Phylogenetic analyses reveal the shady history of C₄ grasses

Erika J. Edwards and Stephen A. Smith

Grasslands cover more than 20% of the Earth’s terrestrial surface, and their rise to dominance is one of the most dramatic events of biome evolution in Earth history. Grasses possess two main photosynthetic pathways: the C₃ pathway that is typical of most plants and a specialized C₄ pathway that minimizes photorespiration and thus increases photosynthetic performance in high-temperature and/or low-CO₂ environments. C₄ grasses dominate tropical and subtropical grasslands and savannas, and C₃ grasses dominate the world’s cooler temperate grassland regions. This striking pattern has been attributed to C₄ physiology, with the implication that the evolution of the pathway enabled C₄ grasses to persist in warmer climates than their C₃ relatives. We combined geospatial and molecular sequence data from two public archives to produce a 1,230-taxon phylogeny of the grasses with accompanying climate data for all species, extracted from more than 1.1 million herbarium specimens. Here we show that grasses are ancestrally a warm-adapted clade and that C₄ evolution was not correlated with shifts between temperate and tropical biomes. Instead, 18 of 20 inferred C₄ origins were correlated with marked reductions in mean annual precipitation. These changes are consistent with a shift out of tropical forest environments and into tropical woodland/savanna systems. We conclude that C₄ evolution in grasses coincided largely with migration out of the understory and into open-canopy environments. Furthermore, we argue that the evolution of cold tolerance in certain C₃ lineages is an overlooked innovation that has profoundly influenced the patterning of grassland communities across the globe.

C₄ photosynthesis | climate niche evolution | cold tolerance | phylogeny

The term “C₄ photosynthesis” refers to a suite of biochemical and anatomical modifications to the standard plant C₃ photosynthetic pathway that work to concentrate CO₂ around the carbon-fixing enzyme Rubisco. The C₄ pathway greatly improves photosynthetic performance in situations that promote photorespiration, typically high-temperature and low-CO₂ environments (1). The pathway also promotes more efficient photosynthetic water use, because the CO₂ concentration mechanism allows C₄ plants to maintain a lower stomatal conductance for a given photosynthetic rate. C₄ photosynthesis is estimated to have evolved at least 50 times in terrestrial plants (2, 3) and is most prominent in grasses, where roughly half the species (~5,000) are C₄ (4), including economically important species such as maize, sugarcane, sorghum, and switchgrass. C₄ grasses currently dominate wide regions of the Earth, and it is estimated that they account for up to 25% of global annual terrestrial primary production (5).

Recent studies suggest that C₄ photosynthesis is a relatively recent innovation in plants, with the earliest appearances coinciding with plummeting atmospheric CO₂ levels during the mid-Oligocene and many origins occurring much later (4, 6). In grasses, the evolutionary history of C₄ photosynthesis is complex, with multiple origins, probable reversals, and a general lag-time between the evolution of the pathway and the formation of C₄-dominated ecosystems (3, 6–8). Although low atmospheric CO₂ certainly was a prerequisite for C₄ evolution, it is thought that multiple stressors worked in concert to promote the pathway. C₄ plants are notably prevalent in arid, high-light, saline, and disturbed environments, and it is largely accepted that water stress has provided a strong selection pressure for C₄ evolution in eudicots (4). Grasses have long been viewed as an interesting exception to this pattern (9). Significant positive correlations between C₄ grass abundance and growing season temperature have been documented at both continental and regional scales (10–13); C₄ grasses dominate tropical grasslands and savannas but are virtually absent from cool-temperate grasslands and steppes. Furthermore, both experimental measurements of photosynthetic light use efficiency (termed “quantum yield”), and predictions of leaf models of C₃ and C₄ photosynthesis provide strong evidence that C₄ grasses outperform C₃ grasses at higher temperatures (5, 14–16). Most studies concerning C₄ grasses and precipitation have focused on C₄/C₃ mixed temperate grassland systems, where the timing of C₄ growth often is restricted to periods with significant rainfall (12, 13, 17, 18).

The C₄ pathway thus has been largely dismissed as an adaptation to water stress in grasses, with all data indicating that C₄ evolution allowed grasses to invade and diversify successfully into hot climates. However, few studies have compared C₄ grasses with their closest living C₃ relatives. C₃ origins are not distributed uniformly across Poaceae but are clustered in one major grass lineage, informally named the “PACMAD” clade (3, 6). Most of the C₃ grasses that dominate cool-climate grasslands belong to the Pooideae, a lineage that last shared a common ancestor with PACMAD grasses ~65–50 Mya (19). It thus is possible that differences between Pooideae and PACMAD grasses that have nothing to do with photosynthetic pathway variation are driving the apparently strong sorting of C₃ and C₄ species along temperature gradients.

We employed an explicitly phylogenetic approach to assess the evolutionary history of climate niche space for grasses on a worldwide scale. We used two public archives of data to build the most inclusive phylogeny for Poaceae that permitted analysis of a climate dataset for all taxa (Materials and Methods). This analysis resulted in a 1,230-taxon tree with broad coverage of all of the major Poaceae lineages and, importantly, good sampling of known C₃ and C₄ transitions. Our phylogenetic tree includes roughly 10% of all Poaceae species and is the largest grass phylogeny built to date. We identified 21 nodes representing evolutionary transitions between photosynthetic types and used these nodes to generate phylogenetically independent C₃/C₄ pairwise comparisons (Tables S1 and S2) (20). Nearly all identified photosynthetic transitions were reconstructed as C₄ origins, with one purported reversal. Fifteen of the 21 transitions occurred within the Panoideae, a major PACMAD lineage containing more than 3,000 species. To account for topological uncertainty in this area of the phylogeny, we performed Bayesian analyses for a 299-taxon Panoideae dataset and ran all diver-

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To whom correspondence should be addressed. E-mail: erika.edwards@brown.edu.

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gence analyses across the Bayesian posterior distribution of trees. We also used this reduced dataset to reconstruct environmental niche evolution in the Panicoideae and tested whether shifts between photosynthetic types corresponded with significant changes in temperature and precipitation niche optima under a stabilizing selection model of evolution.

**Results**

Climate data extracted from all available geo-referenced herbarium material provided clear evidence that certain grass lineages have specialized in certain habitats (Fig. 1). Importantly, two strictly C3 grass lineages, the Pooidae and the Danthonioideae, stood apart as inhabiting much cooler environments, measured either by mean annual temperature (MAT) (Fig. 1) or temperature of the wettest, coldest, or warmest month (Fig. S1). All other C3 lineages were indistinguishable from C4 lineages with respect to their temperature profiles. Distinctive sorting of precipitation variables was less apparent, although Pooidae occupied the drier end of the spectrum alongside the C4 lineages Aristidoideae and Chloridoideae (Fig. S2).

Our phylogenetic analyses concurred with these general temperature and precipitation profiles for the different grass clades, particularly the observation that, with the exception of Pooidae and Danthonioideae, grasses are warm-climate specialists (Fig. 2). The evolution of C4 photosynthesis appears to have had little influence on gross temperature niche: Only 10 of 21 photosynthetic shifts resulted in increases in MAT in the C4 lineage, with a mean divergence of −0.13 °C. Likewise, there was no significant relationship between photosynthetic pathway and mean temperature of the hottest, coldest, or warmest months (Table 1). In sharp contrast, there were large and predictable shifts in mean annual precipitation (MAP) between C3 and C4 sister taxa: 18 of the 21 divergences resulted in C4 taxa living in areas with lower MAP (P < 0.01), with a mean difference of 546 mm year−1. Fifteen of those divergences also resulted in shifts toward increased seasonality of precipitation. Results were statistically similar using a more conservative, reduced dataset generated by removing three focal taxa with questionable placement in the phylogeny and recoding a C3–C4 intermediate species as C3 (Table 1) and also were robust to testing against a wide range of alternative tree topologies within Panicoideae (Table S3). These results are highly consistent with an earlier analysis of the Hawaiian grass flora that used a similar approach but was based on very limited geographic and phylogenetic sampling (21).

Analyses of niche evolution within Panicoideae provided further support that C4 evolution was associated with shifts into drier, but not warmer, environments. For most climate variables, two-optimum stabilizing selection Ornstein-Uhlenbeck (OU) models were preferred over Brownian motion or single-optimum models, implying that C3 and C4 lineages have experienced divergent selection over time (Table 2). However, the mean C3 and C4 temperature optima were not largely different, and in all cases, the model inferred lower temperature optima for C4 Panicoideae lineages than for C3 Panicoideae lineages. MAP received the strongest support for a two-optimum model, with the C4 optimum inferred to be drier than the C3 optimum by more than 500 mm year−1.

**Discussion**

These analyses provide clear evidence that C4 origins in grasses coincided with ecological shifts into drier environments. However, the actual values of precipitation divergences across our datasets largely occurred squarely within what generally is considered a mesophytic moisture niche (22). No reconstructed C4 ancestral focal node had a MAP of less than 500 mm year−1, across all C3/C4 contrasts, the C3 MAP average was 1,839 mm year−1, and the C4 average was 1,229 mm year−1. In the tropics, such differences in annual precipitation values often lead to major changes in biome structure, because a MAP of 1,500 mm year−1 generally is considered the minimum amount of rainfall required to support a closed-canopy forest (23). Considering the relative values of MAT, MAP, and precipitation seasonality, one-third of our focal nodes present climate profiles that are highly consistent with a shift from a closed-canopy tropical moist forest to an open tropical woodland/savanna biome (Fig. 3) (24). This finding also is consistent with the unexpected and significant increase in temperature seasonality in C4 lineages, suggesting geographic movements north and south from the equator and into the savanna belt. This general picture of a shift from closed to open canopies mirrors results of an independent dataset that focused on qualitative ecological assessments for 117 genera of grasses (22) and suggests a direct link between C4 evolution and the establishment of the savanna biome.

![Species accumulation curves for mean annual temperature and precipitation, sorted by major grass lineage. These data represent 1,584,351 independent collection localities spread across 10,469 taxa. Each point in the curve is a species’ mean value.](image-url)
Although our data provide strong evidence that C₄ origins were not correlated with movements between cooler and warmer climates at a macroecological scale, these results do not preclude high temperature as a potential factor in driving the evolution of the pathway. Ultimately, C₄ photosynthesis is an evolutionary response to debilitating levels of photorespiration, so any hypotheses regarding the ecological drivers of C₄ evolution must include an explanation of how that new environment translates into increased photorespiration at the leaf level. In the case of a shift from tropical moist forests to tropical open-canopy systems, we propose two alternatives. First, water limitation caused by lower rainfall could promote photorespiration directly via increased stomatal resistance and lowered internal leaf CO₂ levels. However, it is possible that these initial reductions in precipitation had little effect on plant water status, particularly if species shifted their timing of growth to coincide with the rainy season (as most C₄ grasses do today). Instead, changes in precipitation may have had an indirect effect on C₄ evolution by limiting canopy growth, thereby creating a high-irradiance environment that simultaneously would raise leaf temperatures and provide the high light needed to drive the CO₂-concentrating mechanism. Thus, high temperatures still could be an important driving force of C₄ evolution at the leaf level. Discerning between these two hypotheses requires more information than can be gleaned from the macroecological approach presented here. In fact, this uncertainty is a fine illustration of the limitations inherent in interpreting ecological function from geographical distribution data; what we ultimately need are more field studies comparing important eco-physiological parameters of different organisms. In this case, we currently lack critical information about the seasonal water status, gas exchange, and phenology of grass species spanning these C₃/C₄ focal nodes.

Finally, it is clear that the C₄ pathway, although a complex and ecologically important innovation, is not an evolutionarily difficult feat for grasses. Certain aspects of grass leaf anatomy and genetics probably precondition these species toward developing the syndrome whenever the right environmental conditions arise.

Fig. 2. The evolution of photosynthetic pathway and temperature niche in grasses. (A) Green lines indicate C₃ photosynthesis; black lines indicate C₄ photosynthesis. Maximum likelihood methods reconstructed 20 origins of C₄ photosynthesis and one reversal to C₃ photosynthesis. (B) Maximum likelihood reconstructions of mean annual temperature (MAT), using species’ mean values that were generated from 1,146,612 geo-referenced herbarium specimens.
These factors have been reviewed extensively elsewhere (25, 26) and include traits such as high leaf vein densities, enlarged bundle sheaths, and a propensity for whole-genome duplications. Diversification into cold climates, on the other hand, has happened much more rarely. Thus grasses lend further support to the “tropical conservatism” hypothesis, generated by the observation that relatively few lineages have diversified successfully outside of the tropics, perhaps because low temperature is one of the more difficult environmental barriers for organisms to overcome (27). The Pooidae stand out as an exceptional lineage of grasses in that they occupy both the coldest and the driest climate space in Poaceae. They should not be thought of as typical C3 grasses nor as a starting point for comparative C3/C4 physiology and ecology. Traits that promote chilling and frost tolerance in Pooidae have received attention with regard to understanding the physiology of winter crop cereals (28–30) but have not really been considered in the context of global grassland ecology. It is likely that the evolution of cold tolerance in the Pooidae has been just as relevant as C4 photosynthesis in shaping current global patterns of grass distribution.

Materials and Methods

Climate Dataset. We extracted all geo-referenced herbarium specimens housed in herbaria and natural history collections that have been made available via the Global Biodiversity Information Facility (GBIF) web portal (http://www.gbif.org/). For each set of coordinates we extracted monthly temperature and precipitation values from the Climate Research Unit 10° global gridded climate layers (31). After we purged duplicate records and outliers, our global gridded climate dataset consisted of 1,584,351 independent collection points spanning 10,469 taxa, including subspecies, varieties, and hybrids. From these collection points, we excluded all hybrid taxa as well as taxa that were represented by fewer than 10 independent localities, reducing the dataset to 1,146,612 collection points spanning 4,309 taxa. The mean number of collections per species was 932, although this value was heavily influenced by several taxa (e.g., Nardus stricta, Danthonia decumbens, Phragmites australis, Poa pratensis) with very large numbers of collections. After the 5% most heavily collected taxa were removed, the mean number of collections per species for the remaining 95% was 89.

Estimating geographical ranges from herbarium specimens is prone to inherent biases in sampling; for instance, in our dataset, Europe and North America are far more heavily sampled than tropical regions (Fig. S3). However, most of this imbalance results from repeated collecting of a small handful of widespread taxa that represent a very small number of tips on our phylogeny. In general, with the exception of India, the GBIF coverage of the major tropical and subtropical grassland areas (e.g., Eastern Africa, Northern Australia, Northern South America) is quite good.

Phylogenetic Tree Construction. We assembled a 1,230-taxon DNA sequence matrix that consisted of chloroplast regions atpB (59 taxa), matK (266 taxa), ndhF (437 taxa), rbcL (251 taxa), rpl16 (176 taxa), and trnL-trnF (810 taxa), and nuclear regions phyB (93 taxa) and the internal transcribed spacer (ITS; 753 taxa), using the Phylogeny Assembly with Databases (PHLAWD) tool (http://code.google.com/p/phlawd) (32). Before matrix building, we filtered all sequences retrieved from the National Center for Biotechnology Information (NCBI) through our 4,309-taxon list to ensure a complete climate dataset across the tips of the tree. All sequence alignments were conducted using MUSCLE (v. 3.6) (33). The phylogeny was constructed using RAXML 7.1.0 (34) with all genes partitioned, allowing each gene region to have independent parameter estimates for molecular rate matrices. We employed a GTRGAMMA model of nucleotide substitution, with a GTR substitution model and a I* model of among-site rate heterogeneity. All matrices and trees can be obtained from the Dryad repository (199122).

Table 1. Independent contrast analyses for temperature and precipitation variables

<table>
<thead>
<tr>
<th>Variable</th>
<th>Contrast (C4 relative to C3)</th>
<th>Number of positive contrasts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full dataset (n = 21)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MAT (°C)</td>
<td>−0.13</td>
<td>10</td>
</tr>
<tr>
<td>Tmax (°C)</td>
<td>1.24</td>
<td>14</td>
</tr>
<tr>
<td>Tmin (°C)</td>
<td>−1.71</td>
<td>11</td>
</tr>
<tr>
<td>Tseasonality (°C)</td>
<td>1.07</td>
<td>16*</td>
</tr>
<tr>
<td>MAP (mm year⁻¹)</td>
<td>−546**</td>
<td>3**</td>
</tr>
<tr>
<td>Pseasonality</td>
<td>0.20**</td>
<td>15</td>
</tr>
<tr>
<td>Reduced dataset (n = 17)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>MAT (°C)</td>
<td>0.60</td>
<td>9</td>
</tr>
<tr>
<td>Tmax (°C)</td>
<td>1.30</td>
<td>11</td>
</tr>
<tr>
<td>Tmin (°C)</td>
<td>−0.202</td>
<td>10</td>
</tr>
<tr>
<td>Tseasonality (°C)</td>
<td>1.14</td>
<td>9</td>
</tr>
<tr>
<td>MAP (mm year⁻¹)</td>
<td>−541**</td>
<td>2**</td>
</tr>
<tr>
<td>Pseasonality</td>
<td>0.21**</td>
<td>13*</td>
</tr>
</tbody>
</table>

MAP, mean annual precipitation; MAT, mean annual temperature; Pseasonality, coefficient of variation of monthly average precipitation; Tseasonality, average temperature of the warmest month; Tmin, average temperature of the coldest month; Tseasonality, SD of monthly average temperature; Tseasonality, average temperature of the wettest month; *, P < 0.05; **, P < 0.01.

Table 2. Mean modeled selection optima for temperature and precipitation variables in Panicoideae, enforcing separate optima for C3 and C4 lineages

<table>
<thead>
<tr>
<th>Variable</th>
<th>OU attraction strength (± SD)</th>
<th>C3 mean (± SD)</th>
<th>C4 mean (± SD)</th>
<th>ΔAICc*</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAT (°C)</td>
<td>5.94 (2.29)</td>
<td>21.97 (0.14)</td>
<td>20.12 (0.11)</td>
<td>5.46</td>
</tr>
<tr>
<td>Tmax (°C)</td>
<td>4.66 (0.68)</td>
<td>25.45 (0.14)</td>
<td>25.46 (0.16)</td>
<td>−0.05</td>
</tr>
<tr>
<td>Tmin (°C)</td>
<td>6.08 (2.50)</td>
<td>18.11 (0.15)</td>
<td>14.27 (0.12)</td>
<td>7.91</td>
</tr>
<tr>
<td>Tseasonality (°C)</td>
<td>5.31 (0.93)</td>
<td>23.27 (0.10)</td>
<td>22.68 (0.10)</td>
<td>1.14</td>
</tr>
<tr>
<td>MAP (mm year⁻¹)</td>
<td>6.23 (2.63)</td>
<td>2.60 (0.04)</td>
<td>4.02 (0.03)</td>
<td>11.65</td>
</tr>
<tr>
<td>Pseasonality</td>
<td>5.33 (1.00)</td>
<td>1781 (25.51)</td>
<td>1269 (16.94)</td>
<td>34.06</td>
</tr>
</tbody>
</table>

* Larger ΔAICc numbers indicate stronger support for a two-optimum model.
the authors. We rooted Poaceae by including known close relatives Flagellaria indica, Edeiocolea monostachya, and Joinvillea ascendens, which subsequently were pruned from the tree for character analyses. Rapid bootstrap analyses were conducted on the dataset before a final maximum likelihood analysis, which used 10 random bootstrap resamples as starting trees. The Panicoideae were found to be monophyletic in the constructed phylogeny and therefore were used for phylogenetically informed trait analyses. This dataset was excised from the larger dataset and resulted in 299 taxa and 6128 sites. A second phylogenetic analysis on Panicoideae was conducted using MrBayes v. 3.1.2 (35) with a GTR +I+Γ model. The MrBayes analysis was conducted with two runs of four chains each for 3,000,000 generations of burn-in. Convergence and burn-in were determined by examining time-series plots of likelihood scores and parameter estimates, as well as by examining the effective sample size. Trees were used after removing 3,000,000 generations of burn-in.

**Dating.** Methods for creating a time-calibrated phylogeny for Poaceae are limited because of the size of the dataset. To overcome this problem, we first constructed a pruned Poaceae tree of 300 tips and ran the nonparametric rate smoothing procedure as implemented in r8s (36). We used the Powell algorithm and restarted three times to verify convergence. We extracted the estimated ages for all internal nodes in the reduced dataset and used these ages as calibrations for the full dataset. To date the remaining nodes in the full dataset, we used the nonparametric dating method PATHd8 (37). For the initial r8s run, we fixed the age of the node subtending crown Poaceae (BEP + PACMAD lineages) to 65 Ma (19). We also set minimum stem ages of 34 Ma for Stipeae and Chusquea (7) and set a minimum stem age of Dicantherium to 8 Ma (38). Using fossil phytolith data to date the Poaceae tree is black dots indicate C4 value. Background shading indicates gross climate delineations between a closed-canopy tropical forest (white) and open woodland/savanna system (gray); transition occurs around MAP ∼ 1500 mm year−1, coefficient of variation of precipitation ∼ 0.75. Gray bars highlight C3/C4 transitions that are consistent with a movement of the C4 lineage from tropical forest open into open woodland/savanna.

**Character Evolution Analyses.** We used the Analysis of Traits module in Phylocode (40) to identify 21 phylogenetically independent contrasts between C3 and C4 taxa. We also used LASRdisc (v. 1.0) (41) to reconstruct the evolution of photosynthetic shifts using maximum likelihood. Both approaches converged on the same focal nodes. Despite missing several known evolutionary transitions from our dataset (e.g., Sartidia, a C3 lineage in C4 Aristidoideae; Merxmuellera rangei, the C3 sister taxon of C4 Centropodia; and Neurachne munroi, which represents a recent C3 origin within the C4 Neurachne (3)) we have recovered more shifts in pathway than previously had been reported for grasses. Of the 21 nodes, 20 were reconstructed as a C3 origin rather than as a reversal to C3; the one exception is Homopholis proluta, which now stands as a putative reversal to C3 alongside Eragrostis walteri (42) and Allotropis semialata subsp. ecksonii (43) (neither sampled here). Three of our focal nodes result from the addition of taxa that have not been included in previous analyses: Homopholis proluta, Panicum amarum, and Panicum decompositum. These taxa were represented by archived sequence data from NCBI that is not published, and there is no way to confirm that the samples were correctly identified. Another potentially misleading node is the divergence between Steinchisma laxa (C3) and Steinchisma hians (a C3/C4 intermediate). In the 21-focal node analysis we coded S. hians as C3. To account for these uncertainties, we ran analyses on a reduced dataset, removing H. proluta, P. amarum, and P. decompositum and recoding S. hians as C3. This revision resulted in 17 rather than 21 C3/C4 divergences. We tested that the magnitude of change in climate variables at focal nodes was significantly different from zero by using a one-sample t test and assessed whether the overall directionality of change was significantly positive or negative against the binomial expectation, as recommended in (40).

To determine whether C3/C4 species have experienced divergent selection for temperature and precipitation variables, we tested the fit of multiple models of trait evolution using the noncensored approach in BROWNIE (v. 2.1) (44). The noncensored approach requires ancestral reconstructions for C3/C4 at internal nodes, so we used a procedure in BROWNIE that reconstructs the most likely state at each internal node before testing for model fit. We then reconstructed an OU model (45), allowing the C3 branches and the C3 branches to have optimal mean values for climate variables from which values could deviate according to the attraction parameter (OU2). To test for the significance of these results, we compared the Akaike Information Criterion scores corrected for small sample size (AICc), comparing the OU2 model described above with a model with a global mean value (OU1) and a Brownian motion model with a single rate of evolution and with no attraction parameter (BM). We made this comparison for each of the 100 dated trees drawn from the Bayesian posterior distribution and calculated the mean and standard deviation of each estimated parameter. A difference <2 between the AICc of the OU2 model and the OU1 model was taken as evidence for the OU1 model, whereas a difference >4 suggested considerable evidence for the OU2 model. In all cases, the difference between AICc of the OU2 model and the BM model was >200, suggesting that Brownian motion is a very poor model of trait evolution in this case.

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