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Review



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What do we know about the global methane budget? Results from four decades of atmospheric CH₄ observations and the way forward

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Atmospheric CH₄ is arguably the most interesting of the anthropogenically influenced, long-lived greenhouse gases. It has a diverse suite of sources, each presenting its own challenges in quantifying emissions, and while its main sink, atmospheric oxidation initiated by reaction with hydroxyl radical (OH), is well-known, determining the magnitude and trend in this and other smaller sinks remains challenging. Here, we provide an overview of the state of knowledge of the dynamic atmospheric CH₄ budget of sources and sinks determined from measurements of CH₄ and $\delta^{13}C_{CH4}$ in air samples collected predominantly at background air sampling sites. While nearly four decades of direct measurements provide a strong foundation of understanding, large uncertainties in some aspects of

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1. Introduction

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Atmospheric CH₄ absorbs terrestrial infrared radiation in a band at approximately 7.6 μ m, so it affects Earth's radiation balance and is a primary driver of the human impact on climate. Based on measurements of air extracted from ice cores and firn, we know that atmospheric CH₄ increased from approximately 360 ppb (see note about units in electronic supplementary material) during glacial periods to approximately 700 ppb during interglacial periods [1] as receding high-northern latitude glacial ice and changes to monsoon circulation resulted in more wetland area. This change was initiated by small variations in Earth's orbital parameters that affected the seasonal distribution of solar radiation across Earth's surface, but most of the increase in atmospheric CH₄ likely resulted from feedbacks in the carbon cycle [2]. Natural emissions in the southern tropics may have driven a sustained atmospheric CH₄ increase starting 5000 years before present [3]. These Holocene and earlier paleo records demonstrate the strong sensitivity of atmospheric CH₄ to changing climate.

By contrast, the atmospheric CH₄ growth from pre-industrial (approx. 720 ppb) to 2020 (1879.2 \pm 0.6 ppb) was predominantly caused by human activity; it results in 0.52 W m⁻² additional direct heating (see: e.g. https://www.esrl.noaa.gov/gmd/ccgg/ghgpower/, references therein, and electronic supplementary material), behind only CO₂ (with 2.13 W m⁻²). There is an additional approximately 0.3 W m⁻² from indirect effects of CH₄'s atmospheric chemistry [4], specifically the production of tropospheric O₃ and stratospheric H₂O.

Developing sensible mitigation strategies for methane [5] requires a quantitative understanding of individual source and sink terms. Strong climate feedbacks similar to those that drove changes in atmospheric CH_4 abundance in the Holocene have the potential to greatly enhance CH_4 emissions, both in the Arctic, where current warming might release huge stores of carbon (greater than 1 PgC in the top 3 m of permafrost [6]) as CH_4 or CO_2 , depending on hydrology, and also in the tropics, where wetlands, including flooded land used for rice agriculture, and cattle may respond to warmth and rainfall.

2. Observations of atmospheric CH₄ abundance and $\delta^{13}C_{CH4}$ as budget constraints

(a) Experimental methods

Blake *et al.* [7] showed that CH₄ in the surface atmosphere varied spatially and seasonally across the globe. Recognizing the importance of improved understanding of the global CH₄ budget for climate, NOAA began measurements of atmospheric CH₄ from discrete air samples collected in its existing Cooperative Global Air Sampling Network in 1983, expanding available high-quality data both in frequency and spatially [8] to better constrain the global CH₄ budget. Since 1998, $\delta^{13}C_{CH4}$ has been measured in a subset of the same air samples by the Institute of Arctic and Alpine Research (INSTAAR), University of Colorado [9]. Currently, many laboratories around the world monitor atmospheric CH₄ abundance and a few measure $\delta^{13}C_{CH4}$. To help ensure the availability of comparable, high-quality observations for global CH₄ budget studies, these efforts are organized under the umbrella of the World Meteorological Organization (WMO) Global Atmosphere Watch (GAW) program, and data are reported to the World Data Center for Greenhouse Gases hosted by the Japan Meteorological Agency (https://gaw.kishou.go.jp/). Even with this structure in place, there are still 'ease of use' issues; not all laboratories report data in a timely fashion, they report data on different standard scales without providing conversion to the WMO GAW CH₄ mole fraction scale, uncertainties are not reported, etc. Isotopic data are further hindered by a lack of CH₄-in-air reference materials and different methods of tracing $\delta^{13}C_{CH4}$ to calcite reference materials [10]. To circumvent these issues, some research programs have collected data and made packages of observations available (see e.g. https://www.esrl. noaa.gov/gmd/ccgg/obspack/). Future effort is required to ensure high-quality measurements can be traced back to common standard scales, because our understanding of the global CH₄ budget relies on precise spatio-temporal differences from these measurements. The data should also be publicly accessible in a timely manner, fully quality controlled and ready for scientific analysis; data users should not have to add uncertainties nor ensure traceability to a common standard scale.

We do not cover column CH_4 abundances retrieved from measurements of radiance by satellite or surface IR sensors, which are not direct, calibrated measurements of CH_4 abundance and have potential biases similar in magnitude to the large-scale signals we are trying to determine. While they are still useful, especially in the tropics, where direct measurements remain sparse, and in regional studies, they will always require calibrated direct measurements of CH_4 for evaluation.

Here, we focus on NOAA CH₄ and INSTAAR δ¹³C_{CH4} measurements in air samples collected from the Global Monitoring Laboratory's Cooperative Global Air Sampling Network. The network and sampling procedures were described in Dlugokencky et al. [11]. Briefly, weekly discrete whole-air samples are collected in pairs of borosilicate glass flasks with PTFE O-ring seals using a portable air sampler from a globally distributed network of air sampling sites (https://www.esrl.noaa.gov/gmd/ccgg/flask.html). Air samples were analysed for CH4 by gas chromatography with flame ionization detection through mid-2019. Since then, samples have been analysed for CH₄ using a cavity ring-down spectrometer (CRDS). Responses of the GC/FID and CRDS were calibrated for CH₄ abundance against a CH₄ standard scale maintained at NOAA ([12]; this scale serves as the WMO CH₄ mole fraction scale, X2004A). These data provide a spatially and temporally consistent set of observations. INSTAAR began analysing a subset of air samples for $\delta^{13}C_{CH4}$ in 1998. CH₄ from the sample is pre-concentrated, separated from other remaining components in a GC, combusted to CO₂ and the CH₄-derived CO₂ analysed by continuous flow isotope ratio mass spectrometry. CH₄ and $\delta^{13}C_{CH4}$ data are available from NOAA's data server: ftp://aftp.cmdl.noaa.gov/data/trace_gases/xxx/flask/, where xxx = 'ch4' or 'ch4c13'.

(b) Documenting CH₄'s global burden

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Observations of the spatial and temporal distribution of atmospheric CH₄ abundance provide key constraints on the global CH₄ budget. In 2020, preliminary globally averaged CH₄ determined by smoothing data as described below was 1879.2 ± 0.6 ppb, with an increase of 14.8 ± 0.5 ppb from January 1, 2020 to January 1, 2021 (https://gml.noaa.gov/ccgg/trends_ch4/; downloaded 8 July 2021). This implies an imbalance between emissions and sinks in 2020 of 41.0 Tg CH_4 (based on 1 globally averaged ppb = 2.77 Tg CH_4 [13]). The time series of these observations from 1983.5 through 2019 is shown in figure 1*a*. Global (and zonal) means are calculated by first smoothing data from a subset of sampling sites that capture well-mixed, background air, extracting values from the smoothed curve fits at synchronized time steps, then smoothing again as a function of latitude to define a matrix of CH₄ abundance as a function of time (48 values per year) and latitude (intervals of sine (latitude) = 0.05). The quality of the data ensures that the variations in growth rate are significant, as indicated by the uncertainties (dashed lines in figure 1*b*) determined with a combination of resampling of the network (bootstrap; [14]) and Monte Carlo (to assess measurement uncertainty) methods (see electronic supplemenatry material for details).



(c) Long-term changes

From the start of NOAA's measurements in 1983 through 2006, the CH₄ growth rate was decreasing with significant interannual variability (IAV) superimposed on top of that. Then, globally averaged CH₄ began increasing again in 2007.

For 1983–2006, if we assume no trend in the atmospheric CH_4 lifetime, then the atmospheric CH_4 burden was approaching steady state [15]. Starting from a global mass balance and rearranging to solve for emissions we have:

$$E = \frac{d[CH_4]}{dt} + \frac{[CH_4]}{\tau},$$
 (2.1)

where *E* is emissions, [CH₄] is the global annual mean CH₄ burden and d[CH₄]/d*t* its annual increase, both determined from our surface global means, and τ is the atmospheric CH₄ lifetime. After calculating emissions using $\tau = 9.1 \text{ yr}$ [16] and fitting a straight line to emissions from 1984 through 2006, we find no trend in emissions (electronic supplementary material, figure S1; slope = 0.0 ± 0.3 Tg CH₄ yr⁻¹; uncertainty is 68% C.I.). It may seem surprising that atmospheric CH₄ continues to increase while emissions and lifetime are constant, but this is expected from a chemical system with pseudo-first-order loss approaching steady state,

$$[CH_4]_t = [CH_4]_{ss}(1 - \exp[-t/\tau]).$$
(2.2)

Accounting for the atmospheric CH₄ abundance at the start of our measurements gives

$$[CH_4]_t = [CH_4]_{ss} - ([CH_4]_{ss} - [CH_4]_0) \exp[-t/\tau]),$$
(2.3)

where subscripts 'ss' and '0' are for steady state and initial conditions (start of measurements). Equation (2.3) is fitted to the global means from 1983.5 through 2006, solving for $[CH_4]_{ss}$, $([CH_4]_{ss} - [CH_4]_0)$, and τ to give a steady-state value of 1794 ppb and a lifetime of 9.2 yr, in agreement with IPCC's Fifth Assessment Report CH₄ budget lifetime of 9.1 yr [16]. A curve calculated using the coefficients of the fit, extrapolated to 2020, is plotted in figure 1*a* (green), and the difference between it and the deseasonalized trend (green, in ppb) is shown with the growth rate (red) in figure 1*b*.

Since 2007, when this difference starts increasing dramatically, there has been a significant change in the global CH₄ budget that will be explored later. Our conclusion of no trend in



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emissions from 1983 to 2006 is true only if there has been no trend in τ_{CH4} , which is the largest source of uncertainty in this simple global mass balance. For changes in global CH₄ emission rates to be consistent with the observed global means, τ_{CH4} would have had to change in a way that still coincidentally fits this steady-state model.

(d) Interannual variability

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In addition to the long-term changes in atmospheric CH₄, IAV is also evident in the growth rate and the residuals (figure 1*b*), and it has inherent information about budget processes. When the process, or processes, responsible for IAV can be identified, it allows us to quantitatively test our understanding. An example is a large increase in CH₄ growth rate in 1991 (red line in figure 1*b*) and the positive anomaly in the difference between the trend and steady-state fit in 1991 (green line in figure 1*b*). In this smoothed representation, which is similar to a 12-month running mean, the anomaly peaked at approximately 5 ppb; at weekly time resolution, it was approximately 9 ppb (see electronic supplementary material, figure S2). The subsequent decrease in 1992 growth rate is a result of the trend adjusting back to the long-term behaviour as indicated by the difference between the trend and steady-state fit returning to near-zero (green line in figure 1*b*).

Dlugokencky *et al.* [17] attributed this positive anomaly to the impact of the eruption of Mt. Pinatubo on the atmospheric CH₄ sink. SO₂ and ash from the eruption, and subsequent sulfate aerosol produced from the oxidation of SO₂, decreased transmission of UV radiation, which decreased production of O(¹D) and, ultimately, primary production of hydroxyl radical (OH). Qualitatively, this made sense, since anomalies in CO abundance were coincident and consistent in magnitude with the changes in CH₄. Bândă *et al.* [18] investigated this change with a threedimensional global chemistry model and found SO₂ and sulfate reduced the tropospheric CH₄ sink from reaction with OH by 17.8 Tg CH₄ from June, 1991 to June, 1993, which only explained approximately 40% of the change in atmospheric CH₄ burden from June, 1991 to May, 1992. Bândă *et al.* [19] identified additional factors that contributed significantly to changes in atmospheric CH₄ growth rate including: changes in stratospheric O₃, which also affected UV flux; atmospheric temperature, which affected CH₄ emission rates from wetlands and the rate of oxidation of CH₄ (k_{OH+CH4} is strongly temperature-dependent, approximately 2% K⁻¹); biomass burning; and changes in rates of emission of other compounds that affect OH concentration (e.g. CO and NMHCs).

(e) Spatial patterns as budget constraints

Because atmospheric mixing is not instantaneous, signals of sources and sinks exist in the atmospheric CH_4 distribution that are useful for quantifying emissions. Fung *et al.* [13] compared the time-averaged latitude gradient calculated with a chemical transport model (CTM) to NOAA CH_4 observations (electronic supplementary material, figure S3) to rule out specific emission scenarios that were inconsistent with atmospheric observations. They showed, for example, that early estimates of CH_4 emissions from northern wetlands were too large to be consistent with the observed latitude gradient in surface CH_4 .

We exploited the gradient further by analysing temporal changes in spatial patterns to identify changes in the distribution of emissions, although this analysis cannot quantitatively determine the magnitude of the emission changes. Following the economic collapse in the former Soviet Union, Dlugokencky *et al.* [20] showed that a change in inter-polar CH₄ difference (IPD; the difference between zonal averages calculated for 53–90°N and 53–90°S) was consistent with a decrease in CH₄ emissions from the former Soviet Union of approximately 10 Tg between 1991 and 1992. Agreement between observations and a forward simulation with an atmospheric transport model (TM3) using emission estimates from Emissions Database for Global Atmospheric Research (EDGAR) was quite good; both showed a decrease of approximately 12 ppb in annual means calculated for high-northern latitude sites relative to the South Pole between the mid-1980s and the late-1990s (electronic supplementary material, figure S4).

(f) Other tracers: $\delta^{13}C_{CH4}$

We get additional information about processes responsible for emitting CH₄ from other tracers. For example, anomalies in CO abundance measured in the same air samples measured for CH₄ were the right magnitude to be consistent with a decreased atmospheric OH sink after the eruption of Mt. Pinatubo. CO is also used as a tracer for biomass burning, where its molar emission ratio with CH₄ is significantly larger than for the sink process. Ethane (C₂H₆) has been used as a tracer for fossil fuel emissions of CH₄. Both are also emitted from other sources, which can complicate their interpretation. For example, Lan *et al.* [22] analysed long-term observations of CH₄ and a suite of NMHCs from samples collected at a site significantly influenced by oil and gas production. Because emission ratios of C₂H₆/CH₄ from fossil fuel production vary with time, predicting trends in CH₄ enhancements from trends in C₂H₆ (and other NMHCs) enhancements and an assumed constant C₂H₆/CH₄ emission ratio results in significant over-estimates of those trends.

A much better tracer than CO or C_2H_6 to identify processes affecting atmospheric CH₄ is the isotopic composition of CH₄, which is emitted and removed by exactly the same processes as CH₄ itself, though with isotope signatures that vary by source, and fractionation factors that are unique to various sink pathways. Here, we focus on $\delta^{13}C_{CH4}$, which can be used to partition the relative fraction of emissions from broad categories: microbial (mic), fossil (fe; including natural fossil emissions)) and biomass burning (bb). Globally averaged atmospheric $\delta^{13}C_{CH4}$ is equal to the average $\delta^{13}C_{CH4}$ of sources mass-weighted by their emissions (equation (2.4)) and corrected for fractionation by sinks [23] (equation (2.5)).

$$\delta^{13}C_{Q} \cdot Q_{\text{Total}} = \delta^{13}C_{\text{mic}} \cdot Q_{\text{mic}} + \delta^{13}C_{\text{fe}} \cdot Q_{\text{fe}} + \delta^{13}C_{\text{bb}} \cdot Q_{\text{bb}},$$
(2.4)

and, at steady-state,

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$$\delta^{13}C_Q = (\varepsilon + 1)\delta^{13}C_{Atm} + \varepsilon, \qquad (2.5)$$

where the subscript 'Q' refers to sources. An isotopic fractionation factor, α , is defined as the ratio of rate coefficients ($k^{13}C/k^{12}C$) for the reaction of ${}^{13}CH_4$ relative to that for ${}^{12}CH_4$ for a specific sink (e.g. reaction with OH). It is related to ε by $\varepsilon = (\alpha - 1)$ In equation (2.5), ε is weighted by the relative contributions of each sink process and commonly reported in per mille (%). Interpretation of the isotope data requires knowledge of the mass-weighted isotopic composition of the sources that emit CH₄ to the atmosphere. An isotope source signature database compiled by Sherwood et al. [24] greatly increased the sample size and reduced the uncertainty of source signatures for a global CH₄ isotope mass balance. Within each source category, a range of $\delta^{13}C_{CH4}$ values has been observed as a result of systematic spatial differences in these source signatures. These differences help constrain the spatial distribution of CH₄ emissions. This database was complemented by other studies on the spatial patterns in $\delta^{13}C_{CH4}$ of sources ([25] for oil, natural gas, coal, biomass burning and ruminants; [26] for wetlands; [27] for geological seeps; and [28] for ruminants), which can help to better leverage the spatial information in observed $\delta^{13}C_{CH4}$. Adjustment for fractionation by sinks, as with emissions, is weighted by the magnitude of each sink process. Since atmospheric observations of CH₄ isotopes are sensitive to CH₄ sources and sinks with their distinct fingerprints, they are an extremely powerful constraint on the CH4 budget (see the following section for the $\delta^{13}C_{CH4}$ constraints on global CH₄ budget).



Figure 2. (*a*) The same as figure 1, but for 1998 through 2019. (*b*) Globally averaged δ^{13} C(CH₄) calculated from a subset of the samples used in (*a*). The dashed green line is from a model simulation where the OH component of τ CH₄ is adjusted to drive the increase in atmospheric CH₄ burden. (Online version in colour.)

In figure 2, CH₄ abundance (2a) and $\delta^{13}C_{CH4}$ (2b) are plotted. The start of the decline in $\delta^{13}C_{CH4}$ is nearly coincident with the increase in atmospheric CH₄ budget. Unfortunately, INSTAAR measurements of $\delta^{13}C_{CH4}$ started after the large El Niño of 1997/1998, when there were large biomass burning signals, which $\delta^{13}C_{CH4}$ should be very sensitive to. But based on analysis of measurements of CH₄ abundance and $\delta^{13}C_{CH4}$ at Ny Ålesund, Spitzbergen, Morimoto *et al.* [29] found significant enhancements of CH₄ emissions from both wetlands and biomass burning in 1998, highlighting that the interpretation of the observations can be complicated by multiple processes changing nearly simultaneously. From measurements of air extracted from ice cores, we know that $\delta^{13}C_{CH4}$ was increasing for approximately 200 yr [30] before the recent decline began. This suggests a rather significant change in the global budget of CH₄ sources and sinks that will be explored below, although the timing of the change in emissions is difficult to match directly with the $\delta^{13}C_{CH4}$ observations because of the slow response of $\delta^{13}C_{CH4}$ to changes in CH₄ emissions or average source signatures [31].

3. The global CH₄ budget, past and present

Based on measurements of CH₄ abundance from ice cores and firn, and assuming its lifetime was similar to today (approx. 9 yr), pre-industrial emissions were approximately 220 Tg CH₄ yr⁻¹. The dominant source then was emissions from wetlands, but there is substantial uncertainty with regard to emissions from natural fossil sources, which impacts estimates of anthropogenic fossil emissions in today's budget. $\delta^{13}C_{CH4}$ can be used in an isotope mass balance to assess the fraction of total CH₄ emissions from fossil sources [32], but it cannot distinguish between natural seeps and anthropogenic emissions from current fossil fuel exploitation.

A potential tracer for that is ${}^{14}C_{CH4}$ in pre-industrial air extracted from ancient ice, because all ${}^{14}C$ has decayed in fossil CH₄ (lifetime approximately 8000 yr). While the number of observations of ${}^{14}C_{CH4}$ in air extracted from ice cores remains very small (in large part because of the enormous amount of ice needed for sufficient air for a measurement), two studies now indicate that emissions from fossil sources prior to industrialization were small. Petrenko *et al.* [33] found emissions were less than approximately 15.4 Tg CH₄ yr⁻¹ based on ice with air age from 11.6 kyr ago. In a study representative of the atmosphere just prior to industrialization, Hmiel *et al.* [34] reported a fossil component of less than 5.4 Tg CH₄ yr⁻¹. These conclusions were based on very few ${}^{14}C_{CH4}$ measurements, but if we assume natural fossil emissions have not changed since pre-industrial, then there is a large discrepancy between this direct quantification and

aggregating assessments that suggest much greater natural geological fossil emissions with a range of 43–50 Tg CH₄ yr⁻¹ [27]. This large, important discrepancy will only be resolved with far more measurements of ¹⁴C_{CH4} from ancient pre-industrial ice complemented by systematic measurement of fluxes from seeps.

The main sources of CH_4 to the atmosphere today are known, and its budget of emissions and sinks has been summarized periodically by the Global Carbon Project, most recently by Saunois *et al.* [35]. While these budget assessments use *in situ* observations and satellite retrievals of CH_4 , they do not apply the constraints imposed by measurements of CH_4 isotopic composition.

Briefly, from comparison with a pre-industrial atmospheric CH_4 mass balance assuming the same lifetime as today, we know approximately two-thirds of current CH_4 emissions are anthropogenic and come from agriculture (ruminants and their manure, and rice agriculture), waste management (landfills and waste treatment), fossil fuel use (coal, oil and natural gas) and biomass burning (including biofuels). The remaining emissions are from natural processes, predominantly tropical and high-northern latitude wetlands. There remains a great disparity between bottom-up and top-down estimates of natural emissions [35]. If natural fossil emissions are indeed small, as Hmiel *et al.* [34] infer, then to balance the isotopic budget, anthropogenic fossil fuel emissions are correspondingly larger.

The largest atmospheric loss process in the global CH_4 budget is mostly initiated by reaction with OH, especially in the tropical mid-troposphere [36], but also by Cl and O(¹D) (stratosphere only). Oxidation by microbes in soils is likely a small sink, but uncertainty in its magnitude and trend remain large [37].

Atmospheric observations of spatial gradients and their changes with time are used in atmospheric CTM to infer emissions from specific regions. In figure 3, contours of CH₄ zonally averaged growth rate determined from NOAA's observations are plotted; warm colours show where the combined effects of source/sink imbalance and transport result in greater than zero growth rate and cool colours where it is less than zero. Data assimilation systems (also known as inverse modelling) start with an initial distribution of atmospheric CH₄ and a first-guess distribution and magnitude of emissions. This first-guess is used to calculate atmospheric CH₄ at the next model time step after winds and sinks have acted. Based on a comparison of the model and measured CH_4 , the emissions are optimized to get the best fit to the data. North-south gradients in growth rate, to first order, are qualitative indicators of where emissions have likely changed, but to be quantitative, the use of a CTM is absolutely necessary, except perhaps when global totals are estimated. This method has a limitation; it depends on the *a priori* first-guess for emissions. Optimized emissions can only be varied within the uncertainty limits specified for the first-guess, and they are also confined to the prior's spatial distribution. If that first-guess is grossly biased, the assimilation system cannot correct it enough, so emissions from another source are adjusted, perhaps erroneously, to match the observations. Improvements to bottom-up inventories and process models that supply priors would improve top-down assessments of the CH₄ budget.

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The renewed long-term increase in atmospheric CH₄ and near-coincident decrease in $\delta^{13}C_{CH4}$ in 2007 suggests a significant change in the global CH₄ budget with potentially important implications for climate. Many scenarios have been proposed to explain the increase; Lan *et al.* [25] investigated some of them to determine which are consistent with observations of atmospheric CH₄ abundance and $\delta^{13}C_{CH4}$. Most scenarios look at a single process or combination of competing processes acting from 2007 to present, but the reality is likely more complicated than that.

Dlugokencky *et al.* [38] attributed the initial rise to increased CH₄ emissions from tropical wetlands responding to increased precipitation in tropical land regions during an intense La Niña phase of ENSO in 2007 and 2008. But this change was not sufficient to sustain the increase; other factors, some perhaps that had acted over the previous decade must have contributed.

An isotope mass balance for CH_4 requires that emissions from fossil sources (natural seeps and exploitation of fossil fuels) are a larger fraction of total emissions than captured in emission

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Figure 3. Contours of zonally averaged CH₄ growth rate from 2000 through 2019. Use of sine latitude on the y-axis weights each latitude band by its atmospheric mass. (Online version in colour.)

inventories [25,32,39], although with no significant trend [32]. Further studies using $\delta^{13}C_{CH4}$ as a constraint on emission changes after 2006 are not consistent with increased fossil emissions dominating the increase in atmospheric CH₄ burden [25,39]. A large increase in U.S. shale gas production contributing to the increase in global CH₄ burden is not consistent with continental observations of CH₄ abundance [22] nor with the heavy isotopic signature of U.S. shale gas [40].

Another possible explanation of rising methane is that [OH] has had a sustained decline, thereby increasing τ_{CH4} . But this hypothesis is also inconsistent with the decade-long decreasing trend in $\delta^{13}C_{CH4}$. Lan *et al.* [25] used a three-dimensional atmospheric transport model to investigate a scenario where the OH component of τ_{CH4} was adjusted to match the increase in CH₄ burden. The impact on $\delta^{13}C_{CH4}$ through 2016 is shown in figure 2*b* (green). As the proportion of oxidation by non-OH sinks becomes a larger fraction of the total sink, fractionation due to loss increases and the net effect is an increase in $\delta^{13}C_{CH4}$, opposite to what is observed. More generally, there is little evidence for a profoundly decreasing trend in [OH] (see e.g. [41]), but [OH], particularly its trend, remains a large uncertainty in quantifying the global CH₄ budget and, especially, how it is changing.

Lan *et al.* [25] tested competing hypotheses explaining the post-2006 global CH₄ increase in a common three-dimensional tracer model framework. The scenarios most consistent with both CH₄ and $\delta^{13}C_{CH4}$ observations and most plausible in regard to other evidence involve increased emissions from microbial sources (agriculture and wetlands) in the tropics with a small contribution from fossil sources [39,42–44], with a small decrease in biomass burning to maintain isotope mass balance.

4. Uncertainties that limit our understanding of the global CH₄ budget

There are many processes for which our ability to assess IAV and trends is limited, and this imposes uncertainty in assessing atmospheric CH_4 's global budget. Here, we highlight some of the larger uncertainties in our understanding.

(a) Sink processes

Oxidation of atmospheric CH₄ by OH is the largest term in the global CH₄ budget, and small trends, much less than $1\% \text{ yr}^{-1}$, are important. We cannot accurately evaluate magnitudes and trends in global total CH₄ emissions (nor determine which particular sources are changing), without an accurate understanding of the magnitude and trend in [OH]. To date, [OH] and

its trend have been assessed indirectly through observations of 1,1,1-trichloroethane (CH₃CCl₃; common name: methyl chloroform, which we abbreviate to MC), an anthropogenic tracer predominantly removed from the atmosphere by reaction with OH. To be effective, the method required accurate estimates of MC emissions. When MC production was phased out by the Montreal Protocol on Substances that Deplete the Ozone Layer, it allowed Montzka *et al.* [45] to exploit the greatly decreased emissions of MC to estimate [OH] averaged over multiple years based on the rate of decay in the atmospheric burden of MC. However, as the abundance of MC in the atmosphere has decreased closer to its analytical detection limit, this approach has become more sensitive to small remaining emissions. Moreover, while this approach was useful for assessing the lifetimes of CH₄ and MC averaged over some years and their IAV, it was less suitable to assess small trends in [OH].

Recent reports of a decreasing trend in [OH] driving the renewed growth in atmospheric CH₄ since 2007 may be artefacts of the box-modelling methods used ([46] and references therein) and are inconsistent with models of atmospheric chemistry and transport [41]. This scenario is also inconsistent with trends in $\delta^{13}C_{CH4}$ as discussed above. While the reaction rate coefficient for OH + CH₄, including its temperature dependence, has been greatly studied [47], uncertainty in the fractionation of the reaction on atmospheric $\delta^{13}C_{CH4}$ is still substantial, and that uncertainty affects our ability to quantitatively partition emissions among broad source categories. Saueressig *et al.* [48] reported a significantly smaller KIE for fractionation of ¹³C by OH (1.0039 ± 0.0004) than reported by Cantrell *et al.* [49] (1.0054 ± 0.0009). Uncertainty in this factor muddles interpretation of measurements of atmospheric $\delta^{13}C_{CH4}$ imposing a 20% uncertainty on emissions from the fossil fuel sector (approx. 30 Tg CH₄ yr⁻¹) [25].

Oxidation of CH₄ initiated by Cl has little impact on atmospheric CH₄ total loss, but strongly fractionates $\delta^{13}C_{CH4}$. Using a CTM (TOMCAT) and sources of Cl from natural and anthropogenic halocarbons and dechlorination of sea-salt aerosol, Hossaini *et al.* [50] estimated a sink for atmospheric CH₄ by reaction with Cl of 12–13 Tg CH₄ yr⁻¹. Based on analysis of $\delta^{13}C_{CO}$ measurements in the extra-tropical Southern Hemisphere, Gromov *et al.* [51] concluded that the loss of CH₄ by reaction with Cl is no greater than 1% of the total atmospheric CH₄ sink (less than half the estimate of Hossaini *et al.* [50]), and even 1% makes it difficult to balance the ¹³C(CO) budget. Wang *et al.* [52] calculated a similar size sink for CH₄ from reaction with Cl, 5.3 Tg CH₄ yr⁻¹. The impact of Cl on the interpretation of $\delta^{13}C_{CH4}$ observations is still clear; Strode *et al.* [53] compared multiple tropospheric Cl spatial and seasonal distributions and found the choice impacts the north-south gradient and seasonal cycle of $\delta^{13}C_{CH4}$. And Lan *et al.* [25] found that the magnitude of the Cl sink chosen (none or 13 Tg CH₄ yr⁻¹) impacts the amount of fossil emissions that can be consistent with observations by approximately 20%.

Oxidation of CH₄ in soils is intermediate in magnitude, much smaller than loss initiated by reaction with OH and likely much larger than reaction with Cl. Saunois *et al.* [35] reported a bottom-up range for soil loss of 12 to 49 Tg CH₄ yr⁻¹ based on a literature review. Fractionation of CH₄ carbon isotopes by soil loss is reported to be approximately -20%, with a range of measured values of -16 to 27‰ (e.g. [54]). Since the sink mostly occurs in forested soils, which are susceptible to land use, the magnitude [55] and distribution of the sink may be changing. Given that the fluxes of soil loss are very small and occur over large areas of forested and Savannah soils, better estimates of global soil loss will remain challenging.

(b) Wetlands

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CH₄ emissions from wetlands are the largest natural source of atmospheric CH₄. Emissions from wetlands vary greatly over small spatial scales and in time; this variability, combined with relatively few flux measurements, makes up-scaling uncertain. Prior estimates of emissions, therefore, rely on process models, but the range of emissions estimated among different wetland process models is quite large, approximately 40% [56]. One limitation is the accurate assessment of time-varying wetland area and inundation extent. While space-based methods seem ideal to

Using the isotopic constraint effectively requires that the source signatures of different wetland ecosystems are accounted for. Recently, Ganesan *et al.* [26] developed a map of spatially varying $\delta^{13}C_{CH4}$ source signatures for wetlands, which vary by approximately 10% between high-northern latitudes and the tropics. The $\delta^{13}C_{CH4}$ of CH₄ emitted from a wetland will depend on factors such as the mechanism of CH₄ production (CO₂ versus acetate reduction), the source of the carbon (the proportion of plants with C3 versus C4 photosynthetic pathways), and oxidation between production and emission to the atmosphere, which depends on the transport route. Further efforts to identify the underlying processes that control wetland isotopic signature can improve the spatial mapping and thus improve the accuracy of attributing wetland emissions spatially.

5. Conclusion

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Changes to the global CH₄ budget are recorded in the atmosphere in CH₄ abundance and $\delta^{13}C_{CH4}$ spatial and temporal gradients. High-quality, calibrated observations of atmospheric CH₄ and its stable isotopic composition are crucial for decreasing uncertainty in quantifying the evolving global CH₄ budget, yet the world is still greatly under-sampled. If we want to track future changes in the global CH₄ budget, including potential climate feedbacks, we need to expand these observations to under-sampled regions, such as the tropics, Siberia, Africa and South America. Most studies using CH₄ isotope composition as a constraint use stable carbon isotopes, but more extensive observations of stable isotopes of hydrogen would further constrain the budget [39], especially for constraining the OH sink, which has a large fractionation factor for $\delta^2 H_{CH4}$. However, the use of $\delta^2 H_{CH4}$ would require both an extended time series from an adequate global network of air sampling sites and detailed work to define source signatures by type of source and geography.

Explaining the renewed and accelerating increase in atmospheric CH_4 burden since 2007 remains challenging, and the exact causes are not yet clear. But, the observations we describe suggest that increased emissions from microbial sources are the strongest driver, with a relatively smaller contribution from other processes, e.g. fossil fuel exploitation. A more difficult question to answer is the one posed by this special issue: is warming feeding the warming? We cannot say for certain, but we cannot rule out the possibility that climate change is increasing CH_4 emissions. The strong signals from the tropics combined with the isotopic data are consistent with increased emissions from natural wetlands, but large IAV and inter-decadal variability in wetland drivers like precipitation make it difficult to identify small trends. Observations are needed that will help process models capture this variability. The size of the IAV illustrates the potential scope of uncontrollable near-future change and emphasizes the urgency of reducing the global methane burden by mitigating the methane emissions that we can control, from the fossil fuel and agricultural sectors.

Data accessibility. GML and INSTAAR data are available from NOAA's data server: ftp://aftp.cmdl.noaa.gov/data/trace_gases/xxx/flask/, where xxx = 'ch4' or 'ch4c13'.

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