 24 07 N 123 49 E 5 CO2 South China Sea (05N) N/A SCS903N00 30 0 N 107 00 E 15< <td>¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (05N) N/A SCS903N00 90 0N 109 00 E 15¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS903N00 100 N 113 00 E 15¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 18 00 N 113 00 E 15¹³CO2, C¹⁴O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 36 0N 137 05 E 140 07 E 26 South China Sea (12N) N/A SCS913N00</td> <td>Nagova</td> <td>Japan</td> <td>NGY235N00</td> <td>35 08 N</td> <td>136 58 E</td> <td>35 N2O</td>	¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (05N) N/A SCS903N00 90 0N 109 00 E 15 ¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS903N00 100 N 113 00 E 15 ¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 18 00 N 113 00 E 15 ¹³ CO2, C ¹⁴ O, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS913N00 36 0N 137 05 E 140 07 E 26 South China Sea (12N) N/A SCS913N00	Nagova	Japan	NGY235N00	35 08 N	136 58 E	35 N2O
Ryori Japan RY0239N00 39 01 N 141 49 E 260 CCl, CFCs, CH ₅ CCl, CH ₄ , CO, CO ₂ , N ₂ O, O Sary Taukum Kazakhstan KZD244N00 44 27 N 75 34 F 12 ¹ CO ₂ , CPO ₃ , CH ₄ , CO, CO ₂ , N ₂ O, O Shangdianzi China SD2240N00 40 38 N 117 06 E 287 CH ₄ , CO ₂ Island Subit China Sea (03N) N/A SCS905N00 3 00 N 105 00 E 15 ¹⁴ CO ₂ , CPO ₂ , CH ₄ , CO, CO ₂ , H ₂ South China Sea (03N) N/A SCS905N00 6 00 N 107 00 E 15 ¹⁴ CO ₂ , CPO ₂ , CH ₄ , CO, CO ₂ , H ₂ South China Sea (12N) N/A SCS905N00 100 N 110 00 E 15 ¹⁴ CO ₂ , CPO ₂ , CH ₄ , CO, CO ₂ , H ₂ South China Sea (18N) N/A SCS915N00 13 00 E 15 ¹⁴ CO ₂ , CPO ₂ , CH ₄ , CO, CO ₂ , H ₂ South China Sea (18N) N/A SCS915N00 36 08 N 132 1E 63 CO ₂ ¹⁴ CO ₂ , CPO ₂ , CH ₄ , CO, CO ₂ , H ₂ South China Sea (18N) N/A SCS915N00 36 08 N 137 25 E 120 ¹⁵ C		-					
CO2, NO, 0.3 Sary Taukum Kazakhstan KZD244N00 44 27 N 75 34 E 412 ¹¹ CO2, C ¹⁸ O2, C	Ryori						
Shangdianzi China SDZ240N00 40 38 N 117 06 E 287 CH4, CO2 Ship between Ishigaki Japan SIH224N00 24 07 N 123 49 E 5 CO2 Island and Hateruma Island and Hateruma South China Sea (03N) N/A SCS903N00 3 00 N 105 00 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (03N) N/A SCS903N00 9 00 N 109 00 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS912N00 12 00 N 11 00 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS912N00 12 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 21 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 21 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 31 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 31 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 31 00 N 11 30 0 E 15 ¹³ CO2	2	1					
Shangdianzi China SDZ240N00 40 38 N 117 06 E 287 CH4, CO2 Ship between Ishigaki Japan SIH224N00 24 07 N 123 49 E 5 CO2 Island and Hateruma Island and Hateruma South China Sea (03N) N/A SCS903N00 3 00 N 105 00 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (03N) N/A SCS903N00 9 00 N 109 00 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS912N00 12 00 N 11 00 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (12N) N/A SCS912N00 12 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 21 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 21 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 31 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 31 00 N 11 30 0 E 15 ¹³ CO2, C ¹⁴ O2, CH4, CO, CO2, H2 South China Sea (21N) N/A SCS912N00 31 00 N 11 30 0 E 15 ¹³ CO2	Sary Taukum	Kazakhstan	KZD244N00	44 27 N	75 34 E	412 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H	
Island and Hateruma Island Island South China Sea (03N) N/A SCS903N00 3 00 N 105 00 F. 15 ¹³ CO ₂ , C ¹⁴ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (08N) N/A SCS909N00 9 00 N 109 00 E 15 ¹³ CO ₂ , C ¹⁴ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (12N) N/A SCS912N00 12 00 N 111 00 E 15 ¹³ CO ₂ , C ¹⁴ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (12N) N/A SCS912N00 12 00 N 113 00 E 15 ¹³ CO ₂ , C ¹⁴ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (12N) N/A SCS912N00 21 00 N 114 00 E 15 ¹³ CO ₂ , C ¹⁴ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (21N) N/A SCS912N00 21 00 N 114 00 E 15 ¹³ CO ₂ , C ¹⁴ O ₂ , CH4, CO, CO ₂ , H2 Suita Japan TKP236N00 36 02 N 140 07 E 20 ¹³ CH4, CO ₂ CH4, CO ₂ Tsukuba Japan TKB236N10 36 02 N 140 07 E 25 0.3 111 04 E 14 140 07 CO ₂ , CH ₄ , CO, CO ₂ , H2 112 ava Japan UM235N00 35 2 N 139 35 E 10 CO ₂ CH	Shangdianzi	China	SDZ240N00	40 38 N	117 06 E	287 CH ₄ , CO ₂	
Island South China Sea (03N) N/A SCS903N00 3 00 N 105 00 E 15 ¹³ CO ₂ , C ¹³ O ₂ , C ¹³ O ₂ , C ¹⁴ O, CO, CD, H ₂ South China Sea (09N) N/A SCS909N00 9 00 N 109 00 E 15 ¹³ CO ₂ , C ¹³ O ₂ , C ¹⁴ O, CO, CD, H ₂ South China Sea (12N) N/A SCS912N00 12 00 N 111 00 E 15 ¹³ CO ₂ , C ¹³ O ₂ , C ¹⁴ O, CO, CD, H ₂ South China Sea (13N) N/A SCS912N00 18 00 N 113 00 E 15 ¹³ CO ₂ , C ¹⁴ O, CO, H ₂ CO ₂ South China Sea (18N) N/A SCS918N00 18 00 N 113 00 E 15 ¹³ CO ₂ , C ¹⁴ O, CO, H ₂ CO ₂ C ¹⁴ O, CO ₂ , H ₂ CO ₂ ¹³ CO ₂ , C ¹⁴ O, CO, H ₂ South China Sea (18N) N/A SCS918N00 36 43 N 126 07 E 26 O ₂ ¹⁴ CO ₂ , C ¹⁴ O, CO, H ₂ South China Sea Japan TKP236N00 36 02 N 140 07 E 25 CO ₂ ¹⁴ SU ₂ SO ₂ C ¹⁴ A, ¹⁵ CO ₂ , C ¹⁴ O, CO ₂ , C ¹⁴ A, ¹⁵ CO ₂ , ¹⁴ SU ₂ CO ₂ ¹⁴ SU ₂ SO ₂ C ¹⁴ A, ¹⁵ CO ₂ , C ¹⁴ A, ¹⁵ CO ₂ , ¹⁴ CO, ¹⁴ CO, ¹⁴ CO CO ₂ ¹⁴ SU ₂ SO ₂ C ¹⁴ A, ¹⁵ CO	Ship between Ishigaki	Japan	SIH224N00	24 07 N	123 49 E	5 CO ₂	
South China Sea (03N) N/A SCS903N00 3 00 N 105 00 E 15 ¹³ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (06N) N/A SCS906N00 60 0 N 107 00 E 15 ¹³ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (12N) N/A SCS9012N00 12 00 N 111 00 E 15 ¹³ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (15N) N/A SCS915N00 15 00 N 113 00 E 15 ¹³ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (15N) N/A SCS912N00 21 00 N 114 00 E 15 ¹³ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (15N) N/A SCS921N00 21 00 N 114 00 E 15 ¹³ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea (12N) N/A SCS921N00 36 04 N 126 07 E 20 ¹⁴ CH4, ¹⁴ CO ₂ , C ¹⁸ O ₂ , CH4, CO, CO ₂ , H2 South China Sea Japan TKY236N00 36 02 N 140 07 E 25 03 Takayama Japan TKB236N10 36 02 N 140 07 E 25 03 CH4, CO ₂							
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Easter Island Chile EIC327S00 27 07 S 109 27 W 50 13CO2, C18O2, CH4, CO, CO2, H2, VOCs Huancayo Peru HUA312S00 12 04 S 75 31 W 3313 CO2 Tierra del Fuego Argentina TDF354S00 54 52 S 68 28 W 20 13CO2, C18O2, C2Cl4, CBrClF2, CFCs, CH2Cl2, CH3Br, CH3CCl3, CH3Cl, CH3Cl, CH3Cl, CH4, CO, CO2, H2, HCFCs, HFCS, VOCS Ushuaia Argentina USH354S00 54 49 S 68 17 W 18 CO, O3 REGION IV (North and Central America) ALT482N00 82 27 N 62 31 W 210 13CO2, CH4, CO, CO2, H2, N2O		-					
Huancayo Peru HUA312S00 12 04 S 75 31 W 3313 CO2 Tierra del Fuego Argentina TDF354S00 54 52 S 68 28 W 20 ¹³ CO2, C ¹⁸ O2, C ₂ Cl ₄ , CBrClF2, CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH4, CO, CO2, H2, HCFCs, HFCS, VOCs Ushuaia Argentina USH354S00 54 49 S 68 17 W 18 CO, O ₃ REGION IV (North and Central America) ALT482N00 82 27 N 62 31 W 210 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O		Northern Ireland					
Huancayo Peru HUA312S00 12 04 S 75 31 W 3313 CO2 Tierra del Fuego Argentina TDF354S00 54 52 S 68 28 W 20 ¹³ CO2, C18O2, C2Cl4, CBrClF2, CFCs, CH2Cl2, CH3Br, CH3CCl3, CH3Cl, CH4, CO, CO2, H2, HCFCs, HFCS, VOCS Ushuaia Argentina USH354S00 54 49 S 68 17 W 18 CO, O3 REGION IV (North and Central America) ALT482N00 82 27 N 62 31 W 210 ¹³ CO2, CH4, CO, CO2, H2, N2O	Easter Island	Chile	EIC327S00	27 07 S	109 27 W		
Tierra del Fuego Argentina TDF354S00 54 52 S 68 28 W 20 ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, VOCs Ushuaia Argentina USH354S00 54 49 S 68 17 W 18 CO, O ₃ REGION IV (North and Central America) Alert Canada ALT482N00 82 27 N 62 31 W 210 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O	Huancayo	Peru	HUA312S00	12 04 S	75 31 W		
CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, VOCs Ushuaia Argentina USH354S00 54 49 S 68 17 W 18 CO, O ₃ REGION IV (North and Central America) Alert Canada ALT482N00 82 27 N 62 31 W 210 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O							
Ushuaia Argentina USH354S00 54 49 S 68 17 W 18 CO, O3 REGION IV (North and Central America) Alert Canada ALT482N00 82 27 N 62 31 W 210 13CO2, CH4, CO, CO2, H2, N2O	C C	C				CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ ,	
Alert Canada ALT482N00 82 27 N 62 31 W 210 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O	Ushuaia	Argentina	USH354S00	54 49 S	68 17 W		
	REGION IV (North an	d Central America)					
	Alert	Canada	ALT482N00	82 27 N	62 31 W	210 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O	
	Alert	Canada	ALT482N00	82 27 N	62 31 W		

Station	Country/Torritory	Inday Number	Latituda	Location	Altitude Daramatar
Station	Country/Territory	Index Number	(° ')	(° ')	Altitude Parameter (m)
Alert	Canada	ALT482N00	82 27 N	62 31 W	210 ¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, N ₂ O, SF ₆ , VOCs
Algoma	Canada	ALG447N00	47 01 N	84 22 W	411 O3
Argyle	United States of America	AMT445N00	45 01 N		50 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , VOCs
Barrow	United States of America	BRW471N00	71 19 N	156 35 W	 ¹³CH₄, ¹³CO₂, C¹⁸O₂, C₂Cl₄, CBrClF₂, CBrF₃, CCl₄, CFCs, CH₂Cl₂, CH₃Br, CH₃CCl₃, CH₃Cl, CH₄, CO, CO₂, H₂, HCFCs, HFCs, N₂O, O₃, SF₆, VOCs
Bratt's Lake	Canada	BRA450N00	50 12 N	104 42 W	595 O ₃
Candle Lake	Canada	CDL453N00	53 52 N	104 39 W	489 CH ₄ , CO, CO ₂
Cape Meares	United States of America	CMO445N00	45 28 N	123 58 W	30 CCl ₄ , CFCs, CH ₃ CCl ₃ , CH ₄ , N ₂ C
Cape Meares	United States of America	CMO445N00	45 28 N	123 58 W	30 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Cape St. James	Canada	CSJ451N00	51 55 N	131 01 W	89 CO ₂
Chalk River	Canada	CHA446N00	46 04 N	77 24 W	184 O ₃
Chapais	Canada	CPS449N00	49 49 N	74 58 W	381 O ₃
Chibougamau	Canada	CHM449N00	49 40 N	74 20 W	393 CH ₄ , CO, CO ₂
Churchill	Canada	CHL458N00	58 45 N		35 ¹³ CO ₂ , C ¹⁸ O ₂
Cold Bay	United States of America	CBA455N00	55 12 N	162 43 W	25 ¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
East Trout Lake	Canada	ETL454N00	54 21 N	104 59 W	492 CH ₄ , CO, CO ₂
Egbert	Canada	EGB444N00	44 13 N		253 O ₃
Egbert	Canada	EGB444N01	44 13 N		253 CH ₄ , CO, CO ₂ , VOCs
Estevan Point	Canada	ESP449N00		126 32 W	39 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Estevan Point	Canada	ESP449N00		126 32 W	39 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , N ₂ O, SF ₆
Esther	Canada	EST451N00		110 12 W	707 O ₃
Experimental Lakes Area	Canada	ELA449N00	49 40 N	93 43 W	369 O ₃
Fraserdale	Canada	FSD449N00	49 52 N		210 CH ₄ , CO, CO ₂
Grifton	United States of America	ITN435N00	35 21 N	77 22 W	505 ¹³ CO ₂ , C ¹⁸ O ₂ , CCl ₄ , CFCs, CH ₄ , CO, CO ₂ , H ₂ , N ₂ O, SF ₆
Harvard Forest	United States of America	HFM442N00	42 53 N	72 17 W	340 C ₂ Cl ₄ , CBrClF ₂ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, HCFCs, HFCs, N ₂ O, SF ₆
Kejimkujik	Canada	KEJ444N00	44 25 N	65 12 W	127 O ₃
Key Biscayne	United States of America	KEY425N00	25 40 N	80 12 W	3 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ VOCs
Kitt Peak	United States of America	KPA431N00	31 58 N	111 35 W	2083 CH ₄
La Jolla	United States of America	SIO432N00	32 49 N	117 16 W	14 CH ₄
La Palma	Cuba	PLM422N00	22 45 N	83 31 W	47 NO ₂
Lac La Biche	Canada	LLB454N00		112 27 W	540 CH ₄ , CO ₂ , VOCs
Lac La Biche (Alberta)	Canada	LLB454N01	54 57 N	112 27 W	540 CH ₄ , CO, CO ₂
Longwoods	Canada	LON442N00	42 52 N	81 28 W	239 O ₃

Station	Country/Territory	Index Number	Latitude (° ')	Location Longitude (° ')	Altitude (m)	e Parameter
Mex High Altitude Global Climate Observation Center,	Mexico	MEX419N00	19 58 N	97 10 W	4560	CH4, CO2, VOCs
Mexico						
Moody	United States of America	WKT431N00	31 19 N	97 19 W	708	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄
Mould Bay	Canada	MBC476N00	76 15 N	119 19 W	58	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Niwot Ridge (C-1)	United States of America	NWR440N00	40 02 N	105 32 W	3021	C ₂ Cl ₄ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, HCFCs, HFCs, N ₂ O, O ₃ , SF ₆
Niwot Ridge (Saddle)	United States of America	NWR440N02	40 03 N	105 35 W	3528	
Niwot Ridge (T-van)	United States of America	NWR440N01	40 03 N	105 35 W	3523	¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Olympic Peninsula	United States of America	OPW448N00	48 15 N	124 25 W	488	CH4, CO2, H2
Pacific Ocean (15N)	N/A	POC915N00	15 00 N	145 00 W	10	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (20N)	N/A	POC920N00	20 00 N	141 00 W	10	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (25N)	N/A	POC925N00	25 00 N	139 00 W		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (30N)	N/A	POC930N00	30 00 N	135 00 W	10	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (35N)	N/A	POC935N00	35 00 N	137 00 W	10	¹³ CO ₂ , C ¹⁸ O ₂ , CO, H ₂
Pacific Ocean (40N)	N/A	POC940N00	40 00 N	136 00 W	10	¹³ CO ₂ , H ₂
Pacific Ocean (45N)	N/A	POC945N00	45 00 N	131 00 W		¹³ CO ₂ , H ₂
Park Falls	United States of America	LEF445N00	45 55 N	90 16 W	868	 ¹³CO₂, C¹⁸O₂, C₂Cl₄, CBrClF₂, CCl₄, CFCs, CH₂Cl₂, CH₃Br, CH₃CCl₃, CH₃Cl, CH₄, CO, CO₂, H₂, HCFCs, HFCs, N₂O, SF₆, VOCs
Point Arena	United States of America	PTA438N00	38 57 N	123 43 W	17	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂
Ragged Point	Barbados	RPB413N00	13 10 N	59 25 W	45	C ₂ Cl ₄ , C ₂ HCl ₃ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CHCl ₃ , HCFCs, HFCs, N ₂ O, PFCs, SF ₆ , SO ₂ F ₂
Ragged Point	Barbados	RPB413N00	13 10 N	59 25 W	45	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , O ₃
Sable Island	Canada	WSA443N00	43 55 N	60 01 W	5	CH4, CO, CO2, N2O, SF6
Saturna	Canada	SAT448N00	48 46 N	123 07 W	178	
Shemya Island	United States of America	SHM452N00	52 43 N	174 04 E		¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Southern Great Plains	United States of America	SGP436N00	36 46 N	97 30 W	314	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , N ₂ O, SF ₆ , VOCs
St. Croix	United States of America	AVI417N00	17 45 N	64 45 W	3	CH ₄ , CO ₂
St. David's Head	United Kingdom of Great Britain and Northern Ireland	BME432N00	32 22 N	64 39 W	30	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
	Canada					

				Location	
Station	Country/Territory	Index Number	Latitude (° ')		Altitude Parameter (m)
Trinidad Head	United States of America	THD441N00	41 02 N	124 09 W	120 C ₂ Cl ₄ , C ₂ HCl ₃ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CHCl ₃ , HCFCs, HFCs, N ₂ O, PFCs, SF ₆ , SO ₂ F ₂
Trinidad Head	United States of America	THD441N00	41 02 N	124 09 W	120 ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , HCFCs, HFCs, N ₂ O, O ₃ , SF ₆ , VOCs
Tudor Hill	United Kingdom of Great Britain and Northern Ireland	BMW432N00	32 16 N	64 52 W	30 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , O ₃ , VOCs
Wendover	United States of America	UTA439N00	39 52 N	113 43 W	1320 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
West Branch	United States of America	WBI441N00	41 43 N	91 21 W	241.7 ¹³ CO ₂ , C ¹⁸ O ₂
REGION V (South-We	est Pacific)				
Baring Head	New Zealand	BHD541S00	41 24 S	174 52 E	85 ¹³ CH ₄ , ¹⁴ CO ₂ , CH ₄ , CO, CO ₂ , N ₂ O, O ₃ , VOCs
Baring Head	New Zealand	BHD541S00	41 24 S	174 52 E	85 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂
Bukit Koto Tabang	Indonesia	BKT500S00	0 12 S		864.5 NO ₂ , SO ₂
Bukit Koto Tabang	Indonesia	BKT500S00	0 12 S		864.5 CO, O ₃
Bukit Koto Tabang	Indonesia	BKT500S00	0 12 S		864.5 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O, SF ₆ , VOCs
Cape Ferguson	Australia	CFA519S00	19 16 S	147 03 E	2 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Cape Grim	Australia	CGO540S00	40 40 S	144 40 E	94 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Cape Grim	Australia	CGO540S00	40 40 S	144 40 E	 94 C₂Cl₄, C₂HCl₃, CBrClF₂, CBrF₃, CCl₄, CFCs, CH₂Cl₂, CH₃Br, CH₃CCl₃, CH₃Cl, CH₄, CHCl₃, CO, H₂, HCFCs, HFCs, N₂O, PFCs, SF₆, SO₂F₂
Cape Grim	Australia	CGO540S00		144 40 E	94 CO ₂ , O ₃
Cape Grim	Australia	CGO540S00	40 40 S	144 40 E	94 ¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, N ₂ O, SF ₆ , VOCs
Cape Kumukahi	United States of America	KUM519N00	19 31 N	154 49 W	3 ¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, N ₂ O, SF ₆ , VOCs
Christmas Island	Kiribati	CHR501N00	1 42 N	157 10 W	3 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Danum Valley GAW Baseline Station	Malaysia	DMV504N00	4 58 N	117 49 E	426 CO ₂ , O ₃
Guam	United States of America	GMI513N00	13 25 N	144 46 E	2 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Jakarta	Indonesia	JKR506S00	6 10 S	106 49 E	7 NO ₂ , SO ₂
Kaitorete Spit	New Zealand	NZL543S00	43 49 S	172 37 E	3 CH4
Lauder	New Zealand	LAU545S00	45 01 S	169 40 E	370 O ₃
Macquarie Island	Australia	MQA554S00	54 28 S	158 58 E	12 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O

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Mauna Loa	United States of America	MLO519N00	19 32 N	155 34 W	3397 ¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Mauna Loa	United States of America	MLO519N00	19 32 N	155 34 W	 3397 ¹³CH₄, ¹³CO₂, C¹⁸O₂, C₂Cl₄, CBrClF₂, CBrF₃, CCl₄, CFCs, CH₂Cl₂, CH₃Br, CH₃CCl₃, CH₃Cl, CH₄, CO, CO₂, H₂, HCFCs, HFCs, N₂O, O₃, SF₆, VOCs
Pacific Ocean (00N)	N/A	POC900N00	0 00 N	155 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (05N)	N/A	POC905N00	5 00 N	151 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (05S)	N/A	POC905S00	5 00 S	159 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (10N)	N/A	POC910N00	10 00 N	149 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (10S)	N/A	POC910S00	10 00 S	161 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (15S)	N/A	POC915S00	15 00 S	171 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (20S)	N/A	POC920S00	20 00 S	174 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (25S)	N/A	POC925S00	25 00 S	171 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (30S)	N/A	POC930S00	30 00 S	176 00 W	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific Ocean (35S)	N/A	POC935S00	35 00 S	180 00 E	10 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Sand Island	United States of America	MID528N00	28 11 N	177 22 W	7.7 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Tanah Rata	Malaysia	TAR504N00	4 28 N	101 22 E	1545 O ₃
Tutuila (Cape Matatula)	United States of America	SMO514S00	14 14 S	170 34 W	42 C ₂ Cl ₄ , C ₂ HCl ₃ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CHCl ₃ , HCFCs, HFCs, N ₂ O, PFCs, SF ₆ , SO ₂ F ₂
Tutuila (Cape Matatula)	United States of America	SMO514S00	14 14 S	170 34 W	42 ¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, N ₂ O, O ₃ , SF ₆ , VOCs
REGION VI (Europe)					
Ähtäri	Finland	AHT662N00	62 34 N	24 11 E	180 NO ₂ , O ₃ , SO ₂
Adrigole	Ireland	ADR651N00	51 40 N		50 CCl ₄ , CFCs, CH ₃ CCl ₃ , N ₂ O
Angra do Heroismo	Portugal	ANG638N00	38 40 N		74 O ₃
BEO Moussala	Bulgaria	BEO642N00	42 10 N		2925 CO, CO ₂ , NO, NO ₂ , NOx, O ₃ , SO ₂
Baltic Sea	Poland	BAL655N00	55 21 N	17 13 E	28 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Begur	Spain	BGU641N00	41 58 N	3 13 E	13 CH ₄ , CO ₂
Beja	Portugal	BEJ638N00	38 01 N	7 52 W	246 O ₃
Black Sea	Romania	BSC644N00	44 10 N	28 40 E	3 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Bragança	Portugal	BRG641N00	41 47 N	6 43 W	690 SO ₂
Brotjacklriegel	Germany	BRT648N00	48 49 N	13 13 E	1016 VOCs
Brotjacklriegel	Germany	BRT648N00	48 49 N	13 13 E	1016 CO ₂ , O ₃
Burgas	Bulgaria	BUR642N00	42 28 N	27 28 E	16 NO ₂ , SO ₂
Campisabalos	Spain	CAM641N00	41 16 N	3 08 W	1360 VOCs
Castelo Branco	Portugal	CAS639N00	39 49 N	7 28 W	386 O ₃
Danki	Russian Federation	DAK654N00	54 53 N	37 47 E	140 O ₃
Deuselbach	Germany	DEU649N00	49 46 N	7 02 E	480 CH ₄ , CO ₂ , O ₃

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Doñana	Spain	DON637N00	37 02 N	6 32 W	5	NO ₂ , O ₃ , SO ₂
Dobele	Latvia	DBL656N00	56 22 N	23 11 E	42	O ₃
Donon	France	DNN648N00	48 30 N	7 07 E	775	VOCs
Dwejra Point	Malta	GOZ636N00	36 02 N	14 10 E	30	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Eskdalemuir	United Kingdom of Great Britain and Northern Ireland	EDM655N00	55 19 N	3 12 W	242	O ₃
Finokalia	Greece	FIK635N00	35 20 N	25 40 E	150	CH ₄ , CO ₂
Fundata	Romania	FDT645N00	45 28 N	25 18 E	1383.5	CO ₂ , NO ₂ , O ₃
Fundata	Romania	FDT645N00	45 28 N	25 18 E	1383.5	NO ₂ , SO ₂
Giordan Lighthouse	Malta	GLH636N00	36 04 N	14 13 E	167	CO, O_3
Hegyhatsal	Hungary	HUN646N00	46 57 N	16 38 E	248	CO ₂
Hegyhatsal	Hungary	HUN646N00	46 57 N	16 38 E	248	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Heimaey	Iceland	ICE663N00	63 23 N	20 16 W	100	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , O ₃ , VOCs
Hohe Warte	Austria	HHE648N00	48 15 N	16 22 E	202	NO, NO ₂ , SO ₂
Hohe Warte	Austria	HHE648N00	48 15 N	16 22 E		NO, NO ₂ , SO ₂
Hohenpeissenberg	Germany	HPB647N00	47 47 N	11 01 E	985	²²² Rn, CO, H ₂ O ₂ , NO, NO ₂ , NOx , NOy, O ₃ , PAN, ROOH, SO ₂ , VOCs
Hohenpeissenberg	Germany	HPB647N00	47 47 N	11 01 E	985	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , VOCs
Ile Grande	France	LPO648N00	48 48 N	3 35 W	10	CH ₄ , CO ₂
Iskrba	Slovenia	IRB645N00	45 34 N	14 52 E	520	NO ₂ , O ₃ , SO ₂
Ivan Sedlo	Bosnia and Herzegovina	IVN643N00	43 46 N	18 01 E	970	NO ₂ , SO ₂
Jarczew	Poland	JCZ651N00	51 49 N	21 58 E	180	NO_2 , SO_2
Jungfraujoch	Switzerland	JFJ646N00	46 32 N	7 59 E	3580	CO ₂
Jungfraujoch	Switzerland	JFJ646N00	46 32 N	7 59 E	3580	CH4, CO, CO2, N2O, NO, NO2, NOx, NOy, O3, PAN, SF6, SO2
Jungfraujoch	Switzerland	JFJ646N00	46 32 N	7 59 E	3580	C ₂ Cl ₄ , C ₂ HCl ₃ , CBrClF ₂ , CBrF ₃ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CHCl ₃ , HCFCs, HFCs, PFCs, SF ₆ , SO ₂ F ₂
K-puszta	Hungary	KPS646N00	46 58 N	19 33 E	125	CO ₂ , NO ₂ , O ₃ , SO ₂
Kamenicki Vis	Serbia	KAM643N00	43 23 N	21 56 E	813	NO_2 , SO_2
Kloosterburen	Netherlands (the)	KTB653N00	53 23 N	6 25 E	0	CO, NO, NO ₂ , NO _x , SO ₂
Kollumerwaard	Netherlands (the)	KMW653N00	53 19 N	6 16 E		CH ₄ , CO, CO ₂ , NO, NO ₂ , NO _x , O ₃ , SO ₂
Kosetice	Czech Republic	KOS649N00	49 34 N	15 04 E		VOCs
Kosetice	Czech Republic	KOS649N00	49 34 N	15 04 E	534	CH ₄ , CO, NO, NO ₂ , O ₃ , SO ₂
Kovk	Slovenia	KVK646N00	46 07 N	15 05 E	600	O3
Krvavec	Slovenia	KVV646N00	46 17 N	14 31 E		CO, O ₃
La Cartuja	Spain	CAR637N00	37 12 N	3 36 W	720	NO_2 , SO_2
La Tardiere	France	LAT646N00	46 38 N	0 45 W	133	VOCs
Lampedusa	Italy	LMP635N00	35 31 N	12 37 E	45	CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Br ₂ , CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₃ I, CH ₄ , CHCl ₃ , CO ₂ , HCFCs, HFCs, N ₂ O SF ₆
Lampedusa	Italy	LMP635N00	35 31 N	12 37 E	45	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂
Lazaropole	The former Yugoslav Republic of Macedonia	LZP641N00	41 31 N	20 41 E		NO ₂ , SO ₂

Station	Country/Territory	Index Number	Latitude (° ')	Location Longitude (° ')	Altitude Parameter (m)	
Leba	Poland	LEB654N00	54 45 N	17 31 E	2 NO ₂ , SO ₂	
Lisboa / Gago Coutinho	Portugal	LIS638N00	38 46 N	9 07 W	105 O ₃	
Logroño	Spain	LOG642N00	42 27 N	2 30 W	370 NO ₂ , SO ₂	
Mace Head	Ireland	MHD653N00	53 19 N	9 54 W	8 O3	
Mace Head	Ireland	MHD653N00	53 19 N	9 54 W	8 CH ₄ , CO ₂	
Mace Head	Ireland	MHD653N00	53 19 N	9 54 W	8 C ₂ Cl ₄ , C ₂ HCl ₃ , CBr CCl ₄ , CFCs, CH ₂ Cl ₂ CH ₃ CCl ₃ , CH ₃ Cl, C CO, H ₂ , HCFCs, HF PFCs, SF ₆ , SO ₂ F ₂	2, CH3Br, H4, CHCl3,
Mace Head	Ireland	MHD653N00	53 19 N	9 54 W	8 ¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ CBrClF ₂ , CBrF ₃ , CC CH ₂ Cl ₂ , CH ₃ Br, CH CH ₃ Cl, CH ₄ , CO, C HCFCs, HFCs, N ₂ O	Cl4, CFCs, 3CCl3, O2, H2,
Mahón	Spain	MHN639N00	39 52 N	4 19 E	78 NO ₂ , O ₃ , SO ₂	
Monte Cimone	Italy	CMN644N00	44 10 N	10 41 E	2165 CH ₄ , CO, H ₂ , N ₂ O,	O ₃ , SF ₆
Monte Cimone	Italy	CMN644N00	44 10 N	10 41 E	2165 CO ₂	
Monte Velho	Portugal	MVH638N00	38 04 N	8 48 W	43 O ₃	
Neuglobsow	Germany	NGL653N00	53 10 N	13 01 E	65 CH ₄ , CO, CO ₂ , O ₃	
Noia	Spain	NIA642N00	42 43 N	8 55 W	685 NO ₂ , O ₃ , SO ₂	
Ocean Station "M"	Norway	STM666N00	66 00 N	2 00 E	5 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ ,	CO, CO ₂ , H ₂
Ocean Station Charlie	Russian Federation	STC652N00	52 45 N	35 30 W	5 CO ₂	
Ocean Station Charlie	United States of America	STC654N00	54 00 N	35 00 W	6 CO ₂	
Ochsenkopf	Germany	OXK650N00	50 01 N	11 48 E	1185 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , VOCs	CO, CO ₂ ,
Oulanka	Finland	OUL666N00	66 19 N	29 23 E	310 NO ₂ , O ₃ , SO ₂	
Pallas-Sammaltunturi	Finland	PAL667N00	67 58 N	24 07 E	560 VOCs	
Pallas-Sammaltunturi	Finland	PAL667N00	67 58 N	24 07 E	560 CH ₄ , CO ₂ , O ₃	
Pallas-Sammaltunturi	Finland	PAL667N00	67 58 N	24 07 E	560 ¹³ CO ₂ , C ¹⁸ O ₂ , CBrF CO ₂ , VOCs	3, CH4, CO,
Payerne	Switzerland	PAY646N00	46 49 N	6 57 E	490 CO, NO, NO ₂ , NOx	, O3, SO2
Penhas Douradas	Portugal	PEN640N00	40 25 N	7 32 W	1380 O ₃	
Peyrusse Vieille	France	PVI643N00	43 37 N	0 10 E	200 VOCs	
Pic du Midi	France	PDM642N00	42 56 N	0 08 E	2877 CO, O ₃	
Pic du Midi	France	PDM642N00	42 56 N	0 08 E	2877 CH ₄ , CO ₂	
Plateau Rosa	Italy	PRS645N00	45 55 N	7 42 E	3480 CH ₄ , CO ₂ , O ₃	
Pleven	Bulgaria	PLV643N00	43 25 N	24 36 E	64 NO ₂ , SO ₂	
Plovdiv	Bulgaria	PLD642N00	42 07 N	24 45 E	179 NO ₂ , SO ₂	
Puszcza Borecka/Diabla Gora	Poland	DIG654N00	54 08 N	22 04 E	157 CO ₂ , NO ₂ , O ₃ , SO ₂	
Puy de Dome	France	PUY645N00	45 46 N	2 57 E	1465 CH ₄ , CO ₂	
Rigi	Switzerland	RIG646N00	46 04 N	8 26 E	1031 CO, NO, NO ₂ , NOx VOCs	, O ₃ , SO ₂ ,
Roquetes	Spain	ROQ640N00	40 49 N	0 28 E	50 NO ₂ , O ₃ , SO ₂	
Rucava	Latvia	RCV656N00	56 09 N	21 10 E	18 NO ₂ , O ₃ , SO ₂	
San Pablo de los Montes	Spain	SPM639N00	39 32 N	4 20 W	917 NO ₂ , O ₃ , SO ₂	
Schauinsland	Germany	SSL647N00	47 55 N	7 55 E	1205 CH4, CO, CO ₂ , N ₂ O O ₃ , PAN, SF ₆	, NO, NO ₂ ,
Schmuecke	Germany	SCH650N00	50 38 N	10 46 E	937 VOCs	
Sede Boker	Israel	WIS631N00	31 07 N	34 52 E	400 ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ ,	CO, CO ₂ , H ₂

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Semenic	Romania	SEM645N00	45 07 N	21 58 E	1432	NO ₂ , SO ₂
Shepelevo	Russian Federation	SHP659N00	59 58 N	29 07 E		O ₃
Shetland	United Kingdom of Great Britain and Northern Ireland	SIS660N00	60 04 N	1 15 W	30	¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Site J	Denmark	GRL666N00	66 30 N	46 12 W	2030	CH ₄
Sniezka	Poland	SNZ650N00	50 43 N	15 43 E	1603	NO ₂ , SO ₂
Sofia	Bulgaria	SOF642N00	42 38 N	23 22 E	586	NO ₂ , SO ₂
Sonnblick	Austria	SNB647N00	47 02 N	12 56 E	3106	CO, CO ₂ , NO, NO ₂ , NOy, O ₃
Stîna de Vale	Romania	STN646N00	46 40 N	22 37 E	1116	NO ₂ , SO ₂
Starina	Slovakia	STA649N00	49 02 N	22 16 E	345	VOCs
Stephansplatz	Austria	STP648N00	48 13 N	16 22 E	171	NO, NO ₂ , SO ₂
Stephansplatz	Austria	STP648N00	48 13 N	16 22 E	171	NO, NO ₂ , SO ₂
Summit	Denmark	SUM672N00	72 34 N	38 28 W	3238	¹³ CO ₂ , C ¹⁸ O ₂ , CBrClF ₂ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO ₂ , HCFCs, HFCs, N ₂ O, O ₃ , SF ₆ , VOCs
Suwalki	Poland	SWL654N00	54 07 N	22 56 E	184	NO_2 , SO_2
Terceira Island	Portugal	AZR638N00	38 46 N	27 22 W		¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Teriberka	Russian Federation	TER669N00	69 12 N	35 06 E	40	CH4, CO ₂
Utö	Finland	UTO659N00	59 46 N	21 22 E		VOCs
Utö	Finland	UTO659N00	59 46 N	21 22 E	7	NO_2, O_3, SO_2
Varna	Bulgaria	VRN643N00	43 12 N	27 55 E	41	NO ₂ , SO ₂
Viana do Castelo	Portugal	VDC641N00	41 42 N	8 48 W	16	SO_2
Vindeln	Sweden	VDL664N00	64 15 N	19 46 E	271	O ₃
Virolahti	Finland	VIR660N00	60 31 N	27 40 E		NO_2, O_3, SO_2
Waldhof	Germany	LGB652N00	52 47 N	10 46 E	74	VOCs
Waldhof	Germany	LGB652N00	52 47 N	10 46 E		CO ₂ , O ₃
Wank Peak	Germany	WNK647N00	47 31 N	11 09 E		CO ₂ , NO _x , SO ₂
Westerland	Germany	WES654N00	54 55 N	8 19 E		CO ₂ , O ₃
Zabljak	Montenegro	ZBL643N00	43 08 N	19 07 E		NO_2 , SO_2
Zavodnje	Slovenia	ZRN646N00	46 25 N	15 00 E	770	
Zeppelinfjellet (Ny-Alesund)	Norway	ZEP678N00	78 54 N	11 52 E		CCl ₄ , CFCs, CH ₃ CCl ₃ , N ₂ O, O ₃ , SO ₂
Zeppelinfjellet (Ny-Alesund)	Norway	ZEP678N00	78 54 N	11 52 E		CO ₂
Zeppelinfjellet (Ny-Alesund)	Norway	ZEP678N00	78 54 N	11 52 E	475	C ₂ Cl ₄ , C ₂ HCl ₃ , CBrClF ₂ , CBrF ₃ , CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CHCl ₃ , HCFCs, HFCs
Zeppelinfjellet (Ny-Alesund)	Norway	ZEP678N00	78 54 N	11 52 E	475	¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ , VOCs
Zingst	Germany	ZGT654N00	54 25 N	12 43 E	1	VOCs
Zingst	Germany	ZGT654N00	54 25 N	12 43 E		CH4, CO ₂ , O ₃
Zoseni	Latvia	ZSN657N00	57 04 N	25 32 E		NO ₂ , O ₃ , SO ₂
Zugspitze	Germany	ZUG647N00	47 25 N	10 58 E	2960	
Zugspitze	Germany	ZUG647N00	47 25 N	10 58 E	2960	CH4, CO, CO ₂ , NO, NOx, NOy, O ₃
Zugspitze / Schneefernerhaus	Germany	ZSF647N00	47 25 N	10 58 E	2656	SO ₂
Zugspitze / Schneefernerhaus	Germany	ZSF647N00	47 25 N	10 58 E	2656	CH4, CO, CO2, N2O, NO, NO2, NOy, O3, PAN, SF6

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ANTARCTICA						
Arrival Heights	New Zealand	ARH777S00	77 47 S	166 40 E	184	¹³ CH ₄ , CH ₄ , CO, N ₂ O
Arrival Heights	New Zealand	ARH777S00	77 47 S	166 40 E	184	O ₃
Casey Station	Australia	CYA766S00	66 16 S	110 31 E	60	¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Halley Bay	United Kingdom of Great Britain and Northern Ireland	HBA775S00	75 34 S	26 30 W	33	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ VOCs
Jubany	Argentina	JBN762S00	62 13 S	58 40 W	15	CO ₂
Mawson	Australia	MAA767S00	67 37 S	62 52 E	32	¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
McMurdo Station	United States of America	MCM777S00	77 49 S	166 34 E		CH4, O3
Mizuho	Japan	MZH770S00	70 42 S	44 17 E	2230	CH4
Neumayer	Germany	NMY770S00	70 39 S	8 15 W		O ₃
Palmer Station	United States of America	PSA764S00	64 55 S	64 00 W		¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ H ₂ , HCFCs, HFCs, N ₂ O, SF ₆ , VOCs
South Pole	United States of America	SPO789S00	89 58 S	24 48 W	2810	¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
South Pole	United States of America	SPO789S00	89 58 S	24 48 W	2810	¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , C ₂ Cl ₄ , CBrClF ₂ , CBrF ₃ , CCl ₄ , CFCs, CH ₂ Cl ₂ , CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, CH ₄ , CO, CO ₂ , H ₂ , HCFCs, HFCs, N ₂ O, O ₃ , SF ₆ , VOCs
Syowa Station	Japan	SYO769S00	69 00 S	39 34 E	16	CO ₂
Syowa Station	Japan	SYO769800	69 00 S	39 34 E	16	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂ VOCs
Syowa Station	Japan	SYO769S00	69 00 S	39 34 E	16	O ₃
MOBILE STATION						
Aircraft (over Bass Strait and Cape Grim)	Australia	AIA999900				¹³ CO ₂ , CH ₄ , CO, CO ₂ , H ₂ , N ₂ O
Aircraft Observation of Atmospheric trace gases by JMA	Japan	AOA9999900				CH ₄ , CO, CO ₂ , N ₂ O
Aircraft: Orleans	France	ORL999900			150	CH ₄ , CO ₂
Akademik Korolev, R/V	United States of America	AKD999900				CH4
Alligator liberty, M/V	Japan	ALG999900				CO ₂
Atlantic Ocean	United States of America	AOC9XXX00			10	CH4, CO ₂
BACPAC 99	United States of America	BAC999900				CCl ₄ , CFCs, CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, HCFCs
BLAST1	United States of America	BLA999900				CCl ₄ , CFCs, CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, HCFCs
BLAST2	United States of America	BLA999901				CCl ₄ , CFCs, CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, HCFCs
BLAST3	United States of America	BLA999902				CCl ₄ , CFCs, CH ₃ Br, CH ₃ CCl ₃ , CH ₃ Cl, HCFCs

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CLIVAR 01	United States of America	CL1999900				CCl4, CFCs, CH3Br, CH3CCl3, CH3Cl, HCFCs
Comprehensive Observation Network for TRace gases by AIrLiner (CONTRAIL)	Japan	EOM999900				CH4, CO2
Discoverer 1983 & 1984, R/V	United States of America	DIS999900				CH4
Discoverer 1985, R/V	United States of America	DSC999900				CH ₄
Drake Passage	United States of America	DRP999900				¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO ₂
Gas Change Experiment	United States of America	GAS999900				CCl4, CFCs, CH3Br, CH3CCl3, CH3Cl, HCFCs
HATS Ocean Projects	United States of America	HOP999900				HFCs
INSTAC-I (International Strato/Tropospheric Air Chemistry Project)	Japan	INS999900				¹³ CO ₂ , CH ₄ , CO ₂
John Biscoe, R/V	United States of America	JBS999900				CH ₄
Keifu Maru, R/V	Japan	KEF999900				CO ₂
Kofu Maru, R/V	Japan	KOF999900				CO ₂
Korolev, R/V	United States of America	KOR999900				CH4
Long Lines Expedition, R/V	United States of America	LLE9999900				CH4
MRI Research, 1978-1986, R/V	Japan	MRI999900				CH4
MRI Research, Hakuho Maru, R/V	Japan	НКН999900				CO ₂
MRI Research, Kaiyo Maru, R/V	Japan	KIY999900				CO ₂
MRI Research, Mirai, R/V	Japan	MMR999900				CO ₂
MRI Research, Natushima, R/V	Japan	NTU999900				CO ₂
MRI Research, Ryofu Maru, R/V	Japan	RFM999900				CO ₂
MRI Research, Wellington Maru, R/V	Japan	WLT999900				CO ₂
Mexico Naval H-02, R/V	United States of America	MXN999900				CH ₄
NOPACCS - Hakurei Maru -	Japan	HAK999900				TIC
Observation of Atmospheric Chemistry Over Japan	Japan	OAJ999900				CFCs, N ₂ O
Oceanographer, R/V	United States of America	OCE9999900				CH4
PHASE I-04	United States of America	PHA999900				CCl4, CFCs, CH3Br, CH3CCl3, CH3Cl, HCFCs
Pacific Ocean	New Zealand	BSL999900				¹³ CH ₄ , CH ₄ , VOCs

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Pacific Ocean	United States of America	POC9XXX00			10	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Pacific-Atlantic Ocean	United States of America	PAO999900				CH4
Polar Star, R/V	United States of America	PLS999900				CH4
Ryofu Maru, R/V	Japan	RYF999900				CFCs, CH4, CO2, N2O, TIC
Santarem	Brazil	SAN999900				CH4, CO, CO2, N2O, SF6
South China Sea	United States of America	SCS9XXX00			15	¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO, CO ₂ , H ₂
Soyo Maru, R/V	Japan	SOY999900				CO ₂
Surveyor, R/V	United States of America	SUR999900				CH4
The Observation of Atmospheric Methane Over Japan	Japan	OAM999900				CH ₄
The Observation of Atmospheric Sulfur Hexafluoride Over Japan	Japan	OAS9999900				SF ₆
WEST COSMIC - Hakurei Maru No.2 -	Japan	HAK9999901				TIC
Wakataka-Maru	Japan	WAK999900				CO ₂
Western Pacific	United States of America	WPC9XXX00			10	¹³ CH ₄ , ¹³ CO ₂ , C ¹⁸ O ₂ , CH ₄ , CO ₂
northern and western Pacific	Japan	NWP999900				N ₂ O

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Ascension Island St. David's Head			
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Atlantic Ocean			
St. Croix			
Barrow Cold Bay			
Cold Bay Cape Meares			
Discoverer 1983 & 1984,			
R/V			
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GLOSSARY

ATMOSPHERIC SPECIES:

CCl ₄	tetrachloromethane (carbon tetrachloride)
CFC-11	chlorofluorocarbon-11 (trichlorofluoromethane, CCl ₃ F)
CFC-11 CFC-12	chlorofluorocarbon-12 (dichlorodifluoromethane, CCl_3F)
CFC-12 CFC-113	
	chlorofluorocarbon-113 (1,1,2-trichlorotrifluoroethane, CCl ₂ FCClF ₂) chlorofluorocarbons
CFCs	
CH ₃ Cl	chloromethane (methyl chloride)
Halon-1211	chlorodifluorobromomethane (CBrClF ₂)
Halon-1301	bromotrifluoromethane (CBrF ₃)
HCFC-141b	hydrochlorofluorocarbon-141b (1,1-dichloro-1-fluoroethane, CH ₃ CCl ₂ F)
HCFC-142b	hydrochlorofluorocarbon-142b (1,1-difluoro-1-chloroethane, CH ₃ CClF ₂)
HCFC-22	hydrochlorofluorocarbon-22 (chlorodifluoromethane, CHClF ₂)
HCFCs	hydrochlorofluorocarbons
HFC-134a	hydrofluorocarbon-134a (1,1,1,2-tetrafluoroethane, CH ₂ FCF ₃)
HFC-152a	hydrofluorocarbon-152a (1,1-difluoroethane, CHF ₂ CH ₃)
HFCs	hydrofluorocarbons
CH ₄	methane
CH ₃ CCl ₃	trichloroethane (methyl chloroform)
CO	carbon monoxide
CO_2	carbon dioxide
N_2O	nitrous oxide
NO	nitrogen monoxide
NO_2	nitrogen dioxide
NO _X	nitrogen oxides
O ₃	ozone
SF ₆	sulphur hexafluoride
SO_2	sulphur dioxide
UNITS:	
ppb	parts per billion
ppm	parts per million
ppt	parts per trillion
Others:	

ENSO	El Niño-Southern Oscillation
M/V	merchant vessel
R/V	research vessel

LIST OF WMO WDCGG PUBLICATIONS

DATA REPORTING MANUAL:

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March	1999
March	2001
	March March

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WDCGG No.34	March	2010
WDCGG No.35	March	2011

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CD-ROM No. 5	March	1999
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CD-ROM No. 7	March	2001
CD-ROM No. 8	March	2002
CD-ROM No. 9	March	2003

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October	1990	~	August	1992
October	1990	~	August	1992
September	1992	~	March	1993
April	1993	~	September	1993
September	1993	~	March	1994
April	1994	~	December	1994
January	1995	~	June	1995
July	1995	~	December	1995
January	1996	~	June	1996
July	1996	~	November	1996
December	1996	~	June	1997
July	1997	~	December	1997
January	1998	~	June	1998
July	1998	~	December	1998
January	1999	~	June	1999
July	1999	~	December	1999
January	2000	~	June	2000
July	2000	~	December	2000

(period of data accepted)

(period of data accepted)

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October	1990	~	December	1994
October	1990	~	June	1995
October	1990	~	June	1996
October	1990	~	December	1997
October	1990	~	December	1998
October	1990	~	December	1999
October	1990	~	December	2000
October	1990	~	January	2002
October	1990	~	December	2002

CD-ROM No.10	March	2004	October	1990	~	December	2003
CD-ROM No.11	March	2005	October	1990	~	December	2004
CD-ROM No.12	March	2006	October	1990	~	December	2005
CD-ROM No.13	March	2007	October	1990	~	November	2006
CD-ROM No.14	March	2008	October	1990	~	November	2007
CD-ROM No.14	March	2008	October	1990	~	November	2007

WMO WDCGG DVD:

WMO WDCGG DVD:			(period	(period of data accepted)			
DVD No. 1	March	2009	October	1990	~	November	2008
DVD No. 2	March	2010	October	1990	~	November	2009
DVD No. 3	March	2011	October	1990	~	November	2010
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