

# The impact of Miocene atmospheric carbon dioxide fluctuations on climate and the evolution of terrestrial ecosystems

Wolfram M. Kürschner\*<sup>†</sup>, Zlatko Kvaček<sup>‡</sup>, and David L. Dilcher<sup>§</sup>

\*Faculty of Science, Institute of Environmental Biology, Department of Palaeoecology, Laboratory of Palaeobotany and Palynology, Utrecht University, Budapestlaan 4, 3584 CD, Utrecht, The Netherlands; <sup>‡</sup>Faculty of Natural Sciences, Charles University, Albertov 6, 128 43 Prague 2, Czech Republic; and <sup>§</sup>Florida Museum of Natural History, University of Florida, Gainesville, FL 32611-7800

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**The Miocene is characterized by a series of key climatic events that led to the founding of the late Cenozoic icehouse mode and the dawn of modern biota. The processes that caused these developments, and particularly the role of atmospheric CO<sub>2</sub> as a forcing factor, are poorly understood. Here we present a CO<sub>2</sub> record based on stomatal frequency data from multiple tree species. Our data show striking CO<sub>2</sub> fluctuations of ≈600–300 parts per million by volume (ppmv). Periods of low CO<sub>2</sub> are contemporaneous with major glaciations, whereas elevated CO<sub>2</sub> of 500 ppmv coincides with the climatic optimum in the Miocene. Our data point to a long-term coupling between atmospheric CO<sub>2</sub> and climate. Major changes in Miocene terrestrial ecosystems, such as the expansion of grasslands and radiations among terrestrial herbivores such as horses, can be linked to these marked fluctuations in CO<sub>2</sub>.**

atmospheric CO<sub>2</sub> | fossil plants | paleoclimates | stomata | C<sub>4</sub> plants

The Miocene is distinguished by extreme climatic optima alternating with major long-term climatic cooling, which together mark the founding of the modern late Cenozoic cold mode and the evolution of modern terrestrial biomes (1). Grass-dominated ecosystems became established in the low and middle latitudes of many parts of the world, such as North America, Eurasia, Africa, and Australia (2). Major radiations in large mammalian herbivores have been attributed to changes in the distribution of vegetation and terrestrial primary productivity (3–5). A significant change in dental morphology from low- to high-crowned toothed horses occurs during the middle Miocene, whereas a transition from a C<sub>3</sub> plant to C<sub>4</sub> plant diet did not take place before the late Miocene (6).

Both Cenozoic climate trends and changes in terrestrial ecosystems have been thought to be influenced by long-term CO<sub>2</sub> fluctuations (6–8). Before marine pCO<sub>2</sub> proxy records were available, Cenozoic CO<sub>2</sub> trends were inferred from carbon-isotope records of paleosols (9) and from carbon cycling models (10), which indicated a long-term decrease from ≈1,000 to <500 parts per million by volume (ppmv) throughout the Cenozoic. Approximately a decade later, CO<sub>2</sub> reconstructions based on marine geochemical proxies indicated consistently low late Pleistocene (glacial-like) CO<sub>2</sub> values of ≈200–280 ppmv (11, 12). Consequently, the Miocene has been regarded as a geological period in which climate and the carbon cycle were essentially decoupled. Because of this alleged decoupling, the role of atmospheric CO<sub>2</sub> as a climate forcing factor has been disputed (13–15). However, a permanently low CO<sub>2</sub> scenario has been challenged because photosynthetic models predict that plant life would not have thrived under such conditions (16). Climate models showed the importance of atmospheric CO<sub>2</sub> as a fundamental boundary condition for Cenozoic climate change (17). In fact, a coupling between atmospheric CO<sub>2</sub> and glacial–interglacial cycles over the past 600,000 years is well documented by ice core analysis (18). Understanding the long-term perspective beyond the Pleistocene is essential because climate and CO<sub>2</sub>

fluctuations on progressively shorter time scales are ultimately dependent on the evolution of the global carbon cycle and earth's climate on longer time scales. Recently, refined geochemical studies of marine and terrestrial sedimentary records suggest a coupling between atmospheric CO<sub>2</sub> and temperature over Phanerozoic time scales (19–21).

In addition to geochemical CO<sub>2</sub> proxies, stomatal frequency analysis on fossil leaf remains represents a terrestrial proxy for CO<sub>2</sub> that is based on the inverse relationship between atmospheric CO<sub>2</sub> and stomatal frequency (22). In the present study, stomatal frequency is expressed as the stomatal index (*SI*), which is calculated as  $SI (\%) = [SD/(SD + ED)] \times 100$ , where *SD* is the stomatal density and *ED* is the epidermal cell density. Because *SI* normalizes for leaf expansion, it is largely independent of plant water stress and is primarily a function of CO<sub>2</sub> (22, 23). Calculation of *SI* provides a robust method for estimating CO<sub>2</sub> levels on short (24) and geologically long time scales (25, 26). Because the stomatal frequency response to CO<sub>2</sub> is species-specific, quantitative estimates of CO<sub>2</sub> are limited to extant species. Here we present a CO<sub>2</sub> reconstruction based on a multiple-species stomatal frequency record from leaf remains of two extant lineages of laurel species (the *Laurus abchasica* and *Laurus nobilis* lineage and the *Ocotea hradekensis* and *Ocotea foetens* lineage), maidenhair tree (*Ginkgo biloba*), and an extinct laurel species (*Laurophyllum pseudoprinceps*). For each extant species, the stomatal frequency response has been independently calibrated based on historical sets of herbarium leaf material, using standard protocols (27). The training datasets and CO<sub>2</sub> inference models, respectively, for *L. nobilis* and *O. foetens* are shown in Figs. 1 and 2 (for details, see *Material and Methods*). The *SI* calibration for the extinct species *L. pseudoprinceps* has been established by cross-calibration with known CO<sub>2</sub> levels from those stratigraphic units in which it occurs together with the extant laurel species and *Ginkgo*.

We address two main questions in this article: (i) What was the relationship between the long-term CO<sub>2</sub> fluctuations and climate evolution during the Miocene? and (ii) Was CO<sub>2</sub> an environmental stress factor that influenced the evolution of modern terrestrial ecosystems?

## Results and Discussion

Fossil *SIs* of all four species (*L. pseudoprinceps*, *L. nobilis*, *O. foetens*, and *G. biloba*) show pronounced fluctuations over the

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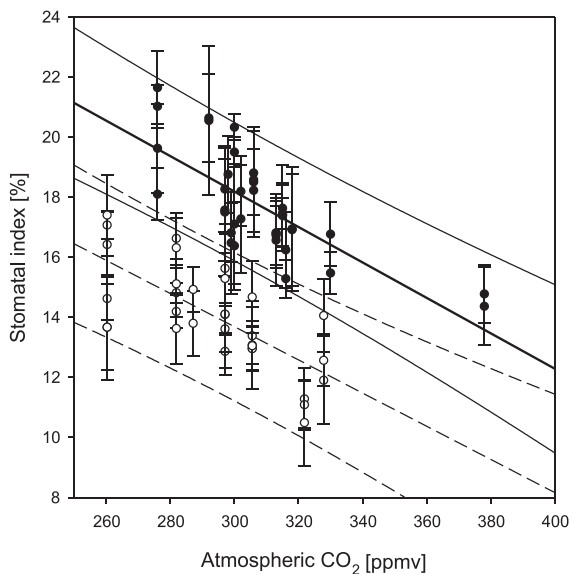
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<sup>†</sup>To whom correspondence should be addressed. E-mail: w.m.kuerschner@uu.nl.

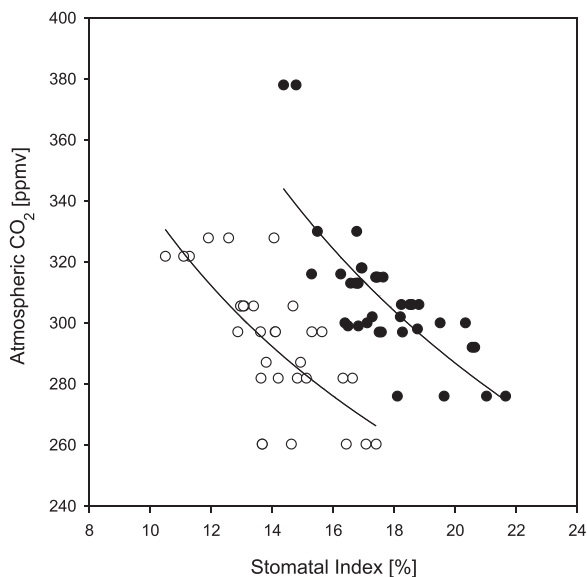
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**Fig. 1.** Training dataset of mean *SI* values of *L. nobilis* (filled circles) and *O. foetens* (open circles) vs. atmospheric  $\text{CO}_2$  levels with 95% prediction interval. Error bars show standard deviation.

time interval studied (Fig. 3A). Three periods can be recognized in the present record. During the oldest period—the late Oligocene to early Miocene (25–20 Ma)—the *SI* of *L. pseudoprinceps* shows a significant increase from  $\approx 9\%$  to  $15\%$  and remains high during the early Miocene. During the middle period—the late early to early middle Miocene (20–16 Ma)—*L. pseudoprinceps* shows a decrease in *SI* from  $15\%$  to  $12\%$ . A simultaneous decline in *SI* is found for two other laurel species, *L. nobilis* and *O. foetens*, from  $12\%$  to  $10\%$  and from  $12\%$  to  $8\%$ , respectively. In the youngest period of the present study—the middle Miocene (16–12 Ma)—*L. nobilis* shows a marked increase from  $10\%$  to  $20\%$ , followed by a decrease to  $14\%$  during the late middle Miocene. The pronounced increase in the *SI* of *L. nobilis* during the early middle Miocene is confirmed by a sharp shift in the *SI* of *G. biloba* from  $8\%$  to  $15\%$ .

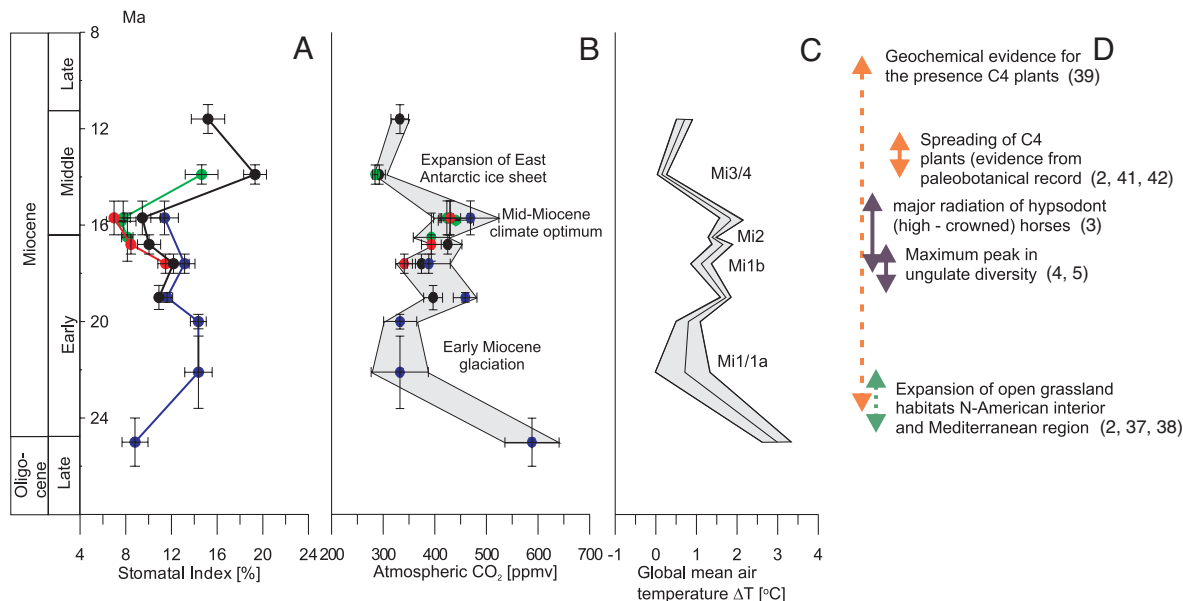


**Fig. 2.**  $\text{CO}_2$  inference model based on historical series of herbarium sheets for *L. nobilis* (filled circles) and *O. foetens* (open circles).

The *SI* record of the present study reveals prominent changes in  $\text{CO}_2$  since the late Oligocene (Fig. 3B).  $\text{CO}_2$  declines drastically, from  $\approx 600$  to  $\approx 340$  ppmv, during the Oligocene–Miocene transition. It remains low (at about this level) until the middle early Miocene. From  $\approx 20$  Ma,  $\text{CO}_2$  increases again to reach a maximum of  $\approx 400$ – $500$  ppmv in the early middle Miocene (15.5 Ma). Superimposed on this rising trend are several smaller, temporary  $\text{CO}_2$  oscillations with amplitude of  $\approx 50$  ppmv. A second major  $\text{CO}_2$  decrease to 280 ppmv occurs during the middle Miocene (15.5–14 Ma), followed by a slight increase to  $\approx 340$  ppmv during the late Miocene. A previous  $\text{CO}_2$  reconstruction based on North American leaf remains (26) fit perfectly into the present study, but it represented only a few measurements at  $\approx 15$  Ma.

It should be noted that because of the nonlinear nature of stomatal frequency response, the sensitivity of this proxy decreases at high  $\text{CO}_2$  levels (greater than or equal to  $\approx 400$  ppmv). Consequently, the high  $\text{CO}_2$  levels during the climate optima at  $\approx 22$  Ma and 15.5 Ma may represent a conservative estimation rather than the average  $\text{CO}_2$  level. Despite the applied log-transformation, the  $\text{CO}_2$  inference models (Eqs. 3 and 4 in *Materials and Methods*) underestimate the actual  $\text{CO}_2$  levels at extrapolation far beyond the calibration interval. To estimate this uncertainty, the  $\text{CO}_2$  levels predicted by the inference models are plotted vs. the input  $\text{CO}_2$  values used to establish the transfer functions (Fig. 4). At  $\approx 500$  ppmv  $\text{CO}_2$ , the inference models of *L. nobilis* and *O. foetens* underestimate the  $\text{CO}_2$  value by  $\approx 150$  and 200 ppmv, respectively. Consequently, the atmospheric  $\text{CO}_2$  levels during the middle Miocene climatic optimum would have been closer to  $\approx 650$ – $700$  ppmv than to the uncorrected value of  $\approx 500$  ppmv. During the late Oligocene, climate optimum  $\text{CO}_2$  levels would have been 800–900 ppmv, rather than the uncorrected 600 ppmv, which is intriguingly close to the estimates provided by the alkenone-based  $\text{CO}_2$  proxy record (14). Moreover, well preserved leaf material is rare, and the number of observations per individual sample is (at the present stage) in some cases rather low. However, the largest portion, 70%, of the  $\text{CO}_2$  record is based on stomata counts from the three extant species that are independently calibrated. Different species show the same trends, which reinforces the fidelity of the reconstructed  $\text{CO}_2$  trends through the Miocene. All in all, our data do not show any signs of a prolonged Miocene low- $\text{CO}_2$  scenario, as suggested by geochemical proxy records (11, 12, 14). If such a scenario had existed, it should also be evident in the present study because the botanical proxy used is sensitive to low  $\text{CO}_2$  levels (22).

The present leaf-based  $\text{CO}_2$  reconstruction shows a similar  $\text{CO}_2$  decline at the Oligocene–Miocene transition as in the marine proxy records (12, 14), but it deviates significantly throughout the course of the Miocene. The alkenone-based  $\text{CO}_2$  record indicates consistently low Pleistocene-like  $\text{CO}_2$  levels (200–280 ppmv) and shows hardly any covariation with Miocene climate fluctuations. The boron isotope-based  $\text{CO}_2$  reconstruction shows a decrease in atmospheric  $\text{CO}_2$   $\approx 15$  Ma, which is in the order of the present study ( $\approx 150$  ppmv), although the absolute values are significantly lower (by  $\approx 200$  ppmv) than in our study. However, our results provide evidence from the terrestrial fossil record that  $\text{CO}_2$  had a profound influence on the Miocene long-term climate evolution as recorded in marine records (1) (Fig. 3C). Despite the limited temporal resolution of the present study,  $\text{CO}_2$  fluctuations correlate with some of the climate events (Mi1/1b, Mi2, and Mi3/4) known from the marine oxygen isotope curve (1). In the future, leaf-based  $\text{CO}_2$  reconstructions with a finer time resolution may could potentially reveal patterns of higher order fluctuations that are coupled to orbital cycles, as suggested by marine proxy records (28). The global mean temperature model shows a 2–3°C decline during the early Miocene and middle Miocene climate events (Fig. 3C).

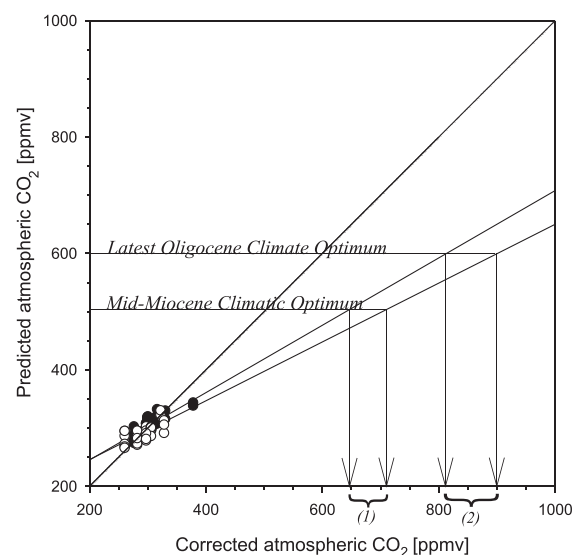


**Fig. 3.** Late Oligocene–Miocene stomatal index records, inferred atmospheric CO<sub>2</sub> fluctuations, and effects on global temperature compared with major events in terrestrial ecosystems. (A) *SI* of fossil leaf remains between 25 and 12 Ma (late Oligocene until late middle Miocene. For a list of locations and their age assessment, see *SI Table 3*). The lines represent trends in *SI*: blue, *L. pseudoprinceps*; black, *L. nobilis*; red, *O. foetens*; green, *G. biloba*. The values represent means per stratigraphic unit, with error bars indicating the standard deviation of the *SI*. The age error bars indicate the minimum and maximum ages of the sample. The stratigraphic framework is established by vertebrate biostratigraphy and magnetostratigraphy (see also *SI Table 3*). (B) Reconstructed late Oligocene–middle Miocene CO<sub>2</sub> levels based on individual independently calibrated tree species. The error bars of the species-specific CO<sub>2</sub> estimates are based on the standard deviation of the *SI* measurements on individual fossil leaf samples. The gray band indicates the envelope as determined by the minimum and maximum CO<sub>2</sub> levels inferred from all individual samples per stratigraphic unit. (C) Modeled temperature departure of global mean surface temperature from present day, calculated from mean CO<sub>2</sub> estimates by using a CO<sub>2</sub>–temperature sensitivity study (46). Also indicated are the major Miocene climate key events and the position of the Miocene cooling events Mi1/1a, Mi1b, Mi2, and Mi3/4 known from the marine oxygen isotope record (1). The effects of pCO<sub>2</sub> level changes on the global mean land surface temperature were estimated assuming a radiative relationship between CO<sub>2</sub> mixing and global air temperature inferred from climate–CO<sub>2</sub> sensitivity models expressed as  $\Delta T = 4 \ln(C/CO)$ , where *C* is the mixing ratio and CO is the preindustrial CO<sub>2</sub> mixing ratio of 278 ppmv (46). (D) Major events in the terrestrial ecosystems in response to the Miocene CO<sub>2</sub> trends, such as changes in terrestrial herbivore communities (3–5), the expansion of Miocene grasslands (2, 37, 38), and evidence for C<sub>4</sub> biomass from paleosols (43).

Our data confirm modeling experiments that show that Cenozoic CO<sub>2</sub> must decline below a threshold of ≈500 ppmv to induce significant buildup of Antarctic ice sheets (17). Unlike the

geochemical proxy records (11, 12, 14), our data indicate that elevated CO<sub>2</sub> levels contributed to the middle Miocene climatic optimum. Several lines of evidence from the fossil and geological record (15, 29–32) suggest that this time interval (≈14.5–17 Ma) was the warmest period of the past 35 Ma. Our results demonstrate that this climate optimum was forced significantly by elevated CO<sub>2</sub> levels similar to those, for example, during the early Eocene (12, 14, 26). A likely source of the late, early, and middle Miocene CO<sub>2</sub> increases was extensive volcanic activity during the Columbia River Flood Basalt volcanism and the Central European volcanism (33). The marked CO<sub>2</sub> drop during the Miocene, in turn, may be the result of increased C<sub>org</sub> burial resulting from the Himalayan uplift (34) and/or of enhanced marine productivity in the Pacific ocean (7) and the global occurrence of vast brown-coal-forming basins (35).

The marked Miocene CO<sub>2</sub> variations may have directly impacted the structure and productivity of terrestrial biomes by affecting plant photosynthetic performance. In sensitivity tests at 280 and 560 ppmv for Miocene global vegetation models (36), CO<sub>2</sub> shows pronounced changes related to vegetation distribution with regard to the degree of tree coverage for the seasonal dry tropics. However, evidence from the fossil record for the global distribution pattern of these biomes in the Miocene is rather scattered because fossil plant assemblages are highly influenced by taphonomical processes. Phytolith studies from the North American continental interior have shed new light on Miocene vegetational history (37). They reveal that pronounced changes in vegetation took place at the Oligocene–Miocene transition. Late Oligocene vegetation was a closed forest with palm and bamboo understory, whereas early Miocene plant communities were characterized by a mix of C<sub>3</sub> grasses and



**Fig. 4.** Predicted CO<sub>2</sub> concentrations calculated from Eqs. 3 and 4 for *L. nobilis* (filled circles) and *O. foetens* (open circles), plotted vs. the actual atmospheric CO<sub>2</sub> levels with a linear extrapolation to elevated CO<sub>2</sub>. The arrows indicate the value of the corrected CO<sub>2</sub> level for the climatic optima in the middle Miocene (1) and latest Oligocene (2).



herbs, forming savannas or open woodland habitats (37). Recently, phytolith studies from the eastern Mediterranean region reveal that relatively open, grass-dominated habitats were established by at least the early Miocene (38). Intriguingly, these examples from the fossil record are similar to changes in vegetation distributions that occur in the Miocene global vegetation models as the result of CO<sub>2</sub> sensitivity runs (36). Growth experiments and vegetation models with modern mixed tree/grass ecosystems show that changes in atmospheric CO<sub>2</sub> directly affect tree cover by modifying water relationships in herbaceous vs. woody plants (39) or by modifying the (re)growth rates of plants recovering after a disturbance such as wildfire (40). Therefore, the opening of the forest vegetation and proliferation of open grassland habitat may have been influenced by the initial CO<sub>2</sub> drawdown to ≈300 ppmv during the early Miocene (Fig. 3D). After a continuous CO<sub>2</sub> increase up to ≈550 ppmv between 20 and 16 Ma, a second major drop in CO<sub>2</sub> at ≈12 Ma down to ≈280 ppmv may have increased the environmental stress induced by CO<sub>2</sub> starvation. We hypothesize that under these CO<sub>2</sub>-limiting conditions, in concert with an open habitat, exposure of grasses to high light intensity and water-stressed environments, combined with the coevolution of herbivores, may have facilitated the first radiation of C<sub>4</sub> grasses. To date, the fossil record of C<sub>4</sub> plants is still rather enigmatic. A few paleobotanical studies reveal evidence for C<sub>4</sub> plants in the middle Miocene (41, 42), whereas geochemical data suggest their presence since the early Miocene (43). Another line of evidence on the origin of C<sub>4</sub> plants comes from the molecular clock approach, which estimates an approximate maximum age for the origin of modern C<sub>4</sub> plants of 25 Ma (44). These data coincide approximately with the CO<sub>2</sub> drop in the early Miocene. Carbon isotope records of paleosols and fossil tooth enamel indicate, however, that the major proliferation of C<sub>4</sub> grasslands did take place during the late Miocene, ≈8 Ma ago (6). The later vegetation change has been attributed to an increased seasonality and increased fire frequency (45).

Changes in the productivity and species richness of terrestrial vegetation must have affected herbivore communities. Hoofed herbivorous large mammals on the North American continent show a maximum diversity during the middle Miocene climatic optimum (4, 5) (Fig. 3D). This maximum in both local and regional diversity greatly exceeds the diversity of ungulates in any present-day habitat, which implies a greater primary productivity than is seen today. Preliminary review data suggest that the pattern of elevated ungulate diversity is a global phenomenon, and, therefore, a global driving force is the most likely explanation. CO<sub>2</sub> fertilization during the middle Miocene climatic optimum may have made possible the expansion of high-productivity terrestrial biomes that supported high-diversity browser communities.

As a result of climatic changes such as increased seasonality and cooling, as well as the decline in primary productivity due to the marked post-middle Miocene CO<sub>2</sub> crisis, woodland biomes retreated, the species richness of browsing mammals declined, and grazers increased their diversity (3). A large number of terrestrial mammalian herbivores, such as equids, camelids, antilocaprids, rhinos, and proboscideans show dramatic morphological changes of their dentitions during the Miocene (3–5). In particular, early Cenozoic horses are characterized by brachydont (short-crowned) dentitions but show, from the late middle Miocene on (≈15 Ma), a rapid diversification of hypsodont (high-crowned) taxa, which has been attributed as an adaptation to include more fibrous and abrasive material, such as grasses, in the diet (3). This development implies the coevolution of large herbivores and plants in response to Miocene climate and atmospheric CO<sub>2</sub> fluctuations.

## Materials and Methods

Standardized computer-aided determinations of epidermal parameters were performed on an AnalySIS image analysis system (Soft Imaging Systems) software. All statistics and graphs were made with Sigmaplot version 9.1 (Systat Software). On average, 5–10 stomata-bearing alveoles per leaf sample were measured for ED (n/mm<sup>2</sup>) and SD (n/mm<sup>2</sup>). From SD and ED, the area-independent *SI* was calculated:  $SI (\%) = [SD/(SD + ED)] \times 100$ . Most of the cuticle preparations (stored at Charles University) were collected from several brown-coal basins in the Czech Republic (Most and Zitava basins, South Bohemia), supplemented with some material from Austria (Parschlug Basin, Styria) and Germany (Lower Rhine Embayment, Lausitz Basin). References for the age assessments of the individual fossil leaf samples are given in [supporting information \(SI\) Table 3](#).

A total of 68 herbarium leaf samples (see [SI Tables 1 and 2](#)) and 36 fossil leaf samples ([SI Table 3](#)) were studied for their epidermal cell properties. The data shown in [SI Tables 1–3](#) represent means with standard deviation per herbarium leaf or fossil leaf remains, respectively. The species included are the extant taxa *L. abchasica*, *O. hradekensis*, and *G. biloba*, as well as one extinct species, *L. pseudoprinceps*. To convert the *SI* values from the extant species *L. abchasica* and *O. hradekensis* to atmospheric CO<sub>2</sub> levels, CO<sub>2</sub> inference models were established from their living equivalents, *L. nobilis* and *O. foetens* (Figs. 1 and 2), from historical herbarium sheets collected over the 19th and 20th centuries, covering the CO<sub>2</sub> increase since the industrial revolution. Historical atmospheric CO<sub>2</sub> concentrations used for calibration are annual means as measured on Mauna Loa, HI, since 1952 (47), supplemented by CO<sub>2</sub> measurements from Antarctic ice cores (Siple Station) (48). The taxonomical relationships have been established by extensive comparative studies on leaf morphology and cuticle anatomy (49–51). CO<sub>2</sub> estimates from *G. biloba* were based on calibration data by Royer *et al.* (26). In the absence of a modern equivalent for the extinct species *L. pseudoprinceps*, the *SI* response of this species has been cross-calibrated by using Miocene CO<sub>2</sub> levels inferred from the three extant species.

The *SI* calibration of *L. nobilis* to atmospheric CO<sub>2</sub> concentrations results in a linear relationship, with

$$SI = -0.059 CO_2 + 35.876 \quad [1]$$

and a coefficient of determination (*R*<sup>2</sup>) of 0.56.

The *SI* calibration of *O. foetens* to atmospheric CO<sub>2</sub> concentrations results in a linear relationship, with

$$SI = -0.0552 CO_2 + 30.245 \quad [2]$$

and an *R*<sup>2</sup> of 0.5.

To account for the nonlinear response of *SI* to changing CO<sub>2</sub> concentrations, both herbarium *SI* data and the historical CO<sub>2</sub> concentrations are log-transformed before fitting a linear response curve through the datasets. For *L. nobilis*, this results in a relationship of

$$CO_2 = 10^{3.173} - [0.5499 \times \log(SI_{fossil})], \quad [3]$$

with an *R*<sup>2</sup> of 0.78 between measured and inferred CO<sub>2</sub> values and a root mean square error (RMSE) of 13.5 ppmv CO<sub>2</sub>.

For *O. foetens*, the regression curve (Fig. 2) and statistics are

$$CO_2 = 10^{2.9567} - [0.4284 \times \log(SI_{fossil})], \quad [4]$$

with an *R*<sup>2</sup> of 0.5 between measured and inferred CO<sub>2</sub> values and an RMSE of 16 ppmv CO<sub>2</sub>.

In the absence of a modern equivalent for the extinct species *L. pseudoprinceps*, the *SI* response of this species has been cross-calibrated by using Miocene CO<sub>2</sub> levels inferred from the three extant species. This results in a relationship of

$$CO_2 = -46.011 \times SI_{fossil} + 993.37, \quad [5]$$

with an *R*<sup>2</sup> of 0.68.

The multiple-species *SI* record based on three extant species allows that CO<sub>2</sub> values are independently inferred from the individual species-specific CO<sub>2</sub> inference models and verified for interspecific coherence.

*O. foetens* is an endemic tree from Madeira and the Canary Islands. It occurs over a wide range of altitudes in valleys and hills of the interior of the islands and is the highest tree of the laurel forest (Larissilva). Because the historical herbarium leaves were collected at different altitudes from approximately sea level up to 1,000 m, the stomatal frequency response is expressed against the

CO<sub>2</sub> partial pressure and corrected for the altitude at which leaves were collected (see SI Table 2).

The atmospheric CO<sub>2</sub> partial pressure was calculated with Eqs. 6 and 7 (52, 53). Atmospheric pressure decreases with altitude in a predictable manner,

$$P_{\text{air}, z} = 101,325^{[MW(\text{air}) \cdot g \cdot z / RT]}, \quad [6]$$

where MW<sub>(air)</sub> is the molecular weight of air (28.964 × 10<sup>-3</sup> kg/mol), *z* is altitude in meters, *g* is acceleration due to gravity in meters per second, *R* is the gas constant (8.3144 J/mol), and *T* is mean July temperature in kelvin.

The CO<sub>2</sub> partial pressure depends on the altitude, with

$$P_{\text{CO}_2, z} = [P_{\text{air}, z} / 101,325] \cdot P_{\text{CO}_2, \text{sea level}}, \quad [7]$$

where *P*<sub>CO<sub>2</sub>, *z*</sub> is the CO<sub>2</sub> partial pressure at altitude *z*, and *P*<sub>CO<sub>2</sub>, sea level</sub> is the CO<sub>2</sub> partial pressure at sea level.

To calculate CO<sub>2</sub> estimates in mixing ratios (ppmv), the altitude-corrected CO<sub>2</sub> partial pressure values of the *O. foetens* training set were reconverted into mixing ratios at sea level with

$$C_{\text{CO}_2} = (P_{\text{CO}_2, \text{sea level}} / 101,325) \cdot 10^6, \quad [8]$$

where *C*<sub>CO<sub>2</sub></sub> is the mixing ratio in ppmv, and *P*<sub>CO<sub>2</sub>, sea level</sub> is the partial pressure at sea level.

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