Multidecadal variability of atmospheric methane, 1000–1800 C.E.

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[1] We present a new high-precision, high-resolution record of atmospheric methane from the West Antarctic Ice Sheet (WAIS) Divide ice core covering 1000–1800 C.E., a time period known as the late preindustrial Holocene (LPIH). The results are consistent with previous measurements from the Law Dome ice core, the only other high-resolution record of methane for this time period, and confirm most of the observed variability. Multidecadal variability in methane concentrations throughout the LPIH is weakly correlated or uncorrelated with reconstructions of temperature and precipitation from a variety of geographic regions. Correlations with temperature are dominated by changes in Northern Hemisphere high latitude temperatures between 1400 and 1600 C.E. during the onset of the Little Ice Age. Times of war and plague when large population losses could have reduced anthropogenic emissions are coincident with short periods of decreasing global methane concentrations.


1. Introduction

[2] Atmospheric methane, the second most important greenhouse gas directly impacted by anthropogenic activities, accounts for 18% of the total increase in radiative forcing by all long-lived greenhouse gases and varied in the past on timescales ranging from seasons to hundreds of thousands of years [e.g., Bousquet et al., 2006; Forster et al., 2007; Loulergue et al., 2008]. Systematic direct atmospheric measurements since 1983 from the NOAA (E. J. Dlugokencky et al., Atmospheric methane dry air mole fractions from the NOAA ESRL Carbon Cycle Cooperative Global Air Sampling Network, 1983–2009, Version 2010-08-12, 2010, available at ftp://ftp.cmdl.noaa.gov/ccg/ch4/flask/event/) and GAGE/AGAGE [Cunnold et al., 2002] sampling networks reveal that methane sources and sinks are perturbed by seasonal climate cycles, atmospheric chemistry, large-scale atmospheric patterns such as El Niño Southern Oscillation (ENSO), volcanic eruptions, and anthropogenic activities [Bousquet et al., 2006; Steele et al., 1987]. These direct measurements also documented a surprising decrease in the methane growth rate during the 1990s and a recent resumption of growth [Dlugokencky et al., 2009]. Understanding methane variability on decadal to multidecadal timescales and discriminating between changes in natural and anthropogenic sources is difficult because of spatial and temporal variability of sources and sinks. Furthermore, direct atmospheric records are not yet long enough to assess variability on multidecadal timescales and ice core records have rarely achieved the precision and time resolution needed to observe changes on such short timescales. These limitations have made it difficult to place the decadal-scale variability observed in the record of direct atmospheric measurements in a longer term context.

[3] Air occluded in polar ice sheets provides a unique archive that enables us to extend our record of methane into the past. Accumulation rate (i.e., annual accumulation of snowfall) is one of the primary factors that limit the temporal resolution of ice core paleoatmospheric reconstructions. Long records from low accumulation sites (≤50 kg m⁻² yr⁻¹) such as Vostok and Dome C have revealed orbital-scale variability over the last 800,000 years [e.g., Delmotte et al., 2004; Loulergue et al., 2008] but do not preserve the detail needed to examine short-term variability because diffusion in the firm smoothes the records [Spahni et al., 2003]. The degree of smoothing is much smaller at sites where accumulation rates are high, and the most detailed history of methane over the last few millennia is captured in ice cores from Law Dome in East Antarctica [Etheridge et al., 1998; MacFarling Meure et al., 2006], where the very high accumulation rate (600–1100 kg m⁻² yr⁻¹) allows for large sample sizes yielding precise measurements, high temporal resolution, and excellent chronological control. Furthermore, the Law Dome ice core and firn air records overlap with direct atmospheric measurements and confirm that polar ice faithfully preserves past atmospheric concentrations of methane.
[4] In this study we present a new, precise, decadally resolved ice core methane record spanning 1000–1800 C.E., a period known as the late preindustrial Holocene (LPIH) [Etheridge et al., 1998]. The results come from the West Antarctic Ice Sheet (WAIS) Divide ice core site (core WDC05A). We confirm the magnitude and timing of multidecadal variability observed in the Law Dome methane record over this time period. We also investigate correlations between our methane concentration record and paleoclimate records relevant to methane sources to investigate the processes which likely controlled LPIH methane variations.

2. Methods

2.1. WDC05A Core Recovery

[5] A deep ice core, the WAIS Divide Core (WDC), is located on the West Antarctic Ice Sheet divide at 79.467°S, 112.085°W, surface elevation of 1759 m. The modern annual accumulation rate of \( \sim 200 \pm 34 \) kg m\(^{-2}\) yr\(^{-1}\) [Banta et al., 2008], mean temperature of \(-31^\circ\)C, and the simple ice flow regime at the site are ideal for high-resolution analysis of greenhouse gases over the past \( \sim 110,000 \) years [Morse et al., 2002]. These characteristics are similar to high-resolution deep ice cores in central Greenland (GISP2, GRIP, and NGRIP) simplifying interhemispheric comparisons. The samples used here came from a 298 m shallow core that was extracted in the austral summer of 2005/2006 (WDC05A) 1.3 km northwest of the main borehole (WDC06A). It was drilled with a 10 cm electromechanical drill without drilling fluid and core quality was excellent.

2.2. Analytical Procedures

2.2.1. Methane Measurements

[6] Samples were processed using a wet extraction technique similar to that described by Grachev et al. [2007, 2009] and further elaborated on here. The outer 1–2 mm of ice samples was removed with a band saw in a \(-25^\circ\)C freezer to produce fresh ice that was not recently exposed to the modern atmosphere. This ice was divided in half along the vertical axis to produce a pair of samples with the same depth and age. Each sample had a cross-sectional area of \( \sim 2.5 \) cm\(^2\), height of \( \sim 10 \) cm, and weight of 50–63 g. Samples were weighed to the nearest 0.1 g and placed in precooled cylindrical glass vacuum flasks joined to stainless steel CF flanges with a glass to metal transition, manufactured by Larson Electronic Glass (Redwood City, CA, USA). Each flask was bolted to a stainless steel vacuum line assembly manufactured by Areva. The vacuum line accommodated eight flasks. The flasks were submerged in an ethanol bath maintained at \(-60^\circ\)C to \(-70^\circ\)C and ambient air was removed by pumping with a turbo molecular pump (Alcatel ATP80) backed with a dry (scroll) pump for 1 h. The valves to the flasks were then closed and flasks were submerged in a warm water bath at \(-50^\circ\)C to completely melt the ice and release the gas into the headspace above the water. Melting was usually complete in \( \sim 15 \) min. The flasks were then resubmerged in the ethanol bath for 1 h to freeze the samples and prevent water vapor from interfering with the measurements. The air in the headspace of the flasks was expanded to a 10 cm\(^3\) sample loop on a six-port Valco gas sampling valve. Samples were injected on to a packed column (6 foot, 1/8 inch Hayesep D 80/100 mesh) in an Agilent Technologies 6890N Gas Chromatograph (GC) equipped with a flame ionization detector, using ultrapure nitrogen as a carrier gas. The pressure in the sample loop was measured using an MKS Baratron capacitance manometer (0–100 torr, 0.15% accuracy) installed in the GC oven for temperature stability. The air from each sample was analyzed four times with the pressure of the first expansion \( P_{\text{headspace}} \) typically between 35 and 45 torr and the last 10–20 torr. The peak area over loop pressure ratio was measured for each air sample and compared to a linear regression line fitted to the ratios from a working air standard (500.22 ppb methane on the NOAA04 methane scale [Dlugokencky et al., 2005]) covering the entire range of sample pressures. Concentrations from the four measurements from each sample were then averaged to produce a mean methane concentration for each sample. Mean concentrations were averaged for each pair of samples to produce a mean concentration for each depth/age (Figure 1).

[7] Calibration of daily measurements was maintained throughout the experiment by analyzing our working air standard tank 12 times each day over the pressure range of 10–50 torr. The working air standard tank was a high-pressure cylinder of synthetic air prepared by Scott–Marrin Inc. that was calibrated to primary laboratory standard tanks with concentrations ranging from 380 to 1853 ppb, which were calibrated by the NOAA GMD Carbon Cycle Group on the NOAA04 methane scale [Dlugokencky et al., 2005]. The response of the measurement system is linear to better than 1.5 ppb over the standard tank concentration range. This implies a negligible correction over the sample concentration range so we have assumed a linear response.

[8] Over the course of this study 294 samples from 147 depths were measured. The results from two depths were rejected due to leaks. In addition, we further investigated the 5% of samples (7 depths) with the greatest disagreement between the sample pairs (>5.9 ppb). We measured additional duplicate samples from the same or adjacent depths and in the three depths with the greatest disagreement (10.5–11.8 ppb) we observed that of the four samples measured there was good agreement between three of them and one outlier. In these three cases we suspected that the outlier was a result of contamination or a leak and rejected that pair of samples. Between 21 September and 23 October 2007 we noticed slightly higher methane levels than expected in 34 samples, and traced the contamination to a change in flask cleaning procedures after a personnel change. We detected this contamination when measurement of air–free ice (see below) indicated a contamination of 5–14 ppb. To eliminate the contamination, we cleaned the flasks daily (i.e., after every use) with Alconox® detergent soap, whereas prior to 23 October we had cleaned the flasks weekly. Additional samples from identical or adjacent depths to the contaminated samples were measured after daily cleaning of flasks was implemented. Duplicate measurement of the samples analyzed prior to 21 September and after 23 October 2007 yielded consistent results. All results, including leaks, outliers, and contaminated samples, are shown in Figure 1.
The average standard error of the four analyses for individual samples was 1.4 ppb and the pooled standard deviation between the means of the sample pairs was 1.9 ppb. These statistics exclude the rejected results described above. To quantify the long-term reproducibility of the methane analysis, we measured 16 duplicate pairs of ice samples with the time between measurements ranging from days to months. These duplicates had the same depths as the original samples and should therefore have identical methane concentrations. The pooled standard deviation between the mean of the original and duplicate pairs of samples is 2.8 ppb which is the value we use to represent the long-term analytical uncertainty ($1\sigma$) for this data set. The greater variability observed in these duplicates most likely comes from subtle changes in the solubility of gases in the meltwater, discussed in greater detail below.

2.2.2. Corrections to Measurements

2.2.2.1. Blanks

To constrain the influence of leaks or other contamination in our analysis, we routinely measured air-free ice (AFI). To create AFI, we boiled ultrapure, 18 MΩ water in a cylindrical stainless steel vacuum flask with a Conflat flange seal and metal bellows seal valve welded to the top flange. We affixed a ~30 cm piece of 1/8 inch stainless steel tubing to the outlet, and during boiling the valve remained open. The boiling drives air from the water, which is swept from the chamber by the released steam. We boiled the water for 30 min then sealed the bellows and slowly froze the remaining liquid from the bottom up in an ethanol bath kept at −20°C. Sample preparation and analysis of the artificial ice was identical to ice core samples with the exception that before the artificial ice was melted in the flasks, sufficient standard air from our working air standard tank was added to the flasks so that when expanded to the sample loop it produced pressures equivalent to those from ice core samples. In this way any errors resulting from leaks or contamination of the flasks could be quantified and corrected for. Average AFI corrections were linearly interpolated between days when AFI was analyzed to create a time-dependent correction to the data. The average AFI correction was 1.1 ± 0.5 ($1\sigma$) ppb.

2.2.2. Solubility of Methane in Water

Gases dissolve in liquid water with the partitioning between the air and water described by Henry’s law. Methane is ~2.5X as soluble as nitrogen and therefore the headspace methane concentration decreases when air is exposed to liquid water. As the water is refrozen a small fraction of the air is trapped in the ice leading to a depletion of methane in the headspace after freezing. We express a methane solubility correction factor as

$$\text{Correction Factor} = \frac{[\text{CH}_4]_{\text{Total}}}{[\text{CH}_4]_{\text{Headspace}}}$$

Figure 1. Atmospheric methane concentrations over the past 2 centuries from the Law Dome compilation (0–1995 C.E., black line) [Etheridge et al., 1998; MacFarling Meure et al., 2006], WDC05A on the WDC05A:2 chronology (1019–1814 C.E., blue line), and direct atmospheric measurements from the South Pole (1983–2010 C.E., red line) (E. J. Dlugokencky et al. Atmospheric methane dry air mole fractions from the NOAA ESRL Carbon Cycle Cooperative Global Air Sampling Network, 1983–2006, version 2007-09-19, report, 2007). Inset shows methane data from WDC05A on the WDC05A:2 chronology with the mean at each depth/age (blue line), individual measurements (light blue circles), leaks (red diamonds), outliers (pink squares), and contaminated measurements (brown triangles). All data are plotted on the NOAA04 calibration scale [Dlugokencky et al., 2005].
where \([\text{CH}_4]_{\text{Headspace}}\) is the concentration in the headspace which we measure during our typical analysis and \([\text{CH}_4]_{\text{Total}}\) can be expressed as the molar ratio

\[
[\text{CH}_4]_{\text{Total}} = \frac{[\text{CH}_4]_{\text{Headspace}}}{[\text{Ar} + \text{O}_2 + \text{N}_2 + \text{CH}_4]_{\text{Headspace}}}
\]

where \([\text{CH}_4]_{\text{Headspace}}\) and \([\text{Ar} + \text{O}_2 + \text{N}_2 + \text{CH}_4]_{\text{Headspace}}\) are the total number of moles that were in the original ice sample. Inclusion of trace atmospheric gases has a negligible effect and was ignored. At equilibrium the distribution of air constituents between the headspace and the meltwater is dictated by Henry’s law constants at 273.2 K [Fogg and Sangster, 2003] because the majority of dissolution happens as bubbles escape from the melting ice sample and rise through the meltwater next to it. Using typical values for flask and ice volumes, the headspace air would have a methane concentration depleted by 2.1–2.2% relative to the original concentration. For example, a sample which has a measured \([\text{CH}_4]_{\text{Headspace}}\) of 700 ppb, 60 mL of water in a 134 cm³ flask, at 273.2 K, would have a \([\text{CH}_4]_{\text{Total}}\) of 715 ppb and therefore a correction factor of 1.021.

[12] We examined this correction factor empirically by measuring the concentration of methane in the frozen sample water (\([\text{CH}_4]_{\text{Refreeze}}\)) after our typical analysis was completed for 32 samples. After the initial melt-refreeze-\(\text{CH}_4\) measurement was completed we evacuated the headspace for 1 h then another melt-refreeze-\(\text{CH}_4\) measurement was done. \([\text{CH}_4]_{\text{Total}}\) was then calculated as follows:

\[
[\text{CH}_4]_{\text{Total}} = \frac{[\text{CH}_4]_{\text{Headspace}} [\text{P}_{\text{Headspace}}] + [\text{CH}_4]_{\text{Refreeze}} [\text{P}_{\text{Refreeze}}]}{[\text{P}_{\text{Headspace}}] + [\text{P}_{\text{Refreeze}}]}
\]

where \(P\) is pressure in torr. Mean and standard deviations (1σ) from the 32 measurements were \(P_{\text{Headspace}} = 0.50 \pm 0.06\) torr and \([\text{CH}_4]_{\text{Refreeze}} = 1698 \pm 169\) ppb. The correction factor from these experiments is therefore 1.0170 ± 0.0031%. Since \(P_{\text{Refreeze}}\) is ~1% of \(P_{\text{Headspace}}\), we expect the amount gas trapped in the ice during the second melt-refreeze cycle to be ~1% of \(P_{\text{Refreeze}}\), or ~0.005 torr, and therefore negligible.

[13] The uncertainty in the correction factor is probably caused by a variety of subtle differences between samples such as sample size, air content, and the refreezing rate. This uncertainty is inherently incorporated into our estimate of the long-term uncertainty of our measurements because it is derived from duplicate measurements from different days and should therefore represent the full range of possible variability. Given these considerations we have increased our final methane concentration values by the average empirically derived solubility correction factor of 1.0170.

[14] The difference between the calculated and empirical correction factors is likely caused by the headspace air not reaching a solubility equilibrium with the meltwater when the samples are refrozen. This has been observed in greater detail in other studies with much larger samples [Petrenko et al., 2008]. The values given above indicate that the samples reach ~80% of solubility equilibrium, presumably because of exclusion of gases during freezing. Slower refreezing may reduce the effect at the expense of longer processing times.

### 2.2.2.3. Gravitational Fractionation

[15] Gases within the firm undergo mass dependant fractionation due to gravity [Craig et al., 1988; Schwander, 1989; Schwander et al., 1997; Sowers et al., 1989]. The magnitude of gravitational fractionation is controlled by the thickness of the diffusive air column in the firm and can be estimated by measuring the \(^{15}\text{N}/^{14}\text{N}\) ratio of \(\text{N}_2\) and reported using standard delta notation as \(\delta^{15}\text{N}_2\). Because the turnover time of the atmospheric \(\text{N}_2\) reservoir is longer than a million years, we can assume that the \(^{15}\text{N}/^{14}\text{N}\) ratio of atmospheric \(\text{N}_2\) has remained constant over ice core timescales [Sowers et al., 1992]. Measurement of \(\delta^{15}\text{N}_2\) along the main core (WDC06A) at WAIS Divide (100–300 m) revealed \(\delta^{15}\text{N}\) values of 0.303 ± 0.006‰ (J. Severinghaus, personal communication, 2010). Gravitational fractionation of methane concentrations results from the mass difference between methane (\(M = 16.04\) g mol⁻¹) and dry air (\(M = 28.96\) g mol⁻¹). The gravitational correction is therefore \(\Delta M \times \delta^{15}\text{N}_2\) where \(\Delta M = 12.92\) g mol⁻¹. This results in a gravitational fractionation correction (increase of the measured concentration) of 0.39% which we have applied to all of our ice core measurements. The \(\delta^{15}\text{N}_2\) is not expected to be significantly different between the WDC05A and WDC06A cores.

### 2.3. WDC05A Chronology

[16] Convective and diffusive processes move air through the firm faster than the annual accumulation of ice causing air to be younger than the surrounding ice at a given depth within the firm [Schwander, 1989]. The age difference between the air and the enclosing ice is termed “delta age” (\(\Delta\)age). The original WDC05A chronology [Mischler et al., 2009], was based on annual layer counting of the non-sea-salt sulfur to sodium ratio (nssS/Na) between the surface and 70 m [Banta et al., 2008] and alternative current electric conductivity measurement (ACECM) measurements from 70 to 298 m to establish the age of the ice. A 1-D firm air diffusion model following Battleg et al. [1996] and Trudinger et al. [1997] was used to estimate the mean age of the air at the lock in depth (LID) which is then subtracted from the age of the ice at the LID to estimate \(\Delta\)age [Mischler et al., 2009]. The ice chronology from 70 to 298 m was recently improved by optimizing correlations between the monthly resolved mineral acidity measurements from the main borehole (WDC06A) which are dominated by fallout from volcanic emissions and the WDC05A ACECM measurements. Annual layer counting of the high-resolution chemical records from the upper 70 m of WDC05A and WDC06A from ~1300 C.E. to ~2000 C.E. was confirmed by comparison with the volcanic sequence of nssS from Law Dome, adapted from Palmer et al. [2001]. This generated 71 unique tie points between the WDC05A and WDC06A records. The annual accumulation rate from WDC06A was then mapped onto WDC05A assuming a linear change between tie points. To evaluate the uncertainty in the estimated ice age for WDC05A, we compared the estimated ice ages in the upper 70 m with those determined from high-resolution chemistry measurements on the same core. For the 231 years common to both records, the average difference between the two depth-age scales is 0.056 mweq (water equivalent), with a maximum difference of 0.283 mweq. The mean annual accumulation rate is ~0.20 mweq, so the mean difference is 3–4 months. The maximum difference corresponds to
1.4 years and the standard deviation between the estimated and observed depth-age scales is ∼0.07 mweq or ∼4 months. The difference between our final ice chronology (WDC05A:2) and the original ice chronology (WDC05A:1) [Mischler et al., 2009] is negligible between 0 and 70 m, increasing to 13 years at 148 m, and then decreasing to ∼36 years at 298 m.

A visible volcanic ash layer observed in both cores was also used to confirm the dating of the two cores. The ∼5 mm volcanic ash layer was observed at 190.83 m (1248.1 C.E.) in WDC05A and 190.39 m (1248.5 C.E.) in WDC06A (ages determined using the revised chronologies). Electron microprobe analysis indicates that the chemical composition, grain size and particle morphology of the two layers are nearly indistinguishable, indicating that both are from the same eruption and therefore deposited contemporaneously (N. Dunbar, personal communication, 2010). The ash layer has abundant ash particles and is relatively coarse grained with particles up to 20 μm suggesting that it has an Antarctic source. Positive chemical correlations with ash layers in the Siple Dome and Taylor Dome ice cores indicate that this ash layer is a major regional time stratigraphic marker [Dunbar et al., 2003, 2007].

A 1-D firn air diffusion model estimated that the mean age of the air within the open porosity at the LID (65.5 m at WDC05A) is 9.9 years for CO2 and 7.2 years for CH4 (M. Battle, personal communication, 2009). The difference between the mean age of CH4 and CO2 arises from the different diffusivities of the two gases in the firn air. The width of the CH4 age distribution at half height is 5.9 years. The age of the ice at the LID is 215 years which makes Δage 205.1 years for CO2 and 207.8 years for CH4. Since temperature (E. Steig, personal communication, 2010) and accumulation at WDC05A have remained relatively constant in the LPIH we do not expect that Δage has changed significantly and for the purpose of creating a chronology we have held it constant. To construct the final gas chronology, we have subtracted 207.8 years from the ice chronology. The estimated uncertainty of the chronology is ±10 years based on a detailed comparison between our record and the Law Dome methane record, discussed below. This chronology is designated WDC05A:2.

3. Results and Discussion

3.1. Comparison With Previous High-Resolution Antarctic Data From Law Dome

Replication of paleoclimate records is an important means of verifying their reliability. The methane record from Law Dome, Antarctica, a well-known data set covering the last 2000 years, is a compilation of data from three different ice cores (DSS, DE08, DE08–2) and until now has been the only high-resolution, high-precision record covering the past 1000 years [Etheridge et al., 1998; MacFarling Meure et al., 2006]. The Law Dome record from 1000 to 1800 C.E. comes from the DSS ice core and the data are plotted with the WDC05A results in Figure 2, after conversion of Law Dome results to the NOAA04 calibration scale [Dlugokencky et al., 2005]. Error bands are ±2.8 ppb for WDC05A, ±5 ppb from Etheridge et al. [1998], and ±4.1 ppb from MacFarling Meure et al. [2006]. The Law Dome data were produced using a dry extraction technique with large samples (500–1500 g) while we use a wet extraction technique with small samples (two samples, ∼60 g each) which requires a correction for solubility effects (discussed above).

Law Dome is located at 66.733°S, 112.833°E, with a surface elevation of 1390 m and is over 3500 km away from the WAIS Divide site. Since there are essentially no sources of methane in the high latitude Southern Hemisphere, the atmospheric concentration around Antarctica is homogenous [Dlugokencky et al., 1994] and should have been so in the past. The absolute methane concentrations in both records are very consistent (Figure 2). The only time interval where the records differ beyond the 1-s envelope of analytical and temporal uncertainty is 1410–1470 C.E. where they diverge by 10–15 ppb. This is within the 2-σ level and despite the
divergence both records show a peak in concentrations during this time period. We observe that the amplitude of high-precision variability in the WDC05A core is ~10–20% smaller than in Law Dome. The exact mechanisms behind the slight amplitude reduction are not currently known but we speculate that it is a result of greater mixing of air within the lock in zone during the bubble closure process, which is longer at WDC05A because of the lower accumulation rate.

[21] The data from WDC05A and Law Dome show that atmospheric methane in the high latitude southern hemisphere averaged ~690 ppb (NOAA04 scale) and experienced multidecadal variability during the LPIH. During the 18th century anthropogenic activities increased methane emissions and caused global concentrations to increase rapidly [Etheridge et al., 1998; MacFarling Meure et al., 2006]. The general trends in methane concentration during the 19th and 20th centuries have been described in greater detail elsewhere and preliminary measurements from WDC05A (not shown) support those previous conclusions [Etheridge et al., 1998; MacFarling Meure et al., 2006]. Here we focus on a detailed comparison with previously published high-resolution records of methane during the LPIH.

[22] A statistical comparison of the two methane records shows a high degree of correlation (r = 0.87 for linear regression). The WDC05A record appears to slightly lag the Law Dome record. A maximum correlation (r = 0.91) between the records is obtained by shifting the WDC05A record 9 years older. While it is not possible to determine which record is “correct,” the highest correlation occurs within the stated uncertainty of ±10 years for the WDC05A gas chronology [Mischler et al., 2009], which is largely due to uncertainty in Δage. One possible explanation for the offset is that the firn air model used by Mischler et al. [2009] to deduce Δage for WDC05A assumes that all bubbles close off below the LID in the WAIS Divide ice core, but the large accumulation rate at this site reduces the magnitude of this effect [Trudinger et al., 1997]. Qualitatively, bubble closure above the LID combined with a low to moderate accumulation rate would decrease Δage and cause additional smoothing of the gas records, consistent with our observations.

[23] Alignment of large rapid changes in methane concentrations have been used to establish chronostatigraphic tie points between ice cores in the past because methane has a relatively short lifetime and variations recorded in polar ice cores are expected to represent global signals [Blunier and Brook, 2001; Brook et al., 2005; EPICA Community Members, 2006; Lemieux-Dudon et al., 2010]. This approach has generally not yet been utilized for small-scale variations (~50 ppb) because methane records have, until now, lacked sufficient temporal resolution and analytical precision to uniquely identify small-scale variability. The high degree of correlation between the Law Dome and WDC05A methane records demonstrate that this technique could be a viable way to establish chronostatigraphic tie points between high-precision methane records from moderate to high accumulation rates.

3.2. Multidecadal Methane Variability From 1000 to 1800 C.E.

3.2.1. Implications for the Global Methane Budget

[24] The global methane budget can be expressed in its simplest form as dB/dt = S − B/τ, where B is the total atmospheric burden (Tg of CH₄), S is the total source in Tg yr⁻¹, and τ is the lifetime of CH₄ in years. The average LPIH methane concentration from WDC05A is ~690 ppb and the interpolated gradient is 43 ± 5 ppb based on preliminary measurements from the Greenland ice core GISP2D (not shown) [Mitchell and Brook, 2009]. The global mean atmospheric concentration, weighted by surface area, has been estimated for the LPIH as the Antarctic concentration plus 37% of the interpolar gradient [Etheridge et al., 1998] and is therefore ~706 ppb from our data. Using the total mass of the dry atmosphere (5.1352 ± 0.0003 × 10¹⁸ kg [Trenberth and Smith, 2005]) the global average atmospheric burden is calculated to be ~2008 Tg (2.844 Tg CH₄ ppb⁻¹) for the LPIH at steady state (dB/dt = 0). Numerous chemical modeling studies have tried to determine likely values for S and τ for the LPIH, but these are sensitive to the concentration of OH, the primary sink for CH₄, and its interaction with the CH₄-CO-NOX chemical system. These models have produced estimates of LPIH τ that range from 17% lower to 16% higher than the present-day range of 8.9–9.2 years [Dentener et al., 2003; Harder et al., 2007; Martinerie et al., 1995; Prinn et al., 2001; Shindell et al., 2003]. For the purposes of the discussion here we choose a LPIH τ of 8 years, implying a steady state flux of ~250 Tg CH₄ yr⁻¹.

[25] We estimated the methane growth rate by linearly interpolating the WDC05A data annually, then determining the annual rate of change. This time series was smoothed with a 30 year Gaussian filter (Figure 3). The most negative growth rates occurred in the early 13th and late 16th centuries while the most positive growth rates occurred at the turn of the 16th century. The interannual variability of methane growth rates may have been greater given that the modern record of direct atmospheric measurements since 1983 record which shows variations of up to ±15 ppb CH₄ yr⁻¹ [Bousquet et al., 2006; Dlugokencky et al., 2009], but such short-term changes would not be captured by the ice core record. The overall decrease of ~10 ppb yr⁻¹ over this
Table 1. LPIH Methane Sources

<table>
<thead>
<tr>
<th>Global Methane Sources</th>
<th>Methane (Tg CH₄ yr⁻¹)</th>
<th>Percent of Total</th>
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<tr>
<td>Animals</td>
<td>18.0</td>
<td>7.2</td>
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<tr>
<td>Termites</td>
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<td>7.7</td>
</tr>
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<td>Ocean</td>
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<td>5.2</td>
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<tr>
<td>Fresh water lakes</td>
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<tr>
<td>Total</td>
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<td>100</td>
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</table>

*LPIH methane sources are after Harder et al. [2007] with the estimate of anthropogenic source after Houweling et al. [2000]; scaled to a total of 250 Tg CH₄ yr⁻¹. See Table 3 for alternate estimates of anthropogenic sources.

It has been common in the ice core literature to interpret past variations as indicators of changes in climate driven methane sources, primarily wetlands, although most work has recognized the possibility that the methane sink can change [Fischer et al., 2008; Kaplan et al., 2006; Valdes et al., 2005]. The wetland centric view is supported by a number of model studies that appear to show that the methane sink has changed relatively little despite the large changes in the global methane burden and sink between preindustrial and modern times [Crutzen and Brühl, 1993; Lelieveld et al., 1998; Martinerie et al., 1995; Shindell et al., 2003; Thompson, 1992; Wang and Jacob, 1998]. Measurements of methyl chloroform (MCF; CH₃CCl₃) since 1978 are used to determine modern OH concentrations. These data show that while there can be significant interannual variability in OH, the longer term trend is small [Bousquet et al., 2005], and rising methane levels. This supports the contention that OH, the main sink for CH₄, has been relatively stable in despite rising methane levels. This supports the contention that OH, the main sink for CH₄, has been relatively stable in.

3.2.2. Methane Source Variations

If we assume that the methane sink has remained relatively constant then the observed multidecadal variability would be a result of variations in source strength. Methane is primarily produced by the anaerobic decomposition of organic material by archaea (see Khalil [2000] for a review). To obtain a source distribution for the LPIH, we combined the “Holocene base” scenario from Harder et al. [2007], which estimates natural methane sources (~230 Tg CH₄ yr⁻¹) with the anthropogenic source estimate (~30 Tg CH₄ yr⁻¹) from Houweling et al. [2000], then scaled the total source to 250 Tg CH₄ yr⁻¹, as shown in Table 1. The magnitude of variability observed in the WDC05A record is ~10–34 ppb, equivalent to 4–12 Tg CH₄ yr⁻¹, or 1.4–4.8% of the total budget (Figure 2). As mentioned previously, ice cores record a smoothed history of atmospheric methane with the degree of smoothing being dependant on the characteristics of the ice core site. The Law Dome record appears to have recorded ~10–20% greater variability than WDC05A, so our record provides a constraint on the minimum amount of variability possible. The variability in the WDC05A record is similar in magnitude to each of the individual nonwetland sources (Table 1). Since it is unlikely that the multidecadal variability comes from very large changes in the smaller budget terms, the most likely explanation is that the variability comes from emissions from wetlands which, given adequate carbonaceous substrate, are predominantly influenced by water table depth and soil temperature [e.g., Allen et al., 2003; Bloom et al., 2010; Christensen et al., 2004; Matthews, 2000; van Hulzen et al., 1999; Walter et al., 2001a, 2001b; Wang et al., 2009; Worthy et al., 2000; Zona et al., 2009]. Sufficient water is required to produce anoxic conditions that are a prerequisite for methanogenesis. Once anoxic conditions are present, increasing temperatures lead to higher emissions with a maximum growth temperature of 37°C–45°C [Boone, 2000]. Temperature and precipitation controls on methanogenesis operate on subannual timescales, so changes in these climatic variables have an immediate impact on annual emissions. The record of methane emissions is then smoothed by the atmosphere and by the firm before being trapped in polar ice sheets. Wetland emissions are thus controlled by temperature and precipitation changes and respond quickly (on subannual timescales) to changes in these variables, so we would expect the ice core methane record to be correlated with temperature and precipitation changes on multidecadal timescales.

In sections 3.2.2.1 and 3.2.2.2 we examine LPIH temperature and precipitation records as well as estimates of anthropogenic emissions for relationships with the WDC05A methane record. Our approach is guided by modeling studies and satellite measurements which show that precipitation exerts a dominant control on tropical (30°S–30°N) methane emission variability through its influence on water table depth and interannual OH concentrations, whereas temperature is the dominant factor in high latitude northern hemisphere (30°N–90°N) variability [Bekki and Law, 1997; Bloom et al., 2010; Bousquet et al., 2006; Khalil and Rasmussen, 1983; Walter et al., 2001a]. Prior to comparison, the paleoclimate records and the methane record were smoothed with a band-pass filter removing periods shorter than 20 years and longer than 500 years, removing variability both higher than the Nyquist frequency of the ice core record and lower than multicentennial frequencies due to slow changes in forcing (e.g., gradual cooling between the Medieval Warm Period and the Little Ice Age), isolating variability on multidecadal timescales. To calculate correlations and their statistical significance (p < 0.05, null hypothesis that r = 0), the smoothed paleoclimate records were then subsampled to match the ages of our methane data points (N = 89).

3.2.2.1. Temperature

Previous studies have documented the striking correlation between the oxygen isotope record of ice (δ¹⁸Oice, a
Figure 4. Comparison between WDC05A methane concentrations on the WDC05A:2 chronology and temperature reconstructions. Greenland: orange, dotted [Johnsen et al., 1997]; light blue, dash-dotted [Vinther et al., 2006]; dark blue, solid [Kobashi et al., 2010]. Extratropical land: green, dotted [D’Arrigo et al., 2006]; light red, dash-dotted [Esper et al., 2002]; dark red, solid [Kaufman et al., 2009]. N. Hemisphere: pink, dotted [Mann et al., 2008]; purple, dash-dotted [Moberg et al., 2005]; brown, solid [Hegerl et al., 2007]. Tropical: blue, dash-dotted [Oppo et al., 2009]; black, solid [Black et al., 2007]. All records were smoothed with a band-pass filter with a period of 20–500 years.

proxy for local temperature) from Greenland ice cores and methane during the past glacial cycle [e.g., Brook et al., 2000; Chappellaz et al., 1993; Huber et al., 2006; Severinghaus and Brook, 1999; Severinghaus et al., 1998]. See Table 2 and Figure 4 for comparisons between methane and temperature proxies. The correlation coefficient between our methane record and the GRIP [Johnsen et al., 1997] δ18Oice record is statistically significant, but not high (r = 0.24, p = 0.03). The correlation coefficient is even lower and not statistically significant with the NGRIP [Vinther et al., 2006] δ18Oice record (r = 0.06, p = 0.56). A record of temperature reconstructed from δ40Ar and δ15N isotopes from GISP2 [Kobashi et al., 2010] also does not have a statistically significant correlation with methane (r = 0.18, p = 0.10). Sliding the chronologies of these records relative to our methane record by ±50 years does not greatly increase the correlation coefficients. We infer from this analysis that the close relationship between methane and large temperature changes during the last ice age apparently does not extend to the very small temperature variability in Greenland during the LPIH, at least to the extent that these proxies are actually recording site temperature.

[30] We have also compared our record to three northern hemispheric land temperature reconstructions [Mann et al., 2008] EIV Land (r = −0.10, p = 0.37) [Moberg et al., 2005] (r = −0.13, p = 0.23) [Hegerl et al., 2007] (r = 0.03, p = 0.81) (Table 2 and Figure 4). The weakly negative correlation coefficients and lack of statistical significance indicates that hemispheric temperature variability during the LPIH did not directly control global methane concentrations. Hemispheric to global temperature reconstructions have been used to scale methane emissions in model reconstructions for the late Holocene using the argument that emissions are temperature sensitive and because of a lack of other constraints [Houweling et al., 2008]. Our results suggest that this approach will not yield accurate results.

[31] Local- to regional-scale temperature reconstructions that are specific to methane source regions might be expected to have a greater correlation with global methane concentrations. Analysis of modern methane emissions indicates that Northern Hemisphere extratropical wetlands are more sensitive to temperature than tropical wetlands [Bloom et al., 2010]. We examined two extratropical Northern Hemisphere temperature reconstructions [D’Arrigo et al., 2006] (r = 0.04, p = 0.73) and [Esper et al., 2002] (r = 0.16, p = 0.13) that utilize similar data sets and found that they both have low correlation coefficients with methane that lack statistical significance. A multiproxy Arctic temperature reconstruction [Kaufman et al., 2009] (r = 0.24, p = 0.03) has a low but statistically significant correlation with methane which increases to r = 0.34 if the record is interpolated annually and shifted by −30 years, within the uncertainty in their chronology (~2–10%).

[32] We also examine correlation coefficients with tropical sea surface temperature records from the Indo Pacific Warm Pool (IPWP) [Oppo et al., 2009] (r = 0.26, p = 0.01) and Cariaco Basin [Black et al., 2007] (r = 0.35, p < 0.01) and find statistically significant correlation coefficients that are slightly higher than with other temperature reconstructions. The record from the Cariaco basin has a high correlation (r = 0.77) when the record is interpolated annually and shifted forward in time by 52 years. The magnitude of this
Table 2. Linear Correlation Coefficients (r) Between Temperature Reconstructions, Precipitation Proxy Records, and the WDC05A Methane Record

<table>
<thead>
<tr>
<th>Source</th>
<th>Region</th>
<th>r</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Temperature Proxy Records and Reconstructions</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Johnsen et al. [1997]</td>
<td>Greenland (GRIP)</td>
<td>0.24</td>
<td>0.03</td>
</tr>
<tr>
<td>Vinther et al. [2006]</td>
<td>Greenland (NGRIP)</td>
<td>0.06</td>
<td>0.56</td>
</tr>
<tr>
<td>Kobashi et al. [2010]</td>
<td>Greenland (GISP2)</td>
<td>0.18</td>
<td>0.10</td>
</tr>
<tr>
<td>D’Arrigo et al. [2006]</td>
<td>Extratropical N.H. 40°N-90°N</td>
<td>0.04</td>
<td>0.73</td>
</tr>
<tr>
<td>Esper et al. [2002]</td>
<td>Extratropical N.H. 30°N-90°N</td>
<td>0.16</td>
<td>0.13</td>
</tr>
<tr>
<td>Kaufman et al. [2009]</td>
<td>Extratropical N.H. 60°N-90°N</td>
<td>0.24</td>
<td>0.03</td>
</tr>
<tr>
<td>Mann et al. [2008] EVI Land</td>
<td>Northern Hemisphere</td>
<td>−0.10</td>
<td>0.37</td>
</tr>
<tr>
<td>Moberg et al. [2005]</td>
<td>Northern Hemisphere</td>
<td>−0.13</td>
<td>0.23</td>
</tr>
<tr>
<td>Hegerl et al. [2007]</td>
<td>Northern Hemisphere</td>
<td>0.03</td>
<td>0.81</td>
</tr>
<tr>
<td>Oppo et al. [2009]</td>
<td>Tropical SST (Indo-Pacific Warm Pool)</td>
<td>0.26</td>
<td>0.01</td>
</tr>
<tr>
<td>Black et al. [2007]</td>
<td>Tropical SST (Cariaco Basin)</td>
<td>0.35</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Trouet et al. [2009]</td>
<td>North Atlantic Oscillation</td>
<td>0.03</td>
<td>0.76</td>
</tr>
<tr>
<td>MacDonald and Case [2005]</td>
<td>Pacific Decadal Oscillation</td>
<td>0.35</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td><strong>Precipitation Proxy Records</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hu et al. [2008]</td>
<td>China Speleothem δ¹⁸O (Heshang)</td>
<td>0.24</td>
<td>0.02</td>
</tr>
<tr>
<td>Wang et al. [2005]</td>
<td>China Speleothem δ¹⁸O (Dongge)</td>
<td>0.02</td>
<td>0.88</td>
</tr>
<tr>
<td>Zhang et al. [2008]</td>
<td>China Speleothem δ¹⁸O (Wanxiang)</td>
<td>0.28</td>
<td>0.01</td>
</tr>
<tr>
<td>Reuter et al. [2009]</td>
<td>Peru Speleothem δ¹⁸O (Cascayunga)</td>
<td>−0.19</td>
<td>0.09</td>
</tr>
</tbody>
</table>

*All records were smoothed with a band-pass filter removing periods shorter than 20 years and longer than 500 years and then subsampled to match the ages of the WDC05A methane data (N = 89). Correlation coefficients are statistically significant when p < 0.05 (null hypothesis is that r = 0).
timescales La Niña conditions (when SSTs are anomalously cold in this region) are associated with greater precipitation over tropical land areas and greater tropical methane emissions [Dlugokencky et al., 2009; Gu et al., 2007]. We also examined a proxy for the North Atlantic Oscillation (NAO) [Trouet et al., 2009] (r = 0.03, p = 0.76); however, this proxy has no correlation with methane on multidecadal timescales.

[35] The positive correlations we observe between temperature reconstructions in some regions and the methane record appear to be driven in large part by temperature variations in the latter part of the record, particularly between 1400 and 1600 C.E. The largest feature in our methane record for the LPIH is a large increase from 1470 to 1520 and a subsequent decrease from 1560 to 1600 C.E. A similar feature is seen in many temperature reconstructions; however, the temperature decline is seen most prominently in records from higher latitudes: Sweden [Grudd et al., 2002]; north (55°–70°N) and eastern hemisphere regions [Cook et al., 2004]; northern Siberia [Briffa et al., 2001]; Yukon, Central Northwest Territories, Jaamtland, Tromntraesk, and Mongolia [D’Arrigo et al., 2006]; modeled Arctic temperatures calibrated to Arctic temperature proxies [Crespin et al., 2009]. The fall in methane is also coincident with the start of the “classical” climatological Little Ice Age (LIA) [Matthews and Briffa, 2005]. Using the temperature field reconstruction discussed above [Mann et al., 2009], we used the annually interpolated records to calculate correlation coefficients in 200 year moving windows and found the highest correlation during the time period 1400–1600 C.E. (Figure 5b). During this time period the land region with the greatest spatially consistent statistically significant positive correlations is extratropical Eurasia. These observations suggest that temperature perturbations in this time period, particularly in the high latitude northern hemisphere, may have impacted global methane concentrations as has been noted previously by other workers [Etheridge et al., 1998; MacFarling Meure et al., 2006].

3.2.2. Precipitation

[36] The largest areas of methane emissions from natural wetlands are the monsoon regions of East Asia, India, and South America [Bergamaschi et al., 2009]. High temporal resolution records of rainfall variability in specific areas of monsoon regions have been inferred from the oxygen isotope composition (δ18O) of speleothems (cave deposits). While the correlation between speleothems from monsoon regions to Greenland temperature [e.g., Wang et al., 2006; Wang et al., 2001] and Greenland temperature to methane [e.g., Brook et al., 2000; Chappellaz et al., 1993] have been widely reported for the last ice age, these relationships have not been explored for the late Holocene.

[37] Speleothem δ18O records that cover the LPIH with enough temporal resolution to observe multidecadal variability have been recovered from the East Asian monsoon region and Peru. Of the East Asian speleothem records, the Dongge [Wang et al., 2005] and Heshang [Hu et al., 2008] chronologies have uncertainties of ± 50 years which reduces the confidence in the timing of multidecadal variability. The Wanxiang [Zhang et al., 2008] chronology is much better (< ± 5 years) and is highly correlated on decadal timescales with another speleothem from the Dandak cave in east central India [Berkelhammer et al., 2010] which supports...
the interpretation that these records represent regional precipitation. On their stated chronologies, the Heshang ($r = 0.24, p = 0.02$) and Wanxiang ($r = 0.28, p = 0.01$) speleothem $\delta^{18}O$ records have statistically significant correlation with methane, but the Dongge ($r = 0.02, p = 0.88$) record does not (Table 2 and Figure 6). The speleothem $\delta^{18}O$ record from the Cascayunga cave in Peru is inversely correlated with tropical SSTs in the Cariaco basin on multidecadal timescales [Reuter et al., 2009] and other South American speleothems have shown this same relationship on millennial timescales [Cruz et al., 2005; Lea et al., 2003].

Correlation between methane and the Cascayunga speleothem is negative and is not statistically significant on its stated chronology ($r = -0.19, p = 0.09$). The correlation increases to $r = -0.35$ when the annually interpolated chronology is shifted forward in time by 19 years, although this shift is larger than the uncertainty of their chronology ($\pm 4$–9 years) [Reuter et al., 2009]. This possible correlation on multidecadal timescales is intriguing because a negative correlation has been noted between South American speleothems and methane on millennial timescales [Cruz et al., 2005].

A recent spatial reconstruction of the Asian monsoon region Palmer Drought Severity Index (PDSI) [Cook et al., 2010] extends back to 1300 C.E. and offers a significant advancement in that the records are annually resolved and have a broad distribution over a large wetland region. Spatial correlation coefficients and analytical uncertainty from this reconstruction were constructed in the same manner as the correlation with the temperature field reconstruction and cover 1300–1800 C.E. (Figure 7).

Positive PDSI values indicate wetter conditions and the highest positive correlation coefficients are centered on the eastern Tibetan Plateau. This area is the source of many major Asian rivers (Yangtze, Mekong, Yellow, Pearl, and Salween rivers) which feed much of the Asian monsoon region. Modern satellite observations show that this area is also a major source of methane emissions which are positively correlated with groundwater depth and temperature on interannual and shorter timescales [Bergamaschi et al., 2009; Bloom et al., 2010]. Our results suggest that the relationship between methane and drought in this area holds for multidecadal timescales and may have been an important factor contributing to global methane variability during the LPIH.

3.2.2.3. Anthropogenic Methane Sources

[39] The dramatic increase in atmospheric methane concentrations at the start of the industrial revolution in the mid-18th century is a result of increasing anthropogenic emissions. Before this time the contribution of anthropogenic emissions are not well constrained [Ferretti et al., 2005; Houweling et al., 2008; Mischler et al., 2009]. A range of published LPIH anthropogenic methane emissions are shown in Table 3. These estimates are generally constructed by scaling modern anthropogenic emissions down with population and then making educated guesses about how anthropogenic activities (land use, farming practices, etc.) would have altered per capita emissions. The process involves many difficult to verify assumptions and we thus caution that the exact value of the following semiquantitative calculations is less important than the approximate magnitudes.

[40] The “early anthropogenic” hypothesis argues that human activity began altering atmospheric methane concentrations as early as 5000 years ago with the biggest contribution coming from agricultural activities, particularly rice farming in China [see Ruddiman, 2003, 2007, and references therein]. The hypothesis presumes that early rice farming techniques were inefficient resulting in disproportionately large methane emissions per capita relative to modern times. Following this line of reasoning, any large reduction in human population or agricultural production during the LPIH caused by plagues or wars, especially in areas of rice cultivation, should have reduced methane emissions on the timescales of those events. Here we use the historic record to explore the possibility that reductions in human populations, agricultural production, or land use patterns were large enough to have a demonstrative impact on global methane concentrations.

[41] The two biggest wars in Asia during the LPIH were the Mongol invasion beginning in 1211 C.E. lasting for about three decades, and the overthrow of Ming dynasty and the establishment of the Qing dynasty in the mid-17th century. These events were associated with large losses of population estimated at 35 million ($\sim 30\%$ or $\sim 15\%$ of the total Chinese or Asian population, respectively) during the Mongol invasion and 25 million ($\sim 15\%$ or $\sim 7\%$ of the total Chinese or Asian population, respectively) during the transition between the Ming and Qing dynasties [McEvedy and Jones, 1978; Pongratz et al., 2008], Zhang et al. [2007] examined agricultural production associated with wars in China from 1500 to 1800 C.E. and concluded that the war and loss of human life associated with the transition between
the Ming and Qing dynasties resulted in a sharp decrease in agricultural production. The invasion of the Mongol armies would have resulted in a similar or larger decrease in agricultural production due to the greater percentage of the population that was killed.

Assuming that essentially all preindustrial rice production occurred in Asia and given that ~70% of the total world population lived in Asia [McEvedy and Jones, 1978], we can add 100% of the rice emissions and 70% of all other anthropogenic emissions (Table 3) to estimate Asian anthropogenic emissions. Here we will focus on the high and low estimates of anthropogenic emissions that are commonly cited in the literature. This leads to estimates of Asian anthropogenic emissions of ~24 Tg CH₄ yr⁻¹ based on the work by Houweling et al. [2000] and ~56 Tg CH₄ yr⁻¹ based on the work by Ruddiman [2007]. Assuming a linear scaling between Asian population change and anthropogenic methane emissions, the invasion of the Mongol armies during ~1211–1241 C.E. would have resulted in a reduction of ~3.3–7.9 Tg CH₄ yr⁻¹ or ~10–23 ppb for Houweling et al.’s [2000] and Ruddiman’s [2007] estimates, respectively. The WDC05A record shows that methane decreased by ~25–30 ppb from 1218 to 1235 C.E. Using the same argument, the transition from the Ming to Qing dynasties (1618–1662 C.E.) would have led to a reduction of ~1.6–3.7 Tg CH₄ yr⁻¹ or ~5–11 ppb for Houweling et al.’s [2000] and Ruddiman’s [2007] estimates, respectively. The WDC05A record has a small gap during this time period, but there is a decrease of ~13 ppb observed in the Law Dome record. Preliminary measurements from the main borehole (WDC06A) have a decrease similar to Law Dome (not shown) in this time interval. Therefore, the timing and magnitude of putative reductions in anthropogenic rice emissions resulting from war is within the chronology uncertainty of our record and represents a possible cause for these reductions in methane.

There is further anecdotal evidence that the invasion of the Mongol empire could have influenced methane emissions. After razing villages and cities, the Mongol army laboriously dismantled the irrigation systems and used their horses to churn up the soil [Weatherford, 2004]. This prevented people from immediately resettling after the Mongol army left and also allowed the land to revert to grasslands which have significantly lower methane emissions than irrigated farmland. Following the invasion, the governing Mongols encouraged scientific innovations which led in 1261 C.E. to the establishment of the Office for the Stimulation of Agriculture which sought to increase the agricultural output of farmlands by the diversification of crops and improvement of farming practices [Weatherford, 2004]. At this same time, global methane concentrations began to increase. These land use changes would have changed the areal extent of methane emissions, and we speculate that it could have contributed to the rapid changes in global methane concentrations during this time period.

In the mid-14th century, plague broke out and spread rapidly across Asia and Europe along the extensive Mongol trade network. Population losses in Asia and Europe were estimated to be in excess of 60 million and 20 million, respectively [McEvedy and Jones, 1978; Weatherford, 2004]. Use of the linear scaling argument above leads to a reduction of Asian anthropogenic emissions of ~6–14.3 Tg CH₄ yr⁻¹ and European anthropogenic emissions of ~1.5–3.3 Tg CH₄ yr⁻¹ for Houweling et al.’s [2000] and Ruddiman’s [2007] estimates, respectively. Thus the total reduction by linear scaling caused by plague would have been ~7.5–17.6 Tg CH₄ yr⁻¹ or ~21–50 ppb. The WDC05A record shows that methane decreases by ~16 ppb from 1314 to 1359 C.E. A possible explanation for the smaller than estimated reduction in methane is that plague would not have caused the land use changes that occurred during wars. This suggests that the land use changes could have had a greater impact than population changes and might be an explanation for why the reduction in methane during the Mongol invasion is slightly greater than that estimated with a linear scaling to population.

Changes in the δ¹³CH₄ record can help identify methane sources which have divergent isotopic signatures. The δ¹³CH₄ record over the LPIH shows a large, gradual decrease from 1400 to 1700 C.E.; however, a large portion of the decrease occurs during 1560–1600 C.E. [Ferretti et al., 2005; Mischler et al., 2009]. This is coincident with the largest decrease in methane concentrations during the LPIH, ~32 ppb in the WDC05A record. Ferretti et al. [2005] proposed that the reduction in δ¹³CH₄ was caused by the decrease of biomass burning (an isotopically heavy source) in the Americas after the arrival of European settlers introduced disease to Native American populations causing a widespread pandemic [Cook, 1998; Ruddiman, 2007]. This hypothesis is consistent with a marked decrease in charcoal accumulation in global sedimentary records [Marlon et al., 2008]. Estimates of LPIH biomass burning emissions range from 10 to 38 Tg CH₄ yr⁻¹ which corresponds to 28–108 ppb (Table 3). While these estimates include some anthropogenic biomass burning that is outside of North America, Ferretti et al. [2005] argue that Native American activities provided the greatest contribution to this source. Thus a reduction in biomass burning methane

Table 3. Estimates of Anthropogenic Emissions at ~1500 C.E.⁴

<table>
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<tr>
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<tbody>
<tr>
<td>Rice agriculture</td>
<td>10</td>
<td>~28 (23–32)</td>
<td>~20</td>
<td>30 (26 = biomass; 4 = wood fuel)</td>
<td>38</td>
</tr>
<tr>
<td>Biomass burning</td>
<td>10</td>
<td>20</td>
<td>~20</td>
<td>10</td>
<td></td>
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<tr>
<td>Domestic Ruminants</td>
<td>5</td>
<td>7</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Waste</td>
<td>5</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Climate feedbacks</td>
<td>30</td>
<td>~69 (60–78)</td>
<td>~20</td>
<td>55</td>
<td>53</td>
</tr>
<tr>
<td>Total</td>
<td>120</td>
<td>~28</td>
<td>~8</td>
<td>22</td>
<td>21</td>
</tr>
<tr>
<td>Percent of total budget</td>
<td>12.0%</td>
<td>12.0%</td>
<td>8.0%</td>
<td>22.0%</td>
<td>21.0%</td>
</tr>
</tbody>
</table>

⁴In Tg CH₄ yr⁻¹. Dashes indicate that the study did not estimate that source. Total budget is assumed to be 250 Tg CH₄ yr⁻¹.
emissions during this time period is consistent with three independent lines of evidence: the decrease in methane concentrations, decreasing isotopic $\delta^{13}$CH$_4$ values, and widespread pandemics in the Americas associated with European invasion.

[46] While anthropogenic activities may have had a discernable impact on multidecadal variations in methane concentrations, many uncertainties remain. Past population estimates are highly uncertain, particularly for the precolonial Americas and Asia. Additional work on quantifying the range of emissions from modern and preindustrial rice agriculture techniques as well as the extent of rice agriculture in the LPIH is needed. Longer high-precision methane records can place the LPIH variations in a longer term context. A coupling of methane emissions with anthropogenic activities could be an explanation for the generally low correlations with temperature and precipitation reconstructions over the LPIH discussed earlier.

4. Conclusion

[47] We have presented a new high-resolution, high-precision record of atmospheric methane covering 1000–1800 C.E. from the West Antarctic Ice Sheet Divide ice core (WDC05A). The high correlation between the WDC05A and the Law Dome methane record [Etheridge et al., 1998; MacFarling Meure et al., 2006] confirms the variability observed in both records. We are able to uniquely identify small-scale variability demonstrating that high-resolution methane records can be used to establish chronostatigraphic tie points for ice core gas chronologies on short timescales.

[48] We find that reconstructions of regional to hemispheric temperature are not highly correlated with methane concentrations. Correlation coefficients with a spatially resolved temperature reconstruction are highest in northern Eurasia, consistent with modern satellite observations on interannual timescales. The highest correlation coefficients with the spatial temperature reconstruction as well as many individual extratropical northern hemisphere temperature reconstructions are observed from 1400 to 1600 C.E. during the onset of the Little Ice Age. This suggests that temperature variations during this time period impacted methane variations. The correlation between proxies for East Asian monsoon strength and methane are similarly low on multidecadal timescales, although uncertainty in the age scales prevents a definitive analysis in some cases. A spatial reconstruction of the Asian monsoon region Palmer Drought Severity Index has the greatest correlation with methane at the headwaters of major East Asian rivers, consistent with modern satellite observations. Moderate to high correlations exist with a Peruvian speleothem and tropical SSTs in the Cariaco basin if these records are shifted by the maximum amount allowed by the uncertainty in their chronologies. These possible correlations on multidecadal timescales are compelling because these relationships have been documented on millennial timescales and suggest that they could be a robust feature of the climate system. Possible explanations for the lack of high correlations with temperature and precipitation proxies are that the individual records comprising these reconstructions may not be reflecting conditions in methane source regions or that the variations were not large enough to significantly perturb methane emissions. Anthropogenic activities could have affected methane emissions based on the synchronous timing between large population losses in Asia and the Americas and decreases in methane concentrations. Our work reinforces the need for additional absolutely dated paleoclimate proxies [Jones et al., 2009], particularly from methane source regions.

[49] Future work will involve extending the high-resolution WAIS Divide methane record beyond 1000 C.E. using the main borehole WDC06A. Extension of the record will characterize the frequency of methane variability and changes in multidecadal variability that may be caused by variations in climate or anthropogenic activities. Understanding high-precision methane variability is critical for placing the recent record of atmospheric and satellite measurements in a longer term context, increasing our understanding of the range of variability in the global methane budget, and for prediction of future changes in that budget. Very high resolution records of methane from Greenland are needed to characterize multidecadal and centennial variability in the Inter Polar Gradient which will provide another constraint on the global methane budget.

References


