Dear Kirsten Thonicke,

We thank you for the comments. Below we respond to your points in blue and propose corresponding text in the manuscript in italics.

Please mention this point in the text of the revised manuscript:

Reviewer 1: Page 5, lines 14-18: It is not clear here if step 2 in the synchronization actually does anything. Is it necessary to have this as a separate component? It reads sort of awkwardly. Your response. Yes, we did not find any additional tie points when we tried to align the different datasets within the two hemispheres. However, we think it is important to mention that we could not find any incongruities with the given data resolutions.

We slightly changed passage in the manuscript which now reads as follows: However, within both hemispheres the temporal alignments of all the distinguishable CH₄ features were already within the limitation of the data resolution. Therefore, no additional tie points could be defined at this stage, within each hemisphere.

Please check if you can include your response to the first major point raised by Hinrich Schaefer in his review into the discussion.

After discussing our model calibration using SF_6 data to calculate the polar mixing time (section 2.4.1) we added the following paragraph:

Another valid strategy to cope with the ice core data representing polar tropospheric values would be to correct the data to represent mean hemispheric ($\varphi_{-}ITCZ=5.0^{\circ}N$) values (similarly as it has been done by Etheridge et al. (1998) and by Schaefer et al. (2016) to derive a global average signal from ice core CH₄ and δ^{13} C-CH₄ data) based on the knowledge about the modern CH₄ concentration distribution (e.g. as provided by Dlugokencky et al. (2017)). However, using the corrected data for the box-model inversion also requires the corresponding calculation of θ (to represent the hemispheric mixing time) using mean hemispheric SF₆ concentrations. The lower spatial resolution of the SF₆ concentration data (especially in the low latitudes) biases the mean hemispheric SF₆ values and thus does not allow a satisfying calculation of a hemispheric exchange time. Therefore, we decided not to follow this approach and use polar values in combination with the polar exchange time (as explained above) for our inversion study.

Best wishes, Jonas Beck

Bipolar carbon and hydrogen isotope constraints of the Holocene methane budget

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- Abstract. Atmospheric methane concentration shows a well-known decrease over the first half of the Holocene following the northern hemisphere summer insolation before it started to increase again to preindustrial values. There is a debate about what caused this change in the methane concentration trend, in particular, whether an early anthropogenic influence or natural emissions led to the reversal of the atmospheric CH₄ concentration. Here, we present new methane concentration and stable hydrogen and carbon isotope data measured on ice core samples from both Greenland and Antarctica over the Holocene. With the help of a two-box model and the full suite of CH₄ parameters, the new data allow us to quantify the total methane emissions in the northern and southern hemispheres separately as well as their isotopic signatures, while interpretation of isotopic records
- of only one hemisphere may lead to erroneous conclusions. For the first half of the Holocene our results indicate and asynchronous decrease in northern and southern hemisphere CH_4 emissions by more than 30 Tg- $CH_4/y_{\mp a}$ in total accompanied by a drop in the northern carbon isotopic source signature of about -3. This cannot be explained by a change in the source mix alone, but requires shifts in the isotopic signature of the sources themselves caused by changes in the precursor material
- for the methane production. In the second half of the Holocene global CH₄ emissions increased by about 30 Tg-CH₄/yFa, while preindustrial isotopic emission signatures remained more a less constant. However, our results show that thethis early increase of methane emissions starting in the mid Holocene took place in the southern hemisphere, while northern hemisphere emissions started to increase only about 2000 years ago. Accordingly, natural emissions in the southern tropics appear to be the main cause of the CH₄ increase starting 5000 years ago in contradiction to, not supporting an early anthropogenic influence on the global methane budget by East Asian land use changes.

1 Introduction

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Atmospheric methane (CH₄) is a potent greenhouse gas and its concentrations are strongly coupled to the Earth's climate system. Due to the human influence on the Earth system, the CH₄ concentration ([CH₄]) in the atmosphere has increased by a factor of 2.5 (relative to the preindustrial level) over the last centuries and contributes significantly to the human-induced radiative forcing (Dlugokencky et al., 2005; Etheridge et al., 1998). Today both anthropogenic CH₄ sources (rice agriculture, livestock, fossil fuel production, anthropogenic biomass burning and landfills) and natural CH₄ sources (natural wetlands,

- livestock, fossil fuel production, anthropogenic biomass burning and landfills) and natural CH₄ sources (natural wetlands, wildfires, geologic emissions, wild animals (including termites) and marine CH₄ hydrates) contribute to the global CH₄ emissions (Kirschke et al., 2013). The main mechanism removing CH₄ from the atmosphere is the chemical reaction of CH₄ with OH radicals in the troposphere. Together with the other sink processes (such as stratospheric loss, reaction with Cl radicals
 in the marine boundary layer and soil uptake) the OH sink determines the recent atmospheric life timelifetime of CH₄, which
- is between 8.4 and $10.69.1 \pm 0.9$ years (NaikPrather et al., 20132012).

Despite the fact that the total CH_4 emissions today are well known, there is still considerable debate about the individual contributions of (in particular small) CH_4 source and their variability.sources and their variability, e.g. evidenced by the

mismatch of bottom-up and top-down estimates of the total CH₄ emissions (Crill and Thornton, 2017; Kirschke et al., 2013; Saunois et al., 2017). Moreover, a long-standing debate exists whether long-term Holocene CH4 trends are naturally caused by wetland emissions (Singarayer et al., 2011) or whether an anthropogenic influence started already 4-5000 years ago by early human land-use changes (Ruddiman, 2003) (see below). Accordingly, to To better assess the human influence on the global methane cycle, robust estimates of the natural (pre-anthropogenic) CH4 emissions and their climate-coupled variations are required. Only polar ice cores can provide this information as they represent a direct, albeit (through the bubble enclosure process in the firn) low-pass filtered archive of trapped air and, therefore, offer the possibility to investigate the atmospheric composition of the past. With such ice core measurements the millennial to centennial evolution of the atmospheric [CH₄] has been determined back to 800 thousand years before present (ka BP), where present refers to the year 1950, fromusing the 10 Antarctic EPICA Dome C (EDC) ice core showing glacial/interglacial variations in atmospheric [CH4] by a factor of about 2 (Loulergue et al., 2008). More important for the assessment of the human influence on CH₄ levels are their variations during the Holocene, i.e., our current interglacial that started about 10,000 years ago. In this time period [CH4] decreased by about 100 parts per billion (ppb) until about 5000 yryears ago (Figure 1) after an early maximum during the preboreal period (about 11.5-10 ka BP) (Blunier et al., 1995; Brook et al., 1996; Flückiger et al., 2002; Schilt et al., 2010). CH4 values started to slowly 15 increase again in the mid-Holocene until the fast anthropogenic increase resulting from the industrialisation startingstarted around 200 years ago (MacFarling Meure et al., 2006). High resolution CH4 measurements on the West Antarctic Ice Sheet (WAIS) and Greenland GISP2 (Greenland Ice Sheet Project 2) ice cores (Mitchell et al., 2013) over the last 2800 years show also centennial CH4 variations of 20-30 ppb over the last 2800 years superimposed on the long-term increase, reflecting natural variability of CH4 emissions.

With the trend reversal in the mid-Holocene leading to a "bowl shape" (Sowers, 2010Blunier et al., 1995) of [CH4], the Holocene differs from many previous interglacials, where the [CH4] steadily dropped to glacial levels concurrently with the northern hemisphere summer insolation. A controversialOne hypothesis postulates that CH4 emissions caused by early human land use were responsible for the CH4 turnaround about 5 ka BP (Ruddiman, 2003; Ruddiman and Thomson, 2001). According
to these authors, early farming activities, mainly rice agriculture in the eastern part of China, led to significant releases of CH4 long before the industrial era (Ruddiman et al., 2008) and, thus, the authors propose that the "bowl" was shaped by human activities. Another explanation supported by a model-based study explains the late Holocene CH4 rise by calling upon increased emissions from southern tropical wetlands due to an insolation-driven strengthening of the monsoon in the western Amazon (Singarayer et al., 2011). Accordingly, this model is able to produce a global [CH4] rise in the Holocene driven by changes in the orbital forcing only. Note that due to the different orbital configuration during the last interglacial (MIS5.5), this model is

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also able to reproduce the steadily declining [CH₄] during MIS5.5, when strongly declining boreal CH₄ emissions overcompensate increasing southern tropical wetland emissions.

An important factpoint is that the two scenarios explain the Holocene CH₄ rise with additional emissions taking place in
different geographic regions. By measuring the [CH₄] on ice cores from both polar regions (Greenland and Antarctica), the inter polarinterpolar difference (IPD) can be calculated. This quantity can be used to draw conclusions about the hemispheric distribution of the CH₄ emissions (Brook et al., 2000; Chappellaz et al., 1997; Mitchell et al., 2013; Yang et al., 2017). Another tool to investigate the processes involved in the CH₄ cycle are measurements of the stable isotope signature in ice core CH₄. The CH₄ released by the various sources are associated with typical hydrogen and carbon isotopic signatures δD-CH₄ (D
representing Deuteriumdeuterium or ²H) (Quay et al., 1999; Walter et al., 2008; Whiticar, 1993; Whiticar and Schaefer, 2007) and δ¹³C-CH₄ (Etiope et al., 2008; Quay et al., 1999; Walter et al., 2005; Levine et al., 2011b; Snover and Quay, 2000; Whiticar and Schaefer, 2007), which leaveleaves atmospheric CH₄ being strongly enriched in heavy isotopes relative to the

isotopic emission signatures. Changes of the relative contributions in sources lead to an alteration of the atmospheric CH₄ isotopic composition. CH₄ isotope measurements onin ice cores, therefore, allow us to deduce the role the different processes related to the CH₄ cycle played in the past. By measuring both δ D-CH₄ and δ ¹³C-CH₄ from the Greenland GISP2 ice core over the Holocene, Sowers (2010) was able to provide first evidence of possible changes in the CH₄ precursor material due to a shift in the C3/C4 plant ratio, but could not unambiguously answer the question about the Holocene CH₄ anomaly.

In this study we combined for the first time comprehensive information on the CH₄ IPD and measurements of the <u>bipolar</u> carbon and hydrogen isotopic signature of CH₄ by measuring all three parameters – the [CH₄], δD-CH₄ and δ¹³C-CH₄ – on ice samples from both polar regions with a-high analytical precision and improved resolution. Furthermore, we use a two-box
 model approach, which allows us to deconvolve the atmospheric signal, to calculate the total emissions and their isotopic signatures in each hemisphere.

As mentioned above, changes in the sinks leave an imprint on the isotopic composition of CH₄ in the atmosphere-as well. However, the CH₄ sinks are thought to scale proportionally with the source (not changing the CH₄ life time) and only play a
minor role for the observed atmospheric [CH₄] variation since the last glacial maximum (Levine et al., 2011a; Martinerie et al., 1995). However, while minor changes in the relative importance of the sink processes cannot be ruled out, in general, atmospheric chemistry models point to small changes in total lifetime (Kaplan et al., 2006; Levine et al., 2011a) and we also assume the relative contribution of individual sinks to be rather constant over the Holocene when relatively stable climate conditions prevailed. Therefore, the discussion of this study centres around CH₄ source processes, hence emissions of different 20 CH₄ source categories.

2 Method

2.1 Measurements

The stable isotope data (δD-CH₄ and δ¹³C-CH₄) presented in this study were measured at the University of Bern on samples from the NGRIP (North GReenland Ice core Project) ice core from Greenland. Antarctic samples from the EDML (EPICA
Dronning Maud Land) and the TALDICE (TALos Dome Ice CorE) ice cores were analysed for δD-CH₄ and δ¹³C-CH₄, respectively (Bock et al., 2017). Note that for both parameters the samples from Greenland and Antarctica have been measured during the same time interval with the same measurement system by the same operators and in randomised order. This procedure is crucial to avoid any systematic error (e.g. due to a long-term drift of the system) and ensures unbiased results in our source deconvolutioninversion.

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The two isotopic parameters were measured at the University of Bern on two independent measurement systems designed for this specific purpose as described in detail in previous publications (Bock et al., 2010; Schmitt et al., 2014). They follow the same principal procedure: the air is (i) extracted from the ice sample by purging the melt water <u>with He</u>, (ii) water vapour, CO₂ and the major air compounds (N₂, O₂ and Ar) are removed from the gas sample, (iii) CH₄ is separated from other trace gases using gas chromatography (GC), (iv) CH₄ is pyrolysed to H₂ or combusted to CO₂ in the δ D and δ ¹³C systems, respectively, and (v) the isotopic composition of H₂ and CO₂ is measured using an isotope-ratio mass spectrometer (IRMS). For both systems effects of interfering masses, specifically Krypton, in the mass spectrometer are avoided by technical adaptations (Schmitt et al., 2013). In the case of δ D-CH₄, the focussed H₂ peak is separated from other components using a second GC separation (post-pyrolysis trapping) (Bock et al., 2014). In the δ ¹³C-CH₄ system, the CH₄-derived CO₂ is captured and delayed using a

⁴⁰ fused silica trap (at -196°C) after the combustion oven and thereby separated from Krypton (Schmitt et al., 2014). Note that

many δD -CH₄ and δ^{13} C-CH₄ values based on such a gas chromatography - mass spectrometry method reported in the literature are not corrected for this effect, leading to significant offsets especially for low CH₄ concentrations.

The δD-CH₄ values are determined relative to our primary air standard 'Air Controlé', a recent clean air ([CH₄] = 1971 ± 7
ppb), which has been cross-referenced to -93.6 ± 2.2 ‰ with respect to (wrt) Vienna Standard Mean Ocean Water (VSMOW) using 'Alert 2002/11', an air collected at Alert Station, Canada (Bock et al., 2014). The latter was referenced by Poss (2003) to be -82.15 ± 0.3 ‰ wrt VSMOW, which anchors our δD values to the isotope scale of the University of Heidelberg (Bock et al., 2010, 2014). Note that so far no internationally uniform δD-CH₄ standardisation scale has been established leading to systematic interlaboratoryinter-laboratory offsets as described in Umezawa et al. (2017). The measurement precision (1σ) of
2.3 ‰ was derived by calculating the pooled standard deviation of multiple NGRIP replicate (vertically neighbouring samples) measurements. The reference gas for the δ¹³C-CH₄ measurements is 'Boulder', an ambient air diluted with CH₄-free air ([CH₄] = 1508.18 ± 0.17 ppb) calibrated by the National Oceanic and Atmospheric Administration (NOAA) to a δ¹³C-CH₄ value of -47.34 ± 0.02 ‰ with respect to Vienna Pee Dee Belemnite (VPDB) (Schmitt et al., 2014; Umezawa et al., 2017). The <u>δ¹³C-</u>CH₄ measurement precision (1σ) derived from replicate measurements is better than 0.15 ‰ for ice core samples.

15 2.2 Monte Carlo spline approximation of millennial CH₄ changes

To compare values from different ice core datasets (with different and variable temporal resolutions and measurement precisions) interpolation of the data to a common age scale is needed. Additionally, the records have to be low-pass filtered to the same (millennial) time scales that can be resolved in the isotope records. Therefore, in this study smoothing splines following Enting (1987) are used to calculate a continuous low-pass filtered record from the discretely measured values. The stiffness of the spline curve is characterised by its cutoff period. A Python (http://www.python.org/) routine was developed to calculate splines in a Monte Carlo manner, where the data points are varied within a normal distribution within their standard

deviation defined by the individual uncertainty of the measured values. This procedure allows for the calculation of the average of the splines representing the best guess mean evolution of the data on the time scale resolved by the spline. Furthermore, using the standard deviation of all splines, the spread of the individual splines can be translated to an uncertainty of the mean spline. This spline calculation is used for various purposes in this study. The choice of the cutoff period has a large influence on the outcome of the spline and is, therefore, always documented wherever it is selected manually. For some applications (e.g. in the routines described in the sections 2.3.2 and 2.3.3), however, we apply an objective rulecriterion to define an

appropriate cutoff period for a data set: wherever the cutoff period is not specified explicitly, it is set to eight times the median 30 of the data resolution.

If the sampling rate in some parts of a dataset is much higher than in others, this part has a larger impact on the spline than other coarser resolved sections. For short periods of large signal change (e.g. the 8.2 ka event for CH₄) such an oversampling relative to other parts of the dataset may lead to significant artefacts in the calculated spline. Wherever this might be an issue, an additional step was introduced into the spline calculation routine, where the data were down-sampled in a randomised way within the Monte Carlo loop (i.e. for each individual spline) to end up with a sub set of a prescribed number of data points, where the chance of data points to get picked are inversely proportional to the local data density. The down-sampled data are still unequally distributed, but <u>the localon</u> average the data density is similar over the whole time period studied.

2.3 Data correction and compilation

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40 For further analysis of our new data, we complemented our [CH₄] records with data from other studies from the GRIP (GReenland Ice core Project), GISP2, TALDICE, EDML, WAIS (West Antarctic Ice Sheet) and EDC ice cores shown in

Table 1. However, combining different datasets introduces various challenges. In the following paragraphs we describe how these difficulties have been addressed.

2.3.1 Time scale synchronisation

All ice cores have individually determined gas age scales, which allow us to assign an age to each sample. Since we compare values from different ice cores (and from both hemispheres) small deviations in the age calculations have significant consequences for the interpretation (e.g. artefacts in the <u>deconvolutionbox-model inversion</u> (see section 2.4)). In an attempt to synchronise the different datasets, we linearly stretched and compressed the individual age scales between manually-chosen tie points in three steps:

- 1. Wherever possible, we tied the data sets of each of the cores to the same reference age scale. For CH₄ datasets that reach far enough to the present, the fast CH₄ increase of the last centuries was used to synchronise the time scales. As a reference we used [CH₄] data from Law Dome firm air measurements (MacFarling Meure et al., 2006) and an atmospheric [CH₄] history derived from NEEM firm air (Buizert et al., 2012) representing the CH₄ evolution in Antarctica and in Greenland, respectively. Both records cover the entire anthropogenic CH₄ increase and overlap with recent atmospheric measurements (CSIRO, 2017;
 15 Dlugokencky et al., 2017). For the early Holocene the layer-counted NGRIP δ¹⁸O-H₂O record was used as an absolutereference time markerscale. Where the CH₄ data show strong signals such as at the 8.2 ka event and the Younger Dryas-Holocene transition, these were synchronised to the corresponding δ¹⁸O signal, assuming that there is no significant lag between the
- temperature changes recorded in δ¹⁸O and the response of CH₄ source to this change (Baumgartner et al., 2014).
 20 2. With the use of splines of different cutoff periods and their time derivatives we were looking for typical features and signals
- in the [CH₄] evolution of the different data sets within the same hemisphere to define additional tie points and synchronise the data sets for each hemisphere. However, within both hemispheres the temporal alignments of all the distinguishable CH₄ features were already within the limitation of the data resolution. Therefore, no additional tie points could be defined at this stage, within each hemisphere.
- In the third synchronisation step we aligned the data of both hemispheres after applying the offset corrections (see section 2.3.2). Additional time corrections (for the southern hemisphere data) could be defined by looking at the fine structure of the [CH₄] evolution (e.g. the mid-Holocene [CH₄] minimum (around 5400 a BP) or the local [CH₄] maximum before the early Holocene decrease (around 9850 a BP)). Over the late Holocene period Mitchell et al. (2013) provided already synchronised data for the GISP2 and the WAIS ice core based on decadal to centennial variability; we retained their time scale over this period.

Table 2 lists the original age scales of the different ice cores. The tie points (original age and age correction) used to achieve an optimal temporal synchronisation between the CH₄ datasets are shown in Table 3. Temporal shifts to data with ages younger
than the first tie point and older than the last tie point were chosen to be identical to those of the first and last tie points, respectively. Please note that for the ice cores, where the primary age scales provide an age uncertainty, all but three tie point corrections (GRIP: 273 a; TALDICE: 127 a and 6042 a) lie within the 1*σ*-range of the uncertainty given for the individual age scales.

2.3.2 Offset correction

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40 Another issue that can <u>complicates complicate</u> compiling multiple datasets are concentration offsets. The CH₄ datasets we use to complement our new data were measured with different methods, in different labs, and at different times over the last

decades. If we use the information about the relative evolution of the methane concentration documented by other measurement series, one has to be aware that the absolute level might show an offset to our data due to calibration and extraction issues. Therefore, we correct all datasets for such potential offsets. To do so, we calculated residuals defined as the difference between the spline values of the CH₄ data from other labs evaluated at the age of our data and the CH₄ mixing ratios of our data. A

5 least-squares optimisation routine was used to find the offsets for the spline values of the different data sets. The offset corrections applied to the data are shown in Table 1. The GISP2 and WAIS datasets by Mitchell et al. (2013) are treated as a single dataset, as they were measured in the same lab and during the same measurement campaign, and their difference (Greenland-Antarctica) represents the true IPD. In Figure 1 the synchronised and offset-corrected CH₄ time series as used in this study are shown.

10 2.3.3 Outlier detection

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The GISP2 datasets from Brook et al. (1996) and Brook (2009) show an increased incidence of anomalously high $[CH_4]$ values (relative to other data sets) in the mid-Holocene. As this ice is from the brittle ice zone (the zone where air bubbles and clathrates coexist in situthe ice, and which is prone to bad core quality due to damage during the pressure and temperature relaxation process after the core was brought to the surface) this is probably due to infiltration of modern air into the ice samples in (micro) cracks opened in the ice after core retrieval (Gow et al., 1997; Neff, 2014). To avoid an artificial

- increaseelevation of the splined signal, which would result in a significant blow-upincrease of the IPD, we identifyidentified and removeremoved such outliers within the depth interval 650-1400 m of the GISP2 ice core; (equivalent to the time window 2.7-8.1 ka BP). To do so we used an iterative elimination approach. In each step, first the two GISP2 datasets were corrected for offsets relative to each other (Brook et al. (1996) was shifted to fit with Brook (2009) by 36 ppb) and combined to one data
- 20 set. Thereafter, for each GISP2 data point the difference between its value and the spline of all other GISP2 data in the brittle zone was calculated and divided by the 1σ uncertainty of the Monte Carlo spline at its age. If the largest (positive) relative residuumresidual exceeded the threshold value of 4, the corresponding data point was removed. This procedure was repeated until no outliers could be found any more. Table 4 shows the data points that were identified and eliminated through this method (also shown in Figure 1). In total 15 out of 95 data points (within this depth range) were removed, reflecting the bad
- 25 core quality in this depth interval. Note that <u>in this outlier detection routine</u> we make the assumption<u>here</u> that the [CH₄] evolution in the atmosphere is relatively smooth. As we focus only on millennial changes in the data here, this assumption does not affect our conclusions. <u>In the following discussion we assume that the corrected GISP2 data and the CH₄ data from the GRIP and NGRIP ice cores over this time interval are not affected anymore by modern air intrusion in the brittle ice. If this assumption does not hold, the true northern hemisphere CH₄ concentration would be lower and so would be northern hemisphere emissions at that time.</u>

2.3.4 Isotope data corrections

All stable isotope data have been corrected for gravitational settlingenrichment. This process takes place in the firn column and leads to an enrichment of the heavier isotopes at the depth where the air bubbles are finally closed off and the gas is trapped in the ice (Schwander, 1996). Using δ¹³⁶Xe data, which come as a side product during the δ¹³C-CH₄ measurement of the Bern δ¹³C system, and additional published δ¹⁵N-N₂ data (Capron et al., 2013; Eggleston et al., 2016; Landais et al., 2006), we experimentally derived this gravitational enrichment and directly correct the CH₄ stable isotope data, as this gravitational enrichment is only dependent on the mass difference between the two isotopes considered. The correction for the TALDICE δ¹³C values increasesdecreases from the present to the carly Holocene to the present from 0.2839 ‰ to 0.3928 ‰. The NGRIP δ¹³⁶Xe data do not show a systematic signalchange. Thus, a constant value of 0.32 ‰ is subtracted from the NGRIP isotope
values. Only very few δ¹⁵N-N₂ data points in the early Holocene were available to correct the EDML δD-CH₄ record. Since in the case of δD-CH₄ the measurement error and the observed signal are much larger than the gravitational enrichment, it is

adequate to assume that this correction is also constant over time without any consequences for the interpretation of the data. For the EDML δ D-CH₄ record the<u>a</u> correction of 0.46 ‰ was applied.

Due to the relatively constant climatic conditions and <u>no rapidslow</u> [CH₄] changes occurring over the Holocene, thermal diffusion (Severinghaus et al., 1998) and diffusive fractionation (Buizert et al., 2013) – other processes potentially leading to an artefact in the stable isotope signature – are not considered in this study.

2.4 Deconvolution-Box-model inversion

Our deconvolution-box-model inversion is a tool to disentangle the different processes involved in the atmospheric methane cycle. More precisely, it allows us to obtain quantitative information about the emission processes that led to the local tropospheric composition<u>CH₄ concentration</u> at changes recorded in the ice core sitescores in the past. In other similar studies different approaches of varying complexity and with different boundary conditions have been presented (Baumgartner et al., 2012; Fischer et al., 2008; Mitchell et al., 2013). For this study we use a two-box model where the boxes represent the northern and southern tropospheric hemispheres.

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The following six deconvolution equations allow us to use the six measured parameters (the tropospheric [CH₄] (c_x) and the two stable isotope ratios (${}^{i}R_x$)) for each hemisphere to calculate the emissions (E_x) and their isotope ratios (${}^{i}R_{E_x}$) for both boxes analytically (with x and y representing north and south and *i* the mass numbers 2 or 13 for deuterium and 13 C):

$$E_x = m^* \cdot \left(\frac{r_x \cdot c_x}{\tau_x} + \frac{c_x}{2 \cdot \theta} - \frac{c_y}{2 \cdot \theta} \right) \tag{1}$$

$${}^{i}R_{E_{x}} = \frac{m^{*}}{E_{x}} \cdot \left(\frac{r_{x} \cdot c_{x}}{\tau_{x}} \cdot {}^{i}R_{x} \cdot {}^{i}\alpha_{x} + \frac{c_{x}}{2 \cdot \theta} \cdot {}^{i}R_{x} - \frac{c_{y}}{2 \cdot \theta} \cdot {}^{i}R_{y} \right)$$
(2)

The first term in equations Eq. (1) and Eq. (2) represents the loss due to the CH₄ sinks, and the other two terms represent the hemispherieinterpolar mixing (out of and into the box, with θ being the mixing time between the hemispherespoles as explained in more detail below). The parameter $m^* = \frac{m_0}{c_0}$ translates atmospheric concentrations (ppb) to total atmospheric inventories (in Tg) using the mean atmospheric concentration value $c_0 = 1650 \text{ ppb}$ of the year 1987-and. Here we use the corresponding global CH₄ burden $m_0 = 4800 Tg$ (by Steele et al., (1992), that was also used in previous ice core studies (Baumgartner et al., 2012; Fischer et al., 2008). Note that this value is 7-10% higher than more recent estimates (Dalsøren et al., 2016) and accordingly the derived absolute emissions may be 7-10% overestimated. However, we focus our interpretation on relative emission changes, which are not affected by this scaling factor. The transport between the boxes is quantified by the interpolar exchange time $\theta = 1.56 a$, which is determined through a model calibration using recent sulphur hexafluoride (SF₆) data (described in section 2.4.1).

30 Note that the sizes of the two well-mixed boxes are not equal; the sizes are defined by the global mean annual latitude of the inter-tropical convergence zone (ITCZ) $\varphi_{ITCZ} = 5^{\circ}N$ (Marshall et al., 2014), which acts as a barrier to the hemispheric exchange of air masses. The inequality of the box sizes is reflected in Eq. (1) and Eq. (2) by

$$r_x = \frac{V_x}{V_{atm}} = \frac{1}{2} \left(1 \mp \sin(\varphi_{ITCZ}) \right), \tag{3}$$

the ratio of the box volume relative to the total atmospheric volume. (n = 0.456, r_s = 0.544 for φ_{ITCZ} = 5°N). The global mean life timelifetime of CH₄ quantifies the total methane sink and is set to τ = 8.4 a (as in the 4-box model used in Fischer
et al. (2008) and in Bock et al. (2017)). Again, using other estimates of the atmospheric lifetime (Naik et al., 2013; Prather et al., 2012) changes the absolute emissions estimates but does not affect our conclusions on relative emission changes.

According to the relative importance given inby Kirschke et al. (2013) (average of bottom-up estimates of all three time intervals shown in their study) the sink is proportionally divided into the four different sink processes (s_{OH} , s_{strat} , s_{soil} , s_{Cl}) shown in Table 5. While s_{OH} and s_{strat} are thought to be equally distributed north and south of the ITCZ and, therefore, are set equalthe same in the two boxes, s_{soil} and s_{Cl} are partitioned according to the area ratios of land and open water, respectively, derived from an 0.5° -by- 0.5° resolution land-cover database (Channan et al., 2014). The life timelifetime of CH₄ in each box

$$\tau_{\chi} = \tau \cdot \frac{\sum s_{tot}}{\sum s_{\chi}} \cdot r_{\chi} , \qquad (4)$$

is determined by the north-south distribution of the sinks and the relative box sizes. The chosen values for the atmospheric CH₄ lifetime and the strengths and the distribution of the sinks represent a best guess based on the cited literature. Hence, other studies end up with slightly different values. Note also that we assume the model parameters to remain temporally unchanged over the Holocene.

The different sink processes lead to strong fractionations in both isotopes (Cantrell et al., 1990; Feilberg et al., 2005; Houweling et al., 2000; Quay et al., 1999; Saueressig et al., 2001; Snover and Quay, 2000 and references therein) and yield the hemispheric isotope fractionation factors ${}^{i}\alpha_{x} = 1 + {}^{i}\varepsilon_{x}$, with the fractionations ${}^{i}\varepsilon_{x}$ shown in Table 5. As in our model with only two 15 tropospheric boxes s_{strat} is applied to the total tropospheric CH₄ inventory instead of the isotopically more depleted stratospheric CH₄, the literature values for the stratospheric sink fractionation have to be adjusted to ${}^{2}\varepsilon_{strat} = -170 \%_{0}$ and ${}^{13}\varepsilon_{strat} = -13.1 \%_{0}$. These values have been empirically derived to match the tropospheric isotopic signature of the forward 4-box model described by Fischer et al. (2008) which contains two stratospheric boxes. First, usingUsing the identical CH₄ emission setup in both models, the north-south air mass exchange of the 2-box model has been increased to equal the tropospheric [CH₄] values with those of the 4-box model (IPD). In a second step, the fractionations of the stratospheric sink ${}^{i}\varepsilon_{strat}$ in the 2-box model have been adjusted to match the tropospheric δ D-CH₄ and δ ¹³C-CH₄ values of the two models.

The sink fractionation values used for the 4 box model analysis by Bock et al. (2017) are very similar to the values we used in this study. Note that the numbers for the sink fractionation caused by stratospheric loss and by soils were accidentally swapped
in the printed version of Table S3 by Bock et al. (2017) for both δ¹³C and δD. However, their model runs and therefore the results are based on the correct values.

As the individual data points from ice core [CH₄] and isotope measurements are subject to uncertainty in both time and valuesmagnitude even after aboveour homogenisation stepsprocedure, we only use the millennial variations in CH₄, δD-CH₄
and δ¹³C-CH₄ in our deconvolution box-model inversion. To this end we calculated low-pass filtered versions of the measured data using the Monte Carlo spline approximations described in section 2.2, which allow us to quantify the mean and the uncertainty of the millennial variations. The uncertainty of the splines of the tropospheric values are transmitted to introduced into the emission values using Gaussian error propagation. Due to the mentioned limitations we refrain from analysing shorter-scale variability.

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2.4.1 SF_6 calibration of the two box model

The same approach used to set up the CH_4 deconvolutioninversion described in the previous section (Eq. (1) and (2)) allows us to formulate the general equations for the change of the concentration of any trace gas species in the two hemisphere boxes:

$$\frac{dc_x}{dt} \cdot r_x \cdot m^* = E_x - S_x - \frac{m^*}{2 \cdot \theta} (c_x - c_y) \tag{1}$$

with $\frac{de_x}{dt}$ representing the <u>rate of</u> change-<u>rate</u> of the hemispheric concentration, m^* the translation factor linking a concentration to an atmospheric inventory, E_x and S_x the emissions and the sinks in Box x, and r_x the ratio of the box volumes depending on the mean annual latitude of the ITCZ φ_{ITCZ} as described above, and where x and y standingstand for north and south (or vice versa).

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SF₆ is a strong greenhouse gas with a life timelifetime of more than 1000 years and, is solely of anthropogenic origin and mainly emitted in the northern hemisphere ($r_{E_n} = 95 \%$, $r_{E_s} = 5 \%$) (Geller et al., 1997, p.6; Kovács et al., 2017; Levin et al., 2010). NOAA/ESRL (2017) measurements taken at Alert Station (Canada), Summit (Greenland), Palmer Station (Antarctica) and South Pole (Antarctica) record the SF₆ evolution in both polar regions over the last decades. Since the target in this study is the long-time trend and not seasonal variations, the NOAA/ESRL data are smoothed using a smoothing spline with a cutoff period of 2 years. For every year in the recorded period (1996-2008 CE) the annual SF₆ increase ($\frac{dc_x}{dt}$) and the mean concentration (c_x and c_y) are calculated. The records of the global emissions (E_{tot}), the global atmospheric mean SF₆

concentration (c_x and c_y) are calculated. The feedbals of the global emissions (E_{tot}), the global atmospheric mean SF_6 concentration (c_{tot}) and the total atmospheric inventory (m_{tot}) published by Levin et al. (2010) are used to calculate the hemispheric emissions $E_x = r_{E_x} \cdot E_{tot}$ and the translation factor $m^* = mean\left(\frac{m_{tot}}{c_{tot}}\right) = 25.7 \frac{Gg SF_6}{ppt}$. With the approximation 15 of a negligible sink term (since life timelifetime exceeds the time scale of interest by 2-3 orders of magnitude) Eq. ((1) can be solved for the hemispheric interpolar exchange time:

$$\theta = \frac{m^*(c_x - c_y)}{2\left(\frac{dc_x}{dt} \cdot r_x \cdot m^* - r_{E_x} \cdot E_{tot}\right)}$$
(2)

where c_x and c_y are the SF₆ concentrations at high latitudes. Not that with this calibration we effectively define the exchange time between the northern and southern polar regions, which is somewhat longer than the mean interhemispheric exchange time. As we use θ later for our inversion of polar CH₄ concentrations measured in the ice cores, we effectively correct for the
difference in polar and mean hemispheric CH₄ concentration using this SF₆ calibration. We use the term interpolar exchange time throughout the manuscript. Using Eq. (2) the interpolar exchange time can be calculated for each year-and hemisphere. With φ_{ITCZ} = 5.0°N we get the average value of θ = 1.56 ± 0.17 a.

In reality the air is not perfectly mixed within each hemisphere, which means that our ice core measurements do not perfectly represent the hemispheric average. However, by using the high latitude SF₆ data for the model calibration, we inherently correct for the fact that a signal subject to the inter hemispheric exchange has to be transported to the poles before it is captured in the ice. Accordingly, θ has to be interpreted as an interpolar rather than an interhemispheric exchange time.

DeconvolutionAnother valid strategy to cope with the ice core data representing polar tropospheric values would be to correct the data to represent mean hemispheric (φ_{ITCZ} = 5.0°N) values (similarly as it has been done by Etheridge et al. (1998) and
by Schaefer et al. (2016) to derive a global average signal from ice core CH₄ and δ¹³C-CH₄ data) based on the knowledge about the modern CH₄ concentration distribution (e.g. as provided by Dlugokencky et al. (2017)). However, using the corrected data for the box-model inversion also requires the corresponding calculation of θ (to represent the hemispheric mixing time) using mean hemispheric SF₆ concentrations. The lower spatial resolution of the SF₆ concentration data (especially in the low latitudes) biases the mean hemispheric SF₆ values and thus does not allow a satisfying calculation of a hemispheric exchange

35 time. Therefore, we decided not to follow this approach and use polar values in combination with the polar exchange time (as explained above) for our inversion study.

2.4.2 Inversion of artificial time series

The interpretation of the output of our inversion model is challenging and not always intuitive. To get a better understanding of the model output, some simplified experiments using artificial time series have been performed and are shown in this study for didactic reasons. In a first approach all six parameters are held constant over time. In the panels on the left in Figure 2 the tropospheric [CH₄] (a), and the isotopic signatures δ D-CH₄ (b) and δ ¹³C-CH₄ (c) are shown for the northern box in red and for the southern box in blue. The constant values (solid lines) represent the levels of the splines of our measurements at 2 ka BP and serve as input to the deconvolutioninversion model. The panels on the right show the deconvolutioninversion results, i.e., the integrated emissions (d) and their mean isotopic signatures (e, f) that are emitted into the two hemisphere boxes according to our model. As the input parameters are constant, the emission signal calculated by the deconvolution to achieve the prescribed tropospheric values—must be constant in all parameters as well. However, although the northern hemisphere concentrations are only 6 % larger than the southern ones, a large difference in the emission strengths ($E_n \approx 2.2 \cdot E_s$) is needed to maintain the IPD in the [CH₄] compensating the air mass exchange between the atmospheres. Note also that due to the sink fractionation the isotopic signature of atmospheric CH₄ is strongly enriched in the heavy isotopes compared to the emissions (see Table 5).

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For the second experiment (dashed lines in Figure 2), the IPD of the tropospheric isotope signatures (IPD_{δD} and IPD_{$\delta 13C$}) have been changed. We kept the southern values constant, while we linearly changed-the concentrations remained unchanged. By altering the northern values so that the minimal and stable isotope signal the isotopic IPD's are changed within the maximal IPD_{8D} and IPD_{813C} values matchrange observed in the splined isotope data, respectivelyice core records (dashed lines in (b) 20 and (c)). This leads to changes in the calculated isotope signatures of the emissions, whereas the emission strengths are unaffected. As shown (dashed lines) in panels (e) and (f), the isotopic signature for both the northern and the southern hemisphere emissions need to be altered to yield the given atmospheric values (of constant [CH4] and isotopic signatures changing only in one hemisphere). To achieve the declining isotope signal in the northern troposphere, the isotope signatures of emissions from the northern hemisphere have to become isotopically more depleted in heavy isotopes (further on 25 conveniently referred to as isotopically "lighter", equivalent to lower values δ -values) over time. At the same time the emissions in the southern hemisphere compensate for the isotopically lighter CH4 mixed down-from the northern into the southern box and, therefore, requires the southern box emissions to become more enriched in the heavy isotope (isotopically "heavier", equivalent to higher δ -values) over time. Since the southern source is much weaker than the northern source (more northern-sourced CH4 is mixed from north to south than southern-sourced CH4 is mixed from south to north), the change in 30 the isotopic signatures of the emissions turns out to be even larger in the southern hemisphere.

The dotted line in panel (a) in Figure 2 shows a third scenario, where all tropospheric parameters are kept constant except for the [CH₄] in the northern hemisphere. Similar to the former experiment, the [CH₄] in the northern box is varied between values representing the minimal and the maximal measured IPD_{CH4}. The dotted curves in the right panels (d-f) in Figure 2 indicate that this change has an influence on all six emission parameters of our model. The emission strength in the northern box increases to produce the rising [CH₄]. Declining emission strength in the southern hemisphere compensates for the increasing CH₄ north-to-south flux due to the rising IPD_{CH4}. The CH₄ that enters a box by atmospheric mixing is already fractionated by the sinks and isotopically heavier than the newly emitted CH₄. If the relative amount of CH₄ input by mixing is decreased for the northern (increased for the southern) box, the emission signatures become less (more) depleted in heavy isotopes to

40 maintain the tropospheric CH4 isotopic signatures.

Since the variations for both experiments were done based on a reasonable range (observed changes in IPDs), the span for the calculated values of the emission parameters also provides a range for the changes that are needed to alter the measured

tropospheric IPDs in the Holocene. Note, however, that the variation scenarios (e.g. with a fullyfull decoupling of atmospheric concentration and the isotopic composition of CH₄) do not represent changes we expect to observe in nature and are presented for didacticillustrative reasons only.

3 Results

- All The homogenised tropospheric data (this study complemented with published [CH4] data from Blunier et al., (1995); Brook et al., (1996); Brook, (2009); Chappellaz et al., (1993, 1997); Flückiger et al., (2002); Mitchell et al., (2013); Schilt et al., (2010), see section 2.3, Figure 1 and Table 1) are shown in Figure 3 in the left-hand panels. The smoothing splines, which are all calculated with the same cutoff period of 3000 years, are shown including their $1\sigma \operatorname{and} 2\sigma$ uncertainty bands. In panel (a) the composite of the time-synchronised and offset-corrected CH4 data is shown over the Holocene. The diamond-shaped symbols represent the NGRIP and TALDICE data of this study measured during the δ^{13} C-CH₄ measurement campaign. 10 Overall, our data compilation after the above mentioned corrections supports previous trends: after high concentrations in the Preboreal (ca. 11.5-10 ka BP) the data show a decline of the atmospheric [CH₄] over the first half of the Holocene followed by a reversal of this trend and an accelerating increase from the mid- to the late Holocene. The preindustrial level is comparable to the [CH4] during the Preboreal warm period. The splines emphasise this long-term shape nicely but, deliberately, the splines 15 do not reproduce the fine structure in the data (see for example Mitchell et al. (2013)). The difference between the [CH₄] values in the northern and southern hemispheres (the IPD_{CH4}) is also clearly visibleshown in the data.panel (g). The concentration measured onin ice cores from Greenland (red) is on average 45 ppb higher than the values derived from Antarctic ice cores (blue), which agrees with the result of Chappellaz et al. (1997).
- In panel (b) the measured-\deltaD-CH4 values showmeasured in ice from the NGRIP and the EDML ice cores are shown. As additionally illustrated in panel (h) we find a pronounced difference of -16.3 % between northern and southern \deltaD-CH4 values, which is rather stable over the whole period of investigation. Limited by the coarser resolution of the EDML \deltaD-CH4 data, the smoothed curves and the error bands illustrate the probable evolution of the true tropospheric signals only on millennial timescales and longer. Compared to the large [CH4] changes, the \deltaD-CH4 signal over the Holocene is relatively small and similar in the two datasets. The values become isotopically lighter in the first half of the Holocene and heavier again in the second half. Note that our NGRIP \deltaD-CH4 data differ stronglysignificantly from the dataset from the GISP2 ice core published by Sowers (2010) (see Figure 4). First, there is a difference of the mean level between the two datasets of 10 to 15 %, which can be attributed to an inter-laboratory difference (Bock et al., 2014; Umezawa et al., 2017). Second, the GISP2 data suggested a shift of about 20 % towards heavier values from 5 to 1 ka BP, which is not confirmed by our data. This disagreement, however, may be largely attributed to the much larger scattering of the GISP2 dataset, which doesdid not allow yet to quantify an unambiguous trend in \deltaD-CH4.

In panel (c) the δ^{13} C-CH₄ measured on ice from the NGRIP and TALDICE ice cores are shown. Both datasets show a gradual trend towards lighter values over the first half of the Holocene and almost constant values over the second half. The point of inflection is well-defined at about 5 ka BP. AtAs shown in panel (f) about the same time the δ^{13} C-CH₄ signals measured in the two hemispheres divergeIPD_{613C} changes quickly from a difference of -0.35 ‰ to -0.63 ‰, which is rather constant before and after the point of inflection. Our NGRIP δ^{13} C-CH₄ record is well in line with the GISP2 data by Sowers (2010) (see Figure 4), which have been subsequently corrected for the analytical Krypton interference-during the measurement (Schmitt et al., 2013).

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Our deconvolutioninversion model calculates the emissions that led to the measured tropospheric signals shown in the panels on the left. In panel (d) the strength of the integrated emissions (in Tg-_CH₄ per year) in each of the two troposphere boxes are shown. Additionally, the total CH₄ global emission (sum of north and south) are shown in grey. Note that the uncertainty (1σ and 2σ) in emission fluxes is very small and much smaller than the long-term changes in the emission fluxes over the Holocene indicated by the spline average. This uncertainty reflects only the error in emissions fluxes for changes on millennial time scales that are resolved in our spline. As indicated in high-resolution data (for example Mitchell et al., 2013) the centennial variability of atmospheric CH₄ is high and the uncertainties of reconstructed emission fluxes on such shorter time scales would be significantly higher as well.

10 The following observations can be made from the deconvolutioninversion results: The CH₄ emissions into the northern box are more than two times higher than the emissions into the southern box. After a decrease of roughly 30 Tg-CH₄/a in the early Holocene the northern emissions remain relatively constant between 8 and 2 ka BP before increasing by 15 Tg-CH₄/a. In the southern hemisphere the emissions showsuggest a slight increase in the first 2,000 years of the record followed by a significant 22 Tg-CH₄/a decrease until 4-5 ka BP after which emissions increase by 25 Tg-CH₄/a in the second half of the Holocene.

The weighted averaged δD signature of all the integrated CH₄ emissions is shown in panel (e) for each hemisphere. With values of about -295 ‰ the northern emission δD signature is on average 30 ‰ lighter than the southern hemisphere emissions. TheBefore 7.5 ka BP and around 4.5 ka BP the δD-Ex signal shows long-term variations around the Holocene mean that are in antiphase between the northern and southern hemisphere, however these variations in each hemisphere are not significantly
different from their respective Holocene mean within their 2σ uncertainty. Thewhereas the global mean δD-E (grey line) remains essentially constant. Based on these two observations, we attribute the δD Ex fluctuations in each hemisphere to and the measurement uncertainty of the individual data points of the low resolution δD CH4 EDML record which affect the spline reconstruction.

- 25 Ourknowledge from our experiments with artificial time series (see section 2.4.2) showshowing that a change in the isotopic signature in one hemisphere has a large influence of different sign on the emission isotope signal in the other hemisphere (Figure 2)- Hence, the), we attribute the δD-E_x fluctuations to the measurement uncertainty of the individual data points of the low-resolution δD-CH₄ EDML record which affect the spline reconstruction. The absence of a long-term trend in δD-E_x over the Holocene with the significant changes in the atmospheric [CH₄] is remarkable and a strong constraint on the average
- 30 <u>Holocene</u> CH₄ budget. However, the large uncertainty in δ D-E_x does not allow us to make robust conclusions about millennial variations in the hydrogen isotopic signature of CH₄ emissions.

The δ^{13} C-CH₄ values of the emissions shown in panel (f) indicate a significant shift of the northern emissionemissions to heavier values over the Holocene, which is especially pronounced in the time interval before 4 ka BP. In the south_a δ^{13} C-E_s 35 shows also a slight negative trend over the first half of the Holocene, which however, is still well within the 1 σ deconvolutioninversion uncertainty. A stronger positive trend, however, occurs during the second half of the Holocene, which_a however, is still withwithin the 2 σ uncertainty of the reconstruction. The global mean of emissions the isotopic emission signatures reflects the coherent trends in the measured atmospheric values, with a strong negative trend of $-1.8_{-\infty}$ over the first 5000 years of the Holocene.

40 3.1 Sensitivity studies

The size-ratio of the two boxes of the deconvolution box-model inversion is defined by the annual mean latitude of the ITCZ φ_{ITCZ} . In fact, there is evidence from different proxies of a southward migration of the ITCZ during the Holocene (Haug et al.,

2001; McGee et al., 2014; Zhao and Harrison, 2012). Since it is not possible to deduce robust numbers for the position and the movement of the global mean ITCZ from local studies, φ_{ITCZ} has been kept constant over the time at a reasonable Holocene value in our deconvolutionstandard inversion (described in section 2.4). However, sensitivity runs have been performed to quantify the impact of this assumption on the conclusions drawn from our deconvolutioninversion results.

As described in section 2.4.1, SF₆ calibration has been used to determine the inter hemispheric mixinginterpolar exchange time θ associated to different values of φ_{ITCZ} . For the first sensitivity study these two parameters have always been fitted in parallel to ensure that the different model configurations all fit the SF₆ constraint in an optimaloptimum way. In Figure 5 the splines of the Holocene data (left panels) and the corresponding emission values derived by the deconvolution calculationsinversion (right panels) are shown. In the sensitivity runs, the deconvolutioninversion equations have been solved using other ($\varphi_{ITCZ}, \varepsilon$)-couples with φ_{ITCZ} moving from 0.0°N to 10.0°N. In all six emission parameters, these variations only lead to small changes relative to the total signal variability. For the isotopic signatures of the emissions, all curves laylic well within the 1 σ error of the 5.0°N results.

15 A change of the mean annual position of the ITCZ has <u>also</u> an influence on the box <u>size distributions.sizes</u>. This has a direct impact on the north-south partitioning of the individual CH₄ sinks and thus on all six <u>deconvolutioninversion</u> parameters. Therefore, another sensitivity run was carried out, where only φ_{ITCZ} was varied and θ kept constant at the best guess value. As shown in Figure 6, the sensitivity of the <u>deconvolutioninversion</u> to a change of φ_{ITCZ} is very low if θ is kept constant. Note that the southward migration of the ITCZ over the Holocene is thought to be smaller than 1° (McGee et al., 2014). The results
20 of this second set of experiments where only one parameter was changed (i.e. the constraints of the SF₆ calibration were not fulfilled) also showedshows that on average variations in φ_{ITCZ} and θ contribute by a similar amount to the deviation from the <u>5°N solutionsour standard run (φ_{ITCZ} = 5°N)</u> (see Figure 6).

4 Discussion

The main goal of this study is to profit from the improved CH₄ multi-parameter dataset using our inversion model in order to evaluate the different scenarios and to constrain the global CH₄ budget over the Holocene. To this end, we consider the early, mid- and late Holocene separately.

4.1 Early Holocene (11-8 ka BP)

The significant <u>early</u> drop in the northern emission strength together with the slight shift towards isotopically lighter carbon emissions in the northern hemisphere from 11 to 8 ka BP is challenging to account for. One potential solution to account for
the observed concentration shift would be a general weakening of the boreal CH₄ sources as a result of the decreasing northern summer insolation (Berger and Loutre, 1991). However, this should lead to a trend in both isotope emission signatures towards heavier values in the northern hemisphere, as in general the high-latitude CH₄ emissions from wetlands are isotopically light in both isotopes (Walter et al., 2008; Whiticar and Schaefer, 2007). Accordingly, this conflicts with the stable isotope results from our deconvolution.inversion. For δD-E_n the deconvolutioninversion might be affected by one single measurement point
(EDML δD-CH₄) at 8.19 yr ka BP forcing the emission signature curves of the two hemispheres apart and indicating that δD-E_n becomes lighter. Nevertheless, even if this single data point is removed, the deconvolutioninversion signal would be constant over this period but would not show a trend towardtowards heavier values as expected from a decline in boreal emissions.

Based on ¹⁴C-dating. Walter et al. (2007) estimated the CH₄ emissions from thermokarst lakes to decrease from 26 to 5 40 Tg-CH₄/a in the period from 11 to 8 ka BP in line with the decreasing trend in northern hemisphere emissions in our deconvolution.inversion. To match the result of the isotope deconvolutioninversion, another isotopically heavy northern source (e.g. biomass burning) has to be reduced simultaneously to compensate for the lack of isotopically light thermokarst CH₄ emissions. However, charcoal records from sediments do not indicate a decline of wild fires in the northern hemisphere but rather <u>point to</u> the opposite during the Holocene (Daniau et al., 2012).

Part of the answer of this puzzle may lie in the combined temporal evolution of the strengths and isotopic signature of the boreal wetlands. Due to land ice and permafrost retreat in the early Holocene, minerotrophic fens, characterised by relatively high CH₄ emissions which are not as strongly depleted in ¹³C, turn into ombrotrophic bogs over time with more ¹³C-depleted but also much weaker CH₄ emissions (Ding et al., 2005; Hornibrook, 2013; Yu et al., 2013). This reduction of CH₄ emissions in the course of conversion of fens to bogs requires a very depleted $\frac{13}{2}$ source signature of the bogs to quantitatively explain

- our deconvolutioninversion results and may therefore not suffice to explain the δ^{13} C-E_n shift towards lighter values in the early Holocene. Furthermore, an enrichment trend in δ D-E_n (of about 5_.‰) would be expected if the δ D-depleted high-latitude emissions became relatively less important, while – if at all – we see a 20_.‰ depletion in δ D-E_n from 10 to 8 ka_BP. Also the isotopic signature of the water used in methanogenesis has a direct imprint on the hydrogen isotopic signature of the emitted
- 15 CH₄ (Whiticar and Schaefer, 2007). Speleothem δ¹⁸O records confirm an isotopic shift towards lighter values in the meteoric water in the summer monsoon regions of the northern hemisphere from 11 to 8 ka BP (Wang et al., 2014) and an inverse relationship in the southern hemisphere, in line with our deconvolution.inversion. However, the speleothem records also suggest opposite trends over the rest of the Holocene, which are not seen in our deconvolutioninversion results. In addition, isotopically light melt water from the retreating northern hemisphere ice sheets (for example in proglacial lakes) may contribute to a depletion in the hydrogen isotopic signature of boreal CH₄ source in the early Holocene.

4.2 Mid-Holocene (8-5.5 ka BP)

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Our deconvolutioninversion results show that the decrease in atmospheric CH₄ concentrations over this interval are mainly caused by declining southern hemisphere emissions. This decrease in the southern hemisphere CH₄ emission from 8 to 5.5 ka BP - during the time of the strongest increase of austral summer insolation (Berger and Loutre, 1991) - appears
counterintuitive but is in line with the model results by Singarayer et al. (2011) explaining it with), where this is caused by changes in tropical rainfall location. This is also the period when the ITCZ is thought to have started moving southward as a response to the change in the orbital forcing (Haug et al., 2001; McGee et al., 2014; Zhao and Harrison, 2012), which hashaving an influence on the hemispheric CH₄ source distributions. CH₄ emissions from regions that were attributed to the southern hemisphere box in the early Holocene may now contribute CH₄ to the northern hemisphere due to the shift of the ITCZ. Note that such a change not only has an influence on the spatial emission distribution, but also on other parameters; (such as θ), which are relevant for the deconvolutioninversion model, where the box sizes arewere kept constant over time. Our sensitivity studies (see section 3.1) show, however, that a movement of the ITCZ in the model has only a small effect on the deconvolutioninversion results.

- 35 As discussed above the δD signal of the CH₄ emissions is mostly constant in both hemispheres. This is expected if one hemisphere box gains emissions from tropical wetlands, characterised by stable isotope signatures close to the integrated source mix, at the cost of the other box. The rather constant charcoal wildfire reconstructions for the northern hemisphere in this time window (Daniau et al., 2012) do not support a decline in biomass burning CH₄ emissions being responsible for the 1.5 ‰ depletion in δ¹³C-E_n between 7 and 5.5 ka BP. The δ¹³C shift of the emissions could be related to a shift in the ratio of
- 40 C3 to C4 plants as suggested by Sowers (2010), while a further evolvement of northern fens to bogs is incompatible with the relatively constant emissions in the northern hemisphere. Under Holocene climate conditions, with high availability of humidity and CO₂, C3 plants successfully compete against the relatively ¹³C-depleted C4 plants and thereby alter the carbon

isotopic signature of the precursor material of methanogenesis (Farquhar et al., 1989). This process was also suggested by Möller et al. (2013) to explain the close CO_2 -correlation of the $\delta^{13}C$ -CH₄ signal with atmospheric CO₂ and the decoupling from the [CH₄] in the glacial period.

4.3 Late Holocene (5.5-1 ka BP)

- 5 The early anthropogenic influence hypothesis (Ruddiman and Thomson, 2001) calls upon increasing emissions from rice paddies mainly in Eastern Asia. In our box model such emissions would be located in the northern hemisphere box and, therefore, lead to an increase in northern emissions after 5 ka BP. However, the deconvolutionbox-model inversion shows no significant increase in E_n before 2 ka BP. In contrast at 5.5 ka BP, at the time when all six presented tropospheric records imply a change in the global CH₄ cycle, the deconvolutioninversion indicates an amplification of southern hemisphere CH₄ emissions. This clearly supports the idea of a natural Holocene wetland CH4 emission increase as suggested by Singarayer et al. (2011). 10 In contrast, a southern anthropogenic source can be excluded since the southern hemisphere was barely populated in the mid-Holocene (Kaplan et al., 2011). The observed shift in δD -E_s and especially $\delta^{13}C$ -E_s towards heavier values in this interval cannot be explained simply by increased tropical wetland emissions in the Amazon but requires also an enhancement of a southern hemisphere CH4 source strongly enriched in ¹³C.region but requires also an enhancement of a southern hemisphere 15 CH₄ source strongly enriched in 13 C. Using a simple mass balance approach (E_{total} ${}^{i}\delta_{total} = E_{initial} \cdot {}^{i}\delta_{initial} + E_{additional} \cdot {}^{i}\delta_{additional}$ allows us to determine a δ^{13} C signature of the additional CH₄ emissions of 19 Tg CH₄/a to be about -45.5 ‰ which is isotopically heavier than tropical wetland CH₄ emissions (-56.8 ‰ according to Whiticar and Schaefer (2007)). Note, however, that this quantitative approach is based on the assumption of
- unchanged carbon isotopic signatures of the initial CH₄ emissions and therefore needs to be taken with care. Increasing
 emissions from tropical and subtropical wild fires in the southern hemisphere, which are also documented in charcoal records (Daniau et al., 2012), could have become increasingly important, e.g. due to enhanced fuel production or higher CH₄ emission efficiency (more smoldering fires) both caused by wetter climatic conditions, and would readily explain the joint information derived from deconvolutioninversion results on southern hemisphere CH₄ emissions and its stable isotopic signatures. Other isotopically heavy sources such as geological emissions and marine CH₄ hydrates are unlikely to dramatically gain importance
 in the mid-Holocene and only in one hemisphere and can, therefore, be ruled out as important players driving the observed CH₄ changes.

5 Conclusions

The presentedOur new records of [CH₄], δD-CH₄ and δ¹³C-CH₄ measured on samples from polar ice cores from both hemispheres provide valuable insights into the Holocene CH₄ cycle. A significant IPD for δ¹³C and δD, as already documented
for [CH₄], exists over the entire period of investigation. While the [CH₄] data confirm the well-known Holocene evolution, the two stable isotope records provide additional insights. This shows the value of our multi–jisotope approach needed notin order to drawavoid drawing the wrong conclusions. A two-box model approach has been used to deconvolvetranslate the measured signals into emissions. The calculated hemispheric CH₄ emission signatures, characterised by different trends in the two hemispheres for δ¹³C and no statistically significant variation in δD, show a decoupling of the emission signatures of the two
isotopes. Therefore, we attribute the observedlong-term changes of the atmospheric CH₄ isotopes over the first half of the Holocene to shifts in the isotopic source signatures of individual CH₄ sources rather than to changes in the global CH₄ source mix.

In the early Holocene (11-8 ka BP) we associate the decline of northern hemisphere emissions and the δ^{13} C trend towards 40 lower values with the natural evolution of high–latitude wetlands from fens (high CH₄ emissions) to bogs (lower but strongly 13 C depleted CH₄ emissions). An alternative explanation calling upon a significant decrease of CH₄ emissions from thermokarst lakes requires would require compensation by other processes to fulfil the δ^{13} C constraint, e.g. a significant substantial decline of wild fire activity in the northern hemisphere without such evidence. In the mid-Holocene (8-5.5 ka BP) the deconvolutioninversion shows a decline of the southern hemisphere CH₄ emissions and an ongoing shift of the δ^{13} C emission signatures towards lighter values, which might be related to the southward migration of the ITCZ and the change of the C3to-C4 plant ratio altering the carbon isotopic composition of the precursor material for CH₄ production. During the second half of the Holocene (5.5-1 ka BP), which shows a major increase in CH₄ emissions accompanied by a shift towards higher δ^{13} C signatures in the southern hemisphere, our results favour the 'Holocene anomaly'CH₄ rise to be caused by natural CH₄ emissions (both wetlands and wildfires) in the southern tropics rather than by early rice agriculture in east Asia.

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The CH₄ stable isotope data provide a powerful constraint on the Holocene CH₄ system (e.g. to benchmark CH₄ emission models), however, the relatively large uncertainty ranges in the calculated emission parameters still limit our ability to draw more robust conclusions, on temporal variations. The deconvolution would-probably benefit from an increased temporal resolution in the southern hemisphere δD -CH₄ record. However, since the concentration data have large influence on 15 all six deconvolutioninversion parameters, we see the largest potential to furtherbetter constrain the Holocene CH₄ emissions by actually improving the [CH4] records. Better IPD_{CH4} data measured on the same analytical system during the same measurement campaign in high temporal resolution, which ensures an accurate synchronisation (such as the data by Mitchell et al. (2013) for the last 2,800 years), would bring us a large step forward in further constraining the Holocene CH4 cycle. This is especially true for the Greenland ice cores within the brittle ice zone (e.g. GISP2 records for depths between 650 and 1400 m)., e.g. GISP2 records for depths between 650 and 1400 m, equivalent to the time window 2.7-8.1 ka BP where individual 20 samples subject to modern air intrusion may compromise the Greenland CH4 record. The new EGRIP ice core currently drilled in northeast Greenland with much lower accumulation, hence a brittle ice zone located at older ages may help to improve the CH4 record in this time interval in the future. Alternatively, a high accumulation ice core from Greenland where ice older than 2.7 ka BP is found below the brittle ice zone would circumvent this issue and allow for uncompromised CH4 measurements

Data availability. Data presented in this study are available at (URL Pangaea page)https://www.pangaea.de and https://www.ncdc.noaa.gov

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²⁵ over the Holocene.

6 References

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Figure 1: CH₄ concentration data compilation. Discrete [CH₄] data over the Holocene used for the CH₄ data compilation on synchronised gas ages scales and corrected for offsets between datasets (references are listed in Table 1). The δ¹⁸O-H₂O data from the layer-counted NGRIP ice core have been used as an absolute time marker for fast climatic changes (e.g. the Younger Dryas-Holocene transition and the 8.2 ka event) assuming an essentially synchronous change in rapid temperature variations and atmospheric CH₄. The markers at the bottom of the plot indicate the tie points of the synchronised age scales (see Table 3).





Figure 2: Artificial time series and deconvolutioninversion results. Artificial tropospheric [CH4] (a) and stable isotope time series (b), (c) used as input for the deconvolution box-model inversion. The calculated mean hemispheric CH4 emission strengths (d) and their isotopic signatures (e), (f) show the dependence on the different input parameters. Three experiments are performed: all input values constant (solid lines), all constant but the northern tropospheric CH4 isotope values varied (dashed lines) and all constant but [CH4] varied (dotted lines). To be consistent with the wording in the text, time is running from right to left. Note the inverted y-axes in the panels (b), (c), (e) and (f).





Figure 3: Data and deconvolution. The panels on the left (a-c) show the concentration and the two stable isotope signatures of CH₄ measured in ice cores representing the tropospheric values over the Holocene. The concentration data are complemented with data from other studies (see Table 1). All three parameters were measured on ice cores from Greenland (red) and form Antarctica (blue and cyan) representing the northern and the southern troposphere, respectively. A spline function with a cutoff period of 3000 years has been used to calculate the smoothed evolution of the tropospheric signals represented by the lines and the 1 σ error bands. A deconvolution modeland 2 σ error bands. The panels (g), (h) and (i) show the IPDs of all three parameters also with their 1 σ and 2 σ error bands. A box-model inversion is used to calculate the hemispheric CH₄ emissions shown on the panels on the right. In panel (d) the emission strengths are shown for both hemisphere boxes together with the total CH₄ emissions. Panel (e) and (f) show the mean isotopic signatures of the emission for both stable isotopes of CH₄. All the error bands are present 1 σ and 2 σ uncertainties. Note the inverted y-axes in the panels (b), (c), (e) and), (f), (h) and (j). Red colours refer to northern hemisphere records, blue colours to southern hemisphere trecords throughout the manuscript.



Figure 4: Comparison of stable CH₄ isotope data. This study's stable CH₄ isotope values of the NGRIP ice cores are shown together with the data published by Sowers (2010)-) using the GISP2 ice core. The δD-CH₄ data sets (panel (a)) differ in the absolute level and the variance, whereas the δ¹⁵C-CH₄ data (panel (b)) are in good agreement. Note that all data are corrected for gravitational fractionation in the firm
 column. All data of this study are free of Krypton interference, whereas the GISP2 δ¹⁵C-CH₄ record has been subsequently corrected for it (Schmitt et al., 2013). Note the inverted y-axes.











Figure 6: Sensitivity study "ITCZ position". In the panels (a-c) the data splines with the 1σ uncertainty error bands of the measured tropospheric values are shown. In panels (d-f) the calculated strengths and the isotopic signature of the emissions are shown for the three different φ_{ITCZ} and the best guess value for θ to investigate the sensitivity of the deconvolutioninversion to a shift of the ITCZ. Note the inverted y-axes in the panels (b), (c), (e) and (f).

Table 1: Datasets used for CH4 data compilation. The CH4 offset correction is added to the data to correct for inter-laboratory and time period-related offsets (see section 2.3.2). Additional datasets used for the time matching (e.g. covering the anthropogenic CH4 increase or the Younger Dryas-Holocene transition) are listed. *GRIP data compilation done by J. Flückiger in 2001 containing data from various publications, e.g. Blunier et al. (1995), Chappellaz et al. (1993, 1997).

Data used for CH ₄ compilation						
			CH4 shiftoffset			
Data set name	Data type	Reference	(ppb)			
NGRIP_Be	NGRIP ice core	This study	0			
GRIP_Fl01	GRIP ice core	GRIP data Flückiger, 2001*	17.69			
GISP2_Br96	GISP2 ice core	(Brook et al., 1996)	34.13			
GISP2_Br09	GISP2 ice core	(Brook, 2009)	-1.81			
GISP2_Mi13	GISP2 ice core	(Mitchell et al., 2013)	-6.15			
TALDICE_Be17	TALDICE ice core	This study	0			
TALDICE_Sc10	TALDICE ice core	(Schilt et al., 2010)	-2.09			
EDML_Sc10	EDML ice core	(Schilt et al., 2010)	-4.63			
WAIS_Mi13a	WAIS (WDC05A) ice core	(Mitchell et al., 2013)	-6.15			
WAIS_Mi13b	WAIS (WDC06A) ice core	(Mitchell et al., 2013)	-6.15			
EDC_Fl02	EDC ice core	(Flückiger et al., 2002)	4.23			
Additional data used for time matching						
Data set name	Data type	Reference				
NGRIP_d18O	NGRIP ice core $\delta^{18}O(H_2O)$	(Rasmussen et al., 2006; Vinther et al., 2006)				
NGRIP_Ba12	NGIRP ice core	(Baumgartner et al., 2014)				
NEEM_Bu12	NEEM firn model	(Buizert et al., 2012)				
NOAA_Alert	Alert flasks	(Dlugokencky et al., 2017)				
LawDome_Ma06	Law Dome firn and ice core	(MacFarling Meure et al., 2006)				
CapeGrim_csiro	Cape Grim flasks	CSIRO (27.11.2017)				

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Table 2: Ice cores and age scales. Short names of the ice cores and initial gas age scales used for the CH4 compilation.

Ice core	Initial age scale	References
NGRIP	AICC2012 ice core chronology	(Bazin et al., 2013; Veres et al., 2013)
GRIP	GICC05	(Rasmussen et al., 2014; Seierstad et al., 2014)
GISP2	Mitchell, Meese/Sowers	(Meese et al., 1994; Mitchell et al., 2013)
TALDICE	AICC2012 ice core chronology	(Bazin et al., 2013; Veres et al., 2013)
EDML	AICC2012 ice core chronology	(Bazin et al., 2013; Veres et al., 2013)
EDC	AICC2012 ice core chronology	(Bazin et al., 2013; Veres et al., 2013)
WAIS	Mitchell, Buizert WD2004	(Buizert et al., 2015; Mitchell et al., 2013)

NG	RIP	GR	IP	GIS	SP2	TAL	DICE	EDI	ML	EI	C	WA	IS
294	0	273	-40	2752	-5	127	-134	2747	0	349	-225	2752	-5
11800	-150	8157	30	8172	-15	2835	-88	3754	39	2959	-212	3753	40
		11606	34	11771	-165	3825	-32	4538	63	3961	-168	4532	69
						4596	5	5279	130	4741	-140	5269	140
						5325	84	6008	215	5478	-69	5993	230
						6042	181	6361	129	6204	19	6344	146
						6389	101	8107	36	6555	-65	8079	64
						8106	37	8164	22	8292	-149	8228	28
						8252	4	8254	2	8353	-164	9869	-13
						9876	-20	9875	-19	8438	-182	10391	110
						10393	108	10391	110	9397	-179	11591	49
						11580	60	11576	64	9989	-133		
										10462	39		
										11547	93		

Table 3: Time synchronisation. The points of the reference time scale for the individual ice cores (left column, in a BPa BP) and time corrections (right column, in a) applied with respect to the core dependent time scale listed in Table 2. The correction values are linearly interpolated between the tie points and held constant before the earliest and after the latest tie points.

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Removed data						
age		CH_4				
depth (m)	(aBP<u>a BP</u>)	(ppm<u>ppb</u>)	dataset			
655.1	2731.9	698.8	GISP2_Br09			
736.1	3199.1	680.8	GISP2_Br96			
819.7	3694.8	678.9	GISP2_Br09			
932.7	4411.4	661.8	GISP2_Br09			
943.1	4474.0	643.8	GISP2_Br09			
992.3	4804.6	677.4	GISP2_Br96			
1019.0	4989.5	642.0	GISP2_Br09			
1059.0	5272.2	631.8	GISP2_Br09			
1177.3	6186.3	639.6	GISP2_Br09			
1194.2	6314.4	653.8	GISP2_Br96			
1202.9	6382.8	652.1	GISP2_Br96			
1253.9	6810.3	687.8	GISP2_Br09			
1274.8	6978.6	677.8	GISP2_Br96			
1389.7	7993.8	675.8	GISP2_Br96			
1392.0	8011.9	682.2	GISP2_Br96			

Table 4: GISP2 outliers. Data points that were removed from the two GISP2 data sets by Brook et al. (2009) and Brook (1996) that were removed due to elevated CH₄ values probably due to contamination related to the ice sample quality in the brittle zone. Please note that the age and the concentration values are corrected for the age shift and the CH₄ correction, respectively.

Table 5: Values used in our deconvolution box-model inversion, Relative strengths, isotopic fractionations (${}^{2}\varepsilon$ and ${}^{13}\varepsilon$) (Brenninkmeijer et al., 1995; Cantrell et al., 1990; Feilberg et al., 2005; Gierczak et al., 1997; Irion et al., 1996; Tyler et al., 1994; Quay et al., 1999) and north-south distributionratio of the individual sink processes (ratio n/s) and the parameters for each hemisphere used in our CH₄ deconvolution-box-model inversion.

Formatiert: Schriftart: Nicht Fett

sink processes							
	rel. strength	$^{2}\varepsilon$ (‰)	$^{13}\varepsilon(\%_{0})$	ratio n/s			
S _{OH}	0.820	-231	-5.4	246.0/246.0			
S _{strat}	0.092	-170	-13.1	27.5/27.5			
S _{soil}	0.047	-80	-22.0	19.6/8.4			
s _{cl}	0.042	- <u>315459</u>	- 55<u>58</u>.0	9.6/15.4			
numbers used for deconvolutioninversion equations							
	$\tau_x(a)$	$^{2}\varepsilon_{x}$ (%0)	$^{13}\varepsilon_{x}(\%_{0})$	r_x			
		-					
north box	7.60	218.3 –222.9	- <u>-</u> 7. 52<u>57</u>	0.456			
		-					
south box	9.22	225.5 -232.9	- <u>-</u> 1. 72 80	0.544			