Evolutionary History of Atmospheric CO₂ during the Late Cenozoic from Fossilized Metasequoia Needles

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Abstract

The change in ancient atmospheric CO₂ concentrations provides important clues for understanding the relationship between the atmospheric CO₂ concentration and global temperature. However, the lack of CO₂ evolution curves estimated from a single terrestrial proxy prevents the understanding of climatic and environmental impacts due to variations in data. Thus, based on the stomatal index of fossilized Metasequoia needles, we reconstructed a history of atmospheric CO₂ concentrations from middle Miocene to late Early Pleistocene when the climate changed dramatically. According to this research, atmospheric CO₂ concentration was stable around 330–350 ppmv in the middle and late Miocene, then it decreased to 278–284 ppmv during the Late Pliocene and to 277–279 ppmv during the Early Pleistocene, which was almost the same range as in preindustrial time. According to former research, this is a time when global temperature decreased sharply. Our results also indicated that from middle Miocene to Pleistocene, global CO₂ level decreased by more than 50 ppmv, which may suggest that CO₂ decrease and temperature decrease are coupled.

Introduction

Carbon dioxide (CO₂) is an important greenhouse gas that influences the surface temperature of the Earth [1]. The 5th report of IPCC concluded [2] that the present positive radiative forcing is unequivocally caused by anthropogenic increases in atmospheric CO₂ concentration and that it influences the climate [3,4]. Estimating the impact of high CO₂ concentration on global environmental systems is the first step to propose solutions for the present global climate change. This impact can be unraveled by a better understanding of the relationship between the paleo-atmospheric CO₂ concentration (paleo-[CO₂] atm) and ancient climate change.
A lot of research has involved the estimation of paleo-[CO$_2$]$_{atm}$ to understand the correlation between CO$_2$ and global warming. To obtain the paleo-[CO$_2$]$_{atm}$ values three major approaches have been used: (1) geochemical modeling (GCS) [5–7], (2) composition measurements of air trapped in ice cores [8], and (3) various proxies (reviewed in [9]). Geochemical modeling (GCS) can reconstruct paleo-[CO$_2$]$_{atm}$ but for long geological time scales its resolution cannot be fine enough to show the details of paleo-[CO$_2$]$_{atm}$ fluctuation [10]. Ice core analysis is the most reliable method to measure paleo-[CO$_2$]$_{atm}$ directly, but is only applicable after 0.8 Ma [8]. Several CO$_2$ proxies have been used to estimate paleo-[CO$_2$]$_{atm}$ such as the carbon isotope composition of phytoplankton, the boron (B) isotope composition of fossil foraminifera, the carbon isotope composition of carbonates in paleosol, and the stomatal parameters of fossil leaves [11]. High resolution records for CO$_2$ can be obtained from marine sediments with the two former proxies, but these do not directly show the paleo-[CO$_2$]$_{atm}$. The latter two proxies are terrestrial-based proxies that reflect paleo-[CO$_2$]$_{atm}$ directly, although they rarely provide continuous paleo-[CO$_2$]$_{atm}$ records for a long geological time. Therefore, while there is a consensus on the general tendency of the Cenozoic paleo-[CO$_2$]$_{atm}$ changes, the estimated paleo-[CO$_2$]$_{atm}$ values vary greatly [9]. To understand the paleoclimatic system, it is important to reduce uncertainties in the relationships between paleo-[CO$_2$]$_{atm}$ and past climate [12].

Stomatal parameters (SI (stomatal index) and SD (stomatal density)) are reliable proxies to estimate paleo-[CO$_2$]$_{atm}$. In particular, SI can provide a robust indicator of terrestrial paleo-[CO$_2$]$_{atm}$ as it is independent of other environmental parameters, such as soil moisture supply, atmospheric humidity and temperature [13]. Many studies have already used the SI of different taxa to estimate paleo-[CO$_2$]$_{atm}$ such as Metasequoia Miki ex Hu et Cheng [12,14], Ginkgo Linn. [15,16], Quercus Linn. [17,18], Laurus Linn. [17,19,20], Platanus Linn. [17,21], and Typha Linn. [22]. As the relationship between the SI and paleo-[CO$_2$]$_{atm}$ is species-specific even within a single family [23] and the response sensitivities to CO$_2$ change are different in various taxa [24], it is necessary to select a single modern taxon that has survived for an extended period to reconstruct atmospheric CO$_2$ over a long geological time.

Metasequoia has exhibited an evolutionary stasis since its appearance in the Late Cretaceous [25], and fossilized Metasequoia can be considered to be conspecific with modern Metasequoia based on the morphology, biochemistry and inferred physiology [26]. Therefore, the paleo-[CO$_2$]$_{atm}$ changes over a long geological time can be determined from a correlation between the SI of Metasequoia needles and the paleo-[CO$_2$]$_{atm}$ concentration [14].

In this study, we use Metasequoia needles from seven localities in China and Japan to reconstruct continuous terrestrial paleo-[CO$_2$]$_{atm}$ changes from the middle Miocene to Pleistocene. Based on the reconstructed paleo-[CO$_2$]$_{atm}$ curve, we discuss the interaction between paleo-[CO$_2$]$_{atm}$ evolution and global environment change since the middle Miocene.

Materials and Methods

Materials

The fossilized needles of Metasequoia (Fig 1) were collected from one locality in SW China (Sanzhangtian) and six localities in central Japan (Kumagaya, Sennan, Hachioji, Konan, Tokamachi, and Ikoma sites) (Fig 2, Table 1). We confirm that our field study did not involve endangered or protected species and none of the localities which provided samples for this study are in protected areas. The Sanzhangtian locality belongs to the National land of the People’s Republic of China, and the Land and Resources Bureau of Zhenyuan County gave permission to collect fossils from this locality. The Japanese sites: Kumagaya, Hachioji, Konan, and Tokamachi are on valley floors which are public space, so no permission was required to...
conduct sampling. The Sennan and Ikoma sites belong to private owners, who gave permission for sampling.

*Metasequoia* fossils had previously been reported from all the fossil localities. Their ages were estimated based on stratigraphic studies (Sanzhangtian site), zircon fission-track methods (Kumagaya site), and regional stratigraphic correlation using magnetostratigraphy and calcareous nanoplankton stratigraphy (Sennan, Hachioji, Konan, Tokamachi, and Ikoma sites) (Table 1). For the samples, two were from the Miocene (Sanzhangtian and Kumagaya), three from the upper Pliocene (Sennan, Hachioji, and Konan), and two from the lower Pleistocene (Tokamachi and Ikoma) (Table 1). At least six different needles from different branchlets were used in the studies from each site, and the exact amount depends on the total amount of materials at each fossil site (Table 2).

Voucher specimens from the Sanzhangtian, Kumagaya, Hachioji, Konan, and Tokamachi sites are housed in the Herbarium of Kunming Institute of Botany (KUN), Chinese Academy of Science. Specimens from the Sennan and Ikoma sites are housed in the Graduate School of Horticulture, Chiba University, Japan.

**Methods**

**Pretreatment of the fossilized needles**

To remove the inorganic compounds adhering to the fossilized needles, the material was first immersed in 10%–25% Hydrochloric Acid (HCl) for two hours, then in 40% Hydrofluoric Acid (HF) for 12 hours, and in 10%–25% HCl for at least one hour. The needles were then
Fig 2. Localities where fossilized *Metasequoia* were obtained. Locality map (a) showing the seven fossil sites in China and Japan. Enlarged map (b) illustrating the central area of Japan showing the position of the six localities in Japan: Kumagaya, Sennan, Konan, Hachioji, Tokamachi and Ikoma. Different colors identify the different ages of the localities (Red: Miocene; Green: Pliocene; Blue: Pleistocene).

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rinsed with distilled water and divided into three parts, and the central piece (when available) used to obtain the cuticle.

### Cleaned cuticular membrane maceration

For the material from the Sanzhangtian, Kumagaya, and Konan sites, we followed the methods of Kerp [41] to isolate the lower cuticle of the fossilized needles. (1) The specimens were first macerated with 70% Nitric acid (HNO₃) for between a few minutes to an hour until they turned yellowish-brown. (2) Once it had been rinsed with distilled water several times, (3) the upper and the lower epidermis were separated using a needle. (4) Then, the epidermis was

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### Table 1. *Metasequoia* samples used for reconstructing paleo-CO₂ including fossil sites, ages, latitude, and longitude.

| Fossil site   | Locality                  | Latitude/Longitude | Geologic setting                                      | Epoch                          | Absolute age | Dating method                                | Voucher specimens # | Remark                                      | Reference |
|---------------|----------------------------|--------------------|-------------------------------------------------------|-------------------------------|--------------|----------------------------------------------|---------------------|---------------------------------------------|-----------|
| Sanzhangtian  | Yunnan, China              | 24°06' N, 101°13' E| Dajie Formation in the Matsuyama Group                | middle Miocene                | 10–16 Ma     | Stratigraphic study                          | SZT077, SZT156,     |
|               |                            |                    |                                                       |                               |              |                                              | SZT115, SZT127,     |                                              | [27–29]  |
|               |                            |                    |                                                       |                               |              |                                              | SZT123, SZT126      |                                             |           |
| Kumagaya      | Saitama Prefecture, Japan  | 36°08' N, 139°18' E| Yagii Formation in the Matsuyama Group                | early late Miocene            | 9–10 Ma      | Zircon fission track dating                   | YJ003, YJ005        | Includes marine bed                          | [30]      |
| Sennan        | Osaka Prefecture, Japan    | 34°24' N, 139°28' E| Lower than the Habutaki I Tephra, Osaka Group         | Late Pliocene                 | 2.8–3.0 Ma   | Magnetostratigraphy and calcareous nanoplankton stratigraphy | FT001               | Included in sediments in fluvial backmarsh | [31–33]  |
| Hachioji      | Tokyo, Japan               | 35° 40' N, 139°18' E| Kasumi Formation (below the Gauss and Matuyama Chron boundary) | Late Pliocene                 | 2.6–2.7 Ma   | Magnetostratigraphy and calcareous nanoplankton stratigraphy | BQC001               | Includes marine bed                          | [34,35]  |
| Konan         | Shiga Prefecture, Japan    | 34°59' N, 136°6' E | Horizon correlated with the Kamide I tephra bed in Kobiwako Group (just below the Gauss and Matuyama Chron boundary) | Late Pliocene                 | 2.6 Ma       | Magnetostratigraphy and calcareous nanoplankton stratigraphy | SG001, SG002        | Included in sediments in fluvial backmarsh | [36]      |
| Tokamachi     | Niigata Prefecture, Japan  | 37°07'N, 138°48'E | Middle part of the Unonna Group lower part of Olduvai paleomagnetic chron | middle Early Pleistocene      | 1.85 Ma      | Magnetostratigraphy and calcareous nanoplankton stratigraphy | 156u01               | Includes marine bed                          | [32,37,38]|
| Ikoma         | Nara Prefecture, Japan     | 34°44' N, 135°43'E | Peat layer just below the Ma 2 Marine Clay bed (MIS 25) in the Osaka Group | latest Early Pleistocene      | 0.95 Ma      | Magnetostratigraphy and calcareous nanoplankton stratigraphy | NR001               | Includes marine bed                          | [39,40]  |
treated with a 3%–5% Sodium Hypochlorite (NaClO) solution for around 10 minutes to remove the remnants of the mesophyll, vascular bundle, hypodermal layer, and epidermal cell walls. (5) According to the state of the material, 5%-10% Aqueous Ammonia (NH₃/H₂O) or 30% Hydrogen Peroxide (H₂O₂) can be used instead of the 3%–5% NaClO. (6) Finally glycerol was used to mount the separated cuticles for observation.

Cuticle observation and photography

The separated cuticles of the material from the Sanzhangtian, Kumagaya, and Konan sites were observed using a transmitted light microscope (Zeiss Axio Imager A2) and photographed with a digital camera (Zeiss AxioCam MRc). For the materials from the Sennan, Hachioji, Tokamachi, and Ikoma sites, pretreated fossilized leaves were mounted with water on slides and the lower sides of the needles were directly scanned by a confocal laser scanning microscope (Zeiss LSM710, Imager. Z2, Ar Lasser 488nm). Each field-of-view was larger than 0.03mm² [42]. Photoshop (version CS6, Adobe Systems; Mountain View, CA) was used to merge 6–12 serial images that were taken of the same area but at different focal levels.

Measurement of SI and paleo-[CO₂]atm concentration

Image J (1.43μ, Wayne Rasband, http://rsb.info.nih.gov/ij/) was used to calculate the number of epidermal cells and stomatal complexes (stomatal pore + guard cells). Then, the SI was calculated using Eq 1 [43].

\[
SI = \frac{\text{stomatal complexes number}}{\text{epidermal cell number} + \text{stomatal complexes number}} \times 100\% \\
\text{Equation 1}
\]

The SI data were used to estimate the paleo-[CO₂]atm from the middle Miocene to Pleistocene by using the species-specific, nonlinear negative correlation between atmospheric CO₂ partial pressure and SI (Eq 2) based on Royer et al. [14].

\[
\text{Paleo} - [\text{CO}_2]_{atm} = \frac{SI - 6.672}{0.003883 \times SI - 0.02897} \\
\text{Equation 2}
\]

The significant differences between the mean variance of the SI from different ages were statistically tested using the two tailed one-way ANOVA with the "LSD" option in IBM SPSS Statistics (Version 20.0).

Results

Fossilized Metasequoia needles from the early late Miocene Kumagaya site had the lowest SI value (SI = 9.80 ± 0.65) and those from the middle Miocene Sanzhangtian site had the second
lowest (SI mean = 10.43 ± 0.99). Their calculated paleo-[CO2]atm values were 351 ± 24.8 ppmv and 334 ± 24.8 ppmv, respectively (Fig 3, Table 3, for more details see S1 Table).

The SI of the Pliocene and Pleistocene samples were higher (SI mean = 15.2–17.9) than the SI of the Miocene samples. The SI of the samples from the middle Early Pleistocene Ikoma site had the highest SI value (SI = 17.9 ± 1.9) and give out the lowest CO2 level of 278 ± 3.86 ppmv. SI of the fossilized leaves from the Sennan, Hachioji, Konan, and Tokamachi sites were 17.1, 17.2, 15.2, and 17.1, respectively. The reconstructed paleo-[CO2]atm from the Pliocene and Pleistocene samples in the Sennan, Hachioji, Konan, Tokamachi, and Ikoma sites were 280 ± 5.16, 279 ± 3.74, 285 ± 5.15, 280 ± 6.23, and 278 ± 3.86 ppmv, respectively (Fig 3, Table 3, for more details see S1 Table).

The significant differences between the mean variance of the stomatal index from different fossil localities were statistically tested (F = 54.016, p < 0.001) by one-way ANOVA with the “LSD” option in SPSS Statistics (Version 20.0). The result showed there was no significant difference between the SI data from Sanzhangtian locality (middle Miocene) and Kumagaya locality (late Miocene), but the SI data of these two localities were significantly different from the SI data from late Pliocene and Pleistocene localities. SI data of Konan locality (Late Pliocene) was significantly different from all other localities, but no significant difference has been detected among Sennan (Late Pliocene), Hachioji (Late Pliocene), Tokamachi (middle Early Pleistocene) and Ikoma localities (latest Early Pleistocene) (Table 4).
Discussion

Middle and late Miocene paleo-[CO_2]_{atm} change

The paleo-[CO_2]_{atm} changes reconstructed in previous research generally indicate a peak during the middle Miocene Climatic Optimum (MCO; 17–15 Ma) [44] and a decline during the later stage of the middle Miocene (ca. 15–11.5 Ma), although the reconstructed paleo-[CO_2]_{atm} values and timing of fluctuation were different among proxies (Fig 4B). The most prominent fluctuation was exhibited in the paleosol carbonate records, which showed a spike (ca. 800 ppmv) at 15.6 Ma, drop to 116–310 ppmv at 14.7–13.8 Ma, and increase to 433–519 ppmv around 12.8–13.1 Ma [45]. The stomatal records from fossilized Quercus leaves [23] also indicated a prominent change from the highest value (469–555 ppmv) at 15.7±0.7 Ma to the lower value at 13.0 Ma (ca. 290 ppmv) and 11.6 Ma (ca. 330 ppmv) during the late middle Miocene. Additionally, the stomatal proxies from North America indicate lower paleo-[CO_2]_{atm} values and moderate changes during the earlier stage of the middle Miocene: 396 ppmv from Ginkgo leaves at ca. 16.5 Ma and 310–316 ppmv from Metasequoia needles around 15.2–15.3 Ma [14].

In general, the values of the middle Miocene [CO_2]_{atm} estimated from marine proxies are lower than those from terrestrial records. Boron/Calcium (B/Ca) ratios of surface-dwelling foraminifera give a paleo-[CO_2]_{atm} of ca. 420 ppmv during the MCO that declined gradually to ca. 200 ppmv in the earliest late Miocene [47]. B isotope (δ^{11}B)-based paleo-[CO_2]_{atm} from ODP761 changed from ca. 400 ppmv in the MCO to ca. 280 ppmv in the late middle Miocene [48]. A stable paleo-[CO_2]_{atm} curve with slight changes around 210 ppmv from the MCO to late Miocene was drawn based on phytoplankton δ^{13}C alkene analysis [49, 50].

Table 3. Fossilized Metasequoia stomatal index and paleo-[CO_2]_{atm} concentration estimates during Cenozoic.

| Fossil site     | Epoch           | SI (%) | paleo-[CO_2]_{atm} (ppmv) |
|-----------------|-----------------|--------|---------------------------|
|                 |                 | Mean ± sd | Max | Min | Mean ± sd | Max | Min |
| Sanzhangtian    | middle Miocene  | 10.4±0.99 | 12.5 | 9.09 | 334±24.9  | 382 | 298 |
| Kumagaya        | early late Miocene | 9.80±0.65 | 11.0 | 8.97 | 351±24.8  | 392 | 317 |
| Sennan          | Late Pliocene   | 17.1±2.30 | 19.8 | 14.8 | 280±5.16  | 285 | 274 |
| Hachioji        | Late Pliocene   | 17.2±1.65 | 19.4 | 14.8 | 279±3.74  | 285 | 275 |
| Konan           | Late Pliocene   | 15.2±1.70 | 18.4 | 13.7 | 285±5.15  | 290 | 276 |
| Tokamachi       | middle Early Pleistocene | 17.1±2.52 | 21.7 | 13.3 | 280±6.23  | 293 | 272 |
| Ikoma           | latest Early Pleistocene | 17.9±1.90 | 20.2 | 15.5 | 278±3.86  | 282 | 273 |

Table 4. Mean difference of the least significant different (LSD) on stomatal index of fossil localities.

| Locality       | Sanzhangtian | Kumagaya    | Sennan      | Hachioji    | Konan       | Tokamachi   |
|----------------|--------------|-------------|-------------|-------------|-------------|-------------|
| Kumagaya       | 0.63         |             |             |             |             |             |
| Sennan         | -6.70***     | -7.33***    | -0.10       |             |             |             |
| Hachioji       | -6.80***     | -7.42***    | 1.90*       | 2.00*       |             |             |
| Konan          | -4.80***     | -5.42***    |             |             | 1.88*       |             |
| Tokamachi      | -6.68***     | -7.31***    | 0.02        | 0.12        | -2.63**     | -0.75       |
| Ikoma          | -7.42***     | -8.05***    | -0.73       | -0.63       | -2.63**     | -0.75       |

The sign of the significance is indicated as
* p<0.05
** p<0.01
*** p<0.001.

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[CO$_2$]$_{atm}$ value (334 ppmv) reconstructed from the fossilized leaves of the middle Miocene Sanzhangtian site was similar to the late middle Miocene values based on *Quercus* leaves [51] and between the results based on *Ginkgo* (16.5 Ma) and *Metasequoia* (15.2–15.3 Ma) leaves in the early middle Miocene [14].

The late Miocene stomatal data based on fossilized *Quercus* exhibited a decreasing paleo-[CO$_2$]$_{atm}$ tendency: ca. 370 ppmv at ca. 10.5 Ma, ca. 350 ppmv at ca. 8.5 Ma, and ca. 270 ppmv at ca. 7.2 Ma [51]. This was related to climatic cooling in the later late Miocene [18,52]. When using B/Ca [53] and phytoplankton [49,50] from marine proxies, they showed fluctuating values that were mostly less than 300 ppmv (Fig 4B). The estimated paleo-[CO$_2$]$_{atm}$ values for 10–9 Ma (351 ppmv) from this work are almost the same as the value from ca. 8.5 Ma from *Quercus* leaves [18]. Our data showed little change between the middle Miocene (334 ppmv) and the early late Miocene (351 ppmv) that confirmed the stable paleo-[CO$_2$]$_{atm}$ condition during this time as indicated by the phytoplankton record [49,50].

**Late Pliocene to Pleistocene paleo-[CO$_2$]$_{atm}$ change**

In most of the previous research, paleo-[CO$_2$]$_{atm}$ values are distributed between 200 and 400 ppmv during the Pliocene to Pleistocene (Fig 5A). B/Ca and B data have been used to
determine the paleo-[CO$_2$]$_{atm}$ of this period, as there is a lack of data from the stomatal method. The paleo-[CO$_2$]$_{atm}$ curve based on B/Ca from surface-dwelling foraminifera exhibited a peak of ca. 300 ppmv at ca. 3.4 Ma, this decreased to 181 ppmv at ca. 2.9 Ma, and then increased to 332 ppmv at ca. 1.4 Ma [47]. The downward shift in its fluctuation range was observed in the Early Pleistocene (Fig 5A), and the lowest value of 188 ppmv was recorded in the last glacial maximum (0.02 Ma) [47]. The paleo-[CO$_2$]$_{atm}$ recorded in the B isotopes indicates a higher level than that in B/Ca record during the Late Pliocene, that is, ca. 340 ppmv at ca. 3.4 Ma and ca. 400 ppmv at ca. 3.0 Ma [54]. However, it decreases to the same level (ca.
270 ppmv) as the B/Ca record in the late Late Pliocene and Early Pleistocene (ca. 2.8–1.0 Ma) [54]. The paleo-[CO₂]ₐtm level estimated from Quercus [18] and Cupressaceae [55] stomata indicates a higher level (ca. 350 ppmv) during the early Late Pliocene (3–3.4 Ma) and a lower value (276 ppmv) at 2.7 Ma. While paleo-[CO₂]ₐtm based on the SI of Typha at the Plio-Pleistocene boundary (2.65 Ma) exhibits a much higher value (534 ppmv) than the other results [22].

Our data showed that the paleo-[CO₂]ₐtm was maintained in the range between 280 and 285 ppmv in the Pliocene and Pleistocene (Figs 4B and 5A), which is about 150 ppmv lower than the results estimated from B isotopes [54], and about 70 ppmv higher than the results estimated from Typha from sediment at the Plio-Pleistocene boundary [22]. While different proxies [22,47,54] have recorded fluctuations accompanying climate changes (Fig 5B), the paleo-[CO₂]ₐtm value of this study stabilized at around 280 ppmv. Seiki et al. concluded that the Pliocene CO₂ levels determined by numerous methods agreed well with each other [9,54]. The present research suggests that some disagreements still remain in the results between our stomatal data and B/Ca records in the Pliocene, while the Pleistocene proxies give more consistent CO₂ levels (than the Pliocene).

Paleo-[CO₂]ₐtm change and late Cenozoic climatic deterioration

The overall climate cooling reconstructed for the past 20 Ma has generally been attributed to changes in CO₂ concentration in the atmosphere [46,58]. According to the marine oxygen isotope record, global temperature peaked at around 16 Ma (middle MCO) (Fig 4A), and the later part of the middle Miocene is characterized by climate cooling with expansion of the East Arctic ice sheet [59,60]. However, the middle Miocene paleo-[CO₂]ₐtm reconstructed in this study (around 334 ppmv) was just slightly lower than the present level, which was also the level maintained during the late Miocene (around 354 ppmv). That means that before the global temperature decrease, paleo-[CO₂]ₐtm had already achieved a stable low level. The Miocene paleo-[CO₂]ₐtm estimated based on alkenones also showed that paleo-[CO₂]ₐtm was similar during middle Miocene and late Miocene [61]. The δ¹³C record from foraminifera and B/Ca ratios in the foraminifera suggest that paleo-[CO₂]ₐtm decreases were apparently synchronous with major episodes of glacial expansion during the middle Miocene [53,62, 63], but this synchronization was not observed in our data. This study supports the view that Miocene climate change was not only influenced by paleo-[CO₂]ₐtm changes, but also by increases in seasonality and ocean circulation changes [50,64,65], and these accelerated the cooling in the late middle Miocene that also acted to decrease the paleo-[CO₂]ₐtm [62]. Also, climate sensitivity to paleo-[CO₂]ₐtm may have been greater than previously thought [66]. The impact of high latitude vegetation on Earth's albedo may have also played an important role in the Earth's energy budget in the Miocene [67].

After termination of the mid-Pliocene warmth at ca. 2.9 Ma, cooling trends continued until the onset of major expansion of the Northern Hemisphere ice sheet at ca. 2.7 Ma, which culminated at ca. 2.5 Ma in the earliest Pleistocene [68–70]. However, present results show that the lower paleo-[CO₂]ₐtm level started around 2.8–3.0 Ma and lasted until the late Early Pleistocene. Therefore, we consider that the transition to the icehouse world was possibly induced by a decrease of the paleo-[CO₂]ₐtm, which already dropped to their lowest levels during the complete Cenozoic before the major expansion of the Northern Hemisphere ice sheets. During the Pliocene to Pleistocene, our data are very stable, but the global temperature estimated from the marine oxygen isotope record [56,57] shows drastic fluctuations (Fig 5). However, our middle and late Miocene data are significantly higher than our Pliocene and Pleistocene data. The oxygen isotope record confirms that the temperature in the Pliocene and Pleistocene was much
lower than that of the middle and late Miocene [44,46]. Therefore, we can conclude that the decrease of paleo-[CO₂] atm level is coupled with temperature decrease during middle Miocene to Pleistocene.

Conclusions

We used the stomatal index of *Metasequoia* Miki *ex* Hu *et* Cheng as a proxy to reconstruct the paleo-[CO₂] atm evolution from the middle Miocene to late Early Pleistocene for the first time. Our results indicate that: (1) From middle to late Miocene the atmospheric CO₂ level stabilized around 350 ppmv which is slightly lower than today. (2) The CO₂ level during the Pliocene to Pleistocene was similar to the pre-industrial level and no fluctuation can be detected by this research. (3) The Pleistocene CO₂ level estimated by different proxies agree well with each other. (4) From middle Miocene to Pleistocene, when the global temperature decreased sharply, the global CO₂ level decreased by more than 50 ppmv, which may suggest that CO₂ decrease and temperature decrease are coupled.

Supporting Information

S1 Table. Original paleo-[CO₂] atm results for the seven localities used in this study. (DOC)

S2 Table. Previously reconstructed paleo-[CO₂] atm results based on different proxies over the past 20 Ma. (DOC)

Acknowledgments

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Author Contributions

Conceived and designed the experiments: LW ZKZ. Performed the experiments: YQW LW. Analyzed the data: YQW AM JLA. Contributed reagents/materials/analysis tools: YQW AM LW. Wrote the paper: YQW AM LW JLA ZKZ.

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