



## A new Himalayan ice core CH<sub>4</sub> record

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# A new Himalayan ice core CH<sub>4</sub> record: possible hints on the preindustrial latitudinal gradient

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## Abstract

Two ice cores recovered from the Himalayan East Rongbuk (ER) Glacier on the north-east saddle of Mt. Qomolangma (Everest) ( $28^{\circ} 01' N$ ,  $86^{\circ} 58' E$ , 6518 m a.s.l.) give access to a tentative record of past Himalayan atmospheric mixing ratio of  $CH_4$  spanning the past 1200 yr. The major part of the record is affected by artefacts probably due to in-situ production. After selecting what may represent the true atmospheric mixing ratio, an average of  $749 \pm 25$  ppbv of  $CH_4$  is estimated for the late preindustrial Holocene, which is  $\sim 36 \pm 17$  ( $\sim 73 \pm 18$ ) ppbv higher than the atmospheric levels recorded respectively in Greenland and Antarctic ice cores. A comparison of these new data with model simulations of the  $CH_4$  latitudinal gradient suggests either that the models do not get a correct balance between high and low latitude  $CH_4$  sources, or that the filtered  $CH_4$  profile from the ER cores remains infected by small artefacts.

## 1 Introduction

Methane records from polar ice cores provided a wealth of information about its natural variability over several climatic cycles (Loulergue et al., 2008). Natural  $CH_4$  variations are partly related to orbital and millennial variations in tropical and boreal wetland extent/ $CH_4$  emissions. There is an ongoing debate on the cause for the pre-industrial increase in atmospheric  $CH_4$  mixing ratios, which has started about 5000 yr ago. Early farming has been hypothesized to contribute to this increase in atmospheric  $CH_4$  mixing ratios during the last 5000 yr (Ruddiman et al., 2008), which is also postulated to be of natural origin (Singarayer et al., 2011). High resolution measurements of air samples from the Antarctic Law Dome ice cores and firn air provided a detailed record of  $CH_4$  variability during the late preindustrial Holocene (LPIH), suggesting that LPIH  $CH_4$  changes are mainly a response to changing climate, with small contributions from human activities (MacFarling-Meure et al., 2006). A new high-precision, high-resolution record of atmospheric  $CH_4$  from the West Antarctic Ice Sheet (WAIS) Divide ice core

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covering 1000–1800 AD indicates that times of war and plague when large population losses could have reduced anthropogenic emissions are coincident with short periods of decreasing global CH<sub>4</sub> mixing ratios (Mitchell et al., 2011). Moreover, two ice cores from Greenland provide a high-resolution  $\delta^{13}\text{C}$  record of CH<sub>4</sub> during the last two millennia, confirming changes in pyrogenic and biogenic CH<sub>4</sub> sources due to both natural climate variability and changes in human population and land use (Sapart et al., 2012).

Records of CH<sub>4</sub> mixing ratios in the tropical or sub-tropical atmospheres could provide an additional insight in latitudinal CH<sub>4</sub> gradients and further constrain the attribution of pre industrial variations. So far, few attempts have been made to reconstruct CH<sub>4</sub> levels from tropical glaciers. There, the integrity of the ice core records is indeed exposed to artefacts due to in-situ production by methanogenic bacteria eventually activated by surface melt (Skidmore et al., 2000; Campen et al., 2003). Very few subtropical glaciers may have the potential to provide a reliable CH<sub>4</sub> preindustrial value. Warm summer temperature and elevated concentrations of impurities are the main caveats of most alpine glacier settings, precluding to access an undisturbed CH<sub>4</sub> record in air bubbles. The first record obtained from an Himalayan ice core, i.e., the Dasuopu ice core (28° 23' N, 85° 43' E, 7100 m a.s.l.), gave an average CH<sub>4</sub> preindustrial mixing ratio of 825 ppbv (later modified to 782 ± 40 ppbv; Li et al., 2010) during the last two millennia (Yao et al., 2002). This value is ~ 120 ppbv (or ~ 160 ppbv) higher than its Greenland (or Antarctic) counter parts. While such difference is comparable to the modern inter-polar CH<sub>4</sub> gradient maximum of ~ 140 ppbv (Dlugokencky et al., 1994), simulations of preindustrial CH<sub>4</sub> suggest much lower methane mixing ratios in the subtropical areas than that in the higher northern latitudes (Houweling et al., 2000). The Dasuopu record is also characterized by a very large CH<sub>4</sub> variability at small depth scale, which cannot reflect the true atmospheric evolution as gas diffusion and trapping smooth out any sub-decadal variability (e.g., Buizert et al., 2012). Therefore the average CH<sub>4</sub> mixing ratio deduced from the Dasuopu record is clearly biased. Detailed CH<sub>4</sub> measurements along the last interglacial section of the NEEM core (Greenland) and along specific sections of the Dye 3 core (Greenland) have shown large peaks associated with melt

layers (NEEM community members, 2013). The Dasuopu record is most probably similarly affected by artefacts generating local peaks and artificially increasing the average CH<sub>4</sub> mixing ratios observed along the core. Here we present new CH<sub>4</sub> results spanning the last 1200 yr obtained from two new Himalayan high-elevation ice cores, in an attempt to re-evaluate the CH<sub>4</sub> mixing ratio difference between the Himalayas and polar sites during the LPIH.

## 2 The ice cores

We recovered one ice core to bedrock (117.06 m deep, hereafter “Core2001”) in 2001, and two more cores to bedrock (108.83 m, hereafter “Core2002”, and 95.80 m deep, respectively) in 2002 from the East Rongbuk (ER) Glacier on the northeast saddle of Mt. Qomolangma (Everest) using an electromechanical drill in a dry hole. Diameters are 6.9 cm and 9.4 cm for Core2001 and Core2002, respectively. Ice layers are horizontal along the cores, suggesting negligible ice dynamical deformation. To accommodate gas analyses, we maintained the temperature of the cores below  $-5^{\circ}\text{C}$  from the time of drilling until analysis.

The ER Glacier covers an area of 48.45 km<sup>2</sup> with a length of 14 km. Its equilibrium line is at 6250 m a.s.l., and is amongst the highest on Earth. A repeating survey with a Sokkia GSS1A Global Positioning System (accuracy of  $\pm 5\text{mm}$  over distance up to 10 km) in 1998 and 2002 did not identify horizontal movement at the drill site. The average annual net accumulation is  $\sim 400\text{mm}$  water equivalent as determined by snow pit and ice core studies (Kaspari et al., 2008). Borehole temperatures in the 108.83 m core ranged from  $-8.9^{\circ}\text{C}$  at 10 m to a minimum of  $-9.6^{\circ}\text{C}$  at 20 m then warmed slightly to  $-8.9^{\circ}\text{C}$  at the bottom. However, snow melting occurs at the ER drilling site in summer and percolates downwards. Refreezing of melt-water speeds up the later stages of snow transformation, resulting in low gas content in the summer strata, which is used as an indicator of summer temperature trend (Hou et al., 2007).

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### 3 Measurements and results

The CH<sub>4</sub> measurements were performed at LGGE (Laboratoire de Glaciologie et Geophysique de l'Environnement) using an automated wet extraction method. The analytical method is described in detail by Chappellaz et al. (1997). In brief, an ice sample (~ 40–50 g) is melted under vacuum in a glass container sealed with a viton O-ring and then refrozen from the bottom with an alcohol bath at –50 °C to expel gas dissolved in the water (extraction efficiency of 99 %). The gas sample is then expanded into an evacuated sample loop and injected into the gas chromatograph (Varian 3300) with a multiposition Valco valve. N<sub>2</sub> + O<sub>2</sub> + Ar and CH<sub>4</sub> are separated on a Porapak-N column (3 m, 1.8 mm ID) and detected with a Thermal Conductivity Detector (TCD) and a Flame Ionization Detector (FID), respectively. Each sample was measured three times. Calibration is performed against a standard gas (Air Liquide) with 499 ppbv of CH<sub>4</sub> in air. Linearity of the FID was checked by measuring CSIRO standard gases with CH<sub>4</sub> mixing ratios in air ranging from 394 to 1679 ppbv. Blanks of the experimental procedure, determined by adding the 499 ppbv gas standard to artificial bubble-free ice, amount to 20 ± 13 (2σ) ppbv and are subtracted from all results.

We analyzed 77 and 112 samples from Core2001 and Core2002, respectively. All the samples are taken below the bubble close-off depth (28.4 m and 26.3 m for Core2001 and Core2002, respectively) to the bottom. The CH<sub>4</sub> profiles are shown as a function of depth in Fig. 1, and the CH<sub>4</sub> profile of 75 samples from the Himalayan Dasuopu ice core that were measured by the same set of instrument at LGGE is included for comparison (Yao et al., 2002; Li et al., 2010). Similar features are observed for all the three Himalayan ice cores: CH<sub>4</sub> values measured on neighboring samples exhibit a large scatter that can exceed hundreds of ppbv, incompatible with normal firn air diffusion and gas trapping processes and pointing to local artifacts causing anomalously high CH<sub>4</sub> values, as recently observed on a shallow ice core from the North Greenland Eemian project (NEEM-2011-S1) by continuous CH<sub>4</sub> measurements (Rhodes et al., 2013), NEEM interglacial ice and Dye 3 ice sections (NEEM community members,

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2013). In the case of melting, atmospheric gases diffuse into the meltwater which is subsequently refrozen within the glacier, with a methane/air solubility ratio of 1.9 for water at 0 °C in equilibrium with the atmosphere. Similarly to the NEEM last interglacial ice, a reverse relationship is apparent between CH<sub>4</sub> and gas content measurements that were performed simultaneously by gas chromatography on the same set of samples (Hou et al., 2007; Fig. 2), indicating that the low CH<sub>4</sub> mixing ratios correspond to samples with relatively high gas content i.e., from winter ice layers whose air content was rarely impaired by melting (Hou et al., 2007; Li et al., 2010). Although methanogens that could be cultured oligotrophically at low temperatures (0.3 to 4 °C) were identified from the debris-rich basal ice layers of a high Arctic glaciers (Skidmore et al., 2000), relatively low concentrations of culturable bacteria were observed in the ER ice core (Zhang et al., 2007). On the other hand, Rhodes et al. (2013) have shown that local CH<sub>4</sub> peaks are associated with organic-rich ice layers. The number of bacteria may thus not be the limiting factor to produce CH<sub>4</sub> anomalies, but instead the amount of organic substrate may play a more important role. In the following, we attempt to filter out the ER CH<sub>4</sub> series from artefacts, in order to tentatively reconstruct a pre-industrial and regional atmospheric CH<sub>4</sub> level using the baseline CH<sub>4</sub> profile of ER Core2002.

Corrections may be required to account for changes in firn thermal and gravitational diffusion. The magnitude of this gravitational fractionation is proportional to the thickness of the firn diffusive zone, which can be estimated using <sup>15</sup>N/<sup>14</sup>N measurements in N<sub>2</sub> (δ<sup>15</sup>N<sub>2</sub>). Twenty samples randomly collected in the depth range of 102–117 m of Core2001 yield a δ<sup>15</sup>N<sub>2</sub> maximum of 0.089 ‰, resulting in a negligible correction of about 0.4 ppbv for the LPIH samples (calculation following Sowers et al., 1989). Thermal fractionation is expected to have only a minor impact on the composition of bubble air in tropical glaciers (Campen et al., 2003). We conclude that the magnitude of the gravitational and thermal fractionation is negligible and their effect was ignored.

## 4 Dating

Core2002 was annually dated back to 1534 AD at a depth of 98.0 m using seasonal variations of  $\delta D$  and soluble ions, and the timescale was verified by identifying large volcanic horizons. Below 98.0 m, annual layer counting is not possible due to layer thinning. Thus prior to 1534 AD the core was dated using an ice flow model (Kaspari et al., 2008). Dating uncertainties are estimated negligible at 1963 AD based on a volcanic Bi horizon from the Agung eruption, and  $\pm 5$  yr at 1534 AD based on repeated annual layer counting (Kaspari et al., 2008).

We estimate a present gas-ice age difference of  $\sim 30$  yr based on the firn gas measurements at the Himalayan Dasuopu Glacier and calculation from the present accumulation rate and temperature (Hou et al., 2007). The gravitational signal in the isotopic composition of atmospheric air in the firn can be used to infer information about the past thickness of the diffusive column (Caillon et al., 2001).  $\delta^{15}N_2$  measurements along the ER cores give an average of  $0.035 \pm 0.025\text{‰}$  ( $1\sigma$ ) and a maximum of  $0.089\text{‰}$ , resulting in  $\sim 8$  to  $20$  m of diffusive column thickness with a firn temperature of  $-8.9^\circ C$ . Although there might be limitation in this approach due to the existence of melt layers along the Himalayan ice cores (Hou et al., 2007; Li et al., 2010), the small  $\delta$  values imply a relatively constant firn thickness that controls the gas-ice age difference. Thus we simply assume that the gas age is 30 yr younger than its corresponding ice age for the studied period.

Accurate dating of Core2001 is not available yet. Therefore we only discuss below the  $CH_4$  record of Core2002 covering the last 1200 yr.

## 5 Discussion

A large part of the  $CH_4$  profile of Core2002 (as well as those of Core2001 and Dasuopu) is affected by in-situ artefacts. We thus attempt to define a criterion allowing to exclude the measurements affected by these artefacts and to retain those which may

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reflect the atmospheric composition at the time of gas trapping. We firstly excluded 34 measurements with one standard deviation > 20 ppbv for Core2002, due to unsatisfactory chromatography, system leaks, or extremely low gas content caused by possible partial snow melting. There are 6 samples collected at the depth of 26.3 m, 29.6 m, 48.0 m, 48.1 m, 83.1 m, 90.4 m, respectively, with CH<sub>4</sub> mixing ratios over 1800 ppbv (i.e., the modern CH<sub>4</sub> mixing ratio measured at Mt. Waliguan on the Tibetan Plateau – one of the CMDL Baseline Stations), which are thus removed. Such anomalously high mixing ratios of CH<sub>4</sub> were also observed in a Sajama ice core from the Northern end of the Bolivian altiplano (18° 6′ S, 68° 53′ W, 6048 m a.s.l.), and suggested to be due to metabolizing microorganisms (Campen et al., 2003). Rhodes et al. (2013) identified abrupt (20–100 cm depth interval), high amplitude (35–80 ppbv excess) CH<sub>4</sub> spikes in the Greenland NEEM-2011-S1 ice core, and suggested that these spikes may result from very localised biological in-situ production in polar, glacial ice.

We then filter out the remaining artifact by using a criterion on the deduced CH<sub>4</sub> annual variation rate. The CH<sub>4</sub> record from WAIS Divide, Antarctica, suggested a maximum growth rate of 1 ppbvyr<sup>-1</sup> during the period from the preindustrial back to 1000 A.D. (Mitchell et al., 2011). We thus adopt ±1 ppbvyr<sup>-1</sup> as a maximum acceptable change between adjacent Core2002 samples, possibly reflecting atmospheric composition changes. We then reject the high CH<sub>4</sub> values when the growth rate calculated before or after them overruns this criterion. The procedure obviously lead to selecting low CH<sub>4</sub> values, as we assume that there is no identified process able to consume CH<sub>4</sub> preserved in the ice core bubbles. As to the industrial period, Etheridge et al. (1998) identified high CH<sub>4</sub> growth rates peaking at ~ 17 ppbvyr<sup>-1</sup> in 1981 AD. Thus we adopt this value as a maximum growth rate between adjacent Core2002 samples over the recent increasing part of the record. Finally 15 measurements (i.e., the solid circles of Figs. 1 and 2) are retained in the full record, satisfying the selection criterion and possibly reflecting the atmospheric composition at the ER site (Fig. 3). Among them, nine measurements date from the LPIH, providing an average CH<sub>4</sub> of 749 ± 25 ppbv. This is 76 ppbv (Yao et al., 2002) or 33 ppbv (Li et al., 2010) lower than its counterpart of the



Dasuopu ice cores, albeit  $\sim 36 \pm 17$  ( $\sim 73 \pm 18$ ) ppbv higher than the GRIP (Law Dome) LPIH records. Core2002 and GRIP CH<sub>4</sub> measurements were performed in the same laboratory, with the same extraction and analysis technique and identical standard gas, so that this CH<sub>4</sub> comparison can be considered in terms of absolute values.

Preindustrial CH<sub>4</sub> simulations with a three-dimensional chemistry-transport model (Houweling et al., 2000) provide a preindustrial annual average CH<sub>4</sub> mixing ratio of 711–717 ppbv for the grid box covering the Himalayas, and 733 ppbv for the Greenland grid box. Although problems may arise from the use of a coarse grid model, especially for the high Himalayas, the modeling results are broadly consistent with the Greenland LPIH level observed in the GRIP core, but much lower than our estimation from the ER2002 data selection. Similar results have also been obtained using the LMDz/INCA climate/chemistry model (Kaplan et al., 2006). Present day CH<sub>4</sub> simulations along the surface air sampling sites of the NOAA Global Monitoring Network roughly distributed along latitudes (Fig. 4) perform well in comparison with CH<sub>4</sub> gradient observations (Dlugokencky et al., 1994). Methane retrievals from the Scanning Imaging Absorption Spectrometer for Atmospheric Chartography (SCIAMACHY) suggest that the model underestimates CH<sub>4</sub> mixing ratios over the Himalayas and Tibetan Plateau, potentially resulted from systematic errors related to topography of the high altitudes and model deficiencies in the stratospheric CH<sub>4</sub> mixing ratios (Bergamaschi et al., 2009). The lower LPIH CH<sub>4</sub> mixing ratios than the ER ice core data selection (Fig. 4) may also result from an underestimation of the magnitude and development of boreal CH<sub>4</sub> sources in the LMDz-INCA (Kaplan et al., 2006). Alternatively, the ER 2002 CH<sub>4</sub> profile presented here after data selection may still be affected by local artefacts, although less pronounced than for the rejected data. The only way to ascertain this will be to run continuous-flow CH<sub>4</sub> measurements with repeated analyses along parallel ice sticks (as performed on the NEEM-2011-S1 Greenland ice core by Rhodes et al., 2013) using Himalayan ice cores from different sites. Replicable features in the resulting CH<sub>4</sub> record will bring the required level of confidence for the comparison with polar records.

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Recovering suitable ice cores becomes an emergency, due to the rapid warming of high altitude glaciers (Kehrwald et al., 2008).

## 6 Conclusions

The Himalayan ER ice cores allowed us to tentatively retrieve a background record of High Himalayan atmospheric CH<sub>4</sub> mixing ratios covering the last 1200 yr. Latitudinal gradients are expected to provide an independent observational constraint from subtropical regions for assessing the relative weight of different CH<sub>4</sub> sources with time, in addition to the temporal trend, inter-polar gradient, and stable isotopic ratio evolution. After removing obvious outliers and attempting to select interpretable data based on the CH<sub>4</sub> rate of change, the resulting CH<sub>4</sub> profile depicts a positive gradient between the Himalayan latitude and Greenland, which is not consistent with simulations by climate/chemistry coupled models. Such model/data mismatch may arise from model caveats related to the magnitude of boreal methane fluxes or from the possibility that the selected ER CH<sub>4</sub> data are also affected by less pronounced artefacts.

More ice core CH<sub>4</sub> records from high mountains (e.g., high central Asia, the Alps, Caucasus, Alaska, the Andes) are necessary for investigating latitudinal gradients and obtain robust features. It becomes an urgent need as alpine glaciers are warming fast and become less suitable for trace-gas investigations due to disturbances related with melting or in-situ production. Future measurements should rely on continuous-flow techniques in order to pick up all details regarding the depth/CH<sub>4</sub> trends, the replicability of the records, and to eventually extract a suitable atmospheric reconstruction.

**Supplementary material related to this article is available online at:**  
**<http://www.clim-past-discuss.net/9/2471/2013/cpd-9-2471-2013-supplement.zip>**

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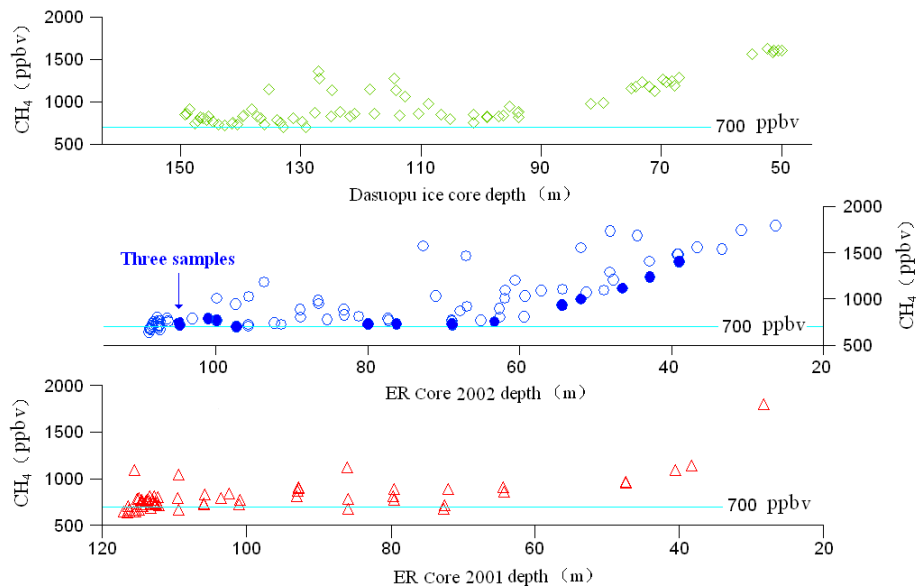
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**Fig. 1.** CH<sub>4</sub> profiles from three Himalayan ice cores. The average 700 ppbv preindustrial CH<sub>4</sub> level depicted by polar ice cores is indicated with a horizontal light blue line. The solid circles in the ER Core2002 panel stand for the filtered 15 samples that are used for the discussion.

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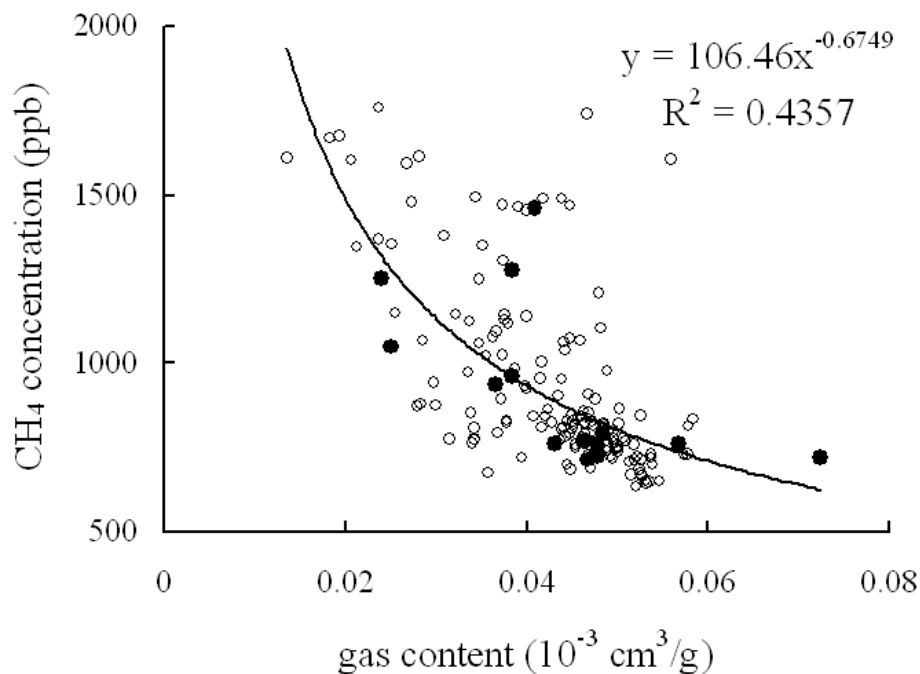
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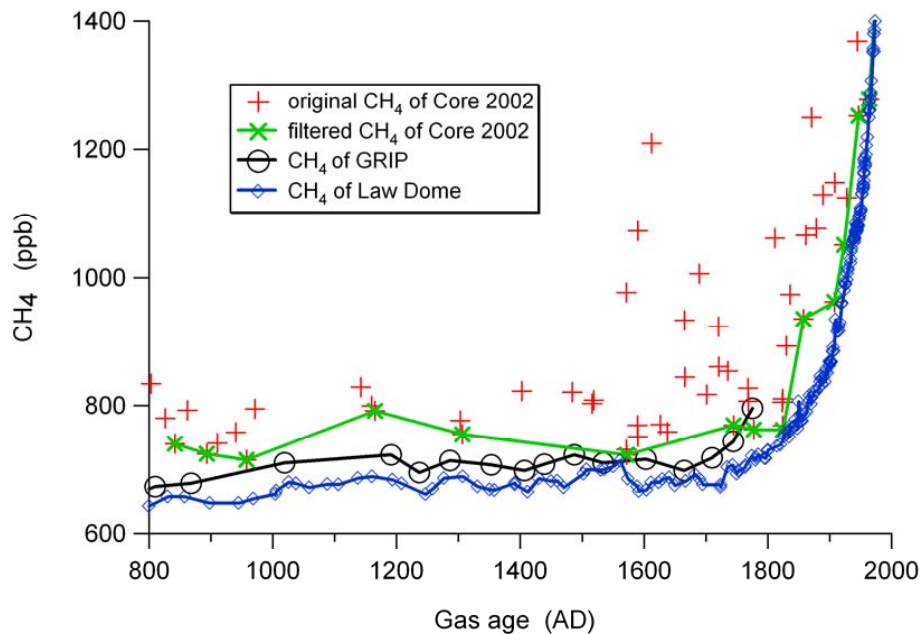
**Fig. 2.** Distribution of CH<sub>4</sub> mixing ratios against gas content of the same samples in ER ice cores, with an exponential fit. Samples with CH<sub>4</sub> mixing ratios over 1800 ppbv are excluded. The solid circles stand for the filtered 15 samples that are used for the discussion.

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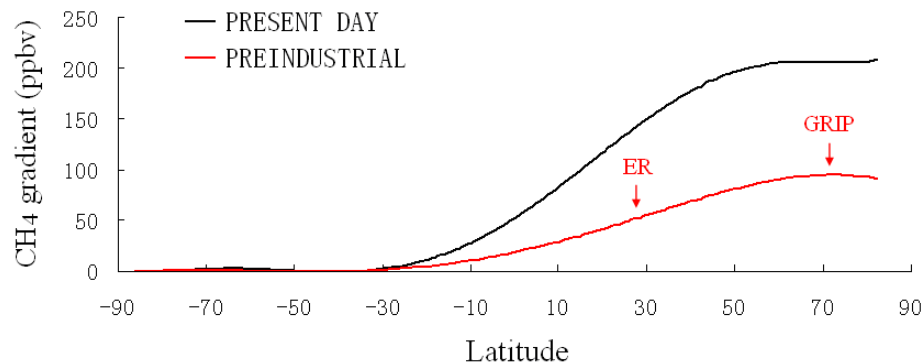


**Fig. 3.** The background CH<sub>4</sub> profiles of Core2002 with comparison to the GRIP (Chappellaz et al., 1997) and Law Dome ice core (MacFarling-Meure et al., 2006) records.

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**Fig. 4.** CH<sub>4</sub> latitudinal gradients for the Present-Day and the Preindustrial periods simulated with the LMDz-INCA climate/chemistry coupled model.

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