

This is a repository copy of *Functional diversification enabled grassy biomes to fill global climate space*.

White Rose Research Online URL for this paper: https://eprints.whiterose.ac.uk/184960/

Version: Submitted Version

# Article:

Lehmann, C.E.R., Griffith, D.M., Simpson, K.J. orcid.org/0000-0001-6673-227X et al. (22 more authors) (Submitted: 2019) Functional diversification enabled grassy biomes to fill global climate space. bioRxiv. (Submitted)

https://doi.org/10.1101/583625

### Reuse

This article is distributed under the terms of the Creative Commons Attribution (CC BY) licence. This licence allows you to distribute, remix, tweak, and build upon the work, even commercially, as long as you credit the authors for the original work. More information and the full terms of the licence here: https://creativecommons.org/licenses/

### Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



eprints@whiterose.ac.uk https://eprints.whiterose.ac.uk/

# Functional diversification enabled grassy biomes to fill global climate space

Caroline E. R. Lehmann<sup>1,2</sup>\*, Daniel M. Griffith<sup>3</sup>, Kimberley J. Simpson<sup>4</sup>, T. Michael Anderson<sup>5</sup>, Sally Archibald<sup>2</sup>, David J. Beerling<sup>4</sup>, William J. Bond<sup>6</sup>, Elsie Denton<sup>7</sup>, Erika J. Edwards<sup>8</sup>, Elisabeth J. Forrestel<sup>9</sup>, David L. Fox<sup>10</sup>, Damien Georges<sup>11</sup>, William A. Hoffmann<sup>12</sup>, Thomas Kluyver<sup>13</sup>, Ladislav Mucina<sup>14,15</sup>, Stephanie Pau<sup>16</sup>, Jayashree Ratnam<sup>17</sup>, Nicolas Salamin<sup>18</sup>, Bianca Santini<sup>4</sup>, Melinda D. Smith<sup>7</sup>, Elizabeth L. Spriggs<sup>8</sup>, Rebecca Westley<sup>4</sup>, Christopher J. Still<sup>3</sup>, Caroline A.E. Strömberg<sup>19</sup> and Colin P. Osborne<sup>4\*</sup>

<sup>1</sup>School of GeoSciences, University of Edinburgh, Edinburgh EH9 3FF, UK

<sup>2</sup>Centre for African Ecology, School of Animal and Plant Sciences, University of Witwatersrand, Johannesburg

<sup>3</sup> Dept. Forest Ecosystems & Society, Oregon State University, Corvallis, OR 97331, USA

<sup>4</sup> Dept. Animal & Plant Sciences, University of Sheffield, Sheffield S10 2TN, UK

<sup>5</sup> Dept. Biology, Wake Forest University, Winston-Salem, NC 27109, USA

<sup>6</sup> Dept. Botany, University of Cape Town, Cape Town, ZA-7701, South Africa

<sup>7</sup> Dept. Biology, Colorado State University, Ft Collins, CO, 80523, USA

<sup>8</sup> Dept. Ecology & Evolutionary Biology, Yale University, 165 Prospect St., New Haven, CT 06520, USA

<sup>9</sup> Dept. Viticulture & Enology, University of California, Davis, CA 95616, USA

<sup>10</sup> Dept. Earth Sciences, University of Minnesota, Minneapolis MN 55455, USA

<sup>11</sup> Laboratoire d'Ecologie Alpine (LECA), Univ. Grenoble Alpes, CNRS, F-38000 Grenoble, France

<sup>12</sup>Dept. Plant Biology, North Carolina State University, Raleigh, NC 27695, USA

<sup>13</sup>Computational Modelling Group, Faculty of Engineering & the Environment, University of Southampton, Southampton SO17 1BJ, UK

<sup>14</sup>Harry Butler Institute, Murdoch University, 90 South Street, Murdoch WA 6150, Perth, Australia

<sup>15</sup>Dept. Geography & Environmental Sciences, Stellenbosch University, Private Bag X1, Matieland 7602, Stellenbosch, South Africa

<sup>16</sup>Dept. Geography, Florida State University, Tallahassee, FL 32306-2190, USA

<sup>17</sup>National Centre for Biological Sciences, Tata Institute of Fundamental Research, Gandhi

Krishi Vignana Kendra, Bellary Road, Bangalore 560 065, India.

<sup>18</sup>Dept. Ecology & Evolution, University of Lausanne, CH-1015 Lausanne, Switzerland

<sup>19</sup>Dept. Biology & the Burke Museum of Natural History & Culture, University of Washington,

Seattle, WA 98195, USA

\* Corresponding Authors

Abstract: Global change impacts on the Earth System are typically evaluated using biome classifications based on trees and forests. However, during the Cenozoic, many terrestrial biomes were transformed through the displacement of trees and shrubs by grasses. While grasses comprise 3% of vascular plant species, they are responsible for more than 25% of terrestrial photosynthesis. Critically, grass dominance alters ecosystem dynamics and function by introducing new ecological processes, especially surface fires and grazing. However, the large grassy component of many global biomes is often neglected in their descriptions, thereby ignoring these important ecosystem processes. Furthermore, the functional diversity of grasses in vegetation models is usually reduced to  $C_3$  and  $C_4$ photosynthetic plant functional types, omitting other relevant traits. Here, we compile available data to determine the global distribution of grassy vegetation and key traits related to grass dominance. Grassy biomes (where > 50% of the ground layer is covered by grasses) occupy almost every part of Earth's vegetated climate space, characterising over 40% of the land surface. Major evolutionary lineages of grasses have specialised in different environments, but species from only three grass lineages occupy 88% of the land area of grassy vegetation, segregating along gradients of temperature, rainfall and fire. The environment occupied by each lineage is associated with unique plant trait combinations, including  $C_3$  and  $C_4$  photosynthesis, maximum plant height, and adaptations to fire and aridity. There is no single global climatic limit where C<sub>4</sub> grasses replace C<sub>3</sub> grasses. Instead this ecological transition varies biogeographically, with continental disjunctions arising through contrasting evolutionary histories.

**Significance statement**: Worldviews of vegetation generally focus on trees and forests but grasses characterize the ground layer over 40% of the Earth's vegetated land surface. This omission is important because grasses transform surface-atmosphere exchanges, biodiversity and disturbance regimes. We looked beneath the trees to produce the first global map of grass-dominated biomes. Grassy biomes occur in virtually every climate on Earth. However, three lineages of grasses are much more successful than others, characterizing 88% of the land area of grassy biomes. Each of these grass lineages evolved ecological specializations related to aridity, freezing and fire. Recognizing the extent and causes of grass dominance beneath trees is important because grassy vegetation plays vital roles in the dynamics of our biosphere and human wellbeing.

# 1 Introduction

The global distribution of terrestrial biomes determines global patterns of carbon storage and 2 3 biodiversity (1). Delineation of biome distributions is crucial because it underpins evaluations of vegetation feedbacks on climate (2), extinction threats for biodiversity (3), and strategies 4 5 for monitoring and reversing land-use change and degradation (4). Global studies of biome 6 distributions typically focus on forests and trees (4-6), following the long-established 7 paradigm in modern ecology of deterministic relationships between forest distributions and 8 environment (7). Within this paradigm, there is a widely held perception that grassy 9 vegetation only occupies semi-arid climates. However, it is increasingly recognized that biome 10 limits are not deterministically linked to climate but arise from multi-directional feedbacks 11 between plant functional traits, environment, and disturbance. These processes operate over 12 evolutionary and ecological timescales (8) creating biogeographic contingencies in biomeenvironment relationships (9). 13

Grassy biomes require open-canopied tree layers (or no tree layer) to permit enough 14 15 light to penetrate for grass photosynthesis. As a result, grasses dominate the ground layer 16 when the rate of woody plant recruitment and growth is limited by climate, soil, drainage, 17 disturbance conditions or light competition (10-12). "Grassy biomes" defined in this way 18 include tropical savannas, montane grasslands, grassy deserts, temperate steppe grasslands, 19 boreal parklands, and many temperate woodlands. The distinction of whether the ground 20 layer is dominated by grasses (Poaceae) is fundamental to understanding global relationships 21 among plants, climate, and disturbance (13). While, both trees and grasses are clearly 22 important in driving vegetation dynamics, grass dominance causes a fundamental shift in 23 disturbance regimes, whereby the consumption of ground layer biomass by fire and grazing 24 reinforces grass dominance and maintains open tree canopies (10). Grass cover and biomass

in the ground layer also affects surface energy, carbon, nutrient and water cycling by, for
example, altering rates of decomposition, water infiltration and absorption of sunlight. Grass
dominance therefore leads to novel ecological processes and properties in the Earth System,
including frequent fire and grazing by mammals (14).

29 During the Cenozoic grasses displaced forests and shrublands by altering disturbance 30 regimes at large scales across tropical and temperate regions (14, 15). The global expansion 31 of grassy vegetation enabled major faunal and floral radiations (14, 16), and is linked to events 32 in human behavioral evolution (17, 18). Today, natural grassy biomes provide grazing lands, 33 water resources and numerous ecosystem services that directly support over a billion people (19). Yet, despite this social and economic significance, and the profound disturbance 34 35 feedbacks engendered by grassy vegetation (20), understanding of grassy biomes is 36 geographically biased towards few regions (e.g., South and East African savannas, North American grasslands), with the global limits of grassy biomes poorly defined. 37

When considering the limits to grassy biomes, the grass diversity present in a system 38 39 is generally reduced to a distinction between species using the  $C_3$  or  $C_4$  photosynthetic 40 pathways. If all else is equal, C<sub>4</sub> grasses should outcompete C<sub>3</sub> grasses under conditions of high 41 light and temperature as well as low CO<sub>2</sub> (21-23). This physiologically based model explains, 42 in general terms, how C<sub>4</sub> grasses dominate tropical regions and C<sub>3</sub> grasses dominate 43 temperate and high-altitude environments under current atmospheric CO<sub>2</sub> levels [ppm  $\approx$  408]. 44 The physiological mechanisms underpinning this model have critical impacts for predicting 45 vegetation trajectories with global climate and atmospheric  $CO_2$  changes (22) yet attempts to 46 parse the consequences of grass physiology for global vegetation is often reliant on sparsely 47 validated modelling (24). Further, a focus on photosynthetic type belies the rich phylogenetic 48 diversity within grasses independent of photosynthetic pathway (14). Grasses are unusual

among vascular plants because C<sub>4</sub> photosynthesis evolved in up to 24 independent lineages
(25), conferring unique ecological characters to each C<sub>4</sub> lineage inherited from its C<sub>3</sub> ancestors
(26). Photosynthetic type therefore interacts with different combinations of other functional
traits to determine plant performance under varied environmental conditions (27, 28), but
the influence of these interactions on the global biogeography of grassy biomes is unknown.

Here, we focus on grass-dominated systems to address three questions. First, what are the global limits of grassy biomes? Second, to what extent is grassy biome structure contingent on evolutionary history, whereby independent phylogenetic lineages characterize grassy biomes on each continent? Finally, how do functional traits of the descendant species of each lineage relate to climate and fire? Our findings have significant implications for the representation of terrestrial vegetation processes in Earth System Models.

60

Identifying grassy biomes. Our dataset provides the first spatially explicit, functional 61 62 classification of grassy vegetation at the global scale (Figs. 1 and S1). Necessarily, our approach 63 that focusses on the ground layer contrasts with efforts to map biomes using remotely sensed 64 tree cover or biomass (4, 29). Such studies generally misclassify extensive areas of tropical 65 savanna as forest or degraded forest (30, 31). Global synthesis of grassy biomes has been 66 prohibited as satellite remote sensing does not see through a tree canopy. Therefore, we 67 mapped grassy formations by integrating and re-classifying 20 existing national and regional 68 vegetation maps produced using botanical data and detailed vegetation descriptions (see 69 Methods and SI).

What is a grassy biome? We defined vegetation units as grassy where the ground layer is characterized by Poaceae and where grasses comprised > 50% of ground layer cover based on descriptions within vegetation maps and associated literature (see Methods and SI). A

relatively small set of species often accounts for the majority of biomass in plant communities,
whether these are communities dominated by trees or grasses, and these species exert major
controls over ecosystem processes (32) and ecosystem services (33). Focusing on dominant
and characteristic species provides one way to explore links between evolutionary history and
ecosystem ecology at large scales (14). Through this process we identified 1,154 grass species
(~10% of the total grass flora) characterizing grassy vegetation.

79

## 80 Results and Discussion

Global limits of grassy biomes. Grasses can dominate ground layer vegetation in all but the 81 82 coldest and driest climates on Earth (Figs. 1 - 2). We estimate that vegetation with a grass-83 dominated ground layer originally covered ~ 41% of the vegetated land surface, although 84 much is now under cultivation. Critically, grasses can dominate the ground layer in every climate where woody vegetation can persist (Figs. 1 - 2). While steppe grasslands and prairies 85 86 occupy a large fraction of the global land area in dry temperate climates (Figs. 2, S2-S3), and 87 savannas and grasslands occupy most of the tropics, grass-dominated ground layers occupy 88 extensive areas in any other part of the vegetated global climate space (Fig. 2 and Fig. S4).

Members of 16 independently derived C<sub>4</sub> grass lineages dominate within at least one vegetation unit worldwide (Fig. 1). However, two C<sub>4</sub> lineages and one C<sub>3</sub> lineage dominate over 88% of the land area of grassy vegetation: C<sub>4</sub> Andropogoneae, 37% (1189 species in the lineage); C<sub>4</sub> Chloridoideae, 14% (1601 species in the lineage); and C<sub>3</sub> BEP, 38% (Fig. 1). The vast majority of C<sub>3</sub> BEP taxa belong to Pooideae (4234 species in the lineage). In contrast, C<sub>3</sub> species of the PACMAD clade dominate only 2% of grassy biomes (Fig. 1); these are the closest relatives of C<sub>4</sub> grasses and are restricted to warm, wet areas (Figs. S2-S4). Of the remaining

area of grassy vegetation, 6.6% is characterised by a mix of lineages, and the rest dominated
by 13 other, independently derived, C<sub>4</sub> lineages (Fig. 1 and Table S1).

98 The three dominant lineages sort in climate space. C<sub>3</sub> Pooideae dominate cooler, drier climates, whereas C<sub>4</sub> Andropogoneae and Chloridoideae dominate grassy biomes in warmer 99 100 climates (Figs. 3, S3-S6). However, precipitation sorts the  $C_4$  lineages, with peak dominance of 101 Andropogoneae occurring at ~ 1200mm MAP (Figs. 4, S3-S5), coinciding almost precisely with 102 the global peak in fire frequency (Fig. 4). This is also the climate space where disturbance-103 driven feedbacks are considered to play a major role in maintaining open (i.e., grassy) or 104 closed (i.e., woody) vegetation (34). In contrast, the peak dominance of Chloridoideae occurs 105 at ~350mm MAP (Figs. 4, S3-S4), within semi-arid climate zones occupied by both dry savannas 106 and shrublands/thickets e.g., (35). Temperature seasonality also differs among the C<sub>4</sub> lineages, 107 with Chloridoideae dominating in regions with strong seasonality, and Andropogoneae 108 dominating in more aseasonal environments (Fig. S5).

109

110 **Continental disjunctions in C<sub>3</sub> and C<sub>4</sub> lineage distributions.** Globally, the mean growing 111 season temperature where dominance of grasses transitions from  $C_3$  to  $C_4$  types varies starkly 112 among continents, from 8.5-26.1 °C, with a global mean of 17.2 °C (Fig. 3). Lineages using  $C_3$ 113 and C<sub>4</sub> photosynthetic pathways are clearly sorted by growing season temperature and mean 114 annual temperature (Figs. 3 and S3-6). The C<sub>3</sub> Pooideae lineage has specialized and radiated 115 in cold environments by evolving physiological cold acclimation to protect tissues from 116 freezing damage, and vernalization to synchronize flowering with the growing season (36, 37). 117 Conversely, in tropical regions, the repeated evolution of C<sub>4</sub> photosynthesis appears vital in 118 expanding the range of grassy biomes, by enabling colonization of hot, high light, and 119 seasonally dry habitats across a wide span of rainfall (38, 39).

120 The C<sub>3</sub> Pooldeae occupy regions with lower winter temperatures and shorter droughts 121 than the C<sub>4</sub> lineages (Fig. S7). C<sub>3</sub> Pooideae dominate grassy biomes to much higher 122 temperatures in the Palearctic than the Nearctic realm, although distributions of C4 123 Andropogoneae and Chloridoideae in these realms are similar (Fig. 3). Conversely, C<sub>3</sub> Pooideae 124 are confined to the geographically restricted colder parts of the Afrotropics and Indo-Malay 125 realms, and C<sub>4</sub> Andropogoneae dominate at much lower temperatures in these regions (Fig. 126 3). The sorting of C<sub>3</sub> and C<sub>4</sub> grass species along local and regional temperature gradients is well 127 established (40, 41), and the crossover temperature can be modified by ecosystem factors 128 (e.g., tree cover) (42). However, our observations are broadly consistent with model 129 predictions of carbon assimilation (22, 23, 43), as modeled crossover temperatures under low 130 light conditions and modern CO<sub>2</sub> levels occurs at ~20-22 °C.

131 In our data, some species of both Andropogoneae and Chloridoideae lineages have adapted to low mean annual temperatures and may persist in grassy vegetation within cool 132 parts of each realm (e.g. Fig. 3). Given equal investment in the carbon-fixing enzyme Rubisco, 133 134 a relatively low canopy leaf area and sunny conditions, a  $C_4$  canopy can theoretically achieve 135 higher total daily rates of photosynthesis than a  $C_3$  at any temperature (37). In this case, the 136 primary limitation on canopy carbon uptake becomes light-mediated damage during low 137 temperature extremes (44), although C<sub>4</sub> photosynthesis is energetically expensive. Low 138 temperature tolerance may be absent from most  $C_4$  species as  $C_4$  photosynthesis evolved in 139 the tropics (38).

140

Trait combinations of each lineage. Chloridoideae are distinguished from Andropogoneae in
 their occupation of regions with lower precipitation, higher daily variation in temperatures
 and longer droughts (Fig. S7). Further, these lineages are differentially associated with fire
 10

where Andropogoneae has the shortest fire return interval of 2 years, the peak occurrence of
Chloridoideae is at an interval of 8 years, while in Pooideae the modal fire return interval
exceeds 20 years (Fig. S7). Maximum plant heights of each lineage sort similarly, with values
peaking at 1.5 m for Andropogoneae and 0.6 m for both Chloridoideae and Pooideae (Fig. S7).
However, annual versus perennial life history is not globally relevant. The only significant areas
dominated by annual grasses occurring at the margins of the Sahara Desert and West Africa,
regions commonly considered as over-grazed.

151 80% of burned area globally occurs in the regions we see dominated by 152 Andropogoneae (20) and differs from other  $C_4$  grass lineages with its greater average height 153 and consequent rapid growth rates. Where rainfall exceeds 800 mm MAP in the tropics, soils 154 are typically leached and infertile (45). Andropogoneae produce leaves with relatively high C:N ratios (46, 47), which resist rapid decomposition. The tall, erect architecture of these 155 156 grasses produces a flammable well-connected fuelbed (48) and productive tropical 157 environments, with an annual dry season of > 5 months (13), are primed to burn as the grass 158 layer senesces. Experimental manipulations demonstrate that fire promotes dominance by 159 Andropogoneae (46) and we see this mirrored at a global scale. Grass persistence in these 160 competitive environments relies on the annual production of a new canopy and, in the 161 absence of woody investment, dead biomass must either rapidly decompose, burn or be 162 consumed by herbivores to avoid self-shading (11, 49). Andropogoneae are known to have 163 morphological adaptations enabling tolerances and persistence to fire that are not commonly 164 present in other grass lineages (49). Fire and other forms of repeated disturbance, such as 165 grazing, are therefore crucial for grass-dominated systems to persist in high rainfall 166 environments. While Andropogoneae appears to be the C<sub>4</sub> lineage most closely associated

with disturbance by fire, multiple lineages in the semi-arid African tropics appear linked to grazing tolerance (Fig S8, (50, 51)), and this may be due to the strength and form of environmental filtering associated with fire versus grazing, as well as the antiquity and biogeography of grazing pressure relative to fire.

171

172 Implications. The Andropogoneae, Chloridoideae and Pooideae grass lineages dominate 173 globally, via mechanisms encompassing plant production and competition, resilience to 174 drought, freezing and disturbance. Why do three of the most diverse grass lineages 175 characterise grassy biomes? Does diversity beget ecological success or does success beget 176 diversity? Early diversification may have enabled ecological success, such that ecological 177 speciation allowed each lineage to radiate across broad environmental envelopes (an 178 ecological mechanism). Alternatively, a neutral mechanism of a long history of diversification 179 may have led to high diversity as Andropogoneae and Chloridoideae are the oldest C<sub>4</sub> lineages. 180 Across our dataset, evidence for this is equivocal. We list 8.8% of all grass species and within 181 lineages: Andropogoneae, 14.5%; Chloridoideae, 6.5%; Pooideae, 10.8%. Perhaps ecological 182 success facilitated diversification, such that large geographical ranges enabled by unique 183 adaptations made the isolation of populations and allopatric speciation more likely (a 184 geographic mechanism). The rapid spread of the cosmopolitan *Themeda triandra* from Asia to 185 Africa in < 500,000 years supports this idea (52). Resolving the relative role of these 186 mechanisms requires comparative phylogenetic analyses of the relationships among ecology, 187 functional traits, range sizes and diversification rates.

188 The biogeographic contingencies described here in crossover temperatures align with 189 emerging evidence that regional evolutionary and environmental histories have been

important modifiers of biome-climate relationships (9, 53). However, the rapid rates of
dispersal observed in grasses (52), along with their short generation times (49), raises critical
questions about whether the biogeographic contingencies observed in woody plants should
be mirrored in grassy communities.

194 Global change will rapidly modify the existing global distribution of grassy biomes. 195 First, environmental change can alter feedbacks between grasses and woody plants via 196 changes in the processes limiting the growth and mortality of woody plants. For example, 197 rising  $CO_2$  is hypothesised to increase tree recruitment in savannas and forest margins (54, 198 55), while extreme drought events and warming may cause forest dieback on large scales (56), 199 where each process has feedbacks with fire leading to ongoing biome shifts (57). Second, 200 environmental changes will shift the community composition of grass communities. Our 201 analysis points to globally important ecotones between  $C_3$  and  $C_4$  likely to be influenced by 202 rising CO<sub>2</sub> and temperature (58), but these are better conceptualised as the boundary 203 between Pooideae and Chloridoideae in arid and semi-arid regions or regions of high grazing 204 pressure, and Pooideae and Andropogoneae in wetter regions. An experimental CO<sub>2</sub> 205 manipulation in dry mixed prairie found elevated CO<sub>2</sub> favoured a Pooideae dominant over a 206 Chloridoideae dominant, with rising temperature having the opposing effect (59). Conversely, 207 in a mesic tallgrass prairie, an Andropogoneae dominant displaced a Pooideae dominant in 208 competition under elevated CO<sub>2</sub> via improved water relations (60). In each case, C<sub>4</sub> 209 photosynthesis was one trait among many that influenced dynamic environmental responses. 210 Finally, the boundary between Andropogoneae and Chloridoideae is more likely to be 211 influenced by changes in rainfall amount and seasonality, along with shifting fire and grazing 212 regimes that can be directly altered by people at small and large scales.

213

Conclusions. The previous lack of synthesis in biome limits between grasses and woody plants constrains our understanding of how ecological and evolutionary processes determine the sensitivity of vegetation to global change. We have shown that divergent evolutionary histories and unique functional trait combinations have enabled three major grass lineages to dominate grassy biomes across global climate space. Local dominance by each lineage brings differing sensitivities to alternative global change drivers.

220

Acknowledgements. This research is a product of the National Evolutionary Synthesis Center 221 222 (NESCent) working group led by CPO, CAES, and CJS. DG was supported by a NESCent graduate 223 fellowship and NSF award 1342703. CPO was supported during the preparation of this 224 manuscript by Natural Environment Research Council grant (NE/I014322/1). Zhiyao Tang 225 helped to obtain the China map. Nikolai Ermakov and Daoud Rafikpoor provided shape files 226 of mapping data for Russia and Afghanistan, respectively. Anita Smyth assisted in obtaining 227 the Aekos data. Les Powrie, and Mike Rutherford assisted in obtaining the plot data from 228 South Africa. The South African National Biodiversity Institute and the South African 229 Biodiversity Facility are thanked for the use of data/information supplied by SANBI from 230 digitized collections. This work forms part of the "The National Vegetation Map" coordinated 231 by the South African National Biodiversity Institute.

Author Contributions. CERL, DMG, KJS, TMA, WB, ED, EJF, WH, LM, SP, JR, BS, MS, ES, RW, and CPO compiled the data. CERL, DMG, KS, DG and TK analysed the data. SA contributed fire data. CERL and CPO designed the study and wrote the paper with text contributions from DMG and KJS. All authors contributed comments on a draft of the paper. DMG and CERL perfected the figures.

# 237 Methods

Classifying grassy biomes. Data from 20 vegetation maps derived from botanical information,
or a combination of botanical and geographic information, were integrated to delineate grassy
biomes (references for these maps are listed in the Supplementary Information). The result
was a global map of grassy biomes resolved into 1,635 discrete vegetation units, each defined
by its characteristic grass species, which formed a list of 1,154 species (accounting for
synonymy) found commonly across global grassy biomes.

244 Vegetation maps are generally based on botanical survey and geographic analysis, 245 combined with expert input, that cluster species composition and vegetation structure to define unique vegetation units. We compiled the ground layer information for the vegetation 246 247 units in each map to identify the grass species considered to characterize a vegetation unit. 248 To determine whether vegetation units were naturally dominated by grasses, we developed 249 a set of criteria. First, artificial vegetation units were defined as those plowed or sown for 250 agriculture and where humans are planting species that would not otherwise occur. We 251 retained data for this analysis of only natural formations. Second, based on the vegetation descriptions we determined whether > 50% of the relative ground cover or biomass was 252 253 derived from grasses. We used this definition in place of 'Is there a continuous grassy ground 254 layer?' because low herbaceous cover in predominantly grassy vegetation would present a 255 problem with the classification of desertic and arid environments. Vegetation units were considered grassy deserts where the total above-ground biomass was considered <50 g m<sup>2</sup>, 256 257 or where total ground cover <25%, throughout the year. Finally, we retained all formations 258 where grasses were the dominant component of the ground layer, irrespective of tree cover. 259 Numerous grassy biomes, such as tropical savannas and woodlands, may be characterised by 260 up to 80% tree cover, but behave functionally as savannas due to a contiguous grassy ground layer (13, 35). Where necessary, we sourced additional information from published vegetation
descriptions and analyses to attribute key grass species to a grassy vegetation unit.
Additionally, vegetation units could be classified as mosaics with patches of closed canopy
vegetation intermingled with open vegetation, e.g. across the Steppe region of Russia.

Mapping grassy biomes. The vegetation maps we used as sources were developed throughout the 20<sup>th</sup> century. While this method provides an incomplete global coverage, we integrated available state-, country- and continent-level mapping to assemble what we consider to be the most robust map possible of the limits of grassy vegetation, where both vegetation characteristics and key constituent species could be identified. We were obliged to use the WWF Ecoregions map (61) where no other mapping was available. We re-assessed this global map to re-define units as grassy or not based on the criteria outlined above.

272 To quantify the global limits of grassy vegetation according to grass lineage, we gridded 273 the mapped data compilation at 0.5 degrees resolution. We calculated the proportion of each 274 0.5-degree grid cell occupied by grassy polygons. Using the grass phylogenetic and trait 275 information compiled, we then calculated the occupancy of grassy polygons by photosynthetic type, annual/perennial life history, grass lineage, and mean maximum grass height. These data 276 277 are not the same as a classic concept of abundance or dominance but are a relative measure 278 of the likelihood of occupancy measured from zero to unity. We undertook a validation of our 279 map compilation described in the Supplementary Information and in Figure S9.

Phylogenetic and plant trait information. We cross-referenced our species list to a taxonomy
of accepted scientific names (GrassBase, <u>http://www.kew.org/data/grasses-syn/cite.htm</u>)
and a recent accepted phylogeny from the Grass Phylogeny Working Group (25) to eliminate
synonymy and link species to descriptions of evolutionary history and functional traits.
Functional traits considered were: C<sub>3</sub>/C<sub>4</sub> photosynthetic pathway, maximum plant height,

285 annual/perennial life history, and tolerance of climatic extremes and fire frequency.  $C_3$  species 286 were divided amongst two groups: a polyphyletic group belonging to the PACMAD clade 287 (including the  $C_3$  sister groups for all  $C_4$  lineages); and the monophyletic BEP clade, a  $C_3$ outgroup to PACMAD, including bamboos, rice relatives and Pooideae species. C<sub>4</sub> grass species 288 289 were attributed to one of 24 independently evolved grass lineages. Maximum plant size is a 290 major axis of plant trait variation at a global scale (62), with maximum culm height in 291 herbaceous grasses reflecting annual rates of height growth, as most grasses annually senesce 292 their canopy (49). Height also describes differences in life history strategies related to light 293 competition and flammability and grazing tolerance (49). We included annual/perennial as 294 while most grasses reach sexual maturity in < 1 growing season, perennial grasses can be long-295 lived. Plant longevity is an effective strategy for occupying space in competitive environments 296 (63). We summarized these data for each grassy vegetation unit based on the grass species 297 listed as characteristic of each unit.

298 For the Poaceae species that we listed, we extracted all available georeferenced 299 occurrence records from the Global Biodiversity Information Facility (GBIF) web portal 300 (http://www.gbif.org/; accessed January 2014) and cleaned these data to ensure longitude 301 and latitude values were viable and to two decimal places. Species distributions were 302 standardised against descriptions of distributions in Grassbase using TDWG regions. For this subset of species produced via distribution records, median fire return intervals were 303 304 calculated at a species level following the methods of Archibald et al. 2010 (64). Information 305 on fire date was extracted for each GBIF location from the MODIS global monthly burnt area 306 (MCD45A1) satellite data product. To calculate climatic extremes for these same species, the 307 WorldClim dataset (www.worldclim.org) was used to obtain species median values of 308 minimum temperature (BIO6 variable) and seasonal drought length (calculated as the number

of successive months where mean annual precipitation was below 30mm). These species level
data were used to construct frequency histograms to examine lineage level variation in fire
regimes and climate extremes (Fig S7).

Environmental data used in global analyses. Our analysis aimed to elucidate lineage, climate 312 313 and disturbance relationships, and whether biogeography impacts the  $C_3$ - $C_4$  crossover 314 temperature. We used the WorldClim dataset at a 0.5 degree resolution to match the 315 vegetation map, and extracted mean annual precipitation (MAP), rainfall seasonality, mean 316 annual temperature (MAT) and temperature seasonality (www.worldclim.org). We used a 317 rainfall concentration index to describe rainfall seasonality based on (35). Growing season temperature (GST) was calculated for each grid cell to quantify regional and global  $C_3$ - $C_4$ 318 319 crossover temperatures. GST was calculated as the mean temperature across months with a 320 greater than or equal to 5 degree mean temperature and at least 25 mm rainfall, and was calculated using WoldClim monthly climate normals (65). 321

322 A median fire return interval (FRI) is the number of years between fire events that 323 represents the time period available for plants to grow. We used fire interval data from the 324 16 year MODIS fire datasets to fit Weibull distributions to 0.5° gridded data for the globe by 325 using the method outlined in Archibald et al. 2010 (64). Tropical grasslands and savannas have 326 the world's shortest fire return times, due to rapid rates of fuel accumulation and a climate 327 that supports frequent fire (annual dry seasons, warm climate and reliably seasonal rainfall) 328 (20). Our dataset of estimated fire return times, while spatially biased, is therefore robust for 329 grassy biomes.

330 Globally consistent data on present or past herbivore pressures are simply not 331 available. We were obliged to restrict our analyses to Africa where efforts have been made to 332 map mammalian herbivore pressures of both wildlife and livestock (66). We combined 18 herbivore and fire data to assess links between lineage composition and disturbance. Soils
data are not of sufficient quality to be meaningfully incorporated in analyses of this scale,
despite being known to mediate local scale vegetation patterns (35). In our global analyses,
we excluded grassy vegetation units defined as flooded, saline or edaphic, where the limits of
these units are generally decoupled from climate.

338 **Analyses.** First, we mapped the distribution of grassy biomes in geographic space according 339 to lineage and photosynthetic type to calculate the land area occupied by different grass 340 lineages in a "rank-abundance" style (Fig. 1). Grassy biome distributions were aligned with 341 MAT (in 1°C intervals) and MAP (in 100 mm intervals) to construct "Whittaker" style plots of 342 the limits of grassy biomes and of C<sub>3</sub> and C<sub>4</sub> photosynthetic types (Figure 2). These data were 343 further decomposed to represent the climate space of 17 grass lineages including Pooideae 344 that dominate grasslands (Figure S4). Data were also analysed by climate intervals of MAT and MAP to calculate the proportion of grassy land area occupied by each grass lineage within 345 346 each climate interval, to consider the potential for deterministic links between climate and 347 biomes (Figs. S3-S4).

Generalised additive models relating the distribution of lineages to growing season temperature, MAT and MAP across continents were fitted using the mgcv R package and the function predict.gam (67). Crossover temperatures plus standard deviations were calculated based on the temperature at which the predicted abundance of C<sub>4</sub> dominance reached 50%.

352 Random forest regressions (<u>https://cran.r-</u>
353 project.org/web/packages/randomForest/randomForest.pdf) were used to examine the
354 climate niche of key grass lineages and to infer correlations between four key climate
355 predictors (MAP, MAT, temperature seasonality and rainfall seasonality). Models were
356 constructed for six groups of interest: the C<sub>3</sub> BEP, PACMAD and Pooideae lineages; and the

357 independent C<sub>4</sub> lineages Andropogoneae, Chloridoideae and MPC (Melinidinae + Panicinae + 358 Cenchrinae) (25). Model fit was checked via a mean of squared residual test. The relative 359 importance of each environmental correlate was computed with a mean decrease of accuracy 360 test. The computed coverage response plots for each grass group was an adaptation of the 361 evaluation strip method developed by (68). These plots demonstrate the non-linear 362 relationships between environmental gradients and the various grass lineages. To produce 363 these plots, an environmental dataset was simulated where the focal environmental variable 364 is varied over its full environmental range and where, for each interval, the observed median 365 of each of other environmental variables (median over areas where the focus environmental 366 variable is within the interval) is returned. The displayed curve in each case is the prediction 367 of our Regression Random Forest model over this simulated dataset. The process used bi-368 variate response curves, where two variables rather than one vary simultaneously. The 90<sup>th</sup> quantile of a kernel density function (function kde2d from the R package ade4) was used to 369 370 plot limits of grass lineages relative to herbivore abundance and fire frequency.

# List of figures in main text Figure 1. Global distributions of grassy biomes and dominant grass lineages. Figure 2. Grassy biomes in global climate space. Figure 3. Continental disjunctions in lineage-growing season temperature relations. Figure 4. Global relationships between rainfall, fire and height.

Figure 1: Global distributions of grassy biomes and dominant grass lineages. A. Grassy biomes coded by the C<sub>4</sub> and C<sub>3</sub> grass lineages dominating each vegetation formation. B. Relationships among the dominant C<sub>4</sub> grass lineages, with colours matching those used on the map. The phylogeny is based on (25) and for simplicity excludes C<sub>3</sub> PACMAD sister clades. C. Rank-abundance curve for C<sub>4</sub> and C<sub>3</sub> grass lineages at the global scale, ordered by the proportion of the terrestrial surface dominated by each lineage.

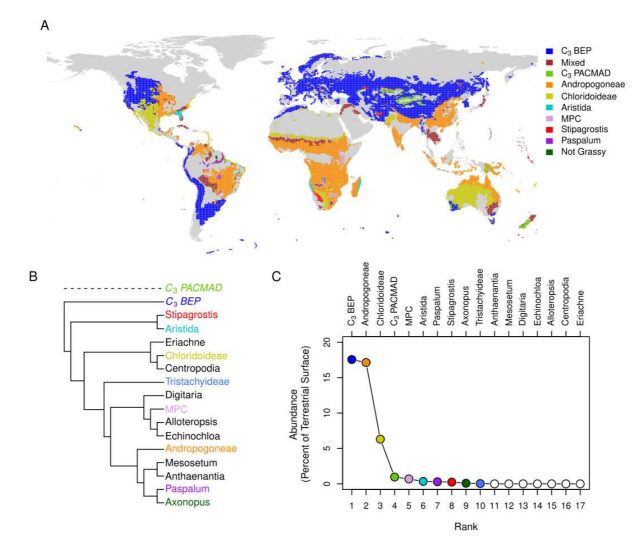
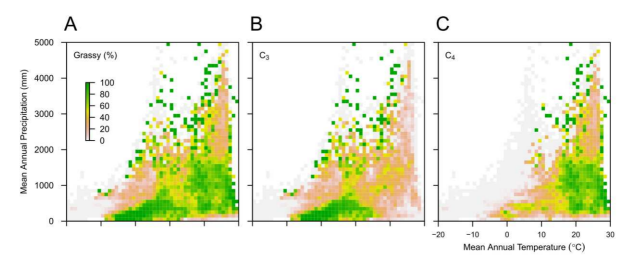


Figure 2: Grassy biomes in global climate space. Based on data at a 0.5-degree resolution, where data has been binned in 1° mean annual temperature (MAT) and 100 mm mean annual precipitation (MAP) intervals. Colour ramp shows the relative proportion of the global climate space for that MAP x MAT bin occupied by grassy biomes. Grey shading represents the vegetated land area. Data shown here link to Figure S1 showing the total vegetated land area within each climate interval.



# Fig. 3. Continental disjunctions in lineage-growing season temperature relationships. The distribution of grass lineages relative to growing season temperature (GST) in degrees Celsius globally (top panel) and then showing the variation in estimated $C_3$ - $C_4$ crossover temperatures by geographic realm. Distributions was fitted using generalized additive models and the crossover temperature calculated as the point where modelled $C_4$ grass abundance is 50%, based on the mapping in Figure 1. The fitted lines and confidence intervals are shown in different colours for each lineage, with the legend on the figure.

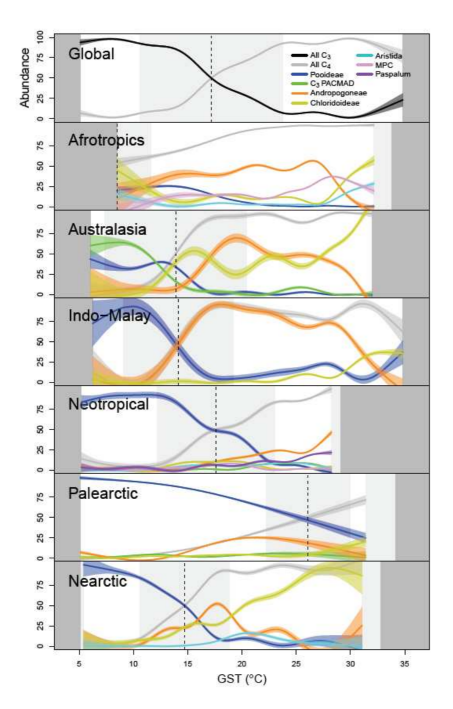
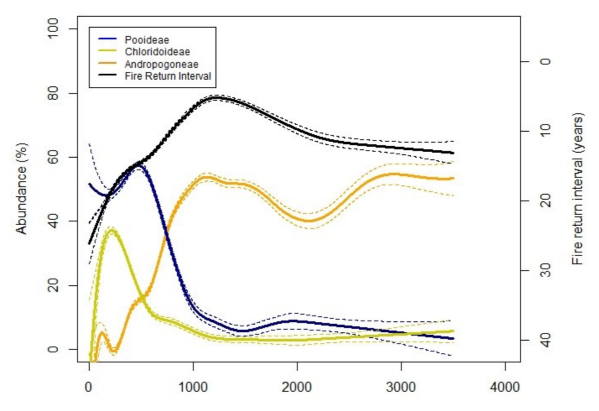




Figure 4. Global relationships between rainfall, fire and grass lineages. The global relationships of grass lineage abundance relative to MAP using a generalised additive model, showing 95% confidence intervals. The right-hand axis is the global relationship between fire return interval and MAP for grassy biomes and is inverted to reflect the inverse relationship with MAP. The global peak in fire activity coincides with the global peak in dominance of Andropogoneae grassy biomes. In contrast longer fire return times are associated with dominance by Pooideae and Chloridoideae.



Mean Annual Precipitation (mm)

410

# 412 References

- Carvalhais N, et al. (2014) Global covariation of carbon turnover times with climate in terrestrial ecosystems. *Nature* 514(7521):213-217.
- 415 2. Randerson JT, *et al.* (2015) Multicentury changes in ocean and land contributions to 416 the climate-carbon feedback. *Global Biogeochemical Cycles* 29(6):744-759.
- 417 3. Searchinger TD, et al. (2015) High carbon and biodiversity costs from converting
  418 Africa's wet savannahs to cropland. Nature Clim. Change 5(5):481-486.
- 419 4. Bastin J-F, *et al.* (2017) The extent of forest in dryland biomes. *Science* 356(6338):635420 638.
- 421 5. Dexter KG, et al. (2015) Floristics and Biogeography of Vegetation in Seasonally Dry
  422 Tropical Regions. International Forestry Review 17(S2):10-32.
- 423 6. Slik JWF, et al. (2018) Phylogenetic classification of the world's tropical forests. PNAS
  424 115(8):1837-1842.
- 425 7. Pausas JG & Bond WJ (2018) Humboldt and the reinvention of nature. *J. Ecology*426 doi.org/10.1111/1365-2745.13109.
- 427 8. Archibald S, et al. (2018) Biological and geophysical feedbacks with fire in the Earth
  428 system. Environmental Research Letters 13(3):033003.
- 429 9. Lehmann CER, et al. (2014) Savanna Vegetation-Fire-Climate Relationships Differ
  430 Among Continents. Science 343(6170):548-552.
- Hoffmann WA, *et al.* (2012) Ecological thresholds at the savanna-forest boundary: how
  plant traits, resources and fire govern the distribution of tropical biomes. *Ecology Letters* 15(7):759-768.
- 434 11. Bond WJ (2008) What Limits Trees in C4 Grasslands and Savannas? *Annual Review of*435 *Ecology, Evolution, and Systematics* 39(1):641-659.
- Lloyd J, et al. (2008) Contributions of woody and herbaceous vegetation to tropical
  savanna ecosystem productivity: a quasi-global estimate. *Tree Physiology* 28(3):451468.
- 439 13. Ratnam J, et al. (2011) When is a 'forest' a savanna, and why does it matter? Global
  440 Ecology and Biogeography 20(5):653-660.
- 441 14. Edwards EJ, Osborne CP, Stromberg CAE, Smith SA, & C4 Grasses Consortium (2010)
  442 The Origins of C4 Grasslands: Integrating Evolutionary and Ecosystem Science. *Science*443 328(5978):587-591.
- 444 15. Strömberg CAE (2011) Evolution of Grasses and Grassland Ecosystems. *Annual Review*445 *of Earth and Planetary Sciences* 39(1):517-544.
- 446 16. Cerling TE (1997) Global vegetation change through the Miocene/Pliocene boundary.
  447 *Nature* 389:153-158.
- 44817.Cerling TE, et al. (2011) Woody cover and hominin environments in the past4496[thinsp]million years. Nature 476(7358):51-56.
- 45018.Wynn JG, et al. (2013) Diet of Australopithecus afarensis from the Pliocene Hadar451Formation, Ethiopia. PNAS 110(26):10495-10500.
- 452 19. Lehmann CER & Parr CL (2016) Tropical grassy biomes: linking ecology, human use and
  453 conservation. *Phil. Trans. R. Soc. B* 371(1703).
- 45420.Archibald S, Lehmann CER, Gomez-Dans J, & Bradstock RA (2013) Defining pyromes455and global syndromes of fire. PNAS 10(16):6442 6447.

- 456 21. Ehleringer JR, Cerling TE, & Helliker BR (1997) C<sub>4</sub> photosynthesis, Atmospheric CO<sub>2</sub>, and
  457 climate. *Oecologia* 112(3):285-299.
- 458 22. Collatz GJ, Berry JA, & Clark JS (1998) Effects of climate and atmospheric CO2 partial
  459 pressure on the global distribution of C4 grasses: present, past, and future. *Oecologia*460 V114(4):441-454.
- 461 23. Still CJ, Berry JA, Collatz GJ, & DeFries RS (2003) Global distribution of C<sub>3</sub> and C<sub>4</sub>
  462 vegetation: Carbon cycle implications. *Global Biogeochem. Cycles* 17(1).
- 463 24. Woodward F, Lomas M, & Kelly C (2004) Global climate and the distribution of plant
  464 biomes. *Phil. Trans. R. Soc. B* 359(1450):1465-1476.
- 465 25. Grass Phylogeny Working G, II (2012) New grass phylogeny resolves deep evolutionary
  466 relationships and discovers C4 origins. *New Phytologist* 193(2):304-312.
- 467 26. Christin P-A & Osborne CP (2013) The recurrent assembly of C4 photosynthesis, an
  468 evolutionary tale. *Photosynthesis Research* 117(1):163-175.
- 469 27. Taub DR (2000) Climate and the US distribution of C<sub>4</sub> grass subfamilies and
  470 decarboxylation variants of C4 photosynthesis. *American Journal of Botany*471 87(8):1211-1215.
- 47228.Liu H, Edwards EJ, Freckleton RP, & Osborne CP (2012) Phylogenetic niche473conservatism in C4 grasses. *Oecologia* 170(3):835-845.
- 474 29. Dixon AP, Faber-Langendoen D, Josse C, Morrison J, & Loucks CJ (2014) Distribution
  475 mapping of world grassland types. *J. Biogeog.* 41(11):2003-2019.
- 476 30. Griffith DM, et al. (2017) Comment on "The extent of forest in dryland biomes".
  477 Science 358(6365):eaao1309.
- Solofondranohatra CL, et al. (2018) Grass Functional Traits Differentiate Forest and
  Savanna in the Madagascar Central Highlands. Frontiers in Ecology and Evolution
  6(184).
- 481 32. Smith MD & Knapp AK (2003) Dominant species maintain ecosystem function with
  482 non-random species loss. *Ecology Letters* 6(6):509-517.
- Winfree R, W. Fox J, Williams NM, Reilly JR, & Cariveau DP (2015) Abundance of
  common species, not species richness, drives delivery of a real-world ecosystem
  service. *Ecology Letters* 18(7):626-635.
- 486 34. Staver AC, Archibald S, & Levin SA (2011) The Global Extent and Determinants of
  487 Savanna and Forest as Alternative Biome States. *Science* 334(6053):230-232.
- 48835.Lehmann CER, Archibald SA, Hoffmann WA, & Bond WJ (2011) Deciphering the489distribution of the savanna biome. New Phytologist 191(1):197-209.
- 490 36. Sandve SR & Fjellheim S (2010) Did gene family expansions during the Eocene–
  491 Oligocene boundary climate cooling play a role in Pooideae adaptation to cool
  492 climates? *Molecular Ecology* 19(10):2075-2088.
- 493 37. McKeown M, Schubert M, Marcussen T, Fjellheim S, & Preston JC (2016) Evidence for
  494 an early origin of vernalization responsiveness in temperate Pooideae grasses. *Plant*495 *physiology* 172(1):416-426.
- 49638.Edwards EJ & Smith SA (2010) Phylogenetic analyses reveal the shady history of C4497grasses. PNAS 107(6):2532-2537.
- 498 39. Osborne CP & Freckleton RP (2009) Ecological selection pressures for C<sub>4</sub>
  499 photosynthesis in the grasses. *Proceedings of the Royal Society B: Biological Sciences*500 276(1663):1753-1760.
- 50140.Teeri J & Stowe L (1976) Climatic patterns and the distribution of C4 grasses in North502America. Oecologia 23(1):1-12.

- 50341.Young HJ & Young TP (1983) Local distribution of C3 and C4 grasses in sites of overlap504on Mount Kenya. *Oecologia* 58(3):373-377.
- 505 42. Griffith DM, et al. (2015) Biogeographically distinct controls on C<sub>3</sub> and C<sub>4</sub> grass
  506 distributions: merging community and physiological ecology. Global Ecology and
  507 Biogeography 24(3):304-313.
- 508 43. Ehleringer JR (1978) Implications of quantum yield differences on the distributions of  $C_3$  and  $C_4$  grasses. *Oecologia* 31(3):255-267.
- 510 44. Long SP (1999) Environmental responses. *C*<sub>4</sub> *plant biology*:215-249.
- 51145.Högberg P (1982) Mycorrhizal associations in some woodland and forest trees and512shrubs in Tanzania. New Phytologist 92(3):407-415.
- Forrestel EJ, Donoghue MJ, & Smith MD (2014) Convergent phylogenetic and
  functional responses to altered fire regimes in mesic savanna grasslands of North
  America and South Africa. *New Phytologist* 203(3):1000-1011.
- 51647.Taylor SH, et al. (2010) Ecophysiological traits in C3 and C4 grasses: a phylogenetically517controlled screening experiment. New Phytologist 185(3):780-791.
- 518 48. Simpson KJ, et al. (2016) Determinants of flammability in savanna grass species. J.
  519 Ecology 104(1):138-148.
- Linder H, Lehmann CE, Archibald S, Osborne CP, & Richardson DM (2018) Global grass
  (Poaceae) success underpinned by traits facilitating colonization, persistence and
  habitat transformation. *Biological Reviews* 93(2):1125-1144.
- 52350.Forrestel EJ, et al. (2017) Different clades and traits yield similar grassland functional524responses. PNAS 201(6)129-09.
- 525 51. Anderson TM, Shaw J, & Olff H (2011) Ecology's cruel dilemma, phylogenetic trait 526 evolution and the assembly of Serengeti plant communities. *J. Ecology* 99(3):797-806.
- 527 52. Dunning LT, *et al.* (2017) The recent and rapid spread of Themeda triandra. *Botany* 528 *Letters*.
- 529 53. Rogers BM, Soja AJ, Goulden ML, & Randerson JT (2015) Influence of tree species on
  530 continental differences in boreal fires and climate feedbacks. *Nature Geoscience*531 8(3):228.
- 53254.Bond WJ & Midgley GF (2000) A proposed CO2-controlled mechanism of woody plant533invasion in grasslands and savannas. Global Change Biology 6(8):865-869.
- 53455.Stevens N, Lehmann CER, Murphy BP, & Durigan G (2017) Savanna woody535encroachment is widespread across three continents. Glob. Change Biol 23(1):235-536244.
- 537 56. Van Mantgem PJ, *et al.* (2009) Widespread increase of tree mortality rates in the 538 western United States. *Science* 323(5913):521-524.
- 53957.Osborne CP, et al. (2018) Human impacts in African savannas are mediated by plant540functional traits. New Phytologist.
- 54158.Ehleringer JR (2005) The influence of atmospheric CO2, temperature, and water on the542abundance of  $C_3/C_4$  taxa. A history of atmospheric CO2 and its effects on plants,543animals, and ecosystems (Springer), pp 214-231.
- 54459.Morgan JA, et al. (2011) C4 grasses prosper as carbon dioxide eliminates desiccation in545warmed semi-arid grassland. Nature 476(7359):202.
- 546 60. Owensby CE, Ham JM, Knapp AK, & Auen LM (1999) Biomass production and species
  547 composition change in a tall grass prairie ecosystem after long-term exposure to
  548 elevated atmospheric CO<sub>2</sub>. *Glob. Change Biol* 5:497.

- 549 61. Olson DM, *et al.* (2001) Terrestrial Ecoregions of the World: A New Map of Life on Earth
  550 A new global map of terrestrial ecoregions provides an innovative tool for conserving
  551 biodiversity. *BioScience* 51(11):933-938.
- 552 62. Díaz S, et al. (2016) The global spectrum of plant form and function. *Nature* 553 529(7585):167.
- 554 63. Raven J & Thomas H (2010) Grasses. *Current Biology* 20(19):R837-R839.
- 55564.Archibald S, Scholes R, Roy DP, Roberts G, & Boschetti L (2010) Southern African fire556regimes as revealed by remote sensing. Int. J. Wildland Fire 19(7):861-878.
- 557 65. Fick SE & Hijmans RJ (2017) WorldClim 2: new 1-km spatial resolution climate surfaces 558 for global land areas. *Int. J. Climatology* 37(12):4302-4315.
- 55966.Hempson GP, Archibald S, & Bond WJ (2015) A continent-wide assessment of the form560and intensity of large mammal herbivory in Africa. Science 350(6264):1056-1061.
- 561 67. Wood S (2011) Mixed GAM Computation Vehicle with GCV/AIC/REML smoothness 562 estimation and GAMMs by REML/PQL.
- 563 68. Elith J, Ferrier S, Huettmann F, & Leathwick J (2005) The evaluation strip: a new and
  564 robust method for plotting predicted responses from species distribution models.
  565 *Ecological modelling* 186(3):280-289.
- 566

# 568 Supplementary Figures, Tables and Information for:

# 569 Functional diversification enabled grassy biomes to fill global climate space

- 570
- 571 Authored by: Caroline E. R. Lehmann, Daniel M. Griffith, Kimberley J. Simpson, T. Michael
- 572 Anderson, Sally Archibald, David J. Beerling, William J. Bond, Elsie Denton, Erika J. Edwards,
- 573 Elisabeth J. Forrestel, David L. Fox, Damien Georges, William A. Hoffmann, Thomas Kluyver,
- 574 Ladislav Mucina, Stephanie Pau, Jayashree Ratnam, Nicolas Salamin, Bianca Santini, Melinda
- 575 D. Smith, Elizabeth L. Spriggs, Rebecca Westley, Christopher J. Still, Caroline A.E. Strömberg
- 576 and Colin P. Osborne
- 577
- 578 Contents
- 579 1. Description of map compilation validation
- 580 2. Supplementary Figures 1 9
- 581 3. Table of land area of different lineages according to different projections
- 582 4. References to dataset development
- a. References to vegetation maps and associated information
- b. References for plot data for map validation

# 586 **Development and validation of map compilation**

587 Rarely, if ever, has this rich body of vegetation mapping research been integrated with Earth 588 system science or evolutionary studies. This is perhaps because vegetation mapping is 589 considered a descriptive natural science in an age of big data. The contiguous land mass 590 covering the countries of China, Mongolia, the former Soviet Union, Afghanistan, Turkey and 591 Europe are represented by detailed botanical data. The regions of Africa, North America, 592 Mexico, Panama, Venezuela, Brazil, Argentina, Papua New Guinea, Indonesia, northern and 593 western Australia are also well documented by botanical data. However, there is a general 594 paucity of adequate vegetation mapping available across India, South-East Asia (Burma, 595 Thailand, Laos, and Vietnam), Central America, and parts of South America (Chile, Peru, 596 Bolivia, Uruguay, Paraguay, Ecuador, and Columbia). It is worth noting that given anticipated 597 impacts of global change on the distribution and dynamics of vegetation, an absence of 598 publicly available vegetation mapping for key regions such as South East Asia and the Andes 599 should be of concern to many.

600

601 We undertook a validation process between plot data describing in situ grass abundance and 602 our global species list. Using publicly available data that intersected with vegetation unit 603 descriptions we found that, at the level of independent evolutionary lineages of grasses (i.e., 604 subfamily), we had strong confidence in the geographic and environmental relationships we 605 elucidate here (Fig. S9). To validate the classification of common grass species across regions, 606 we compared the species list in each vegetation unit to a plot level database developed for 607 validation purposes (Fig. S9). Plot data were sourced from the literature and vegetation 608 databases and assembled by the authors (see references in the Supplementary Information).

609 110 vegetation units contained enough plot level data for validation analyses. To determine 610 what taxonomic levels agree with plot data, the comparison was conducted at the species and 611 subfamily levels. We also examined the agreement of our map and plot datasets at a 612 functional level by comparing the attribution of photosynthetic type. From the 507 common 613 grass species across these vegetation units, 88% of these species were present in the plot 614 dataset of those appropriate vegetation types. This is a very high degree of overlap in species 615 in our mapping classifications and plot data, especially considering the difference in scale 616 between local species plots and large vegetation units. Furthermore, we found that vegetation 617 types generally had similar percentages of characteristic grass species represented in their plot datasets, although the agreement was worse for particularly large and broadscale 618 619 vegetation units. To validate the higher taxonomic classifications and plant functional type 620 classifications of our map units, we compared the proportion each classification in plots 621 (weighted by abundance) to the proportion of that classification in our map. Because these data are on the interval (0,1) we used beta regression to model this relationship. Beta 622 623 regression can be interpreted much like logistic regression, except that it allows continuous 624 values in the dependent variable. Proportions of Poaceae subfamilies and functional types 625 showed that plot values were strongly predictive of classified values in our map (Fig. S9).

627 Figure S1: Global distribution of grassy biomes. The global map was derived as a composite 628 from national and regional maps of vegetation that was gap-filled using the Ecoregions map 629 (see Methods in the main text and references for all maps at the end of the Supplementary 630 Information). Coloured areas show the extent of grassy biomes globally and dominance of 631 these by C<sub>3</sub> grasses and C<sub>4</sub> grasses mapped at the scale of identified vegetation units (i.e., polygons). Red = High proportion of C4 grasses. Blue = High proportion C3 grasses. Datum: 632 WGS84. 633

634

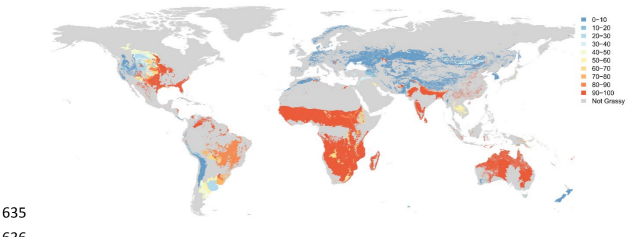
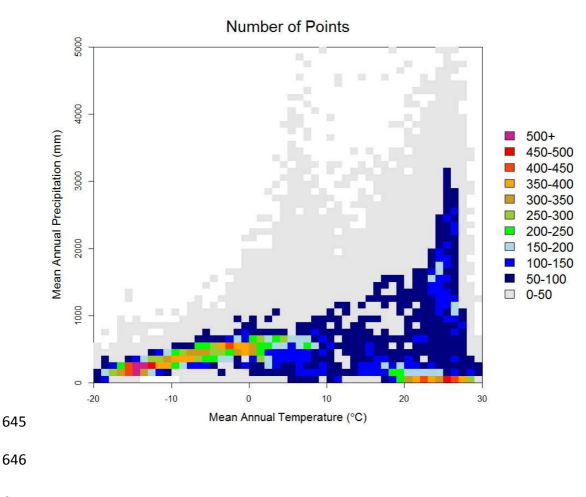


Figure S2. Global vegetated land area as related to Mean annual temperature and mean annual precipitation. Mean annual precipitation is in 100 mm bins, while temperature is in 1oC bins. The color ramp represents the number of 0.5 degree points in 100mm x 1oc unit of climate space. Note the grey background that highlights the global extent of climate space where these temperature – precipitation combinations are essentially rare on the vegetated land surface. The color ramp from dark blue to purple represents an increasing density of points in a given climate bin.

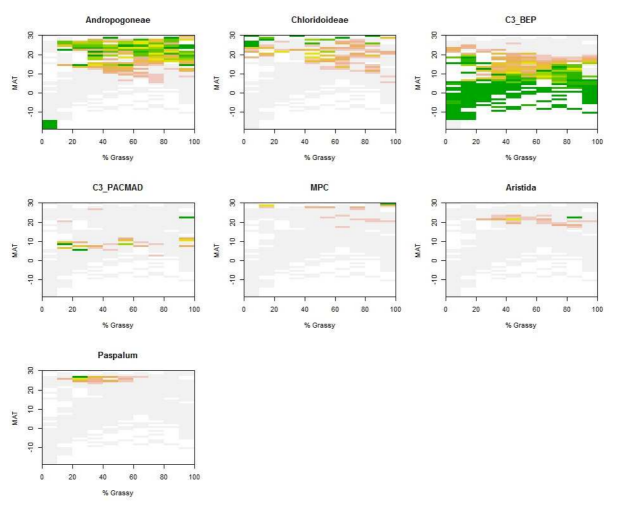




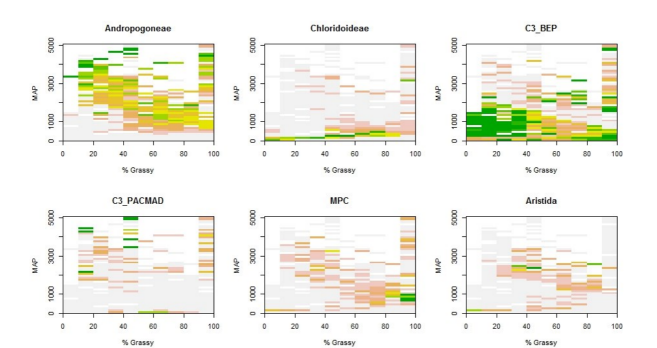
647

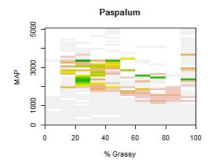
649 Figure S3: Global abundance of the main grass lineages by temperature and rainfall. Colour 650 scale indicates the proportion of 0.5° grid squares dominated by each lineage at the global 651 scale for each of (A) mean annual temperature and (B) Mean annual precipitation. These plots 652 demonstrate that, in cool, dry regions where the C3 Pooideae lineage is concentrated, it tends 653 to be the only grass lineage present, and this lineage dominates that climate space. These can 654 be considered as deterministic grasslands. In contrast, the heterogeneity of the dominance of C4 Andropogoneae and C4 Chloridoideae lineages across climate space could suggest that the 655 grassy biomes where these lineages are found are not deterministic, and dominance may be 656 657 driven by processes other than climate.

#### 658 A. Dominance by mean annual temperature



### 660 **B. Dominance by mean annual precipitation**

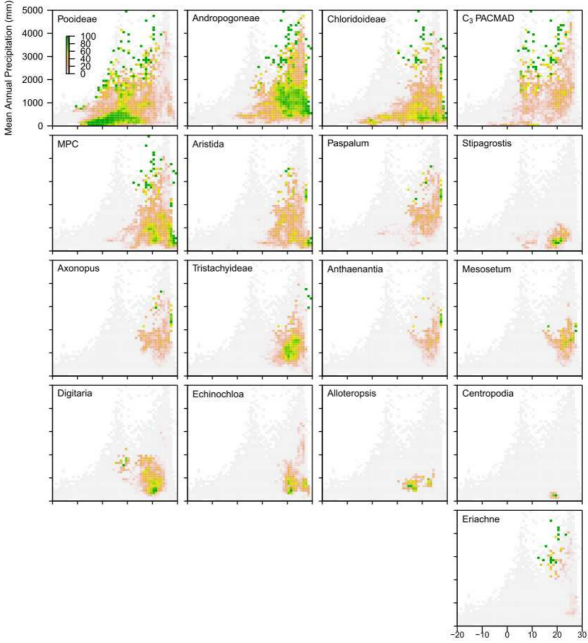




662

661

Figure S4: Concentration of 17 grass lineages in climate space. This figure builds on S1 – S2
by again highlighting the climate space characterised by different grass lineages. It is very clear
that C3 PACMAD dominance is highly restricted to warmer wetter parts of climate and we
know from S1 that geographically these combinations of temperature and precipitation are
limited. These figures also again highlight the wide distribution of Pooideae, Andropogoneae
and Chloridoideae.



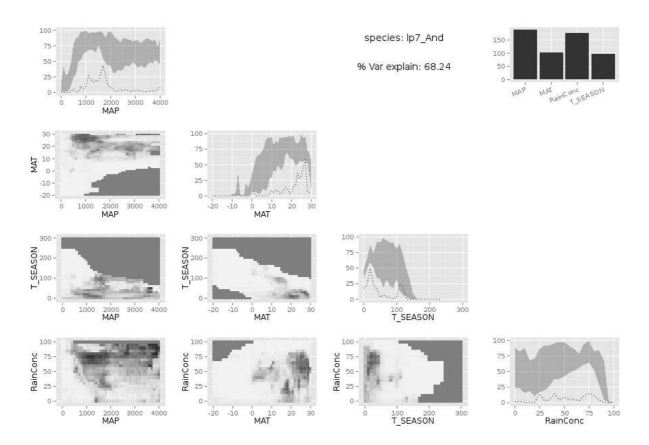
Mean Annual Temperature (°C)

Figure S5: Plots from Random Forest analyses of the relative importance of mean
temperature, mean precipitation, temperature seasonality and rainfall seasonality in the
limits of the three key lineages of grasses. A) C3 BEP, B) C4 Andropogoneae, and C) C4
Chloridoideae. Model fits against data are shown for the land area over which each lineage
dominates against each climate variable.

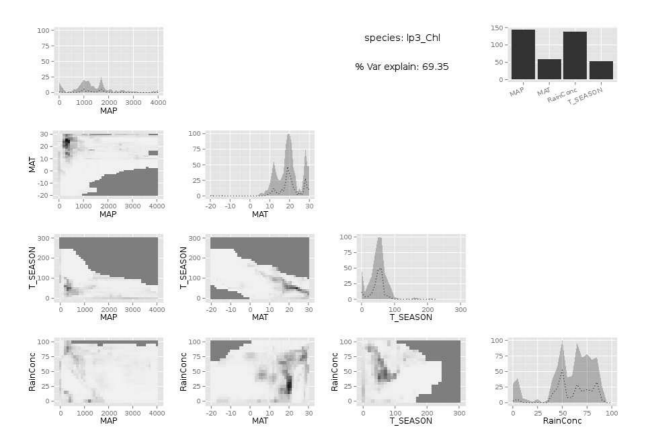
A. C<sub>3</sub> BEP

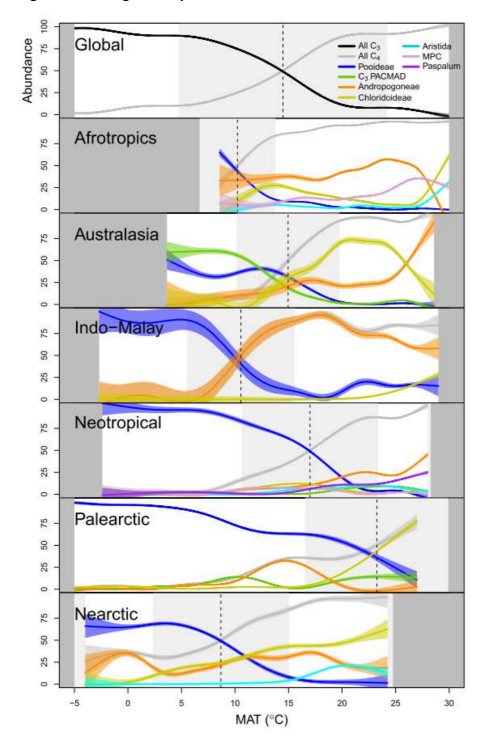
species: lpC3\_BEP % Var explain: 71.56 MAT RainConc LSEASON MARP MAP MAT -10 -20 MAT MAP T SEASON SEASON 3000 4000 -10 -20 MAP 0 10 MAT 100 200 T SEASON RainConc RainConc RainConc 0-RainConc MAP -10 MAT T\_SEASON

#### 679 B. C<sub>4</sub> Andropogoneae



# 681 C. C<sub>4</sub> Chloridoideae





### 683 Figure S6. Lineage – temperature associations.



685 Fig. S7: Trait-environment associations. Density plots of traits that characterise the realised 686 ecological niche of dominant grass lineages: A) Median minimum annual temperature (°C) 687 across the range of each species; B) Median drought length (months); and C) Median fire 688 return interval (years), calculated by mapping GBIF occurrence data for each species onto 689 Earth Observation data layers (see Methods). D) Maximum height of the culm (flowering stem) 690 for each species, as a measure of plant size at maturity (see Methods). In each case, species 691 from each lineage recorded within vegetation units in our dataset were mapped across their whole range (i.e. beyond the area over which they dominate ground cover). 692

693

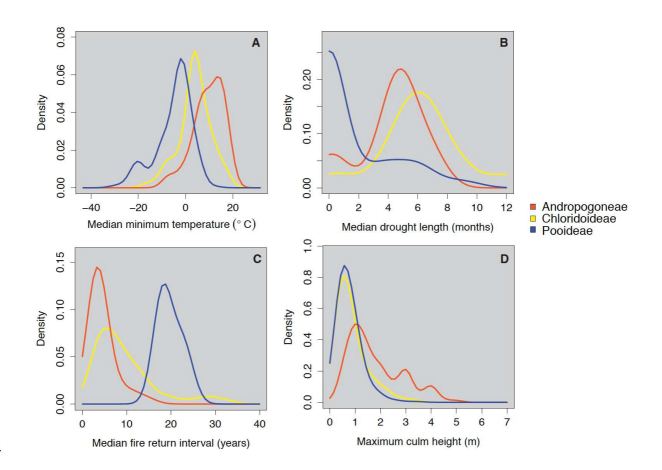
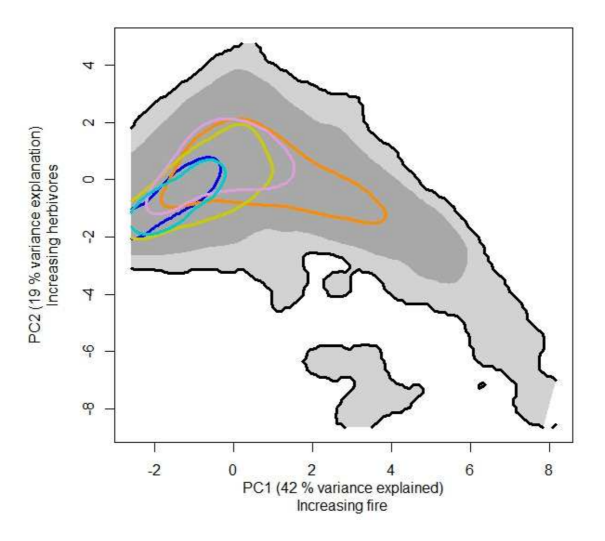
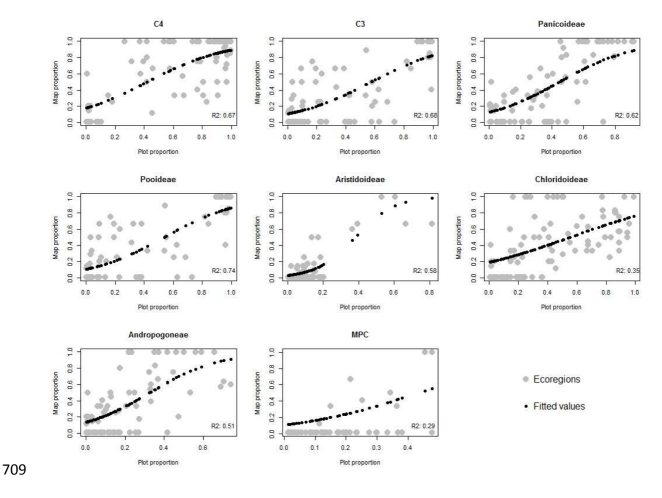


Figure S8. Ordination of fire frequency and herbivore biomass with the 90<sup>th</sup> quantile of 695 696 lineage distributions shown via different colours. This ordination shows the limits of five 697 grass lineages relative to the prevalence of fire and modelled herbivore biomass. Colours 698 representing lineages are consistent with figures in the main text. Key here is that one lineage 699 stretches into environments of more frequent fire (Andropogoneae = orange), while a number 700 of lineages are clustered and overlapping with respect to variation in herbivore biomass. 701 However, it is worth noting the globally poor data on herbivore biomass in contrast to fire that 702 is relatively easily quantified by satellites as changes in surface reflectance and heat.



703

Figure S9. Validation of mapping approach to determine function and lineage level distributions of grassy biomes. Shown are figures relating plot level versus map level estimates of different grass groups (as shown on each plot). Logistic regression was used to quantify relationships and the deviance explained of the analyses are shown on each plot.



- 711 **Table S1: Land area occupied by each grass lineage.** Each column represents a slightly
- 712 different way to calculate the relative coverage of grassy biomes by different grass lineages.
- 713 Polygon calculations are based on the mapped polygons while grid cells represent the
- conversion of data from Both calculations use a WGS84 projection.

LINEAGE	Percentage of grassy (from	Percentage of grassy (from grid
	polygons)	cells)
Andropogoneae	38.99	36.66
C3 BEP	32.51	37.57
Chloridoideae	13.93	13.51
Mixed	10.66	6.56
MPC	1.38	1.48
C3 PACMAD	0.92	2.06
Aristida	0.64	0.73
Stipagrostis	0.62	0.53
Paspalum	0.18	0.64
Tristachyideae	0.09	0.09
Axonopus	0.07	0.17

# 716 Appendix S1: Supplemental Data References

717

## 718 **References to vegetation maps and associated descriptions**

- Barbour, M. G. and Major, J. 2007. Terrestrial vegetation of California, 3rd ed. University of
   California Press, Los Angeles.
- BirdLife International 2004. Important Birds Areas in Asia: key sites for conservation.
   Cambridge, UK: BirdLife International. (BirdLife Conservation Series No. 13).
- Blair, G.J. and Rudolf, W., 1984. Grasslands of Indonesia and the effect of animals on their
   stability. on land evaluation for extensive grazing (LEEG), p.153.
- Blasco, F., Bellan, M.F. and Aizpuru, M., 1996. A vegetation map of tropical continental Asia at
   scale 1: 5 million. Journal of Vegetation Science, 7(5), pp.623-634.
- Bohn U, Neuhäusl R, Gollub G, Hettwer C, Neuhäuslová Z, Schlüter H, Weber H. 2003. Karte
  der natürlichen Vegetation Europas. Maßstab 1:2.500.000 / Map of the Natural
  Vegetation of Europe. Maßstab / Scale 1:2.500.000. Münster, Germany:
  Landwirtschaftsverlag.3 volumes + CD.
- Breckle, S.-W., Ditman, A. & Rafiqpoor, M.D. 2010. Field guide Afghanistan. Flora and vegetation. Scientia Bonnensis, Bonn, Germany. Map: GIS raster (shape files) supplied by Daoud Rafiqpoor and these data correspond to the map in Breckle et al. 2010, Fig. 2.01: natural vegetation of Afghanistan. Description of vegetation types: Breckle et al. 2010, pp 91-115
- Comer, P., D. Faber-Langendoen, R. Evans, S. Gawler, C. Josse, G. Kittel, S. Menard, M. Pyne,
   M. Reid, K. Schulz, K. Snow, and J. Teague. 2003. Ecological Systems of the United
   States: A Working Classification of U.S. Terrestrial Systems. NatureServe, Arlington,
   Virginia.
- Comision Nacional para el Conocim iento y Uso de la Biodiversidad (CONABIO), 1999. "Uso de suelo y vegetacio modificado por CONABIO". Escala 1: 1000 000. Comision Nacional para el Conocimiento y Uso de la Biodiversidad. Ciudad de Mexico, Mexico.
- Djoeroemana, S. and Myers, B., 2000. Fire and sustainable agricultural and forestry
   development in eastern Indonesia and northern Australia. In International workshop
   on Fire and Sustainable Agricultural and Forestry Development in Eastern Indonesia
   and Northern Australia. Australian Centre for International Agricultural Research,
   Canberra.
- Editorial Committee of Vegetation Map of China, 2007. Vegetation map of the People's
   Republic of China (1:1000,000). Geological Publishing House, Beijing, CN.
- 750European Environmental Agency. 2010. Manual for the European Environment Agency's Land751accountsdataviewer2000-2006version1-17/11/2010.

- 752 http://www.eea.europa.eu/data-and-maps/figures/corine-land-cover-2006-by-753 country-1
- Fox, I.D., Neldner, V.J., Wilson, G.W. & Bannink, P.J. 2001. The vegetation of the Australian
   tropical savannas. Brisbane: Environment Protection Agency.
- Garrity, D.P., Soekardi, M., Van Noordwijk, M., De La Cruz, R., Pathak, P.S., Gunasena, H.P.M.,
  Van So, N., Huijun, G. and Majid, N.M., 1996. The Imperata grasslands of tropical Asia:
  area, distribution, and typology. Agroforestry Systems, 36(1-3), pp.3-29.
- Griffith, G.E., Omernik, J.M., Johnson, C.B., and Turner, D.S., In prep., Ecoregions of Arizona
   (color poster with map, descriptive text, summary tables, and photographs): Menlo
   Park, California, U.S. Geological Survey (map scale 1:1,325,000).
- Hannibal, L.W., 1950. Vegetation Map of Indonesia. Planning Department, Forest Service,
  Jakarta. Forest Policies in Indonesia: The Sustainable Development of Forest Lands,
  3.Brookfield, H. and Byron, Y., 1993. South-East Asia's environmental future: the
  search for sustainability.
- Harris, S and Kitchener, A 2005. From Forest to Fjaeldmark: Descriptions of Tasmania's
   Vegetation. Department of Primary Industries, Water and Environment, Printing
   Authority of Tasmania. Hobart.
- Huber, O., 1988. Mapa de vegetación de Venezuela. República de Venezuela, Ministerio del
  Ambiente y de los Recursos Naturales Renovables, Dirección Genreal de Información
  e Investigación del Ambiente, Dirección de Suelos, Vegetación y Fauna, División de
  Vegetación.
- Huber, O., Gharbarran, G. and Funk, V.A., 1995. Vegetation map of Guyana. Centre for the
  Study of Biological Diversity, University of Guyana.
- 775 IBGE, D., 1993. Vegetation Map of Brazil. Rio de Janeiro: IBGE.
- Isachenko, T. I. (ed.) 1990. Vegetation of the Soviet Union. Scale 1:4,000,000. Institute of
   Geography, Siberian Dept. of the USSR Academy of Sciences. Novosibirsk.
- 778 Karamysheva ZV, Khramtsov VN. 1995. The steppes of Mongolia. Braun-Blanquetia 17:1–79.
- Kuchler, A.W. 1964. Potential natural vegetation of the conterminous United States. American
   Geographical Society, University of Minnesota.
- Navarro, G. and Ferreira, W., 2007. Mapa de vegetación de Bolivia a escala 1: 250.000. On CD ROM. The Nature Conservancy (TNC), RUMBOL. Cochabamba, Bolivia.
- Maxwell, A. L. 2004. Fire regimes in north-eastern Cambodian monsoonal forests, with a 9300 year sediment charcoal record. Journal of Biogeography, 31, 225-239.
- Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powell, G. V., Underwood,
   E. C., D'amico, J. A., Itoua, I., Strand, H. E. & Morrison, J. C. 2001. Terrestrial Ecoregions

of the World: A New Map of Life on Earth A new global map of terrestrial ecoregions
provides an innovative tool for conserving biodiversity. BioScience, 51, 933-938.

- Otsamo, A., Ådjers, G., Hadi, T.S., Kuusipalo, J. and Vuokko, R., 1997. Evaluation of
   reforestation potential of 83 tree species planted on Imperata cylindrica dominated
   grassland–A case study from South Kalimantan, Indonesia. New Forests, 14(2), pp.127 143.
- Paijmans K, CSIRO, 1975. Explanatory Notes to the Vegetation map of Papua New Guinea.
   CSIRO, Melbourne, AU.
- Pattiselanno, F. and Arobaya, A.Y.S., 2009. Grazing Habitat of the Rusa Deer (Cervus
  timorensis) in the Upland Kebar, Manokwari. Biodiversitas Journal of Biological
  Diversity, 10(3).
- Shepherd, D.P. 2003. Implementation of the National Vegetation Information System model
  in Western Australia. Milestone 6 Report. Final report on the implementation of the
  National Vegetation System model in Western Australia. Department of Agriculture,
  Perth. Unpublished Report to the Bureau of Rural Sciences, Canberra. Which was later
  replaced by publication of: Beard JS, Beeston GR, Harvey JM, Hopkins AJM, Shepherd
  DP. 2013. The Vegetation of Western Australia at the 1:3,000,000 Scale. Explanatory
  Memoir. 2nd ed. Conservation Science Western Australia 9: 1–152.
- Stone, T.A. and Schlesinger, P. 1993. Digitization of the Map "Vegetation of the Soviet Union,
  1990." A Report to the Northeast Forest Experiment Station, USDA Forest Service,
  Global Change Research Program, Radnor, Pennsylvania. See companion file
  http://daac.ornl.gov/daacdata/russian\_land\_cover/vegetation\_1990/comp/vmap90\_
  method.pdf.
- Stone, T.A. and Schlesinger, P. 2003. RLC Vegetative Cover of the Former Soviet Union, 1990.
  Data set. Available on-line [http://www.daac.ornl.gov] from Oak Ridge National
  Laboratory Distributed Active Archive Center, Oak Ridge, Tennessee, U.S.A.
  doi:10.3334/ORNLDAAC/700.
- Stott, P. 1988. The Forest as Phoenix: Towards a Biogeography of Fire in Mainland South East
  Asia. The Geographical Journal, 154, 337-350.
- Tang Z, Fang J, Chi X, Yang Y, Ma W, Mohhamot A, Guo Z, Liu Y, Gaston KJ. (2012) Geography,
  environment, and spatial turnover of species in China's grasslands. Ecography
  35:1103–1109. doi: 10.1111/j.1600-0587.2012.07713.x
- White, F. 1983. The vegetation of Africa, a descriptive memoir to accompany the
   UNESCO/AETFAT/UNSO vegetation map of Africa.
- Wilson, B. A., Brocklehurst, P. S., Clark, M. J. & Dickinson, K. J. M. 1990. Vegetation survey of
   the Northern Territory, Australia. Darwin: Conservation Commission of the Northern
   Territory.

### 824 Plot data references for validation of map

- ABARES. (2012) Ground Cover Reference Sites Database, Version 9 /2014. ÆKOS Data Portal
   (http://www.portal.aekos.org.au/). Accessed: 16 July 2014.
- Abule E, Smit GN, Snyman HA. (2005) The influence of woody plants and livestock grazing on
- 828 grass species composition, yield and soil nutrients in the Middle Awash Valley of
- 829 Ethiopia. *J Arid Environ* 60:343–358. doi:10.1016/j.jaridenv.2004.04.006.
- Alberti J, Canepuccia A, Pascual J, Pérez C, Iribarne O. (2011) Joint control by rodent
- herbivory and nutrient availability of plant diversity in a salt marsh–salty steppe
  transition zone. *J Veg Sci* 22:216–224. doi:10.1111/j.1654-1103.2010.01240.x.
- 833 Alhamad MN, Alrababah MA, Gharaibeh MA (2012) Impact of burning and fertilization on
- dry Mediterranean grassland productivity and diversity. *Acta Oecol* 40:19–26.
  doi:10.1016/j.actao.2012.02.005.
- Ao M, Ito M, Ito K, Yun JF, Miura R, Tominaga T. (2008) Floristic compositions of Inner
- 837 Mongolian grasslands under different land-use conditions. *Grassl Sci* 54:173–178.
  838 doi:10.1111/j.1744-697X.2008.00121.x.
- Asebrook J. (2003) Grasslands of Glacier National Park. (GRAS) National Park Service
- 840 *Mapping Project in Montana.* http://vegbank.org/cite/VB.Ob.6325.GRAS00005.
- 841 Accessed: 09 January 2013.
- Banyikwa FF, Feoli E, Zuccarello V. (1990) Fuzzy set ordination and classification of Serengeti
  short grasslands, Tanzania. *J Veg Sci* 1:97–104.
- Barnes DL, Rethman NFG, Beukes BH, Kotzé GD. (1984) Veld composition in relation to
  grazing capacity. *J Grassl Soc Southern Afr* 1:16–19.
- 846 doi:10.1080/02566702.1984.9647960.

847 Beckley A. (2012). Honey Hill Vegetation Data.

848 http://vegbank.org/cite/VB.ob.78830.126010504. Accessed: 09 January 2013.

- 849 Belachew G, Demissew S, Woldu Z. (2012) Floristic Composition and Structure of Riverine
- 850 *Vegetation. The Case of Beschillo and Abay (Blue Nile).* Lambert Academic Publishing.
- 851 Bell J, Cogan D, Erixson J, Von Loh J. (2009) Vegetation Inventory Project Report. Craters of
- 852 the Moon National Monument and Preserve. Natural Resource Technical Report
- 853 NPS/UCBN/NRTR—2009/277. National Park Service, Fort Collins, CO.

- 854 Belsky AJ. (1992) Effects of grazing, competition, disturbance and fire on species composition 855 and diversity in grassland communities. J Veg Sci 3:187–200. doi:10.2307/3235679. Berliner D, Kioko J. (1999) The effect of long-term mowing and ungulate exclusion on grass 856 857 species composition and soil nutrient status on the Athi-Kapiti Plains, Central Kenya. 858 *Afr J Range Forage Sci* 16:62–70. doi:10.2989/10220119909485720. 859 Bonyongo MC, Veenendaal E, Bredenkamp GJ. (2000) Floodplain vegetation in the Nxaraga 860 Lagoon Area, Okavango Delta, Botswana. S Afr J Bot 66:15–21. 861 Boyer M, Dellinger B. (2012) Longleaf – Southeastern NC Vegetation Data. 862 http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:1054-%7BF477A5F3-863 D04B-44AC-ADDD-6B15EB709D8F%7D. Accessed: 09/01/2013. Brockett B. (2001) Sampling efficiency for species composition assessments using the wheel-864 865 point method in a semi-arid savanna. *Afr J Range Forage Sci* 18:93–101. doi:10.2989/10220110109485761. 866 867 Brookman-Amissah J, Hall JB, Swaine MD, Attakorah JY. (1980) A re-assessment of a fire 868 protection experiment in north-eastern Ghana savanna. J Appl Ecol 17:85–99. 869 doi:10.2307/2402965. Burns CE, Collins SL, Smith MD. (2009) Plant community response to loss of large herbivores: 870 871 Comparing consequences in a South African and a North American grassland. Biodivers Conserv 18:2327-2342. doi:10.1007/s10531-009-9590-x. 872 Carr S. (2007) Floristic and Environmental Variation of Pyrogenic Pinelands in the 873 Southeastern Coastal Plain: Description, Classification, and Restoration. PhD Thesis, 874 875 University of Florida, Gainesville, FL. 876 Carr SC, Robertson KM, Peet RK. (2010) A vegetation classification of fire-dependent 877 pinelands of Florida. Castanea 75:153-189.
- 878 Carter AJ, O'Connor TG. (1991) A two-phase mosaic in a savanna grassland. *J Veg Sci* 2:231–
  879 236.
- 880 Cecil F (2012) Longleaf Fall-line Sandhills Vegetation Data.
- 881 http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:792-%7BB0F4BECC-
- 882 1771-4A27-89A5-8BFFFD26D232%7D. Accessed: 09/01/2013.
- Chastain RA, Struckhoff MA, Grabner KW, Stroh ED, He H, Larsen DR, Nigh TA, Drake J. (2006)
   Mapping Vegetation Communities in Ozark National Scenic Riverways. Final Technical

885 Report to the National Park Service: U.S. Geological Survey Open-File Report 2006-

886 1354.

Cogan D. (2007) Vegetation Classification and Mapping Project Report, Lyndon B. Johnson
 National Historical Park. Natural Resource Technical Report NPS/SOPN/NRTR—

889 2007/073. National Park Service, Fort Collins, CO.

- 890 Cogan D. (2007) Vegetation Classification and Mapping Project Report, San Antonio Missions
- 891 National Historical Park. Natural Resource Technical Report NPS/GULN/NRTR—
- 892 2007/074. National Park Service, Fort Collins, CO.
- Cogan D. (2007) Vegetation Classification and Mapping Project Report, Washita Battlefield
   National Historic Site. Natural Resource Technical Report NPS/SOPN/NRTR—
- 895 2007/075. National Park Service, Fort Collins, CO.
- 896 Cogan D, Varga K, Kittel G, McCloskey K, Gremer J, Abendroth D, Bolen C. (2005) USGS-NPS-
- 897 USGS Vegetation Mapping Program: Grand Teton National Park 2002–2005.

898 Vegetation Mapping Project Report. U.S. Geological Survey.

- Cogan D, Walker L, Loring H, Jog S, Delisle J. (2007) *Vegetation Classification and Mapping Project Report. Fort Larned National Historic Site.* Natural Resource Technical Report
   NPS/SOPN/NRTR—2007/072. National Park Service, Fort Collins, CO.
- 902 Cogan D, Reid M, Schulz K, Pucherelli M. (2003) USGS-NPS Vegetation Mapping Program:
   903 Zion National Park, Utah. Project Report. U.S. Geological Survey.
- 904 Cogan D, Marriot H, Von Loh J, Pucherelli MJ. (1999) USGS-NPS Vegetation Mapping
- 905 *Program: Wind Cave National Park, South Dakota*. Project Report. U.S. Geological
  906 Survey.
- Dad JM, Khan AB. (2010) Floristic composition of an alpine grassland in Bandipora, Kashmir.
   *Grassl Sci* 56:87–94. doi:10.1111/j.1744-697X.2010.00179.x.

Daly C, Halbleib M, Smith JI, Gibson WP, Doggett MK, Taylor GH. (2008) Physiographically

- 910 sensitive mapping of climatological temperature and precipitation across the
- 911 conterminous United States. *Int J Climatol* 28:2031–2064.
- de Abreu RCR, Durigan G. (2011) Changes in the plant community of a Brazilian grassland
   savannah after 22 years of invasion by *Pinus elliottii* Engelm. *Plant Ecol Divers* 4:269–
- 914 278. doi:10.1080/17550874.2011.594101.

- 915 De Sanctis M, Adeeb A, Farcomeni A, Patriarca C, Saed A, Attorre F. (2012) Classification and
- 916 distribution patterns of plant communities on Socotra Island, Yemen. Appl Veg Sci

917 16:148–165. doi:10.1111/j.1654-109X.2012.01212.x.

- 918 Department of Environment, Water and Natural Resources. (2012) *Biological Survey of South* 919 Australia. Vegetation Survey, *Biological Database of South Australia, Version 11*
- 920 /2014. ÆKOS Data Portal (http://www.portal.aekos.org.au/). Accessed: 16 July 2014.
- 921 Department of Parks and Wildlife, Biogeography Program (2012) Biological Survey of the
- 922 Ravensthorpe Range (Phase 1), Version 1 /2013. ÆKOS Data Portal

923 (http://www.portal.aekos.org.au/). Accessed: 16 July 2014.

924 Department of Primary Industries, Parks, Water and Environment. (2015) *Platypus Survey* 

925 Data, Tasmania (1901–2009), Version 4 /2015. ÆKOS Data Portal

926 (http://www.aekos.org.au/home). Accessed: 16 July 2014.

927 Deshmukh I. (1986) Primary production of a grassland in Nairobi National Park, Kenya. J Appl
 928 Ecol 23:115–123.

Dhaou SO, Abdallah F, Belgacem, AO, Chaieb M. (2010) The protection effect on floristic
diversity in a North African pseudo-savanna. *Pak J Bot* 42:1501–1510.

931 Downing BH, Robinson ER, Trollope WSW, Morris JM. (1978) Influence of macchia

- 932 eradication techniques on botanical composition of grasses in the Döhne Sourveld of
- 933 the Amatole Mountains. *Proc Annual Congr Grassl Soc Southern Afr* 13:111–115.
- 934 doi:10.1080/00725560.1978.9648844.
- 935 du Toit PF, Aucamp AJ. (1985) Effect of continuous grazing in the Döhne Sourveld on species
  936 composition and basal cover. *J Grassl Soc Southern Afr* 2:41–45.

937 doi:10.1080/02566702.1985.9648018.

- 938 Dunham KM. (1989) Vegetation-environment relations of a Middle Zambezi floodplain. *Plant*939 *Ecol* 82:13–24.
- El-Ghareeb RM. (1991) Suppression of annuals by *Tribulus terrestris* in an abandoned field in
  the sandy desert of Kuwait. *J Veg Sci* 2: 147–154.
- Ellery WN, Ellery K, Rogers KH, McCarthy TS, Walker BH. (1990) Vegetation of channels of the
  northeastern Okavango Delta, Botswana. *Afr J Ecol* 28:276–290. doi:10.1111/j.13652028.1990.tb01162.x.

945	Enright NJ, Miller BP, Akhter R. (2005) Desert vegetation and vegetation-environment
946	relationships in Kirthar National Park, Sindh, Pakistan. J Arid Environ 61:397–418.
947	Erixson JA, Cogan D. (2009) Vegetation Classification and Mapping of Hagerman Fossil Beds
948	National Monument. Natural Resource Technical Report NPS/UCBN/NRTR—
949	2009/212. National Park Service, Fort Collins, CO.
950	Erixson JA, Cogan D, Von Loh J. (2011) Vegetation Inventory Project Report: John Day Fossil
951	Beds National Monument. Natural Resource Technical Report NPS/UCBN/NRTR—
952	2011/419. National Park Service, Fort Collins, CO.
953	Erixson JA, Cogan D, Von Loh J. (2011) Vegetation Inventory Project Report: Lake Roosevelt
954	National Recreation Area. Natural Resource Report NPS/UCBN/NRR-2011/434
955	National Park Service, Fort Collins, CO.
956	Evans R, Pyne M. (2012) Angelina National Forest Vegetation Data.
957	http://vegbank.org/cite/VB.ob.26846.ANGE10. Accessed: 09 January 2013.
958	Evans R, Wiseland R. (2012) Mississippi Vegetation Survey Vegetation Data.
959	http://vegbank.org/cite/VB.ob.26967.MSVS4. Accessed: 09 January 2013.
960	Evans R, Teague J. (2012) Sam Houston National Forest Vegetation Data.
961	http://vegbank.org/cite/VB.ob.27063.SAMH13. Accessed: 09 January 2013.
962	Farris, E, Filighedd, R, Deiana P, Farris GA, Garau G. (2010) Short-term effects on sheep
963	pastureland due to grazing abandonment in a Western Mediterranean island
964	ecosystem: A multidisciplinary approach. J Nat Conserv 18:258–267.
965	doi:10.1016/j.jnc.2009.11.003.
966	Felfili JM, Filgueiras TS, Haridasan M, Silva Júnior MC, Mendonça RC, Resende A. (1994)
967	Projeto biogeografia do bioma cerrado: vegetação e solos. Cadernos de Geociêcias
968	12:75–166.
969	Fenton K, Bell JR, Wegner D. (2006) USGS-NPS Vegetation Mapping Program: Lake Meredith
970	National Recreation Area and Alibates Flint Quarries National Monument, Texas.
971	Project Report. U.S. Geological Survey.
972	Ferreira PMA, Müller SC, Boldrini II, Eggers L. (2010) Floristic and vegetation structure of a
973	granitic grassland in southern Brazil. Rev Brasil Bot 33:21–36.

974	Fidelis A. (2008) Fire in Subtropical Grasslands in Southern Brazil: Effects on Plant Strategies
975	and Vegetation Dynamics. Technische Universität München, Wissenschaftszentrum
976	Weihenstephan. http://d-nb.info/991968948/34.
977	Forrestel EJ. (2013) Vegetation Cover Data from 10 Sites in the Great Plains, North America.
978	Unpublished data.
979	Gaddy C. (2012) Congaree Swamp National Monument Vegetation Data.
980	http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:4538-%7BA7C85EE9-
981	4726-401A-B482-82254469D858%7D. Accessed: 09 January 2013.
982	Galal TM, Fahmy AG. (2012) Plant diversity and community structure of Wadi Gimal
983	Protected Area, Red Sea coast of Egypt. <i>Afr J Ecol</i> 50:266–276.
984	http://onlinelibrary.wiley.com/doi/10.1111/j.1365-2028.2012.01320.x/full.
985	Gallimore J. (2012) McCall Outdoor Science School Vegetation Data.
986	http://vegbank.org/cite/VB.ob.78625.JG01. Accessed: 09 January 2013.
987	Glitzenstein J. (2012) St. Francis National Forest Vegetation Data.
988	http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:8996-{A6D80FD4-5777-
989	4B17-A778-08703107E334}. Accessed: 09 January 2013.
990	Grace P. (2010) A Vegetation Survey of the Samford Ecological Research Facility (SERF).
991	https://researchdata.ands.org.au/vegetation-survey-samford-facility-serf/663523.
992	Hall M. (2012) Short Mountain Wildlife Management Area Vegetation Data.
993	http://vegbank.org/cite/VB.ob.27561.SHMO14. Accessed: 09 January 2013.
994	Harkel MJ, van der Meulen F. (1996) Impact of grazing and atmospheric nitrogen deposition
995	on the vegetation of dry coastal dune grasslands. <i>J Veg Sci</i> 7: 445–452.
996	Hayashi I, Kawada K, Akimova A, Nakamura T. (2006) Floristic composition and plant biomass
997	of the grasslands in the vicinity of Pavlodar, Kazakhstan. <i>Grassl Sci</i> 52:141–146.
998	doi:10.1111/j.1744-697X.2006.00059.x.
999	Hejcmanová P, Hejcman M, Camara AA, Antonínová M. (2010) Exclusion of livestock grazing
1000	and wood collection in dryland savannah: An effect on long-term vegetation
1001	succession. Afr J Ecol 48:408–417.
1002	Hop K, Faber-Langendoen D, Lew-Smith M, Aaseng N, Lubinski S. (2001) USGS-NPS
1003	Vegetation Mapping Program: Voyageurs National Park. Project Report. U.S.
1004	Geological Survey.

1005 Hop K, Drake J, Lubinski S, Dieck J, Menard S. (2009) National Park Service Vegetation 1006 Inventory Program: Indiana Dunes National Lakeshore, Indiana. U.S. Geological 1007 Survey, Upper Midwest Environmental Sciences Center, La Crosse, WI. 1008 Hop K, Drake J, Lubinski S, Menard S, Dieck J. (2012) National Park Service Vegetation 1009 Inventory Program: Saint Croix National Scenic Riverway, Minnesota/Wisconsin. 1010 Natural Resource Report NPS/GLKN/NRR—2012/547. National Park Service, Fort 1011 Collins, CO. 1012 Hop K, Pyne M, Foti T, Lubinski S, White R, Dieck J. (2012) National Park Service Vegetation 1013 Inventory Program: Buffalo National River, Arkansas. Natural Resource Report 1014 NPS/HTLN/NRR—2012/526. National Park Service, Fort Collins, CO. Hop K, Reid M, Dieck J, Lubinski S, Cooper S. (2007) USGS-NPS Vegetation Mapping Program: 1015 1016 Waterton-Glacier International Peace Park. U.S. Geological Survey, Upper Midwest Environmental Sciences Center, La Crosse, WI. 1017 Hop K, Lubinski S, Menard S. (2005) USGS-NPS Vegetation Mapping Program: Effiqy Mounds 1018 1019 National Monument, Iowa. U.S. Geological Survey, Upper Midwest Environmental 1020 Sciences Center, La Crosse, WI. Hop K, Menard S, Drake J, Lubinski S, Dieck J. (2010) National Park Service Vegetation 1021 1022 Inventory Program: Apostle Islands National Lakeshore, Wisconsin. Natural Resource 1023 Report NPS/GLKN/NRR—2010/199. National Park Service, Fort Collins, CO. Hop K, Menard S, Drake J, Lubinski S, Dieck J. (2010) National Park Service Vegetation 1024 1025 Inventory Program: Pictured Rocks National Lakeshore, Michigan. Natural Resource 1026 Report NPS/GLKN/NRR—2010/201. National Park Service, Fort Collins, CO. 1027 Hop K, Menard S, Drake J, Lubinski S, Faber-Langendoen D, Dieck J. (2010) National Park 1028 Service Vegetation Inventory Program: Grand Portage National Monument, 1029 Minnesota. Natural Resource Report NPS/GLKN/NRR—2010/200. National Park 1030 Service, Fort Collins, CO. 1031 Jauffret S, Lavorel S. (2003) Are plant functional types relevant to describe degradation in 1032 arid, southern Tunisian steppes? J Veg Sci 14:399–408. 1033 Jennings M. (2012) Composition and Function of Vegetation Alliances in the Interior

1034 Northwest, USA. http://vegbank.org/cite/VB.Ob.10774.INW10717. Accessed: 09
1035 January 2013.

1036 Johnson A. (2012) Apalachicola National Forest Vegetation Data.

1037 http://vegbank.org/cite/VB.ob.26905.APAL74. Accessed: 09 January 2013.

- 1038 Joint Remote Sensing Research Program. (2013) AusCover Supersites SLATS Star Transects,
- 1039 Version 2 /2013. ÆKOS Data Portal (<u>http://www.portal.aekos.org.au/</u>). Accessed: 16
  1040 July 2014.
- Jones E, Pyne M. (2008) Vascular Plant Inventory and Plant Community Classification for
   Abraham Lincoln National Historic Site. NatureServe, Durham, NC.
- 1043 Kawada K, Vovk AG, Filatova OV, Araki M, Nakamura T, Hayashi I. (2005) Floristic
- 1044composition and plant biomass production of steppe communities in the vicinity of1045Kharkiv, Ukraine. *Grassl Sci* 51:205–213. doi:10.1111/j.1744-697X.2005.00026.x.
- 1046 Keeler-Wolf T, Moore PE, Reyes ET, Menke JM, Johnson DN, Karavidas DL. (2012) *Yosemite*
- 1047 National Park Vegetation Classification and Mapping Project Report. Natural
- 1048Resource Technical Report NPS/YOSE/NRTR—2012/598. National Park Service, Fort1049Collins, CO.
- Kelly M, Allen-Diaz B, Kobzina N. (2005) Digitization of a historic dataset: The Wieslander
   California Vegetation Type Mapping Project. *Madroño* 52:191–201.
- 1052 Kindscher K, Kilroy H, Delisle J, Long Q, Loring H, Dobbs K, Drake J. (2011a) Vegetation
   1053 Mapping and Classification of Homestead National Monument of America. Natural
- 1054 Resource Report NPS/HTLN/NRR—2011/345. National Park Service, Fort Collins, CO.
- 1055 Kindscher K, Kilroy H, Delisle J, Long Q, Loring H, Dobbs K, Drake J. (2011b) Vegetation
- Mapping and Classification of Tallgrass Prairie National Preserve. Natural Resource
   Report NRR/HTLN/NRR—2011/346. National Park Service, Fort Collins, CO.
- Kittel G, Reyes E, Evens J, Buck J, Johnson D. (2012) *Vegetation Classification and Mapping Project Report. Pinnacles National Monument.* Natural Resource Report
   NPS/SFAN/NRR—2012/574. National Park Service, Fort Collins, CO.
- 1061 Kittel G, VanWie E, Damm M, Rondeau R, Kettler S, McMullen A, Sanderson J. (1999) A
- 1062 Classification of Riparian Wetland Plant Associations of Colorado: User Guide to the
   1063 Classification Project. Colorado Natural Heritage Program, Colorado State University,
   1064 Fort Collins, CO.
- 1065 Klopfer SD, Olivero A, Sneddon L, Lundgren J. (2002) USGS-NPS Vegetation Mapping Project
   1066 at Fire Island National Seashore. Project Report. U.S. Geological Survey.

1067 Koonamore Research Group. (2015) Kangaroo Transects. Koonamore Vegetation Monitoring 1068 Project (1925–present), Version 12 /2014. ÆKOS Data Portal 1069 (http://www.aekos.org.au/home). Accessed: 16 July 2014. 1070 Koonamore Research Group. (2015) Photopoints. Koonamore Vegetation Monitoring Project (1925–Present), Version 12 /2014. ÆKOS Data Portal 1071 1072 (http://www.aekos.org.au/home). Accessed: 16 July 2014. 1073 Koonamore Research Group. (2015) Myoporum platycarpum Survey. Koonamore Vegetation 1074 Monitoring Project (1925–Present), Version 12 /2014. ÆKOS Data Portal 1075 (http://www.aekos.org.au/home). Accessed: 16 July 2014. 1076 Koonamore Research Group. (2015) Rabbit Activity Monitoring and Control. Koonamore Vegetation Monitoring Project (1925–Present), Version 12 /2014. ÆKOS Data Portal 1077 1078 (http://www.aekos.org.au/home). Accessed 16 July 2014. Koonamore Research Group. (2015) Saltbush Transects. Koonamore Vegetation Monitoring 1079 Project (1925Present), Version 12 /2014. ÆKOS Data Portal 1080 1081 (http://www.aekos.org.au/home). Accessed: 16 July 2014. 1082 Koonamore Research Group. (2015) Senna Populations. Koonamore Vegetation Monitoring Project (1925–Present), Version 12 /2014. ÆKOS Data Portal 1083 1084 (http://www.aekos.org.au/home). Accessed: 16 July 2014. Koonamore Research Group. (2015) Senna Quadrat (Cassia Corner). Koonamore Vegetation 1085 1086 Monitoring Project (1925–Present), Version 12 /2014. ÆKOS Data Portal 1087 (http://www.aekos.org.au/home. Accessed: 16 July 2014. 1088 Koonamore Research Group. (2015). Vegetation Quadrats. Koonamore Vegetation 1089 Monitoring Project (1925–Present), Version 12 /2014. ÆKOS Data Portal (http://www.aekos.org.au/home). Accessed: 16 July 2014. 1090 1091 Kotze DC, O'Connor TG. (2000) Vegetation variation within and among palustrine wetlands 1092 along an altitudinal gradient in KwaZulu-Natal, South Africa. Plant Ecol 146:77–96. 1093 Krueger E, Nordman C. (2012) Francis Marion National Forest Vegetation Data. 1094 http://vegbank.org/cite/VB.ob.78771.068FRMA16. Accessed: 09 January 2013. 1095 Kunz D, Baker G, Peet R. (2012) Croatan National Forest Vegetation Data. http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:7642-{EB625604-8404-1096 1097 4635-8B89-507E699C31E8}. Accessed: 09 January 2013.

- 1098 Kwatrina RT. (2009) Penentuan Kuota Panenan Dan Ukuran Populasi Awal Rusa Timor Di
- 1099 Penangkaran Hutan Penelitian Dramaga.
- 1100 http://repository.ipb.ac.id/handle/123456789/5368.
- 1101 Lenzi-Grillini CR, Viskanic P, Mapesa M. (1996) Effects of 20 years of grazing exclusion in an
- area of the Queen Elizabeth National Park, Uganda. *Afr J Ecol* 34:333–341.
- 1103 doi:10.1111/j.1365-2028.1996.tb00629.x.
- 1104 Lubinski S, Hop K, Grawler S. (2003) USGS-NPS Vegetation Mapping Program: Acadia
- 1105 *National Park, Maine.* Project Report. U.S. Geological Survey.
- Madden M, Welch R, Jordan T, Jackson P, Seavey R, Seavey J. (2004) *Digital Vegetation Maps for the Great Smoky Mountains National Park.* Project Report.
- 1108 Marriot H, Drake J, Curtis A, Grossman D. (1998) USGS-NPS Vegetation Mapping Program:
- 1109 Classification of the vegetation of Fort Laramie National Historic Site. The Nature
- 1110 Conservancy, Minneapolis, MN.
- 1111 Marriot H, McAdams A, Stutzman D, Drake J, Grossman D. (1998) USGS-NPS Vegetation
- 1112 Mapping Program: Classification of the Vegetation of Mount Rushmore National
  1113 Memorial. Project Report. U.S. Geological Survey.
- Marriot H, McAdams A, Stutzman D, Drake J, Grossman D. (2012a) USGS-NPS Vegetation
   Mapping Program: Classification of the Vegetation of Jewel Cave National
- 1116 *Monument.* Project Report. U.S. Geological Survey.
- 1117 Marriot H, McAdams A, Stutzman D, Drake J, Grossman D. (2012b) USGS-NPS Vegetation
- 1118 Mapping Program: Classification of the Vegetation of Devils Tower National
- 1119 *Monument.* Project Report. U.S. Geological Survey.
- 1120 Matthews L. (2012) Cape Fear Riparian Vegetation Data.
- http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:7350-{46DDE438-83DC 4C25-8F03-07CFE23450EE}. Accessed: 09/01/2013.
- 1123 Matthews L. (2012) Yadkin-Pee Dee & Catawba Riparian Vegetation Data.
- 1124 http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:8207-%7B52904CDF-
- 1125 10AC-486D-A492-8CF16BC51CD9%7D. Accessed: 09/01/2013.
- 1126 McKenzie B. (1987) Composition, pattern and diversity of some Transkeian grasslands. J
- 1127 *Grassl Soc Southern Afr* 4:135–138. doi:10.1080/02566702.1987.9648091.

1128 McMillan P, Kjellmark E. (2012) Longleaf – Coastal SC & GA Vegetation Data.

- 1129 http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:2708-%7B72B2B40D-
- 1130 41FE-4436-ABE5-CE4B753C2898%7D. Accessed: 09 January 2013.
- 1131 Michiels B, Babatounde S, Dahouda M, Chabi SLW, Buldgen A. (2000) Botanical composition
- and nutritive value of forage consumed by sheep during the rainy season in a
- 1133 Sudano-Guinean savanna (central Benin). *Trop Grassl Soc* 34:43–47.
- 1134 Miehe G, Bach K, Miehe S, Kluge J, Yang Y, Duo L, Co S, Wesche K. (2011) Alpine steppe plant
- 1135 communities of the Tibetan Highlands. *Appl Veg Sci* 14:547–560. doi:10.1111/j.1654109X.2011.01147.x.
- 1137 Morgan JW. (1998) Composition and seasonal flux of the soil seed bank of species-rich
- 1138 *Themeda triandra* grasslands in relation to burning history. *J Veg Sci* 9:145–156.
- 1139 Moyo CS, Campbell BM. (1998) Grass species composition, yield and quality under and
- outside tree crowns in a semi-arid rangeland in south-western Zimbabwe. *Afr J Range Forage Sci* 15:23–34. doi:10.1080/10220119.1998.9647937.
- 1142 Mucina L, Rutherford MC. (eds) (2010) *The Vegetation of South Africa, Lesotho and*
- 1143 *Swaziland*. South African National Biodiversity Institute, Pretoria, ZA. (CD Set).
- 1144 Mucina L, Rutherford MC. (eds) (2006) *The Vegetation of South Africa, Lesotho and*
- 1145 *Swaziland.* South African National Biodiversity Institute, Pretoria, ZA.
- Mucina L, Rutherford MC, Powrie LW. (eds) (2007) *Vegetation Map of South Africa, Lesotho and Swaziland, edn 2. 1:1 000 000 Scale Sheet Maps.* South African National
  Biodiversity Institute, Pretoria, ZA.
- 1149 Mucina L, Rutherford MC, Powrie LW, van Niekerk A, van der Merwe JH. (eds) (2014)
- 1150 Vegetation Field Atlas of Continental South Africa, Lesotho and Swaziland. South
- 1151 African National Biodiversity Institute, Pretoria, ZA.
- 1152 Müller SC, Waechter JL. (2001) Estrutura sinusial dos componentes herbáceo e arbustivo de
  1153 uma floresta costeira subtropical. *Rev Brasil Bot* 24:395–406.
- 1154 Nachtergaele F, Batjes N. (2012) Harmonized World Soil Database. FAO, Rome, IT.
- 1155 NatureServe. (2004) International Ecological Classification Standard: Terrestrial Ecological
   1156 Classifications. NatureServe, Arlington, VA.

- 1157 Neid S, Stevens JE, Forest K, Fink M. (2007) Sand Creek Massacre National Historic Site:
- 1158 Vegetation Classification and Mapping. Natural Resource Technical Report
- 1159 NPS/SOPN/NRTR—2007/050. National Park Service, Fort Collins, CO.
- 1160 New South Wales Office of Environment and Heritage. (2014) Data from the Atlas of NSW
- 1161 Database: VIS Flora Survey Module, Version 11 /2013. ÆKOS Data Portal
- 1162 (http://www.portal.aekos.org.au/). Accessed: 16 July 2014.
- 1163 Ngwenya P. (2012) Herbaceous plant species richness and composition in Moist Midlands
- 1164 Mistbelt Grasslands in KwaZulu-Natal: Is there a relationship to veld condition? *Afr J* 1165 *Range Forage Sci* 29:75–83. doi:10.2989/10220119.2012.705324.
- 1166 Noor Alhamad M. (2006) Ecological and species diversity of arid Mediterranean grazing land
- 1167 vegetation. J Arid Environ 66:698–715.
- 1168 Nordman C. (2012) Holly Springs National Forest Vegetation Data.
- 1169 http://vegbank.org/cite/VB.ob.27092.HOLL1. Accessed: 09 January 2013.
- 1170 Nordman C, Anglin G. (2012) Osceola National Forest Vegetation Data.
- 1171 http://vegbank.org/cite/VB.ob.27002.OSCE27. Accessed: 09 January 2013.
- 1172 Nordman C, Schultz G. (2012) Osceola National Forest Vegetation Data.
- 1173 http://vegbank.org/cite/VB.ob.26974.OCAL32. Accessed: 09 January 2013.
- 1174 Nordman C, Pierce R. (2012) Delta National Forest Vegetation Data.
- 1175 http://vegbank.org/cite/VB.ob.27106.DELT9. Accessed: 09 January 2013.
- O'Connor TG. (1994) Composition and population responses of an African aavanna grassland
   to rainfall and grazing. *J Appl Ecol* 31:155–171.
- 1178 O'Connor TG, Martindale G, Morris CD, Short A, Witkowski ETF, Scott-Shaw R. (2011)
- 1179 Influence of grazing management on plant diversity of Highland Sourveld Grassland,
- 1180 KwaZulu-Natal, South Africa. *Rangel Ecol Manage* 64:196–207. doi:10.2111/REM-D1181 10-00062.1.
- 1182 Ode D. (2012) LaFramboise Island Management Plan Vegetation Data.
- 1183 http://vegbank.org/cite/VB.ob.28253.LFIPLOT1. Accessed: 09 January 2013.
- Pandey CB, Singh JS. (1991) Influence of grazing and soil conditions on secondary savanna
  vegetation in India. *J Veg Sci* 2:95–102.

- 1186 Parsons DAB, Shackleton CM, Scholes RJ. (1997) Changes in herbaceous layer condition
- 1187 under contrasting land use systems in the semi-arid Lowveld, South Africa. J Arid
- 1188 Environ 37:319–329. doi:10.1006/jare.1997.0283.
- 1189 Patterson K, Pyne M. (2012) Bankhead National Forest Vegetation Data.

1190 http://vegbank.org/cite/VB.ob.26740.BANK24. Accessed: 09 January 2013.

1191 Pattiselanno F, Arobaya AYS. (2009) Grazing habitat of the Rusa Deer (*Cervus timorensis*) in

the Upland Kebar, Manokwari. *Biodiversitas* 10:134–138.

- Peet R, Lee M, Boyle F, Wentworth T, Schafale M, Weakley A. (2012) Vegetation-plot
  database of the Carolina Vegetation Survey. *Biodivers Ecol* 4:243–253.
- 1195 Peet RK. (1975) Forest Vegetation of the East Slope of the Northern Colorado Front Range.
- 1196 PhD thesis, Cornell University, Ithaca, NY.
- 1197 Peet RK. (2012) Maritime Forests Vegetation Data.
- 1198 http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:714-%7B5C325A03-
- 1199 C818-430F-B4C9-EFBED2272AF0%7D. Accessed: 09 January 2013.
- Peet RK, Lee MT, Jennings MD, Faber-Langendoen D. (2012) VegBank A permanent, open access archive for vegetation-plot data. *Biodivers Ecol* 4:233–241
- Peet RK, Lee MT, Jennings MD, Faber-Langendoen D. (eds) (2013) *VegBank: The vegetation plot archive of the Ecological Society of America*. Accessed: 09 January 2013.
- Podniesinski GS, Sneddon LA, Lundgren J, Devine H, Slocumb B, Koch F. (2005) *Vegetation classification and mapping of Valley Forge National Historical Park*. Technical Report
- 1206 NPS/NER/NRTR—2005/028. National Park Service, Philadelphia, PA.
- 1207 Poilecot P, Gaidet N. (2011) A quantitative study of the grass and woody layers of a mopane
- 1208 (Colophospermum mopane) savannah in the mid-Zambezi Valley, Zimbabwe. Afr J
   1209 Ecol 49:150–164.
- Powrie L, Rutherford MC, Mucina L, Mangwale K. (2012) National Vegetation Database of
  South Africa. *Biodivers Ecol* 4:299–299.
- 1212 Pyne M, Schmidt J. (2012) Oconee National Forest Vegetation Data.
- 1213 http://vegbank.org/cite/VB.ob.26571.OCON46. Accessed: 09 January 2013.
- 1214 Pyne M, Evans R. (2012) Sabine National Forest Vegetation Data.
- 1215 http://vegbank.org/cite/VB.ob.26862.SABI8. Accessed: 09 January 2013.

1216 Pyne M, Stewart R. (2012) Tuskegee National Forest Vegetation Data. 1217 http://vegbank.org/cite/VB.ob.26949.TUSK3. Accessed: 09 January 2013. Pyne M, Weakley A, Donaldson J. (2012) Cherokee National Forest Vegetation Data. 1218 1219 http://vegbank.org/cite/VB.ob.26485.CHER29. Accessed: 09 January 2013. Pyne M, Evans R, Spaulding D, Garland B. (2012) Talladega National Forest Vegetation Data. 1220 1221 http://vegbank.org/cite/VB.ob.26732.TALT38. Accessed: 09 January 2013. 1222 Queensland Herbarium. (2012) Queensland CORVEG Database, Version 9 /2013. ÆKOS Data 1223 Portal (http://www.portal.aekos.org.au. Accessed: 16 July 2014. 1224 Reemts C. (2012) Fort Hood Vegetation Map. 1225 http://vegbank.org/cite/VB.ob.27632.FTHD225. Accessed: 09 January 2013. 1226 Regass, T. (2005) An Ecological Study of Vegetation Around Lake Abijata. PhD Thesis, Addis 1227 Ababa University, Addis Ababa, ET. http://localhost:80/xmlui/handle/123456789/8502 1228 1229 Reis AMS., Araújo EL, Ferraz EM, Moura AN. (2006) Inter-annual variations in the floristic and 1230 population structure of an herbaceous community of 'caatinga' vegetation in 1231 Pernambuco, Brazil. Rev Brasil Bot 29:497–508. 1232 Reschke C, Reid R, Jones J, Feeney T, Potter H. (1999) Conserving Great Lakes Alvars: Final 1233 Technical Report of the International Alvar Conservation Initiative. The Nature 1234 Conservancy, Chicago, IL. 1235 Rolfsmeier S, Drake J, Grossman D. (1998) USGS-NPS Vegetation Mapping Program: 1236 Classification of the Vegetation of Agate Fossil Beds National Monument. Project 1237 Report. U.S. Geological Survey. 1238 Rolfsmeier S, Steinauer G, Schneider R, Drake J, Aldrich J, Faber-Langendoen D, Goodin K, 1239 Swinehart C, Grossman D. (1998) USGS-NPS/BRD Vegetation Mapping Program: 1240 *Classification of the Vegetation of Scotts Bluff National Monument.* Project Report. 1241 U.S. Geological Survey. Roy DP, Boschetti L, Justice CO, Ju J. (2008) The collection 5 MODIS burned area product-1242 1243 Global evaluation by comparison with the MODIS active fire product. Rem Sens 1244 Environ 112:3690-3707. 1245 Salama FM, Ahmed MK, El-Tayeh NA, Hammad SA. (2012) Vegetation analysis, phenological patterns and chorological affinities in Wadi Qena, Eastern Desert, Egypt. Afr J Ecol 1246

1247	50:193-
1248	204.http://onlinelibrary.wiley.com/doi/10.1111/j.13652028.2011.01313.x/full.
1249	Salas DE, Folts-Zettner T, Sanders RW, Drake J. (2010) Vegetation Classification and Mapping
1250	at Chickasaw National Recreation Area. National Park Service, Natural Resource
1251	Program Center Fort Collins, CO.
1252	Sasaki T, Okayasu T, Takeuchi T, Jamsran U, Jadambaa S. (2005) Patterns of floristic
1253	composition under different grazing intensities in Bulgan, South Gobi, Mongolia.
1254	<i>Grassl Sci</i> 51:235–242. doi:10.1111/j.1744-697X.2005.00029.x.
1255	Savadogo P, Sawadogo L, Tiveau D. (2007) Effects of grazing intensity and prescribed fire on
1256	soil physical and hydrological properties and pasture yield in the savanna woodlands
1257	of Burkina Faso. Agric Ecosyst Environ 118:80–92. doi:10.1016/j.agee.2006.05.002.
1258	Sawadogo L, Tiveau D, Nygård R. (2005) Influence of selective tree cutting, livestock and
1259	prescribed fire on herbaceous biomass in the savannah woodlands of Burkina Faso,
1260	West Africa. Agric Ecosyst Environ 105:335–345. doi:10.1016/j.agee.2004.02.004.
1261	Schiebout M. (2012) Pawnee Grassland Classification.
1262	http://vegbank.org/cite/VB.ob.78868.001SCHIEB43A. Accessed: 09 January 2013.
1263	Schirokauer D, Keeler-Wolf T, Meinke J, van der Leeden P. (2003) Plant Community
1264	Classification and Mapping Project: Point Reyes National Seashore, Golden Gate
1265	National Recreation Area. San Francisco Water Department Watershed Lands, Mount
1266	Tamalpais, Tomales Bay, and Samuel P. Taylor State Parks. Project Report.
1267	Schmidt M, Janssen T, Dressler S, Hahn-Hadjali K, Hien M, Konaté S, Lykke AM, Mahamane A,
1268	Sambou B, Sinsin B, Thiombiano A, Wittig R, Zizka G. (2012) The West African
1269	Vegetation Database. <i>Biodivers Ecol</i> 4:105–110.
1270	Scott, K, Setterfield S, Douglas M, Andersen A. (2010) Soil seed banks confer resilience to
1271	savanna grass-layer plants during seasonal disturbance. Acta Oecol 36:202–210.
1272	doi:10.1016/j.actao.2009.12.007.
1273	Sexton JO, Song X-P, Feng M, Noojipady P, Anand A, Huang C. (2013) Global, 30-m resolution
1274	continuous fields of tree cover: Landsat-based rescaling of MODIS vegetation
1275	continuous fields with lidar-based estimates of error. Int J Digit Earth 6:427–448.

1276 Seymour S. (2012) Piedmont Nonalluvials Vegetation Data.

- 1277 http://vegbank.org/cite/urn:lsid:cvs.bio.unc.edu:observation:8902-%7BEB5FEBB4-
- 1278 39FE-4565-BFA6-8DB75DF9B44E%7D. Accessed: 09 January 2013.
- 1279 Shelton L, Evans R. (2012) Davy Crockett National Forest Vegetation Data.
- 1280 http://vegbank.org/cite/VB.ob.27046.DAVY17. Accessed: 09 January 2013.
- 1281 Shonhiwa FF. (1998) The Effects of Land-use History on Plant Species Diversity and
- 1282 Abundance in Dambo Wetlands of Zimbabwe. PhD Thesis, West Virginia University,
- 1283 Morgantown, WV. http://wvuscholar.wvu.edu:8881/R/?func=dbin-jump-
- 1284 full&object\_id=6697
- 1285 Short AD, O'Connor TG, Hurt CR. (2003) Medium-term changes in grass composition and
- 1286 diversity of Highland Sourveld Grassland in the Southern Drakensberg in response to
- 1287 fire and grazing management. *Afr J Range Forage Sci* 20:1–10.
- 1288 doi:10.2989/10220110309485792.
- 1289 Sisay A, Baars RMT. (2002) Grass composition and rangeland condition of the major grazing
- areas in the Mid Rift Valley, Ethiopia. *Afr J Range Forage Sci* 19:161–166.
- doi:10.2989/10220110209485789.
- Sneddon L, Lundgren J, Crane E, Salmons S. (1998) USGS-NBS/NPS Vegetation Mapping
   *Program: Vegetation Classification of Rock Creek Park.* Project Report. U.S. Geological
   Survey.
- Soromessa T, Teketay D, Demissew S. (2004) Ecological study of the vegetation in Gamo Gofa
   Zone, Southern Ethiopia. *Trop Ecol* 45:209–221.
- 1297 Sugiyama S, Zabed HM, Okubo A. (2008) Relationships between soil microbial diversity and
- 1298 plant community structure in seminatural grasslands. *Grassl Sci* 54:117–124.
- doi:10.1111/j.1744-697X.2008.00113.x.
- Tanimoto T. (1981) Vegetation of the Alang-alang grassland and its succession in the Benakat
   District of South Sumatra Indonesia. *Bull For & For Prod Res Inst* 314:11–19.
- 1302 TERN AusPlots. (2013) AusPlots Rangelands, Version 6 /2014. ÆKOS Data Portal
- 1303 (http://www.aekos.org.au/). Accessed: 16 July 2014.
- 1304 TERN Australian Transect Network. (2013) South West Australian Transitional Transect
- 1305 (SWATT), Version 11 /2014. ÆKOS Data Portal (http://www.portal.aekos.org.au/).
- 1306 Accessed: 16 July 2014.

1307	TERN Australian Transects. (2013) Transect for Environmental Monitoring and Decision
1308	Making (TREND), Version 6 /2014. ÆKOS Data Portal (http://www.aekos.org.au/).
1309	Accessed: 16 July 2014.
1310	The Association for Biodiversity Information. (2001) International Classification of Ecological
1311	Communities: Terrestrial Vegetation. The Association for Biodiversity Information,
1312	Arlington, VA.
1313	The Nature Conservancy. (1999) USGS-NPS Vegetation Mapping Program: Classification of
1314	the Vegetation of Isle Royale National Park. Project Report. U.S. Geological Survey.
1315	Thompson J. (2012) New York Natural Heritage Program.
1316	http://vegbank.org/cite/VB.Ob.24379.MINNREGPLT47. Accessed: 09 January 2013.
1317	USGS National Gap Analysis Program. (2004) Southwest Regional Gap Analysis Project Field
1318	Sample Database. Version 1.1. RS/GIS Laboratory, College of Natural Resources, Utah
1319	State University.
1320	USGS. (2010) Sequoia and Kings Canyon National Parks Vegetation Mapping Project.
1321	http://www.usgs.gov/core_science_systems/csas/vip/parks/seki.html. Accessed: 09
1322	January 2013.
1323	Verlinden A, Dayot B. (2005) A comparison between indigenous environmental knowledge
1324	and a conventional vegetation analysis in North Central Namibia. J Arid Environ
1325	62:143–175.
1326	Von Loh J, Cogan D, Butler J, Faber-Langendoen D, Crawford D, Pucherelli MJ. (2000) USGS-
1327	NPS Vegetation Mapping Program: Theodore Roosevelt National Park, North Dakota.
1328	Project Report. U.S. Geological Survey.
1329	Von Loh J, Cogan D, Faber-Langendoen D, Crawford D, Pucherelli MJ. (1999) USGS-NPS
1330	Vegetation Mapping Program: Badlands National Park, South Dakota. Project Report.
1331	U.S. Geological Survey.
1332	Walton D. (2012) Shenandoah National Park Vegetation Data.
1333	http://vegbank.org/cite/VB.Ob.25621.SHNP530. Accessed: 09 January 2013.
1334	Weakley A, Patterson K. (2012) Chattahoochee National Forest Vegetation Data.
1335	http://vegbank.org/cite/VB.ob.26587.CHAT135. Accessed: 09 January 2013.
1336	Weakley A, Stewart R. (2012) Oakmulgee National Forest Vegetation Data.
1337	http://vegbank.org/cite/VB.ob.26642.TALO31. Accessed: 09 January 2013.

- 1338 Whelan KRT, Sudalter E, Patterson JM, Vargas RM, Atkinson AJ, Witcher B. (2009) The 2009
- 1339 Vegetation Map of De Soto National Memorial. Natural Resource Technical Report
- 1340 NPS/SFCN/NRTR—2009/240. National Park Service, Fort Collins, CO.
- 1341 White R. (2005) Vascular Plant Inventory and Plant Community Classification for Fort