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Lineage-based functional types: characterising functional diversity to enhance the representation of ecological behaviour in Land Surface Models

Permalink

https://escholarship.org/uc/item/1c12q7dp

Journal

New Phytologist, 228(1)

**ISSN** 

0028-646X

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Publication Date

2020-10-01

DOI

10.1111/nph.16773

Peer reviewed

- 1 **Type:** Viewpoint.
- 2 **Brief Heading:** Lineage Functional Types: Organizing functional diversity around evolutionary history

- 4 **Title:** Lineage Functional Types (LFTs): Characterizing functional diversity to enhance the representation
- 5 of ecological behavior in Land Surface Models

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- 31 Word count: Total: 2620
- 32 Figures: 1
- 33 Tables: 1
- 34 Supporting information: 1

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# **Summary (200/200):**

Process-based vegetation models attempt to represent the wide range of trait variation in biomes by grouping ecologically similar species into plant functional types (PFTs). This approach has been successful in representing many aspects of plant physiology and biophysics but struggles to capture biogeographic history and ecological dynamics that determine biome boundaries and plant distributions. Grass dominated ecosystems are broadly distributed across all vegetated continents and harbor large functional diversity, yet most Land Surface Models (LSMs) summarize grasses into two generic PFTs based primarily on differences between temperate C<sub>3</sub> grasses and (sub)tropical C<sub>4</sub> grasses. Incorporation of species-level trait variation is an active area of research to enhance the ecological realism of PFTs, which form the basis for vegetation processes and dynamics in LSMs. Using reported measurements, we developed grass functional trait values (physiological, structural, biochemical, anatomical, phenological, and disturbance-related) of dominant lineages to improve LSM representations. Our method is fundamentally different from previous efforts, as it uses phylogenetic relatedness to create lineage-based functional types (LFTs), situated between species-level trait data and PFT-level abstractions, thus providing a realistic representation of functional diversity and opening the door to the development of new vegetation models.

**Keywords:** C<sub>4</sub> photosynthesis, Earth system models, land surface models, evolution, grass biogeography, plant functional types, vegetation models

#### Main body:

#### Introduction

Functional trait variation within biomes arises from evolutionary histories that vary biogeographically, leading to plant taxa with differing ecological behavior and differences in ecosystem structure and function across continents (Lehmann *et al.*, 2014; Higgins *et al.*, 2016; Griffith *et al.* 2019). Land Surface Models (LSMs), fundamental components of Earth System Models, typically apply abstracted plant functional types (PFTs; but see Pavlick *et al.*, 2013; Scheiter *et al.*, 2013; Medlyn *et al.*, 2016) to represent physical, biological, and chemical processes crucial for soil and climate-related decision making and policy. However, PFTs must generalize across species, and inevitably encapsulate a wide range of plant strategies and vegetation dynamics, a demand that contrasts with efforts to investigate nuanced and species specific ecological behavior (Cramer *et al.*, 2001; Bonan, 2008; Sitch *et al.*, 2008; Kattge *et al.*, 2011). Furthermore, PFTs account for only a modest degree of variation in a wide array of functional traits, ranging from seed mass to leaf lifespan (LL), in the TRY database (Kattge *et al.*, 2011). For

example, standard PFTs may not generally capture key drought responses in tree species (Anderegg, 2015), although models with a hydraulics module can be specifically applied for this purpose (e.g., *ecosys*; Grant *et al.*, 1995). Oversimplification of the physiognomic characteristics of PFTs can have major unintended consequences when simulating ecosystem function (Griffith et al., 2017), such as highly biodiverse savanna ecosystems (Searchinger *et al.*, 2015). However, studies that explicitly incorporate species-level trait variation into vegetation models (e.g., Grant *et al.*, 1995; Sakschewski *et al.*, 2016; Lu *et al.*, 2017; Grant *el al.*, 2019; Mekonnen *et al.*, 2019) have demonstrated improvements in model performance. Selecting trait data from multi-variate trait distributions for model parameterization (Wang *et al.*, 2012; Pappas *et al.*, 2016) is very challenging for global modeling applications, particularly in hyper-diverse regions like the tropics, and may not be feasible for areas with biased or limited data. Until these data-gaps are filled, a finer-grained representation of the functional diversity among species might be achieved by reorganizing PFTs based on tradeoffs and evolutionary relatedness.

Importantly, in seeking approaches to restructure PFTs, numerous observations over the last decade have shown that both plant traits and biome occupancy are commonly phylogenetically conserved, with closely related species having similar traits and niches (e.g., Cavender-Bares et al., 2009, 2016; Crisp et al., 2009; Liu et al., 2012; Donoghue & Edwards, 2014; Coelho de Souza et al., 2016). The existence of strong evolutionary constraints on plant functioning and distribution suggests that, as an alternative, vegetation types should be organized in a manner consistent with phylogeny. Eco-evolutionary models have increased our mechanistic understanding of ecological patterns in fields ranging from community ecology (e.g., Webb et al., 2002; Cavender-Bares et al., 2009) to global biogeography (e.g., the Latitudinal Diversity Gradient; Pontarp et al., 2019; Visser et al. 2014). We advocate for explicit inclusion of evolutionary history and a consistent framework for integrating traits into global vegetation models. This approach brings a testable method for defining vegetation types, enables the functional traits of uncharacterized species to be inferred from relatives, and allows evolutionary history to be explicitly considered in studies of biome history. Here, we illustrate this approach for grasses and grass-dominated ecosystems, where we use our framework to aggregate species into Lineage-based Functional Types (LFTs) to capture the species-level trait diversity in a tractable manner for large-scale vegetation process models used in LSMs. Capturing the evolutionary history of woody plants is also critical to understanding variation in ecosystems function in savannas (Lehmann et al., 2014; Osborne et al., 2018), and in general we are advocating for the development of LFTs in other vegetation types and in other ecosystems. Grasses provide a tractable demonstration for the utility of LFTs; we also discuss the potential to significantly improve ecological and biogeographical representations of other plants in LSMs.

125 Grasses are one of the most ecologically successful plant types on earth (Linder et al., 2018) and provide 126 great opportunity for increasing understanding of plant functional diversity. Ecosystems containing or 127 dominated by grasses (i.e., temperate, tropical, and subtropical grasslands and savannas) account for 128 a>40% of global land area and productivity, and are a staple for humanity's sustenance (Tilman et al., 129 2002; Still et al., 2003; Asner et al., 2004; Gibson, 2009). The photosynthetic pathway composition (C<sub>3</sub> or 130 C<sub>4</sub>) of grass species is a fundamental aspect of grassland and savanna function, ecology, and 131 biogeography. Of the ~11,000 grass species on Earth, some ~4,500 use the C<sub>4</sub> photosynthetic pathway 132 (Osborne et al., 2014). Although they account for less than 2% of all vascular plant species (Kellogg, 133 2001), C<sub>4</sub> grasses are estimated to account for 20-25% of terrestrial productivity (Still et al., 2003), 134 having risen to such prominence only in the last 8 million years (Edwards et al., 2010). Dominance by 135 C<sub>4</sub> versus C<sub>3</sub> grasses has major influences on gross primary productivity and ecosystem structure and 136 function (Still et al., 2003) and strongly influences interannual variability of the global carbon cycle, due 137 to a combination of ecological and climatic factors (Poulter et al., 2014; Griffith et al., 2015). Dynamic 138 vegetation models largely fail to reproduce spatial patterns of grass cover —both past and present—and 139 productivity at regional to continental scales, limiting ability to predict future plant community changes 140 (Fox et al., 2018; Still et al., 2018). As a consequence, LSMs require significant improvement to 141 adequately represent vegetation responses to increasing CO<sub>2</sub> (Smith et al. 2016; De Kauwe et al., 2016). 142 Many models also miss key transitions between biome states (e.g., Still et al., 2018) that exist as a result 143 of disturbance or biogeographic history (e.g., Staver et al., 2011; Dexter et al., 2018). 144 145 Most LSMs classify grasses into two PFTs based on differences between temperate C<sub>3</sub> grasses and sub-146 tropical and tropical C<sub>4</sub> grasses. However, grass ecological adaptations and physiological properties are 147 highly diverse, ranging from cold-specialized to fire- and herbivore-dependent species. While grasses are 148 often equated functionally, in reality they exhibit a high degree of variation in hydraulic, leaf economic, 149 and phenological traits (Taylor et al., 2010; Liu et al., 2012) that likely explains their broad geographic 150 dominance in different regions (Edwards et al., 2010; Visser et al., 2014). These differences include 151 economically important forest-forming grasses such as bamboos, although here we focus on globally 152 dominant herbaceous lineages. Grasses exhibit strong phylogenetic diversity in leaf economics variation 153 and associations with disturbance (Taylor et al., 2010; Liu et al., 2012; Simpson et al., 2016). 154 Disturbances such as fire and herbivory have large impacts on ecosystem function and distributions, and 155 PFT based approaches are unlikely to capture these differences among lineages. At broad phylogenetic 156 and spatial scales, niche and biome conservatism of major plant lineages is common (Crisp et al., 2009; 157 Cornwell et al., 2014; Donoghue & Edwards, 2014), and we therefore argue that evolution and 158 biogeography provide a framework for aggregating species (across ecosystems and strata) into LFTs that

capture species-level trait diversity in a way that can be feasibly incorporated for use in global vegetation models, and that will improve PFT-based modeling approaches. Focusing on grasses, we developed this approach by collecting grass traits from databases (e.g., Osborne *et al.*, 2011) and literature (e.g., Atkinson *et al.*, 2016; Supplemental Methods S1), for five key categories (physiology, structure, biochemistry, phenology, and disturbance). We summarize these species traits at the lineage level and relate these functional types to their observed global distributions.

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## Methods for establishing lineage-based functional types (LFTs) for grasses

167 There are 26 monophyletic C<sub>4</sub> lineages described in the Poaceae family, yet only two (the Andropogoneae 168 and Chloridoideae) account for most of the areal abundance of C<sub>4</sub> grasses globally (Lehmann et al., 2019 169 ; Fig 1.) (Edwards & Still, 2008; Edwards et al., 2010; Grass Phylogeny Working Group II, 2012). 170 Among C<sub>3</sub> grasses, only the Pooideae are globally dominant today. The Pooideae occupy cooler climates 171 than the C<sub>4</sub> Andropogoneae and Chloridoideae, which dominate in warm and wetter and drier climates, 172 respectively. Therefore, we focused on collecting species-level trait data from the literature and from 173 databases for grass species from these three lineages. The term 'trait' is defined differently across 174 research disciplines (Violle et al., 2007). Our aims necessitate a collection of broad trait space beyond 175 that typically used for the leaf economic spectrum to include morphological and physiological 176 determinants of plant hydraulics, physicochemical controls of photosynthesis, allocation to reproduction, 177 and spectral reflectance. Many traits are highly correlated, reflecting plant functional strategies. Further, a 178 single trait can relate to multiple forms of plant fitness. Here, traits were assigned to groups (Table 1) 179 based on their use in models and how they might be used in future applications (e.g., hyperspectral remote 180 sensing of LFTs, or modeling of fire). We present median and variation in trait values among species for 181 three major grass lineages (LFTs) as per Figure 1, and compare these with commonly used values for C<sub>3</sub> 182 and C<sub>4</sub> PFTs (Table 1).

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# LFTs for grasses differ drastically in key functional traits

Our LFTs demonstrate both the importance of considering lineage to explain ecological patterning, and the need for modification of current LSM PFT approaches. For instance, C<sub>4</sub> plants typically have lower RuBisCO activity (V<sub>cmax</sub>) but higher electron transport capacity (J<sub>max</sub>) than C<sub>3</sub> plants, reflecting both the additional energetic cost of C<sub>4</sub> physiology and the greater efficiency of RuBisCO in higher CO<sub>2</sub> environments (Collatz *et al.*, 1998). The Chloridoideae (C<sub>4</sub>) grasses have intermediate Vcmax and J<sub>max</sub> compared to the Andropogoneae (C<sub>4</sub>) and the Pooideae (C<sub>3</sub>) (Table 1). Furthermore, the Pooideae have evolved to tolerate much colder conditions (reflected in Trange; Sandve & Fjellheim, 2010; Vigeland *et al.*, 2013; McKeown *et al.*, 2016), and our results suggest that C<sub>4</sub> lineages may differ in their thermal

193 tolerances (Watcharamongkol et al., 2018). These differences suggest that macroecological synthesis 194 studies with global implications (e.g., Walker et al., 2014; Heskel et al., 2016) should, at minimum, 195 include more grass species in their datasets, ideally organized as LFTs. 196 197 Trade-offs among adaptations and tolerances in natural systems promote coexistence among plant species 198 (Tilman, 1988; Tilman & Pacala, 1993; Kneitel & Chase, 2004). Specific leaf area (SLA) measures the 199 cost of constructing a leaf, which represents a tradeoff between acquisitive (high relative growth rate) and 200 conservative (high leaf lifespan) plant strategies (Westoby, 1998; Westoby et al., 2002; Wright et al., 201 2004). Model simulations of growth are highly dependent on the value of SLA used (Korner, 1991; Sitch 202 et al., 2003; Bonan, 2008). However, in most of these LSMs, C<sub>3</sub> grass PFTs have higher or similar SLA 203 values as C<sub>4</sub> PFTs likely biasing predictions. In contrast, we found that the C<sub>4</sub> LFTs had higher SLA than 204 the C<sub>3</sub>LFT, but SLA did not differ between the two dominant C<sub>4</sub> grass lineages (Atkinson et al. 2016). 205 SLA can be highly variable within lineages in grasses, likely due to the importance of herbivore pressure 206 as a competing demand on leaf economics (Anderson et al., 2011; Griffith et al., 2017) as well as 207 intraspecific variation. As a result, SLA highlights that some traits are harder to generalize than others 208 using the LFT approach and suggests that a range of values may be appropriate than a single value for 209 constraining LFT parameters. The phylogenetic signal among grass lineages is stronger for stature (Taylor 210 et al., 2010; Liu et al., 2012), with the Andropogoneae being considerably taller on average than the 211 Chloridoideae. This difference suggests that not all traits are oriented along a fast-slow axis at broad 212 taxonomic scales across C<sub>3</sub> and C<sub>4</sub> grass lineages (Reich, 2014; Díaz et al., 2016; Archibald et al. 2019). 213 Furthermore, the C<sub>3</sub>- and eudicot-centric approach in the current leaf economics framework suggests that 214 a higher SLA should also correlate with a higher specific leaf nitrogen content, yet the evolution of C<sub>4</sub> 215 photosynthesis allows for a significant reduction in RuBisCO content, and hence plant nitrogen 216 requirements (Taylor et al., 2010). Thus, grass lineages differ in numerous leaf traits which have 217 consequences that extend from palatability and flammability to hydrological differences. 218 219 Physiological and morphological leaf vascular traits underlie variation in SLA, constrain the hydrology of 220 plants (e.g., Blonder et al., 2014; Sack et al., 2014), and are key traits related to the evolution of C<sub>4</sub> 221 photosynthesis (Sage, 2004; Ueno, 2006). We describe next key hydraulic differences between the two 222 dominant C<sub>4</sub> lineages, which correspond to the C<sub>4</sub> biochemical subtypes (Ueno, 2006; Liu & Osborne, 223 2015). The Chloridoideae have low conductance and high embolism resistance hydraulic traits (Table 1), 224 and tend to inhabit drier sites (Fig. 1). Some Andropogoneae have been described as "water spenders"

(Williams et al., 1998), and their hydraulic traits help to explain their affinity with higher rainfall habitats

where they rapidly expend available soil water (Taub, 2000) and promote fire after curing. These

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hydraulic differences should have large effects in models, especially those that consider tree-grass coexistence (Higgins *et al.*, 2000) and explicit representation of plant hydraulics (Grant *et al.*, 1995; Mekonnen *et al.*, 2019).

Lineages also differ in biogeochemical traits that influence nutrient turnover rates and the reflectance and absorbance properties of vegetation. For example, Andropogoneae have higher C:N than Chlordoideae grasses, likely a result of growth rate differences and the frequent association of Andropogoneae grasses with fire. Similarly, a greater proportion of N in Chloridoideae leaves is allocated to RuBisCO, which is related to Vcmax (Ghannoum et al. 2012). Finally, C<sub>3</sub> and C<sub>4</sub> grasses are distinguishable spectrally at the leaf, canopy, and landscape level based on differences between the functional types in chlorophyll a/b ratio, canopy structure, and seasonality (Foody & Dash, 2007; Siebke & Ball, 2009; Irisarri *et al.*, 2009). C<sub>3</sub> and C<sub>4</sub> grasses are typically given many of the same optical properties in vegetation models, but we show here that Chloridoideae might have considerably higher near infra-red (NIR) reflectance than other lineages, possibly producing interesting optical variation and affecting the surface energy balance and albedo (Ustin & Gamon, 2010)(Table 1). Foliar spectral traits are also correlated with morphological and chemical traits related to nutrient cycling and plant physiology (Dahlin *et al.*, 2013; Serbin *et al.*, 2014).

Grass lineages also show key differences in reproductive traits and the timing of related biological events (e.g., leaf-out times) that should be captured in models, especially those that include demographic predictions (Davis et al., 2010). Chloridoideae grasses have seeds with lower mass than other lineages (Liu et al., 2012; Bergmann et al., 2017), and this may represent a life-history trade-off with higher seed production and other 'fast' growth strategies (Adler et al., 2014). Wind versus animal dispersal strategies might also affect diaspore size in a way not directly related to disturbance (e.g., Westoby 1998; Bergmann et al., 2017), whereas some reproductive traits may also indicate fire and disturbance-related adaptations. Phenological traits, such as flowering and leaf-out times and their cues (which can include disturbance factors) exhibit conservatism across many plant lineages (Davies et al., 2013). Fire and herbivory are two globally important and contrasting disturbances for grass-dominated vegetation (Archibald & Hempson, 2016; Archibald et al., 2019) and adaptations to both can be characterized by phenological and reproductive traits in addition to physiological and leaf traits. It is less clear how herbivory effects can be captured in such models, given that many herbivore-related traits vary greatly in grasses (Anderson et al., 2011). Many fire-related traits show patterns of phylogenetic conservatism, with high flammability clustering into particular lineages such as the Andropogoneae (Simpson et al., 2016). Large-scale vegetation models that have simulated grass fires in Africa have attributed faster curing (becoming dry

fuel) rates to C<sub>4</sub> vegetation (Scheiter *et al.*, 2012), and this behavior appears to be due largely to dominant Andropogoneae grasses.

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We have identified large differences among LFTs, across six trait categories, that are not captured by the standard PFT approach. Many of these trait data have very low sample sizes (from 1 to 1365) and come from non-overlapping species, highlighting the need for systematic data collection for grasses. Such a data collection effort would be an excellent opportunity to test for coordination among trait axes in a phylogenetic context, which has rarely been done in other systems despite the likelihood that relatedness drives patterns of trait covariation (e.g., Salguero-Gómez *et al.*, 2016; Griffith *et al.*, 2016). Furthermore, intra-group (whether PFTs or LFTs) trait variation deserves to be properly estimated (only some traits in Table 1 have enough data to estimate variability) as convergence and adaptation produce meaningful trait variation that should be incorporated into models.

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### Potential for lineage-based functional types in other vegetation types

Many current PFTs implicitly represent groupings of closely related lineages (e.g., pinaceous conifers, grasses). However, even in these cases biogeographic distributions, and the coarseness of the phylogenetic unit, generates a lack of useful resolution. Currently, there are efforts to incorporate specieslevel trait data and methods such as those proposed by Cornwell et al., (2014) could be employed to cluster species into prominent lineage-based groupings representing unique trait combinations. Phylogenies are hierarchical by nature and allow the LFT approach to be scalable and adjustable to the research question being addressed. While many technical challenges still remain, the ability to remotely sense plant lineages adds potential for rapidly developing LFTs from spectral data (e.g., Cavender-Bares et al., 2016). LFTs would be valuable for a wide range of systems. For example, trees in Eurasian boreal forests suppress canopy fires through the structure of their canopies, whereas North American boreal trees enable greater intensity canopy fires (Rogers et al., 2015). These distinctions lead to major differences in CO<sub>2</sub> emissions and function (Rogers et al., 2015) that might be captured in an LFT framework. The boreal tree example is challenging because these communities are comprised of closely related species that are ecologically different, potentially requiring species level parameterization or being better represented by fire-based PFTs. Secondly, LFTs for savanna tree communities could better represent differing climatic responses that are driven by unique evolutionary and biogeographic histories (Lehmann et al., 2014; Osborne et al., 2018). Finally, tropical ecosystems such as the dipterocarp forests in Southeast Asia would be well suited to LFTs which might better represent carbon storage (Brearley et al., 2016).

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Potential challenges with a lineage-based functional approach include the fact that many plant traits do not show strong phylogenetic conservatism (Cadotte *et al.*, 2017), with several being labile. There are likely spatial and phylogenetic scales at which the LFT approach will be most appropriate; for example, at large scales (regional to continental), lineage conservatism is common (Crisp *et al.*, 2009). In contrast, at the scale of local communities, we might expect character displacement and limiting similarity (processes that lead to reduced trait similarity of coexisting species) could obscure phylogenetic patterns and limit the utility of LFTs as proposed here (Webb *et al.*, 2002; Cavender-Bares *et al.*, 2009; HilleRisLambers *et al.*, 2012). However, in grassy ecosystems, there is evidence that the patterns of spatial ecological sorting of lineages would be captured with LFTs also at landscape scales (e.g., within Serengeti National Park, Anderson *et al.*, 2011; Forrestel *et al.*, 2017). Finally, we focus on extant lineages that are functionally important today, but their past interactions with other clades may have shaped the biomes they inhabit (Edwards *et al.*, 2010).

#### Conclusions

We conclude that an LFT perspective captures important variation in functional diversity for grasses (Table 1). Our analysis of current knowledge of grass functional diversity (in terms of physiology, structure, biochemistry, phenology, and disturbance), distributions, and phylogeny indicates that to represent grass ecological behavior, division of today's ecologically dominant grasses into at least two C<sub>4</sub> and at least one C<sub>3</sub>LFT could potentially improve representation in LSMs. These proposed LFTs capture key evolutionary differences in physiological, structural, biogeochemical, anatomical, phenological, and disturbance-related traits. We also highlight the need for systematic trait data collection for grasses, which we show are vastly underrepresented in trait databases, despite their ecological and economic importance. More broadly, we outline the LFT framework which is highly flexible and has the potential for use in a wide range of applications. Here, we speak to incorporating LFTs as groupings in vegetation models, but we also suggest that trait-based models might capture important biogeographic variation (e.g., due to historical contingency) through the inclusion of phylogenetic conservatism. We advocate for the use of phylogeny as a way to help guide and constrain the inclusion of burgeoning plant trait data to expand the range of functional types considered by global vegetation models.

### Acknowledgements

This work was in part inspired by a workshop at the National Evolutionary Synthesis Center. DMG acknowledges support from NASA under the auspices of the Surface Biology and Geology (SBG) Study and from USGS through the National Innovation Center. CJS, SP, BRH, SB, MU, FQ, JBN were supported by National Science Foundation award 1342703, 1926431, and 1856587; CAES and TJG by

- 328 NSF awards 1253713, 1342787, and 1120750. CERL was supported by an award from The Royal
- 329 Society. WJR was supported as part of the RUBISCO Scientific Focus Area in the Regional and Global
- 330 Climate Modeling Program of the U.S. Department of Energy, Office of Science, Office of Biological and
- 331 Environmental Research under contract DE-AC2–5CH11231.

- **Author contributions**
- DMG, CJS, and CPO planned and designed the work. All authors contributed data and writing to the
- 335 manuscript.

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624 **Methods S1** - Description of data and code for Table 1.

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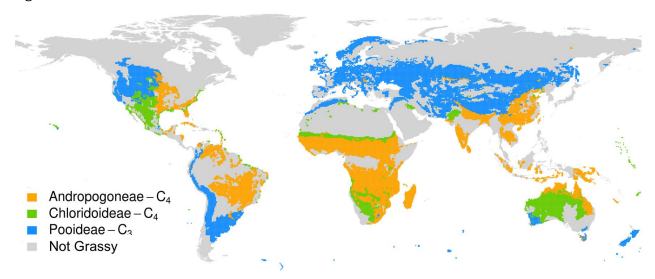
		PFT		LFT*			
Category	Parameter	$C_4$	C <sub>3</sub>	Source	Andropogoneae	Chloridoideae	Pooideae
Physiological	Vcmax ( $\mu mol \ m^{-2} \ s^{-1}$ )	39	90	1, 2	38	45.6 (4.4)	63.6 (28)
	Jmax ( $\mu mol \ m^{-2} \ s^{-1}$ )	400	100	3	180	108.1 (43)	128.8 (45)
	Rd ( $\mu mol \ m^{-2} \ s^{-1}$ )	8.0	1.1	1, 2	0.9a (0.2)	2a (1.4)	0.9a (0.7)
	Phi ( $\mu mol \; \mu mol^{-1}$ )	0.06	0.085	4, 5	0.06	0.06	0.09
	Trange (°C)	< 15.5	> 15.5	6	> 5**	> - 5**	> -30 <sub>5</sub> and <
Structural	$SLA\;(m^2kg^{-1})$	16	33	7	33b (11)	29b (14)	25a (12)
	LDMC $(gg^{-1})$	-	-		-	-	-
	$SRL (m g^{-1})$	-	-		-	-	-
	Culm Height (cm)	-	-		150c (150)	80b (50)	60a (60)
	$R : S \; (g  g^{-1})$	2	2	7	0.4b (0.07)	0.3a (0.07)	0.4b (0.1)
Biogeochemical/	$C:N(gg^{-1})$	10	17	7	66.1b (14.7)	39.9a (22.2)	55.7ab (10)
spectral	Nrubisco (proportion)	0.09	0.137	7	0.05 (0.01)	0.08 (0.03)	0.2
	Reflectance (300-2500 nm)	-	-		-	-	-
	e.g., rNIR (reflectance)	0.35	0.35	7	0.4	0.5	0.3
Anatomical/	IVD (µm)	-	-		85.7a (25.2)	136.8b (40)	242.1c (58)
Hydraulic	Xylem dia. ( $\mu m$ )	-	-	7	21.4b (12.2)	16.8a (10.7)	19.3a (6.7)
	K <i>leaf</i>	-	-		-	-	-
	Vein Hierarchy Primary vein secondary vein		-		0.11a (0.09)	0.29b (0.2)	0.58c (0.6)
	Leaf Width:Length	-	-		0.04b (0.05)	0.03a (0.04)	0.03a (0.02)
Life History	LL (months)	1.68	12	7	2 (0.4)	1.1	2 (1.8)
	1000-seed mass $(g)$	-	-	7	1.4b (2.4)	0.2a (0.4)	1.4b (3.8)
	Life History (% annual)	-	=	7	0.25	0.28	0.15
Disturbance	Curing rate (%)	80	20	8	80	50**	20
	Bud Bank	-	-		-	=	-
	Flammability $(gs^{-1})$	-	-		-	-	-

1 - Farquhar *et al.* (1980), 2 - Collatz *et al.* (1992), 3 - von Caemmerer (2000), 4 - Ehleringer *et al.* (1997), 5 - Collatz *et al.* (1998), 6 - Sitch *et al.* (2003), 7 - Oleson *et al.* (2013), 8 - Scheiter *et al.* (2012); Curing rate is the % cured 30 d after the end of the growing season as

described in Scheiter et al. (2012); \*Published citations for LFT values can be found in Methods S1.

Anatomical data come from Gallaher, T.J. et al. unpublished . \*\*Estimated value. Abbreviations: Vcmax (maximum carboxylation rate), Jmax (light saturated rate of electron transport), Rd (dark Respiration), Phi (quantum efficiency), SLA (Specific Leaf Area), LDMC (Leaf Dry Matter Content), SRL (Specific Root Length), R:S (root to shoot ratio), C:N (Carbon to Nitrogen ratio), IVD (InterVeinal Distance), Kleaf (leaf hydraulic conductance), LL (Leaf Lifespan).

# 649 Figures:



**Figure 1.** Distributions of the three globally dominant grass lineages in the herbaceous layer. These data come from Lehmann et al (2019) and show where each lineage is more abundant than the other two lineages on a 0.5-degree grid.