



Rapid report

Glacial trees from the La Brea tar pits show physiological constraints of low CO₂

Author for correspondence: Joy K. Ward Tel: +1 785 864 5229 Email: joyward@ku.edu

Received: 3 November 2011 Accepted: 24 November 2011

Laci M. Gerhart¹, John M. Harris², Jesse B. Nippert³, Darren R. Sandquist⁴ and Jov K. Ward¹

¹Department of Ecology and Evolutionary Biology, University of Kansas, 1200 Sunnyside Avenue, Lawrence, KS 66045, USA; ²George C. Page Museum at the La Brea Tar Pits, Natural History Museum of Los Angeles County, 5801 Wilshire Blvd, Los Angeles, CA 90036, USA; ³Division of Biology, Kansas State University, 116 Ackert Hall, Manhattan, KS 66506, USA; ⁴Department of Biological Science, California State University, 800 N. State College, Fullerton, CA 92834, USA

New Phytologist (2012) 194: 63-69 doi: 10.1111/j.1469-8137.2011.04025.x

Key words: carbon isotopes, c_i/c_a . interannual variation, Juniperus, last glacial period, low CO₂, tree rings.

Summary

- While studies of modern plants indicate negative responses to low [CO₂] that occurred during the last glacial period, studies with glacial plant material that incorporate evolutionary responses are rare. In this study, physiological responses to changing [CO₂] were compared between glacial (La Brea tar pits) and modern Juniperus trees from southern California.
- Carbon isotopes were measured on annual rings of glacial and modern Juniperus. The intercellular : atmospheric [CO₂] ratio (c_i/c_a) and intercellular [CO₂] (c_i) were then calculated on an annual basis and compared through geologic time.
- Juniperus showed constant mean c_i/c_a between the last glacial period and modern times, spanning 50 000 yr. Interannual variation in physiology was greatly dampened during the last glacial period relative to the present, indicating constraints of low [CO2] that reduced responses to other climatic factors. Furthermore, glacial Juniperus exhibited low c_i that rarely occurs in modern trees, further suggesting limiting [CO₂] in glacial plants.
- This study provides some of the first direct evidence that glacial plants remained near their lower carbon limit until the beginning of the glacial-interglacial transition. Our results also suggest that environmental factors that dominate carbon-uptake physiology vary across geologic time, resulting in major alterations in physiological response patterns through time.

Introduction

The last glacial period began c. 110 000 yr ago and reached a maximum for global ice volume at 18 000-20 000 yr ago. Glacial conditions persisted (except for brief interstadials) until the abrupt transition to the current interglacial period, beginning c. 14 000 yr ago. At the peak of the last glacial period, atmospheric CO₂ concentrations ([CO₂]) ranged between 180 and 200 ppm, which are among the lowest concentrations that occurred during the evolution of land plants (Berner, 2006; also see Pagani et al., 2009 for an account of similarly low concentrations c. 15 million yr ago). When grown at glacial vs modern [CO₂], modern C₃ plants show 40-70% reductions in photosynthesis and biomass production (Polley et al., 1993; Sage & Coleman, 2001), 20-30% lower survival (Ward & Kelly, 2004), and may even fail to reproduce (Dippery et al., 1995).

This is a result of reduced CO₂ substrate concentrations at carboxylation sites, as well as higher photorespiration rates. However, even at reduced paleo-temperatures where photorespiration is decreased, plants are still unable to overcome the severe, negative effects of low [CO₂] (Ward et al., 2008). Such pronounced effects originating at the level of autotrophic physiology have been modeled at the ecosystem scale, and have been predicted to greatly reduce net primary production and carbon storage during glacial periods (Turcq et al., 2002; François et al., 2006). Admittedly, however, modern plants are often grown in glacial conditions for only a single generation, and therefore do not reflect evolutionary responses to low [CO₂]. This realization prompted our recent studies of glacial Juniperus (juniper) trees that were fully preserved within the La Brea tar pits in southern California (Los Angeles), and that had tens of thousands of years to adapt to low [CO₂]. Analysis of stable carbon isotope ratios

of complete wood samples (that integrate all tree rings from a given individual) show severely reduced internal $[CO_2]$ (c_i) during the last glacial period that are unprecedented in modern equivalents, strongly suggesting the existence of major carbon limitations on tree physiology (Ward *et al.*, 2005). Thus, the low $[CO_2]$ of glacial periods likely produced a bottleneck on carbon exchange through reduced transfer of CO_2 from the atmosphere to the biosphere. However, it is still unknown if low $[CO_2]$ presented an overriding limitation on plant physiology relative to other climatic factors (e.g. water, temperature).

The last glacial period represents an ideal time period for addressing this issue, since climate was more variable on an interannual basis and [CO₂] was exceptionally low relative to modern times (Mayewski et al., 2004). More specifically, ice cores from Greenland indicate extreme stability of Holocene climate compared with that of the last glacial period (Dansgaard et al., 1993). In addition, Dansgaard-Oeschger (D-O) cycles, which are periods of rapid and abrupt changes in temperature, dust content, ice accumulation and greenhouse gas concentrations, were more prominent during the last glacial period relative to the Holocene (Broecker, 1994; Roy et al., 1996). These patterns recorded in Greenland ice are also documented in ocean sediment cores from the Santa Barbara Basin (Behl & Kennett, 1996; Heusser, 1998; Hendy & Kennett, 1999; Hendy et al., 2002), c. 100 km northwest of La Brea, our primary research site. Analyses of these cores show a strong teleconnection between atmospheric trends over Greenland and ocean dynamics off the California coast, identifying synchronous climatic events between the two records over the last 60 000 yr (Behl & Kennett, 1996; Heusser, 1998; Hendy & Kennett, 1999; Hendy et al., 2002).

In previous work, we did not have wood specimens that allowed for carbon isotope analysis of individual tree rings. Recent excavations at the La Brea tar pits have now yielded higher-quality *Juniperus* specimens, allowing for discernment of individual tree rings. Thus, these wood specimens make an excellent model system for testing the constraints of low [CO₂] on tree

physiology relative to the effects of other climatic factors during the last glacial period. Here we compare long-term responses of tree physiology, as well as interannual variation within individuals, between the last glacial period and modern times. In doing so, we find the first evidence that low [CO₂] constrained the physiology of glacial trees, as evidenced by a dampened response to interannual climate variability.

Materials and Methods

Site selection

For this study, glacial trees from the Rancho La Brea tar pits (Los Angeles) were sampled and ¹⁴C dated to 14.5–47.6 kyr before present (BP), with the majority of specimens dating to the last glacial period. Juniperus samples from Rancho La Brea cannot be identified to the species level, although analysis by a wood anatomy expert (Ward et al., 2005) and species distributions indicate these samples are either *I. californica* or *I. occidentalis*. Cores of modern trees were collected from three low elevation sites in the Angeles National Forest (J. californica, two trees per site, one core per tree) and three high elevation sites in the San Bernardino National Forest (J. occidentalis, three trees per site, one core per tree), which are close in proximity to La Brea. Only modern trees from natural areas with well-drained, nonirrigated soils were sampled. Low-elevation sites provided a same-site control for glacial La Brea (with the full suite of environmental changes through time), whereas high-elevation sites controlled for lower temperatures and higher precipitation of the last glacial period (see Table 1, Heusser, 1998; Daly et al., 2008), allowing for isolation of CO₂ effects. Note that conditions at glacial La Brea were wetter than modern times, which differs from most regions that were drier during the last glacial period. While [CO₂] does not vary with elevation, CO₂ partial pressure decreases in proportion to total atmospheric pressure. Under modern conditions, partial pressures of CO₂ at high-elevation sites are 10-30% lower than

Table 1 Climate data for glacial and modern Juniperus sampling sites

Site category	Site name and coordinates	Elevation (m)	Mean annual precipitation (mm)	Mean annual temperature (°C)
Glacial	La Brea Tar Pits + 34°3′48″, – 118°21′22″	80	с. 600	c. 7.5–9.5
Modern high elevation (SBNF)	Big Bear Lake + 34°16'12", – 116°55'29"	2830	696	7.3
	Hwy 38 Bend + 34°11'35", – 116°47'6"	2300	452	11.1
	Wildhorse Springs + 34°9'52", - 116°43'14"	2300	572	8.7
Modern low elevation (ANF)	Mt. Emma Rd + 34°28′55″, – 118°4′2″	1340	242	14.7
	Littlerock Reservoir + 34°29'42", – 118°1'36"	1045	230	14.9
	Lyttle Creek + 34°11'22", - 117°26'11"	630	390	18.2

Climate data for glacial La Brea (Heusser, 1998), and modern sampling sites (Daly *et al.*, 2008). Modern samples were collected from San Bernardino National Forest (SBNF) and Angeles National Forest (ANF).

at low-elevation sites, producing an even more conservative comparison between glacial and modern conditions.

Stable isotope measurement

We measured stable carbon isotope ratios on alpha-cellulose from individual tree rings of glacial and modern Juniperus. Whole tree rings were analyzed in order to provide an integrated measure of the full annual response. Ring wood was separated under a dissecting microscope and alpha-cellulose was extracted from each ring using the method described by Ward et al. (2005). Previous work using this method has documented high purity levels of alpha-cellulose with no indication of asphalt contamination from the tar pits. Purity was based on theoretical O: H ratios (weight percent oxygen: weight percent hydrogen) of 7.79-8.08 for alpha-cellulose, with actual values falling well within this range $(8.01 \pm 0.02 \text{ and } 7.97 \pm 0.04 \text{ for modern and glacial samples,}$ respectively, Ward et al., 2005). Because our specific compound reflected high purity levels, we do not believe that diagenetic processes would have influenced our results.

Of the five glacial wood specimens that were available with an adequate number of tree rings, three are trunk specimens while two may be either portions of the trunk or large branch sections. For modern trees, the 10 rings nearest the center were excluded, as is common on dendrochronological work, as the juvenile stage often exhibits altered physiological patterns. Apart from this exception, all available rings in all samples were analyzed. Isotope measurements were performed at the Keck Paleoenvironmental and Environmental Stable Isotope Laboratory (KPESIL) at the University of Kansas. δ^{13} C values were calculated using the following formula:

$$\delta = R_{\text{sample}}/R_{\text{standard}} - 1$$

where R is the ratio of ${}^{13}C$: ${}^{12}C$, using belemnite carbonate from the Pee Dee Formation, Hemingway, SC (PDB) as the standard. Data were converted to 'per mil' (%) notation by multiplying δ values by 1000. $\delta^{13}C_{cell}$ was converted to $\delta^{13}C_{leaf}$ using a constant offset of -3.2% (Leavitt & Long, 1982; Ward et al., 2005). Carbon isotope discrimination was calculated as:

$$\Delta = \frac{\delta^{13} C_{\text{air}} - \delta^{13} C_{\text{leaf}}}{1 + \delta^{13} C_{\text{leaf}}}$$

Conversion to carbon discrimination is necessary as it incorporates changes in $\delta^{13}C_{air}$ through time. $\delta^{13}C_{air}$ was -0.0066(-6.6%) during glacial times, but has decreased in modern times to -0.008 (-8.0%); Leuenberger et al., 1992). From Δ , c_i/c_a was calculated as

$$\frac{c_i}{c_a} = \frac{\Delta - a}{b - a}$$

where a is the fractionation against 13 C as a result of slower diffusion across the stomata $(4.4\%_{00})$ and b is the fractionation against ¹³C as a result of Rubisco (27‰).

For each ring, c_i was also calculated from the c_i/c_a ratio using c_a values. For modern samples, c_a values were obtained from direct atmospheric measurements (Keeling et al., 2009) and the Taylor Law Dome ice core (Etheridge et al., 1996). For glacial trees, c_a values were obtained from the Vostok and EPICA Dome C ice cores (Lüthi et al., 2008). To obtain the appropriate c_a values, ¹⁴C ages of glacial trees were first converted to calendar ages in order to coincide with ice core data (Beck et al., 2001). Since atmospheric [CO₂] showed only minimal changes throughout the latter portion of the last glacial period that is encompassed in our study, we are confident that [CO₂] values corresponding to converted ages are accurate to the actual conditions experienced by glacial trees.

Statistical analyses

Mean c_i/c_a values for high- and low-elevation modern *Juniperus* were not significantly different despite environmental differences between these locations (0.53 \pm 0.05 and 0.53 \pm 0.06, P = 0.1; ANOVA), and therefore, the two modern sets were grouped together for comparison to glacial values. Since the variance in c_i/c_a was significantly different between modern and glacial Juniperus (P < 0.0001), a Welch's ANOVA was used to compare modern and glacial c_i/c_a values that account for lack of equivalence of variance.

The coefficient of variation (CV) was calculated for c_i/c_a in both modern and glacial samples. CV provides a measure of dispersion of data around the mean, allowing us to compare variation between groups. CV was calculated as:

$$CV = \frac{s}{\bar{x}}$$

where s is the standard deviation, and \bar{x} is the mean. Data are shown in percentage notation by multiplying CV by 100. In order to account for differences in chronology length between glacial (shorter) and modern (longer) samples, the following correction (Sokal & Rohlf, 1995) was applied to CV:

$$CV_{corr} = \left(1 + \frac{1}{4n}\right)CV$$

Correlation of modern c_i/c_a with climate

To determine correlations of modern c_i/c_a values with climate, monthly temperature and precipitation data were obtained for each site from PRISM (Daly et al., 2008). The PRISM model is ideal for this comparison as it accurately reflects climatic conditions in mountainous coastal regions with large elevational gradients and complex topography (Daly et al., 2008). Measures of temperature and precipitation alone provided only weak correlations with c_i/c_a , so vapor pressure deficit (VPD) was used for this correlation. VPD is a more integrative climatic parameter that combines water and temperature relationships and is closely linked to evapotranspiration, making this measure more directly related to plant physiology than temperature or precipitation alone. VPD was calculated from monthly average maximum ($T_{\rm max}$), minimum ($T_{\rm min}$) and dewpoint ($T_{\rm dew}$) temperatures using:

$$VPD = \frac{e_s(T_{\text{max}}) + e_s(T_{\text{min}})}{2} - e_s(T_{\text{dew}})$$

where $e_s(T)$ is the saturation vapor pressure at temperature T, calculated as:

$$e_s(T) = 0.6112 \exp\left(\frac{17.67 T}{T + 243.5}\right)$$

In order to correlate ring isotopic composition with VPD, rings of modern trees were associated with specific calendar years. Ring width patterns from trees within the same site were correlated and aligned using marker years of high precipitation and growth.

Results and Discussion

The c_i/c_a ratio is driven by two fundamental processes: stomatal conductance, which controls the rate of CO_2 diffusion from the atmosphere into the intercellular spaces of leaves; and chloroplast demand for CO_2 , which is determined by internal CO_2 diffusion rates to carboxylation sites and photosynthetic biochemistry. Long-term trends in c_i/c_a over evolutionary timescales reflect the degree of coordination between processes affecting CO_2 supply and demand within the leaf. In addition, shorter-term trends in c_i/c_a (e.g. annual rings) reflect integrated shifts in tree physiology in response to changing environmental conditions within the lifespan of a single individual.

We found that mean c_i/c_a of *Juniperus* was similar between glacial and modern trees (Fig. 1a; glacial average, 0.52 ± 0.02; modern average, 0.53 ± 0.05 ; P > 0.2). One possible explanation for this, although one not supported by the literature, is that both stomatal conductance and chloroplast demand for CO₂ remained constant across this expansive period of [CO₂] and climatic change. On the other hand, if only one of these factors predominantly changed through time, there would have been shifts in c_i/c_a , which were not observed here. It is therefore most likely that both stomatal conductance and chloroplast demand for CO₂ were higher during the last glacial period, which would have enhanced CO₂ uptake under limiting carbon conditions. When supply and internal demand for CO₂ covary in the same direction, as has been observed even in highly disparate taxa (Franks & Beerling, 2009a), there are opposing effects on c_i/c_a , likely producing the stabilization effect observed here. When moving into the interglacial period, both stomatal conductance and chloroplast CO₂ demand likely decreased, with the effect of saving water and nitrogen as CO₂ became less limiting. In support of this idea, Ehleringer & Cerling (1995) hypothesized that c_i/c_a represents a metabolic set point that is maintained within species across time. In addition, increases in stomatal conductance are almost always observed in modern C₃ plants grown at low [CO₂] (Gerhart & Ward, 2010), and studies with glacial leaves show

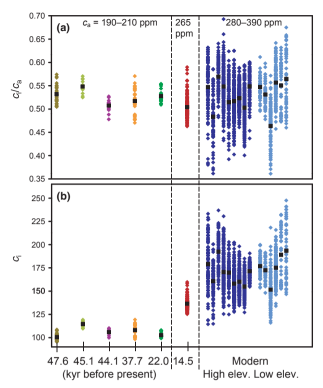


Fig. 1 Results of stable carbon isotope measurements for glacial and modern *Juniperus* tree rings. (a) Intercellular: atmospheric $[CO_2]$ ratio (c_i/c_a) values); (b) corresponding intercellular $[CO_2]$ (c_i values). Each point represents an individual tree ring, and vertical groups represent results from all available tree rings for an individual tree, with values stacked from highest to lowest. Glacial samples are shown in different colors, while modern samples are grouped by elevation (to distinguish two different control groups), with means labeled as black boxes. Atmospheric $[CO_2]$ values (c_a) are provided for each group. The range provided for modern samples reflects a temporal gradient experienced by each tree over its lifetime as a result of rapid changes in atmospheric $[CO_2]$ in the modern period.

evidence for increased stomatal density and decreased stomatal size, which would have increased maximum stomatal conductance in the past (Beerling *et al.*, 1993; Franks & Beerling, 2009b; but also see Malone *et al.*, 1993 for responses of modern plants grown at low [CO₂]). The wetter conditions of the last glacial period may have also provided increased nitrogen availability to support higher leaf nitrogen contents, which may have enhanced photosynthetic capacity.

Despite any physiological adjustments, c_i values remained extremely low in glacial trees relative to modern trees as a result of consistently low c_a throughout the last glacial period (Fig 1b; glacial average, 106 ± 6 ; modern average, 168 ± 20 ; P < 0.0001). Past studies have reported similarly low c_i values in glacial needles of *Pinus flexilis* preserved in packrat middens (Van de Water *et al.*, 1993; Beerling, 1994). When considering all available rings, the vast majority of glacial c_i values fell outside the range of modern values. In fact, no modern trees experienced c_i values below 114 ppm, and no glacial trees experienced values > 120 ppm, leaving only a narrow overlapping range. It is also interesting to note that c_i values of glacial trees never fell below 90 ppm over an integrated annual period. This suggests that this

may be a limiting concentration below which juniper trees may not maintain a positive carbon budget for basic physiological functions for survival (Campbell *et al.*, 2005).

Interannual variation in c_i/c_a , represented by CV_{corr} , was significantly lower in glacial vs modern trees (P < 0.0002; Fig. 2). More specifically, low- and high-elevation modern trees showed CV_{corr} values of 8 ± 2% and 8 ± 3%, respectively. The only available Holocene specimen (14.5 kyr BP) showed an intermediate CV_{corr} value of 5%, while glacial specimens showed the lowest values, averaging 3 ± 1%. Furthermore, although modern trees show occasional, short-term periods of low interannual variation, these periods are rare. Glacial trees show consistently low variation in c_i/c_a in all cases. In fact, the two oldest glacial *Juniperus* samples (45.1 and 47.6 kyr old) correspond to the timing of D–O events recorded in Greenland glaciers

(Blunier & Brook, 2001). The maintenance of low interannual variation in c_i/c_a , even during time periods of rapid and drastic environmental change that are characteristic of D–O cycles, suggests that the maintenance of low variation in glacial *Juniperus* physiology was consistent throughout the last glacial period.

In a plethora of past studies, modern *Juniperus* in southern California and the southwestern US exhibits high interannual variation in c_i/c_a , mainly as a result of changes in soil water availability from year to year (Leavitt & Long, 1989; Feng & Epstein, 1995; Moore *et al.*, 1999; Leffler *et al.*, 2002). In our study, the c_i/c_a of modern trees showed the strongest correlations with monthly or seasonal VPD ($R^2 = 0.06-0.25$; P < 0.05-0.0001), whereby the months showing the strongest correlations were offset between elevations. Although these correlations were relatively low, similar correlations have been reported for modern *Juniperus*

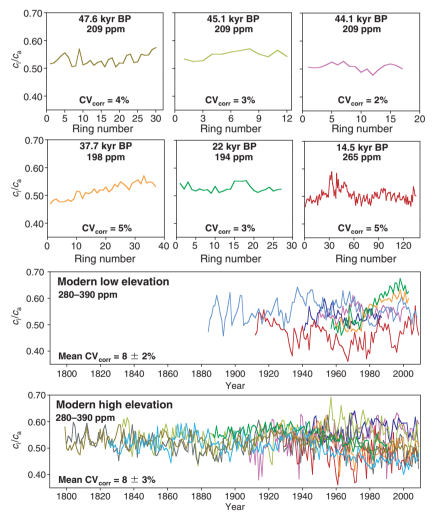


Fig. 2 Annual responses of intercellular: atmospheric $[CO_2]$ ratio (c_i/c_a) for modern and glacial *Juniperus*. These are the same data as in Fig. 1, although in this case, data are placed in chronological order throughout the development of each tree. Full chronologies are not available for glacial trees, and therefore data are arranged from youngest (ring number 1) to oldest. Glacial samples are shown in the same colors as in Fig. 1, although modern samples are given different colors in order to distinguish their responses. ¹⁴C age (thousands of years before present, kyr BP) atmospheric $[CO_2]$, and CV_{corr} (interannual variation in c_i/c_a , see the 'Materials and Methods' section for details) are provided for each sample and/or control group for the sake of comparison. The atmospheric $[CO_2]$ range provided for modern samples reflects a temporal gradient experienced by each tree over its lifetime as a result of rapid changes in atmospheric $[CO_2]$ in the modern period. [Correction added after online publication 20 December 2012: a corrected version of Fig. 2 is now published here, where the following corrections have been applied: Row 1, panel three: the text which previously read as ' $CV_{corr} = 3\%$ ' now reads as ' $CV_{corr} = 5\%$ '].

in other studies (Leavitt & Long, 1989), and in all cases, soil water parameters and/or VPD correlate most closely with tree physiology (Leavitt & Long, 1989; Feng & Epstein, 1995; Moore *et al.*, 1999; Leffler *et al.*, 2002).

Since glacial climate was much more variable than in modern times, one would expect glacial c_i/c_a to also show higher variation if trees were responding to similar climatic factors. To the contrary, we found reduced amounts of interannual variation in c_i/c_a during the last glacial period (Fig. 2), indicating that a stable environmental factor dominated tree physiology. During the last glacial period, [CO2] was extremely stable from year to year (EPICA, 2004), while water availability and temperature were predicted to have been highly variable (Behl & Kennett, 1996; Heusser, 1998; Hendy & Kennett, 1999; Hendy et al., 2002). In our study, extremely low c_i values coupled with reduced variation in c_i/c_a even under a highly fluctuating glacial climate, point strongly to low [CO₂] constraints on tree physiology. While short-term studies with modern plants grown at glacial [CO₂] show major carbon limitations on physiology, our findings highlight the strength and consistency of low CO₂ constraints over evolutionary timescales.

In conclusion, this study has demonstrated that mean c_i/c_a has been maintained in *Juniperus* between the last glacial period and modern times, despite changes in temperature, precipitation and $[CO_2]$; that glacial c_i values were extremely low on an annual basis and occur only rarely in modern trees; that a limiting level for *Juniperus* physiology may exist at or near 90 ppm; and that interannual variation in c_i/c_a was greatly reduced in glacial *Juniperus*, likely as a result of the constraints of low $[CO_2]$ that overrode responses to other climatic factors. This is the first direct evidence from trees that actually lived and evolved under low $[CO_2]$ that carbon limitation persisted on an annual basis during the last glacial period. Moreover, our results suggest that the environmental factors that dominate carbon-uptake physiology can vary across geologic timescales, resulting in major alterations in physiological response patterns through time.

Acknowledgements

This work was supported by the National Science Foundation (NSF) CAREER (0746822) and PECASE awards, and the Wohlgemuth Faculty Scholar Award (endowed by D. Lynch) to J.K.W. The NSF C-CHANGE IGERT fellowship (0801522) and the Madison and Lila Self Graduate Fellowship supported L.M.G. at KU. The Department of Ecology and Evolutionary Biology (KU) also provided assistance for this work. The authors thank G. Cane (KPESIL, KU), E. Duffy (EEB, KU) and K. Colvin (EEB, KU) for assistance with isotope analysis, and J. Nickerman (Angeles National Forest) and M. Lardner (San Bernardino National Forest) for field permit assistance. The authors also thank three anonymous reviewers for their helpful comments.

References

Beck JW, Richards DA, Edwards RL, Silverman BW, Smart PL, Donahue DJ, Hererr-Osterheld S, Burr GS, Calsoyas L, Jull AJT *et al.* 2001. Extremely

- large variations of atmospheric ¹⁴C concentration during the last glacial period. *Science* **292**: 2453–2458.
- Beerling DJ. 1994. Predicting leaf gas exchange and δ^{13} C responses to the past 30,000 years of global environmental change. *New Phytologist* 128: 425–433.
- Beerling DJ, Chaloner WG, Huntley B, Pearson JA, Tooley MJ. 1993. Stomatal density responds to the glacial cycle of environmental change. *Proceedings of the Royal Society of London Series B-Biological Sciences* 253: 53–60.
- Behl RJ, Kennett JP. 1996. Brief interstadial events in the Santa Barbara basin, NE Pacific, during the past 60 kyr. *Nature* 379: 243–246.
- Berner RA. 2006. GEOCARBSULF: a combined model for Phanerozoic atmospheric O₂ and CO₂. *Geochimica et Cosmochiica Acta* 70: 5653–5664.
- Blunier T, Brook EJ. 2001. Timing of millennial-scale climate change in Antarctica and Greenland during the last glacial period. Science 291: 109–112.
- Broecker WS. 1994. Massive iceberg discharges as triggers for global climate change. *Nature* 372: 421–424.
- Campbell CD, Sage RF, Kocacinar F, Way DA. 2005. Estimation of the wholeplant CO2-compensation point of tobacco (*Nicotiana tabacum* L.). Global Change Biology 11: 1956–1967.
- Daly C, Halbleib M, Smith JI, Gibson WP, Doggett MK, Taylor GH, Curtis J, Pasteris PP. 2008. Physiographically sensitive mapping of climatological temperature and precipitation across the conterminous United States. *International Journal of Climatology* 28: 2031–2064.
- Dansgaard W, Johnsen SJ, Clausen HB, Dahl-Jensen D, Gundestrup NS, Hammer CU, Hvidberg CS, Steffensen JP, Sveinbjornsdottir AE, Jouzel J *et al.* 1993. Evidence for general instability of past climate from a 25-kyr ice-core record. *Nature* 364: 218–220.
- Dippery JK, Tissue DT, Thomas RB, Strain BR. 1995. Effects of low and elevated CO_2 on C_3 and C_4 annuals. I. Growth and biomass allocation. *Oecologia* 101: 13–20.
- Ehleringer JR, Cerling TE. 1995. Atmospheric CO₂ and the ratio of intercellular to ambient CO₂ concentrations in plants. *Tree Physiology* 15: 105–111.
- EPICA. 2004. Eight glacial cycles from an Antarctic ice core. *Nature* 429: 623–628.
- Etheridge DM, Steele LP, Langenfelds RL, Francey RJ, Barnola J-M, Morgan VI. 1996. Natural and anthropogenic changes in atmospheric CO₂ over the last 1,000 years from air in Antarctic ice and firn. *Journal of Geophysical Research* 101: 4115–4128.
- Feng X, Epstein S. 1995. Carbon isotopes of trees from arid environments and implications for reconstructing atmospheric CO₂ concentration. *Geochimica et Cosmochimica Acta* 59: 2599–2608.
- François LM, Ghislain M, Otto D, Micheels A. 2006. Late Miocene vegetation reconstruction with the CARAIB model. *Palaeogeography Palaeoclimatology Palaeoecology* 238: 302–320.
- Franks PJ, Beerling DJ. 2009a. CO₂-forced evolution of plant gas exchange capacity and water-use efficiency over the Phanerozoic. *Geobiology* 7: 227–236.
- Franks PJ, Beerling DJ. 2009b. Maximum leaf conductance driven by CO₂ effects on stomatal size and density over geologic time. *Proceedings of the National Academy of Sciences, USA* 106: 10343–10347.
- Gerhart LM, Ward JK. 2010. Plant responses to low [CO₂] of the past. New Phytologist 188: 674–695.
- Hendy IL, Kennett JP. 1999. Latest Quaternary North Pacific surface-water responses imply atmosphere-driven climate instability. Geology 27: 291–294.
- Hendy IL, Kennett JP, Roark EB, Ingram BL. 2002. Apparent synchroneity of submillennial scale climate events between Greenland and Santa Barbara Basin, California from 30–10 ka. *Quaternary Science Reviews* 21: 1167–1184.
- Heusser L. 1998. Direct correlation of millennial-scale changes in western North American vegetation and climate with changes in the California Current System over the past ~ 60 kyr. *Paleoceanography* 13: 252–262.
- Keeling RF, Piper SC, Bollenbacher AF, Walker JS. 2009. Atmospheric CO₂ records from sites in the SIO air sampling network. In: Trends: a compendium of data on global change. Oak Ridge, TN, USA: Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, US Department of Energy doi: 10.334/CDIAC/atg.035
- Leavitt SW, Long A. 1982. Evidence for ¹³C/¹²C fractionation between tree leaves and wood. *Nature* 298: 742–744.

- Leavitt SW, Long A. 1989. Drought indicated in carbon-13/carbon-12 ratios of southwestern tree rings. Water Resource Bulletin 25: 341–347.
- Leffler AJ, Ryel RJ, Hipps L, Ivans S, Caldwell MM. 2002. Carbon acquisition and water use in a northern Utah *Juniperus osteosperma* (Utah juniper) population. *Tree Physiology* 22: 1221–1230.
- Leuenberger M, Siegenthaler U, Langway C. 1992. Carbon isotope composition of atmospheric CO₂ during the last ice age from an Antarctic ice core. *Nature* 357: 488–490.
- Lüthi D, Le Floch M, Bereiter B, Blunier T, Barnola J-M, Siegenthaler U, Raynaud D, Jouzel J, Fischer H, Kawamura K *et al.* 2008. High-resolution carbon dioxide concentration record 650,000–800,000 years before present. *Nature* 453: 379–382.
- Malone SR, Mayeux HS, Johnson HB, Polley HW. 1993. Stomatal density and aperture length in four plan species grown across a subambient CO₂ gradient. *American Journal of Botany* 80: 1413–1418.
- Mayewski PA, Eohling EE, Stager JC, Karlén W, Maasch KA, Meeker LD, Meyerson EA, Gasse F, van Kreveld S, Holmgren K et al. 2004. Holocene climate variability. Quaternary Research 62: 243–255.
- Moore DJ, Nowak RS, Tausch RJ. 1999. Gas exchange and carbon isotope discrimination of *Juniperus osteosperma* and *Juniperus oscidentalis* across environmental gradients in the Great Basin of western North America. *Tree Physiology* 19: 421–433.
- Pagani M, Caldeira K, Berner R, Beerling DJ. 2009. The role of terrestrial plants in limiting atmospheric CO₂ decline over the past 24 million years. *Nature* 460: 85–88.

- Polley HW, Johnson HB, Marino BD, Mayeux HS. 1993. Increase in C₃ plant water-use efficiency and biomass over glacial to present CO₂ concentrations. *Nature* 361: 61–64.
- Roy K, Valentine JW, Jablonski D, Kidwell SM. 1996. Scales of climatic variability and time averaging in Pleistocene biotas: implications for ecology and evolution. *Trends in Ecology and Evolution* 11: 458–463.
- Sage RF, Coleman JR. 2001. Effects of low atmospheric CO₂ in plants: more than a thing of the past. *Trends in Plant Science* 6: 18–24.
- Sokal RR, Rohlf FJ. 1995. Biometry, 3rd edn. New York, NY, USA: Freeman & Co.
- Turcq B, Cordeiro RC, Sifeddine A, Simões Filho FFL, Albuquerque ALS, Abrão JJ. 2002. Carbon storage in Amazonia during the Last Glacial Maximum: secondary data and uncertainties. *Chemosphere* 49: 821–835.
- Van de Water PK, Leavitt SW, Betancourt JL. 1993. Trends in stomatal density and ¹³C/¹²C ratios of *Pinus flexilis* needles during last glacial-interglacial cycle. *Science* 264: 239–243.
- Ward JK, Harris JM, Cerling TE, Wiedenhoeft A, Lott MJ, Dearing M-D, Coltrain JB, Ehleringer JR. 2005. Carbon starvation in glacial trees recovered from the La Brea tar pits, southern California. Proceedings of the National Academy of Sciences, USA 102: 690–694.
- Ward JK, Kelly JK. 2004. Scaling up evolutionary responses to elevated CO₂: lessons from *Arabidopsis. Ecology Letters* 7: 427–440.
- Ward JK, Myers DA, Thomas RB. 2008. Physiological and growth responses of C₃ and C₄ plants to reduced temperature when grown at low CO₂ of the last ice age. *Journal of Integrative Plant Biology* **50**: 1388–1395.



About New Phytologist

- New Phytologist is an electronic (online-only) journal owned by the New Phytologist Trust, a not-for-profit organization dedicated
 to the promotion of plant science, facilitating projects from symposia to free access for our Tansley reviews.
- Regular papers, Letters, Research reviews, Rapid reports and both Modelling/Theory and Methods papers are encouraged.
 We are committed to rapid processing, from online submission through to publication 'as ready' via Early View our average time to decision is <25 days. There are no page or colour charges and a PDF version will be provided for each article.
- The journal is available online at Wiley Online Library. Visit **www.newphytologist.com** to search the articles and register for table of contents email alerts.
- If you have any questions, do get in touch with Central Office (np-centraloffice@lancaster.ac.uk) or, if it is more convenient, our USA Office (np-usaoffice@ornl.gov)
- For submission instructions, subscription and all the latest information visit www.newphytologist.com