

## A multi-species synthesis of physiological mechanisms in drought-induced tree mortality

H.D. Adams, M.J.B. Zeppel, W.R.L. Anderegg, H. Hartmann, S.M. Landhäusser, D.T. Tissue, T.E. Huxman, P.J. Hudson, T.E. Franz, C.D. Allen, et al.

### ► To cite this version:

H.D. Adams, M.J.B. Zeppel, W.R.L. Anderegg, H. Hartmann, S.M. Landhäusser, et al.. A multi-species synthesis of physiological mechanisms in drought-induced tree mortality. Nature Ecology & Evolution, 2017, 1, pp.7. 10.1038/s41559-017-0248-x. hal-02606468

## HAL Id: hal-02606468 https://hal.inrae.fr/hal-02606468

Submitted on 23 Apr 2024  $\,$ 

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



This is a repository copy of A multi-species synthesis of physiological mechanisms in drought-induced tree mortality.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/118137/</u>

Version: Accepted Version

### Article:

Adams, H.D., Zeppel, M.J.B., Anderegg, W.R.L. et al. (59 more authors) (2017) A multispecies synthesis of physiological mechanisms in drought-induced tree mortality. Nature Ecology and Evolution, 1 (9). pp. 1285-1291. ISSN 2397-334X

https://doi.org/10.1038/s41559-017-0248-x

### Reuse

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

### 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/ 1 2

# A multi-species synthesis of physiological mechanisms in drought-induced tree mortality

3	Authors: Henry D. Adams <sup>1*</sup> , Melanie J.B. Zeppel <sup>2,3</sup> , William R. L. Anderegg <sup>4</sup> , Henrik
4	Hartmann <sup>5</sup> , Simon M. Landhäusser <sup>6</sup> , David T. Tissue <sup>7</sup> , Travis E. Huxman <sup>8</sup> , Patrick J. Hudson <sup>9</sup> ,
5	Trenton E. Franz <sup>10</sup> , Craig D. Allen <sup>11</sup> , Leander D.L. Anderegg <sup>12</sup> , Greg A. Barron-Gafford <sup>13,14</sup> ,
6	David J. Beerling <sup>15</sup> , David D. Breshears <sup>16,17</sup> , Timothy J. Brodribb <sup>18</sup> , Harald Bugmann <sup>19</sup> , Richard
7	C. Cobb <sup>20</sup> , Adam D. Collins <sup>21</sup> , L. Turin Dickman <sup>21</sup> , Honglang Duan <sup>22</sup> , Brent E. Ewers <sup>23</sup> , Lucía
8	Galiano <sup>24</sup> , David A. Galvez <sup>6</sup> , Núria Garcia-Forner <sup>25</sup> , Monica L. Gaylord <sup>26,27</sup> , Matthew J.
9	Germino <sup>28</sup> , Arthur Gessler <sup>29</sup> , Uwe G. Hacke <sup>6</sup> , Rodrigo Hakamada <sup>30</sup> , Andy Hector <sup>31</sup> , Michael W.
10	Jenkins <sup>32</sup> , Jeffrey M. Kane <sup>33</sup> , Thomas E. Kolb <sup>26</sup> , Darin J. Law <sup>16</sup> , James D. Lewis <sup>34</sup> , Jean-Marc
11	Limousin <sup>35</sup> , David M. Love <sup>4</sup> , Alison K. Macalady <sup>36</sup> , Jordi Martínez-Vilalta <sup>37,38</sup> , Maurizio
12	Mencuccini <sup>39,40</sup> , Patrick J. Mitchell <sup>41</sup> , Jordan D. Muss <sup>21</sup> , Michael J. O'Brien <sup>42</sup> , Anthony P.
13	O'Grady <sup>41</sup> , Robert E. Pangle <sup>9</sup> , Elizabeth A. Pinkard <sup>41</sup> , Frida I. Piper <sup>43,44</sup> , Jennifer A. Plaut <sup>9</sup> ,
14	William T. Pockman <sup>9</sup> , Joe Quirk <sup>15</sup> , Keith Reinhardt <sup>45</sup> , Francesco Ripullone <sup>46</sup> , Michael G.
15	Ryan <sup>47,48,49</sup> , Anna Sala <sup>50</sup> , Sanna Sevanto <sup>21</sup> , John S. Sperry <sup>4</sup> , Rodrigo Vargas <sup>51</sup> , Michel
16	Vennetier <sup>52</sup> , Danielle A. Way <sup>53,54</sup> , Chonggang Xu <sup>21</sup> , Enrico A. Yepez <sup>55</sup> , Nate G. McDowell <sup>56</sup>
17	
18	Affiliations:

<sup>1</sup>Department of Plant Biology, Ecology, and Evolution, Oklahoma State University, Stillwater,
OK, USA

21 <sup>2</sup>Department of Biological Sciences, Macquarie University, Sydney, NSW, Australia

<sup>3</sup>The Boden Institute, Charles Perkins Centre, University of Sydney, Sydney, NSW, Australia

<sup>4</sup>Department of Biology, University of Utah, Salt Lake City, UT, USA

24	<sup>5</sup> Biogeochemical Processes, Max-Planck Institute for Biogeochemistry, Jena, Germany
25	<sup>6</sup> Department of Renewable Resources, University of Alberta, Edmonton, AB, Canada
26	<sup>7</sup> Hawkesbury Institute for the Environment, Western Sydney University, Penrith, NSW,
27	Australia
28	<sup>8</sup> Ecology and Evolutionary Biology, University of California, Irvine, CA, USA.
29	<sup>9</sup> Department of Biology, University of New Mexico, Albuquerque, NM, USA.
30	<sup>10</sup> School of Natural Resources, University of Nebraska-Lincoln, Lincoln, NE, USA.
31	<sup>11</sup> U.S. Geological Survey, Fort Collins Science Center, Jemez Mountains Field Station, Los
32	Alamos, NM, USA
33	<sup>12</sup> Biology, University of Washington, Seattle, WA, USA
34	<sup>13</sup> B2 EarthScience, Biosphere 2, University of Arizona, Tucson, AZ, USA
35	<sup>14</sup> School of Geography & Development, University of Arizona, Tucson, AZ, USA
36	<sup>15</sup> Department of Animal and Plant Sciences, University of Sheffield, Sheffield, UK
37	<sup>16</sup> School of Natural Resources and the Environment, University of Arizona, Tucson, AZ, USA
38	<sup>17</sup> Department of Ecology and Evolutionary Biology, University of Arizona, Tucson, AZ, USA
39	<sup>18</sup> School of Biology, University of Tasmania, Hobart, TAS, Australia
40	<sup>19</sup> Forest Ecology, Department of Environmental Systems Science, ETH Zurich, Zurich,
41	Switzerland
42	<sup>20</sup> Department of Plant Pathology, University of California, Davis, CA, USA
43	<sup>21</sup> Earth and Environmental Sciences Division, Los Alamos National Laboratory, Los Alamos,
44	NM, USA
45	<sup>22</sup> Institute of Ecology and Environmental Science, Nanchang Institute of Technology, Nanchang,
46	Jiangxi, China

47	<sup>23</sup> Department of Botany and Program in Ecology, University of Wyoming, Laramie, WY,
48	USA <sup>24</sup> Department of Forest Mycology and Plant Pathology, Swedish University of Agricultural
49	Sciences, Uppsala, Sweden
50	<sup>25</sup> Department of Life Sciences, Centre for Functional Ecology, University of Coimbra, Coimbra,
51	Portugal
52	<sup>26</sup> School of Forestry, Northern Arizona University, Flagstaff, AZ, USA
53	<sup>27</sup> Forest Health Protection, R3-Arizona Zone, US Forest Service, Flagstaff, AZ, USA
54	<sup>28</sup> U.S. Geological Survey, Forest and Rangeland Ecosystem Science Center, Boise, Idaho, USA
55	<sup>29</sup> Forest Dynamics, Swiss Federal Research Institute WSL, Birmensdorf, Switzerland
56	<sup>30</sup> Department of Forest Sciences, University of Sao Paulo, Piracicaba, Brazil
57	<sup>31</sup> Department of Plant Sciences, University of Oxford, Oxford, UK
58	<sup>32</sup> Environmental Studies Department, University of California Santa Cruz, Santa Cruz, CA, USA
59	<sup>33</sup> Department of Forestry and Wildland Resources, Humboldt State University, Arcata, CA, USA
60	<sup>34</sup> Louis Calder Center - Biological Field Station and Department of Biological Sciences,
61	Fordham University, Armonk, NY, USA
62	<sup>35</sup> Centre d'Ecologie Fonctionnelle et Evolutive, CNRS, Montpellier, France
63	<sup>36</sup> U.S. Agency for International Development, Washington, DC, USA
64	<sup>37</sup> CREAF, Cerdanyola del Valles, Spain
65	<sup>38</sup> Universitat Autònoma Barcelona, Cerdanyola del Valles, Spain
66	<sup>39</sup> ICREA, ICREA-CREAF, Cerdanyola del Valles, Barcelona, Spain
67	<sup>40</sup> School of GeoSciences, University of Edinburgh, Edinburgh, UK
68	<sup>41</sup> CSIRO Land and Water, Hobart, TAS, Australia

- <sup>42</sup>Estación Experimental de Zonas Áridas, Consejo Superior de Investigaciones Científicas, La
- 70 Cañada, Almería, Spain
- <sup>43</sup>Centro de Investigación en Ecosistemas de la Patagonia, Coyhaique, Chile
- 72 <sup>44</sup>Instituto de Ecología y Biodiversidad, Santiago, Chile
- <sup>45</sup>Department of Biological Sciences, Idaho State University, Pocatello, ID, USA
- <sup>46</sup>School of Agricultural, Forest, Food and Environmental Sciences, University of Basilicata,
- 75 Potenza, Italy
- <sup>47</sup>Natural Resources Ecology Laboratory, Colorado State University, Fort Collins, CO, USA
- <sup>48</sup>Graduate Degree Program in Ecology, Colorado State University, Fort Collins, CO, USA
- <sup>49</sup>USDA Forest Service, Rocky Mountain Research Station, Fort Collins, CO, USA;
- <sup>50</sup>Division of Biological Sciences, University of Montana, Missoula, MT, USA
- <sup>51</sup>Department of Plant and Soil Sciences, University of Delaware, Newark, DE, USA
- 81 <sup>52</sup>Irstea, UR RECOVER, Aix en Provence, France
- <sup>53</sup>Nicholas School of the Environment, Duke University, Durham, North Carolina, USA
- <sup>54</sup>Department of Biology, University of Western Ontario, London, ON, Canada
- <sup>55</sup>Departamento de Ciencias del Agua y Medio Ambiente, Instituto Tecnologico de Sonora,
- 85 Ciudad Obregon, SO, Mexico
- <sup>56</sup>Pacific Northwest National Laboratory, Richland, WA, 99352
- 87 \*Correspondence to: henry.adams@okstate.edu
- 88
- 89 For submission to Nature Ecology and Evolution as an Article
- 90
- 91

92 Widespread tree mortality associated with drought has been observed on all 93 forested continents, and global change is expected to exacerbate vegetation vulnerability. 94 Forest mortality has implications for future biosphere-atmosphere interactions of carbon, 95 water, and energy balance, and is poorly represented in dynamic vegetation models. 96 Reducing uncertainty requires improved mortality projections founded on robust 97 physiological processes. However, the proposed mechanisms of drought-induced mortality, 98 including hydraulic failure and carbon starvation, are unresolved. A growing number of 99 empirical studies have investigated these mechanisms, but data have not been consistently 100 analyzed across species and biomes using a standardized physiological framework. Here 101 we show that xylem hydraulic failure was ubiquitous across multiple tree taxa at drought-102 induced mortality. All species assessed had 60% or higher loss of xylem hydraulic 103 conductivity, consistent with proposed theoretical and modelled survival thresholds. We 104 found diverse responses in non-structural carbohydrate reserves at mortality, indicating 105 that evidence supporting carbon starvation was not universal. Reduced non-structural 106 carbohydrates were more common for gymnosperms than angiosperms, associated with 107 xylem hydraulic vulnerability, and may have a role in reducing hydraulic function. Our 108 finding that hydraulic failure at drought-induced mortality was persistent across species 109 indicates that substantial improvement in vegetation modelling can be achieved using 110 thresholds in hydraulic function.

Increasing forest mortality from global change has been observed in all forested biomes<sup>1,2</sup> and will have profound implications for future energy and element fluxes<sup>3-5</sup>. Predictions of vegetation responses to future climate are uncertain due to the lack of realistic mortality mechanisms in vegetation models<sup>3,6-9</sup>. Recent research supports at least two tightly inter-related

115	physiological mechanisms associated with tree mortality by drought: (a) hydraulic failure
116	through partial or complete loss of xylem function from embolism that inhibits water transport
117	through the vasculature, leading to tissue desiccation; and (b) carbon starvation via imbalance
118	between carbohydrate demand and supply that may lead to an inability to meet osmotic,
119	metabolic, and defensive carbon requirements <sup>3,6,7,10-15</sup> . Hydraulic failure is most typically
120	assessed via percent loss of xylem conductivity (PLC), and carbon starvation via changes in
121	tissue non-structural carbohydrate (NSC) concentrations <sup>12-16</sup> . There has been significant debate
122	over these co-occurring mechanisms of mortality, particularly regarding the prevalence of carbon
123	starvation and whether reduced carbohydrate reserves can be lethal during drought <sup>11,17-22</sup> .
124	Though a number of studies on the mechanism of drought-induced mortality in trees have
125	been conducted for a variety of tree species over the last decade, the prevalence of these
126	mechanisms at a global scale remains uncertain. Differences in approach, variables measured,
127	and species and life stage studied have limited global assessment of drought-induced tree
128	mortality mechanism. Here, we provide the first cross-species synthesis of tree drought
129	mortality mechanisms. We used a standardized physiological framework to analyze drought-
130	induced tree mortality across species and assessed hydraulic function as PLC, and carbohydrate
131	status as NSC normalized relative to controls. We examined data from 19 recent experimental
132	and observational studies on 26 species from around the globe. Most tree species were assessed
133	in only one study, but for several species, data were available from more than one study,
134	resulting in 34 cases (species-study combinations). However, data were not available for all
135	analyses from all cases: more cases had NSC data (31 cases from 24 species) than PLC data (14
136	cases from 9 species) which could be used to compare NSC and PLC at mortality with that of
137	surviving control trees (see Methods below, Supplementary Table 1). In order to make our

138 synthesis comprehensive, we worked with all of the data that were available, including data from 139 studies on a range of tree sizes and ontogenetic life stages (i.e. seedlings, saplings, and large 140 trees), conducted in a variety of settings, including potted plants in greenhouses or growth 141 chambers, and trees grown in the field (Supplementary Tables 1, 2). Given the diversity of 142 studies synthesized, these data were not ideal for a statistical meta-analysis; therefore, we limited 143 our analyses to a standard comparison within each case between plants that died and plants that 144 remained healthy (Supplementary Methods). We also compared differences in degree of 145 embolism and carbohydrate concentrations between plants at mortality and control plants to differences in functional traits<sup>3,24,25</sup>. For each species, we obtained available data for traits that 146 147 are easily measured, widely available, and likely relevant for drought tolerance, including wood density and specific leaf area<sup>26</sup>. We also obtained data for hydraulic traits that are directly related 148 149 to drought tolerance, but harder to measure, including xylem water potential at 50% loss of 150 hydraulic conductivity ( $\Psi_{50}$ ), point of embolism entry ( $\Psi_{e}$ ), and corresponding hydraulic safety margins<sup>24,27</sup> (Supplementary Methods). We used this dataset to address the following 151 152 hypotheses: 1) given the potential role of NSC in the maintenance of water transport during drought<sup>6, 28</sup>, both high PLC and reduced NSC reserves are common at tree death from drought, 153 154 and 2) among species, species-level functional traits that have been positively related to drought 155 tolerance (e.g. low xylem vulnerability to embolism, low SLA, high wood density) are associated 156 with high NSC at tree death. According to this hypothesis we expect that for species with greater 157 xylem vulnerability (quantified by  $\Psi_{50}$ ,  $\Psi_{e}$ , and hydraulic safety margin), NSC at death will be 158 relatively lower. This hypothesis is based on prior proposals that drought-sensitive trees which 159 close their stomata earlier during drought would be more likely to show a reduction in NSC associated with carbon starvation<sup>3,25,29,30</sup>. 160

161

### 162 **Results.**

163 For the cases where PLC data at mortality were available (Supplementary Methods), PLC 164 was 60% or higher (Figure 1A), demonstrating that a high degree of xylem embolism at drought-165 induced death was a universal aspect of mortality physiology in these species. Mean PLC was 166 84.3% at mortality, and PLC was significantly higher at mortality than for control trees in every 167 case (p < 0.05, Student's t-test). For NSC, we focused our analysis on differences in NSC 168 concentration between trees that died from drought and controls which did not die, measured at 169 the same point in time for both groups, although we also considered differences over time for 170 trees that died (Supplementary Discussion). Reductions in NSC at mortality were common 171 among species, but not universal, and no common NSC threshold for mortality was identified. 172 For 48% of cases and 38% of species with available data, NSCs were significantly lower at 173 mortality in dying trees compared to surviving or control trees (for observational and 174 experimental studies, respectively) in at least one tissue (p < 0.05, ANOVA; Figure 1B-D). 175 Among all species, mean NSCs at mortality for leaves, above-ground woody tissues (bole, 176 branch, stem, or twig), and roots were 13, 17, and 35% lower in dying trees than control 177 measurements.

For boreal and temperate angiosperms, lower NSCs at mortality relative to control trees were observed in 56% of cases and 63% of the species for at least one tissue, and NSC reductions exceeded 50% in approximately 33% of these cases and 38% of these species (Figure 18). Higher NSCs at mortality relative to controls were common for tropical angiosperm seedlings<sup>31</sup>, more than 100% higher in some cases, and reduced NSCs were not observed in this group, suggesting different physiological responses to severe drought in non-tropical and tropical 184 tree species (Figure 1B, C). In a similar seedling study with the same tropical species, however, 185 lower pre-drought NSCs were consistently correlated with a shorter time to mortality though NSCs did not decline during drought<sup>14</sup>. Lower NSCs at mortality relative to controls were most 186 common in root tissues<sup>32</sup>, and typically resulted from lower starch concentrations, consistent 187 188 with a starch to sugar conversion to meet metabolic and osmoregulatory demands during drought stress<sup>6</sup> (Supplementary Figure 1). Notably, only a few cases exhibited the hypothesized time-189 190 series trend in NSCs of initial small increase and then a more pronounced decrease in NSCs 191 over time<sup>28</sup> (Supplementary Figures 2, 3, 4).

192 Reductions in NSCs at mortality were more prevalent for gymnosperms than 193 angiosperms (Figure 1, Supplementary Figures 2, 3, 4). Among gymnosperms, 83% of cases and 194 67% of species had lower NSC at mortality relative to controls for at least one tissue (Figure 195 1D). This occurred in at least one tissue for all four species of the Pinaceae, but not for the two 196 species in the Cupressaceae, which is consistent with divergent evolutionary pathways for stomatal control between these families<sup>33</sup>. Relative reductions in NSCs were also generally 197 198 greater in gymnosperms than angiosperms, e.g. Pinus sylvestris had NSC reductions of >80% in 199 some tissues prior to mortality (Figure 1D).

Functional traits related to xylem embolism resistance and stomatal control have been suggested as useful predictors of the physiological causes of drought-induced mortality<sup>3,25,30,34</sup>. For all species, the deviation of NSCs in trees at mortality from their controls was not significantly associated with wood density or specific leaf area (p > 0.05, linear regression), regardless of whether the relationships were assessed for angiosperms, gymnosperms, or all species together. For gymnosperms, reduced NSCs at mortality in aboveground woody tissues (bole, branch, stem, or twig) were associated with lower resistance to xylem embolism (i.e.

207	higher $\Psi_{50}$ and $\Psi_e$ ; $r^2 = 0.88$ and 0.91, respectively, p < 0.001, linear regression; Figure 2),
208	indicating that hydraulic features in gymnosperms associated with drought resistance were
209	related to NSC dynamics during lethal drought. Normalized NSCs in other tissues were
210	positively correlated with embolism resistance at mortality (leaf NSC with $\Psi_{50}$ , root NSC with
211	$\Psi_e$ ; p < 0.05, linear regression), and normalized NSCs in aboveground woody tissue and roots at
212	mortality were also positively correlated with the $\Psi_{50}$ hydraulic safety margin for gymnosperms
213	(p < 0.001, linear regression; Supplementary Figure 5), but these relationships were strongly
214	influenced by one species, Callitris rhomboidea (Supplementary Methods). Variation in PLC at
215	mortality was not related to any functional traits assessed (p > 0.05, linear regression).

216

#### 217 **Discussion.**

218 We found that tree mortality from drought was always associated with substantial loss of 219 hydraulic function, and that lower NSCs at mortality were common but not universal (Figure 1). Our findings for PLC at mortality (Figure 1A) are close to modeling and theoretical predictions 220 of a stem PLC mortality threshold near or above  $60\%^{7,10,35-37}$ . In all cases, we found that PLC at 221 222 mortality was at least 60%, but values were much higher in a number of cases. The studies in 223 our synthesis were not designed to quantify lethal PLC thresholds, which deserve future 224 investigation to determine the duration and intensity of drought required to trigger mortality and 225 the mechanisms underlying such a threshold. The physiological effects of a particular level of 226 PLC likely vary among species, mediated by traits such as the capacity to refill embolism and replace conducting area via new growth<sup>6,38</sup>. Nonetheless, a sustained stem PLC at or above 60% 227 228 provides a generally supported starting point for modeling vegetation response across spatial scales, a point beyond which the probability of mortality increases<sup>7,10,36,37</sup>. 229

230 For the cases where both NSC and PLC data were available at mortality, all trees died 231 with high PLC (100% of cases), but only 62% of cases also had low NSCs at mortality relative to 232 controls (Figure 3). This suggests that trees died from either hydraulic failure alone, or hydraulic 233 failure in combination with reduced NSCs. This finding should help lay to rest the 234 misconception of a dichotomy between hydraulic failure and carbon starvation, which are often mistakenly thought to represent mutually exclusive mechanisms<sup>15</sup>. Clearly, our results 235 236 underscore the importance of maintaining a functional plant hydraulic system for survival, while 237 suggesting a relationship between hydraulic failure and carbon starvation mechanisms in this 238 process. The majority of studies included in our analysis were not designed to distinguish the 239 drivers of mortality from the non-causative symptoms of dying. Thus, it is not possible with our 240 data to conclusively determine if changes in either NSC or PLC facilitated death or were the 241 result of the mortality process (Supplementary Discussion). Results from studies in which light 242 and CO<sub>2</sub> concentration were manipulated to regulate carbon fixation do suggest a role for NSC as 243 a survival mechanism against mortality via hydraulic failure during drought, even when NSC does not decline during drought or is not reduced below control values<sup>12-16</sup> (Supplementary 244 245 Discussion).

Given the diversity of NSC responses found at mortality, there is an obvious need to develop frameworks for the sensitivity of plant metabolism to changes in NSC levels, including the potential for lethal thresholds<sup>22,39</sup>. Specific NSC thresholds for survival or mortality during drought are not well-resolved in our data, nor yet in the literature. Such survival thresholds likely vary with factors including tree species, ontogeny, tree tissue, canopy position, seasonality, environmental conditions, and interactions with other organisms, but empirical investigation of these thresholds is needed<sup>22,40,41</sup>. Determination of these thresholds is hampered by an incomplete understanding of the role of NSC storage in plant function, and its regulatory
mechanism<sup>22,39</sup>. However, significantly lower NSCs at mortality were relatively common for a
variety of species in our analysis, such that reduced NSCs can no longer be considered a rare or
atypical response during tree death.

257 Our finding that reduced NSCs at mortality were more common for gymnosperms, than 258 for angiosperms (Figure 1, Supplementary Figures 2, 3, 4), is consistent with the wider hydraulic safety margins of gymnosperms relative to angiosperms<sup>24,42</sup>. For gymnosperms, our functional 259 260 trait analysis revealed that species with greater xylem embolism resistance had higher NSC at 261 mortality in boles, branches, stems, or twigs than surviving controls, indicating that species' 262 hydraulic traits can affect C balance during lethal drought (Figure 2). As embolism resistance is often associated with an ability to keep stomata open at lower water potentials<sup>30,43</sup>, our results 263 264 suggest that tree species which can maintain stomatal conductance and photosynthesis at higher xylem tension during drought are less likely to have reduced NSC at mortality<sup>29</sup>. These resistant 265 266 tree species would be more likely to die from hydraulic failure alone without reduced NSC — 267 consistent with hypotheses that stomatal regulation and hydraulic transport strategies influence 268 the contribution of carbon starvation and hydraulic failure to mortality mechanism among species<sup>3,25,30</sup>. Caution, however, should be used in assuming stomatal regulation is highly 269 270 coupled with water potential regulation and hydraulic strategy<sup>44</sup>. Importantly, we did not find a 271 relationship between NSC reduction and embolism resistance for angiosperms, nor did any other 272 trait predict mortality physiology in these species.

273 Our synthesis of data from multiple studies on the physiology of drought-induced tree 274 mortality exposes several key knowledge gaps in the field. Our dataset of only 26 species under-275 represents the enormous diversity of tree species found in forests globally, particularly so for

276 tropical forests, where drought-induced mortality can have substantial implications for the global 277 carbon cycle<sup>9</sup>. *Pinus* was relatively over-represented in this synthesis (nine cases from three 278 species), although it is widely distributed and has been widely affected by forest die-off on 279 multiple continents<sup>1</sup>. Also, our dataset is dominated by data from seedlings and saplings, often 280 from studies conducted with potted plants, which may be predisposed to die quickly from 281 hydraulic failure due to limited rooting volume and lack of access to deeper soil water pools 282 (Supplementary Tables 1, 2). Data at mortality for more than one life stage were available for 283 only three species (Figure 1), and the consistency of NSC and PLC responses at mortality across 284 a gradient of size and ontogeny varied in these species. Clearly, more research on the physiology 285 of mortality in large trees in the field and the effect of size and ontogeny on the mortality process 286 is needed. Nonetheless, our overall observation that hydraulic failure was universal, and NSC 287 reduction was not, does not change if we only consider data for each life stage separately. In all 288 cases for which PLC data were available, mean PLC was 60% or greater at mortality, 289 irrespective of life stage (Figure 1A). Our finding that normalized NSC at mortality varied 290 among cases and species also holds when seedlings, saplings, and trees are considered separately 291 (Figure 1B-D, Supplementary Table 1).

Determining whether forests will continue to act as a global carbon sink or transition to a carbon source is a critical uncertainty for the carbon cycle with large ramifications for society and climate policy<sup>8,9,23</sup>. Such a shift largely depends on tree mortality responses which could be anticipated by resolving the relative roles of hydraulic and carbohydrate mechanisms in causing tree death<sup>7,10,45</sup>. We found that hydraulic failure was ubiquitous among the studies we compared, that PLC at mortality in all cases with such data was at least 60%. These results affirm that simulating hydraulic function should be a first priority for development of mechanistic tree 299 mortality algorithms in climate-vegetation models to improve projections of the future terrestrial 300 carbon budget. Hydraulic models that capture drought damage at tree and landscape scales are rapidly developing<sup>7,10,36,37,45-47</sup> and substantial improvement in vegetation model projections may 301 302 be possible with simulation of hydraulic-driven mortality, whether tree carbohydrate status is 303 represented or not. Reduced NSC in tree species dying from drought was common in 304 gymnosperms, but not angiosperms, suggesting an influence of NSC on hydraulic deterioration 305 in some trees that requires further investigation. Yet, the diversity of NSC responses among only 306 26 species and the design limitations of past studies in determining causality demonstrate that we need to further assess the influence of carbon metabolism and storage on mortality<sup>39</sup>. Ultimately, 307 308 an improved representation of the physiology of drought-induced tree mortality that includes 309 both water and carbon relations will be crucial for forecasting the fate of forests in a changing 310 climate.

311

### 312 Methods

313 Data Synthesis. We used literature search and extensive discussion with colleagues to identify 314 data from 19 experimental and observational studies on 26 species, for a total of 34 cases (study and species combinations). Literature search terms included "non-structural carbohydrates", 315 316 "water potential", "tree mortality", and "drought". Our synthesis was not limited to an objective 317 literature search, as we sought to include all published data that fit our criteria for inclusion. 318 Criteria for inclusion were that studies included data on: 1) tree mortality from drought; 2) NSC 319 concentrations of at least one tissue, and/or PLC of aboveground woody tissue, either measured 320 directly, or estimated from plant water potential ( $\Psi_p$ ) measured at mortality, or modeled from hydraulic conductance  $^{48,49}$  (Supplementary Methods); and 3) that data were either: a) 321

322 concurrently collected for trees that died (either at or near mortality) and from trees that either 323 survived the drought or were in a paired control treatment, and/or: b) available prior to drought 324 or pre-treatment from the same trees that later died. We obtained data from each study directly 325 from contributors. Details on the specific studies synthesized can be found in Supplementary 326 Table 1. Determination of the point of mortality in dying trees was defined in each original 327 study, as detailed in Supplementary Table 3, and we relied on data contributors to provide the 328 appropriate data for at- (or near-) mortality assessments.

329 NSC measurements are methodologically challenging and comparisons of absolute 330 concentrations can be problematic across studies due to issues of standards, NSC technique, and lab protocol disagreement<sup>50,51</sup>. However, relative differences (treatment vs. control and changes 331 332 over time assessed with the same technique in the same laboratory) provide robust estimates of NSC dynamics within studies<sup>50,51</sup>. We limited all statistical analyses of absolute NSC data to 333 334 within each case (detailed below) and we only present relative differences in NSC in figures. 335 For studies where data were concurrently available for trees that died and control or surviving 336 trees, we calculated a normalized NSC deviation from the difference between values at or near 337 mortality and those for control or surviving trees divided by the control or surviving tree value. 338 For studies where data were available prior to the drought for the same trees that later died (or 339 seedlings in the same treatment harvested at measurement), normalized values were also 340 calculated as the difference between values at or near mortality and initial pre-treatment or pre-341 drought values divided by the initial or pre-drought values. In both cases, normalized values 342 were expressed as a percent. For comparison of time series trends in NSC, we also calculated 343 normalized, proportional NSCs in trees that died by scaling values relative to the maximum 344 value in each time series, which was defined as a normalized value of 1. When possible,

normalizations were calculated for individual trees, and specifically for each tissue sampled. For
studies 3 and 9 (Supplementary Table 1), only means and standard errors for species and tissues
were available, so normalized values were calculated from these metrics.

348 Note that all types of data were not available for all cases in our synthesis. Among the 34 349 cases in our dataset, PLC measured at mortality was available for nine cases (eight species), PLC 350 was estimated in five cases (two species), NSC deviation from control/surviving trees at 351 mortality was available for 31 cases (24 species), and percent change in NSC was available for 352 28 cases (22 species). Sample sizes for PLC and NSC data are available in Supplementary 353 Tables 4 and 5. Because PLC values are already normalized to the maximum conductivity per 354 sample, no further normalization was conducted with these data. We also acknowledge that 355 direct measurements of PLC and generation of hydraulic vulnerability curves can be challenging, and that method artifacts can effect results<sup>52,53</sup>. Although the majority of hydraulic data we 356 357 report were collected following recommended practices (Supplementary Methods, 358 Supplementary Table 5), we cannot rule out the possibility of such artifacts influencing our data. 359 To compare physiological mortality indicators to tree species traits, we obtained trait data 360 for the species in this synthesis from a variety of sources. We investigated the relationships 361 between physiology at mortality and traits related to drought tolerance that are easily measured 362 and widely available, such as wood density and specific leaf area (SLA). We also included 363 hydraulic traits more directly related to drought tolerance that were measured with morechallenging hydraulic vulnerability curve methods. Wood density data for most species were 364 obtained from the Global Wood Density database<sup>54,55</sup> available through the DRYAD digital 365 366 repository (www.datadryad.org). We obtained SLA data from the TRY database (www.trydb.org)<sup>56-58</sup>, for nearly all non-tropical species. We calculated species means for SLA from all 367

368 data available for each species of interest for our analysis. Data for *Acer pseudoplatanus* were 369 available from the mortality study population<sup>16</sup>. For *Callitris rhomboidea* and *Eucalyptus* 370 smithii, SLA data were not available. Additional sources of wood density data are detailed in 371 Supplementary Methods. Hydraulic trait data for the stem water potential at 50 PLC ( $\Psi_{50}$ ) and hydraulic safety margin  $(\Psi_{50} - \text{minimum } \Psi)^{24,27}$ , were obtained from multiple sources 372 373 (Supplementary Table 5, Supplementary Methods). Data for the embolism entry point ( $\Psi_e$ ) were 374 not available in the literature, so we calculated  $\Psi_e$  from relevant hydraulic vulnerability curve for 375 each case by applying a Weibull fit to the data, and determining the x-intercept of the line tangent to  $\Psi_{50}$  (Supplementary Table 5)<sup>27,59</sup>. Hydraulic trait data were unavailable for 376 377 Eucalyptus radiata, Eucalyptus smithii, and Nothofagus nitida. No trait data were available for 378 the tropical angiosperm species from study 7 (Supplementary Table 1) for any of the traits we assessed<sup>31</sup>. 379

The majority of datasets generated and analyzed during the current study are available from the corresponding author on reasonable request. Trait data obtained for the current study from the TRY Database were used under license and as restrictions apply to the availability of these data, these are not available from the corresponding author, but can be requested from the TRY Database (www.try-db.org).

385

Statistical Analyses. We used MATLAB R2012a (The Mathworks, Inc., Natick, MA, USA) for all statistical analyses, with  $\alpha = 0.05$ . All NSC and PLC comparisons were performed using ANOVA or Student's t-test individually for each case, between dead (or dying) and control/surviving trees or between post-drought dead and corresponding pre-drought values, with tissue as a factor for analysis of NSC. Since our NSC normalization could affect tissue

391 comparisons within the same case, these analyses were performed on non-normalized NSC data 392 to maintain the correct ratio among tissues, a conservative approach. Our within-individual case 393 analysis on relative differences in non-normalized NSC does not bear the risk of error introduced 394 by different NSC techniques or labs, or uncertainty in standards for determining absolute NSC, 395 and furthermore the inferences are based on large effect sizes compared to possible measurement error<sup>50,51</sup>. In experimental cases that included temperature or CO<sub>2</sub> concentration treatments in 396 397 addition to drought, we included these factors in ANOVA tests to determine if PLC and NSC 398 should be pooled or split among levels of these factors (Supplementary Table 1). For NSC, these 399 analyses also included tissue as a factor. Functional trait relationships with normalized NSC data 400 at mortality were analyzed with linear regression. Cook's distance was calculated for all points 401 in significant linear regressions, and a value greater than three times the mean of the Cook's 402 distance was used to identify outliers for exclusion.

403

### 404 **References and Notes**

- 405 1. Allen, C. D. *et al.* A global overview of drought and heat-induced tree mortality reveals
  406 emerging climate change risks for forests. *Forest Ecology and Management* 259, 660-
- 407 684, doi:10.1016/j.foreco.2009.09.001 (2010).
- 408 2. Intergovernmental Panel on Climate Change (IPCC). Impacts, Adaptation, and Vulnerability.
- 409 Contribution of Working Group II to the Fifth Assessment Report of the IPCC.
- 410 Cambridge University Press, Cambridge, UK, 1132 pp. (2014).

411	3. McDowell, N. et al. Mechanisms of plant survival and mortality during drought: why do
412	some plants survive while others succumb to drought? New Phytologist 178, 719-739,
413	doi:10.1111/j.1469-8137.2008.02436.x (2008).
414	4. Adams, H. D. et al. Ecohydrological consequences of drought- and infestation- triggered tree
415	die-off: insights and hypotheses. <i>Ecohydrology</i> 5, doi:10.1002/eco.233 (2012).
416	5. Anderegg, W. R. L., Kane, J. M. & Anderegg, L. D. L. Consequences of widespread tree
417	mortality triggered by drought and temperature stress. Nature Climate Change 3, 30-36
418	(2013).
419	6. McDowell, N. G. et al. The interdependence of mechanisms underlying climate-driven
420	vegetation mortality. Trends in Ecology & Evolution 26, 523-532,
421	doi:10.1016/j.tree.2011.06.003 (2011).
422	7. McDowell, N. G. et al., Multi-scale predictions of massive conifer mortality due to chronic
423	temperature rise. Nature Climate Change 6, 295-300, doi:10.1038/nclimate2873 (2016).
424	8. Friedlingstein, P. et al. Uncertainties in CMIP5 Climate Projections due to Carbon Cycle
425	Feedbacks. Journal of Climate 27, 511-526, doi:10.1175/jcli-d-12-00579.1 (2014).
426	9. Friend, A. D. et al. Carbon residence time dominates uncertainty in terrestrial vegetation
427	responses to future climate and atmospheric CO2. Proceedings of the National Academy
428	of Sciences of the United States of America 111, 3280-3285,
429	doi:10.1073/pnas.1222477110 (2014).

430	10. McDowell, N. G. et al., Evaluating theories of drought-induced vegetation mortality using a
431	multimodel-experiment framework. New Phytologist 200, 304-321, doi:
432	10.1111/nph.12465 (2013).
433	11. Sala, A., Piper, F. & Hoch, G. Physiological mechanisms of drought-induced tree mortality
434	are far from being resolved. New Phytologist 186, 274-281 (2010).
435	12. Hartmann, H., Ziegler, W., Kolle, O. & Trumbore, S. Thirst beats hunger - declining
436	hydration during drought prevents carbon starvation in Norway spruce saplings. New
437	Phytologist 200, 340-349, doi:10.1111/nph.12331 (2013).
438	13. Quirk, J., McDowell, N. G., Leake, J. R., Hudson, P. J. & Beerling, D. J. Increased
439	susceptibility to drought-induced mortality in Sequoia sempervirens (Cupressaceae)
440	trees under Cenozoic atmosphere carbon dioxide starvation. American Journal of
441	Botany 100, 582-591, doi:10.3732/ajb.1200435 (2013).
442	14. O'Brien, M. J., Leuzinger, S., Philipson, C. D., Tay, J. & Hector, A. Drought survival of
443	tropical tree seedlings enhanced by non-structural carbohydrate levels. Nature Climate
444	Change 4, 710-714, doi:10.1038/nclimate2281 (2014).
445	15. Sevanto, S., McDowell, N. G., Dickman, L. T., Pangle, R. & Pockman, W. T. How do trees
446	die? A test of the hydraulic failure and carbon starvation hypotheses. Plant Cell and
447	Environment 37, 153-161, doi:10.1111/pce.12141 (2014).
448	16. Piper, F. I. & Fajardo, A. Carbon dynamics of Acer pseudoplatanus seedlings under drought
449	and complete darkness. Tree Physiology, doi:10.1093/treephys/tpw063 (2016).

450	17. McDowell, N. G. & Sevanto, S. The mechanisms of carbon starvation: how, when, or does it
451	even occur at all? New Phytologist 186, 264-266 (2010).

- 452 18. Sala, A., Woodruff, D. R. & Meinzer, F. C. Carbon dynamics in trees: feast or famine? *Tree*453 *Physiology* 32, 764-775, doi:10.1093/treephys/tpr143 (2012).
- 454 19. Fatichi, S., Leuzinger, S. & Koerner, C. Moving beyond photosynthesis: from carbon source
- 455 to sink-driven vegetation modeling. *New Phytologist* **201**, 1086-1095,
- 456 doi:10.1111/nph.12614 (2014).
- 457 20. Hartmann, H. Carbon starvation during drought-induced tree mortality are we chasing a
  458 myth? *Journal of Plant Hydraulics* 2, e-005 (2015).
- 459 21. Körner, C. Paradigm shift in plant growth control. *Current Opinion in Plant Biology* 25,
  460 107-114, doi:10.1016/j.pbi.2015.05.003 (2015).
- 461 22. Martínez-Vilalta, J., Sala A., *et al.* Dynamics of non-structural carbohydrates in terrestrial
- 462 plants: a global synthesis. *Ecological Monographs* **86**, 495-516. (2016).
- 463 23. Allen, C. D., Breshears, D. D. & McDowell, N. G. On underestimation of global

464 vulnerability to tree mortality and forest die-off from hotter drought in the

- 465 Anthropocene. *Ecosphere* **6**, doi:10.1890/es15-00203.1 (2015).
- 466 24. Choat, B. *et al.* Global convergence in the vulnerability of forests to drought. *Nature* 491,
  467 doi:10.1038/nature11688 (2012).

468	25. Skelton, R. P., West, A. G. & Dawson, T. E. Predicting plant vulnerability to drought in
469	biodiverse regions using functional traits. Proceedings of the National Academy of
470	Sciences of the United States of America 112, 5744-5749, doi:10.1073/pnas.1503376112
471	(2015).

472	26. Poorter, L. & Markesteijn, L. Seedling traits determine drought tolerance of tropical tree
473	species. <i>Biotropica</i> <b>40</b> , 321-331, doi:10.1111/j.1744-7429.2007.00380.x (2008).

- 474 27. Meinzer, F. C., Johnson, D. M., Lachenbruch, B., McCulloh, K. A. & Woodruff, D. R.
- 475 Xylem hydraulic safety margins in woody plants: coordination of stomatal control of
- 476 xylem tension with hydraulic capacitance. *Functional Ecology* **23**, 922-930,
- 477 doi:10.1111/j.1365-2435.2009.01577.x (2009).
- 478 28. McDowell, N. G. Mechanisms linking drought, hydraulics, carbon metabolism, and
  479 vegetation mortality. *Plant Physiology* 155, 1051-1059, doi:10.1104/pp.110.170704
  480 (2011).
- 481 29. Mitchell, P. J., O'Grady, A. P., Tissue, D. T., Worledge, D. & Pinkard, E. A. Co-ordination
  482 of growth, gas exchange and hydraulics define the carbon safety margin in tree species
  483 with contrasting drought strategies. Tree Physiology 34, 443-458,
- 484 doi:10.1093/treephys/tpu014 (2014).
- 30. Mencuccini, M., Minunno, F., Salmon, Y., Martinez-Vilalta, J. & Holtta, T. Coordination of
  physiological traits involved in drought-induced mortality of woody plants. *New Phytologist* 208, 396-409, doi:10.1111/nph.13461 (2015).

488	31. O'Brien, M. J., Burslem, D., Caduff, A., Tay, J. & Hector, A. Contrasting nonstructural
489	carbohydrate dynamics of tropical tree seedlings under water deficit and variability. New
490	Phytologist 205, 1083-1094, doi:10.1111/nph.13134 (2015).
491	32. Landhäusser, S. M. & Lieffers, V. J. Defoliation increases risk of carbon starvation in root
492	systems of mature aspen. Trees-Structure and Function 26, 653-661,
493	doi:10.1007/s00468-011-0633-z (2012).
494	33. Brodribb, T. J., McAdam, S. A. M., Jordan, G. J. & Martins, S. C. V. Conifer species adapt
495	to low-rainfall climates by following one of two divergent pathways. Proceedings of the
496	National Academy of Sciences of the United States of America 111, 14489-14493,
497	doi:10.1073/pnas.1407930111 (2014).
498	34. Anderegg, W. R. L. et al. Meta-analysis reveals that hydraulic traits explain cross-species
499	patterns of drought-induced tree mortality across the globe. Proceedings of the National
500	Academy of Sciences of the United States of America 113, 5024-5029, doi:
501	10.1073/pnas.1525678113 (2016).
502	35. Brodribb, T. J. & Cochard, H. Hydraulic failure defines the recovery and point of death in
503	water-stressed conifers. Plant Physiology 149, 575-584, doi:10.1104/pp.108.129783
504	(2009).
505	36. Anderegg, W. R. L. et al. Tree mortality predicted from drought-induced vascular
506	damage. Nature Geoscience 8, 367-371, doi:10.1038/ngeo2400 (2015).

507	37. Sperry, J. S. & Love, D. M. What plant hydraulics can tell us about responses to climate-
508	change droughts. New Phytologist 207, 14-27, doi:10.1111/nph.13354 (2015).
509	38. Zeppel, M. J. B. et al. Drought and resprouting plants. New Phytologist 206, 583-589,
510	doi:10.1111/nph.13205 (2015).
511	39. Hartmann, H. & Trumbore, S. Understanding the roles of nonstructural carbohydrates in
512	forest trees – from what we can measure to what we want to know. <i>New Phytologist</i> 211,
513	386-403, doi:10.1111/nph.139552016 (2016).
514	40. Oliva, J., Stenlid, J. & Martinez-Vilalta, J. The effect of fungal pathogens on the water and
515	carbon economy of trees: implications for drought-induced mortality. New
516	Phytologist 203, 1028-1035, doi:10.1111/nph.12857 (2014).
517	41. Anderegg, W. R. L. et al. Tree mortality from drought, insects, and their interactions in a
518	changing climate. New Phytologist 208, 674-683, doi:10.1111/nph.13477 (2015).
519	42. Johnson, D. M., McCulloh, K. A., Woodruff, D. R. & Meinzer, F. C. Hydraulic safety
520	margins and embolism reversal in stems and leaves: Why are conifers and angiosperms
521	so different? Plant Science 195, 48-53, doi:10.1016/j.plantsci.2012.06.010 (2012).
522	43. Garcia-Forner, N. et al. Responses of two semiarid conifer tree species to reduced
523	precipitation and warming reveal new perspectives for stomatal regulation. Plant Cell
524	and Environment 39, 38-49, doi:10.1111/pce.12588 (2016).

525	44. Martínez-Vilalta, J. & Garcia-Forner, N. Water potential regulation, stomatal behaviour and
526	hydraulic transport under drought: deconstructing the iso/anisohydric concept. Plant Cell
527	and Environment, doi:10.1111/pce.12846 (2016).
528	45. Adams, H. D. et al. Empirical and process-based approaches to climate-induced forest
529	mortality models. Frontiers in Plant Science 4, doi:10.3389/fpls.2013.00438 (2013).
530	46. Mackay, D. S. et al. Interdependence of chronic hydraulic dysfunction and canopy processes
531	can improve integrated models of tree response to drought. Water Resources Research
532	<b>51</b> , 6156-6176, doi:10.1002/2015wr017244 (2015).
533	47. Sperry, J. S. et al. Pragmatic hydraulic theory predicts stomatal responses to climatic water
534	deficits. New Phytologist, doi: 10.1111/nph.14059 (2016).
535	48. Sperry, J. S., Adler, F. R., Campbell, G. S. & Comstock, J. P. Limitation of plant water use
536	by rhizosphere and xylem conductance: results from a model. Plant Cell and
537	Environment 21, doi:10.1046/j.1365-3040.1998.00287.x (1998).
538	49. Plaut, J. A. et al. Hydraulic limits preceding mortality in a piñon-juniper woodland under
539	experimental drought. Plant Cell and Environment 35, 1601-1617, doi:10.1111/j.1365-
540	3040.2012.02512.x (2012).
541	50. Quentin, A. G. et al. Non-structural carbohydrates in woody plants compared among
542	laboratories. Tree Physiology 35, 1146-1165, doi:10.1093/treephys/tpv073 (2015).
543	51. Germino, M. J. A carbohydrate quandary. Tree Physiology 35, 1141-1145,
544	doi:10.1093/treephys/tpv109 (2015).

545	52. Wheeler, J. K. et al. Cutting xylem under tension or supersaturated with gas can generate
546	PLC and the appearance of rapid recovery from embolism. Plant, Cell & Environment
547	36, 1938-1949, doi:10.1111/pce.12139 (2013).
548	53. Nardini, A., Savi, T. Trifilò, P., Lo Gullo, M. A., Drought stress and the recovery from
549	xylem embolism in woody plants. Progress in Botany, doi:10.1007/124_2017_11
550	(2017).
551	54. Chave, J. et al. Towards a worldwide wood economics spectrum. Ecology Letters 12, 351-
552	366, doi:10.1111/j.1461-0248.2009.01285.x (2009).
553	55. Zanne, A. E., et al., Global wood density database. Dryad Digital Repository. Identifier:
554	http://hdl.handle.net/10255/dryad.235 (2009).
555	56. Kattge, J. et al. TRY - a global database of plant traits. Global Change Biology 17, 2905-
556	2935, doi:10.1111/j.1365-2486.2011.02451.x (2011).
557	57. Niinemets, U. Components of leaf dry mass per area - thickness and density - alter leaf
558	photosynthetic capacity in reverse directions in woody plants. New Phytologist 144, 35-
559	47, doi:10.1046/j.1469-8137.1999.00466.x (1999).
560	58. Niinemets, U. Global-scale climatic controls of leaf dry mass per area, density, and
561	thickness in trees and shrubs. <i>Ecology</i> 82, 453-469, doi:10.1890/0012-
562	9658(2001)082[0453:gsccol]2.0.co;2 (2001).

563	59. Domec, J. C. & Gartner, B. L. Cavitation and water storage capacity in bole xylem segments
564	of mature and young Douglas-fir trees. Trees-Structure and Function 15, 204-214,
565	doi:10.1007/s004680100095 (2001).
566	
567	Acknowledgements. This research was supported by the US Department of Energy, Office of
568	Science, Biological and Environmental Research and Office of Science, Next Generation
569	Ecosystem Experiment-Tropics, the Los Alamos National Laboratory LDRD Program, The EU
570	Euforinno project, the National Science Foundation LTER Program and EF-1340624, EF-
571	1550756, and EAR-1331408, ARC DECRA DE120100518, ARC LP0989881, ARC
572	DP110105102, the Philecology Foundation of Fort Worth, Texas, the Center for Environmental
573	Biology at UC Irvine through a gift from Mr. Donald Bren, and additional funding sources listed
574	in the Supplementary Acknowledgements. We thank Amanda Boutz, Sandra Bucci, Rosie
575	Fisher, Andrew Meador-Sanchez, Rick Meinzer, and Don White for discussions on study design,
576	analysis, and interpretation of results, and Troy Ocheltree for helpful comments on the
577	manuscript. Any use of trade, product or firm names is for descriptive purposes only and does
578	not imply endorsement by the U.S. Government.
579	
580	Author Contributions. ADC, AH, AKM, AS, BEE, CDA, CXU, DAG, DAW, DTT, GBG,
581	HDA, HH, JAP, JDL, JMK, JML, JSS, LDLA, LTD, MJBZ, MJG, MM, NGM, PJH, RCC, RV,
582	SML, SS, TEF, TEH, TEK, UH, WRLA, and WTP designed the study. AH, AOG, BEE, DAG,
583	DDB, DJB, DML, DTT, EAP, EAY, FIP, GBG, HD, HDA, HH, JAP, JDL, JMV, JQ, JSS, KR,

- 584 LDLA, LGP, LTD, MJBZ, MJG, MJO, MLG, NGF, NGM, PJH, PJM, REP, SML, SS, TEH,
- 585 TEK, TJB, UH, WRLA, and WTP contributed data. HDA, MJBZ, PJH, and TEF analyzed the

586	data. ADC, AG, AH, AKM, AOG, AS, BEE, CDA, CXU, DAW, DDB, DJB, DJL, DML, DTT,
587	EAP, FIP, FR, GBG, HB, HD, HDA, HH, JDL, JDM, JMK, JMV, JQ, JSS, KR, LDLA, LGP,
588	LTD, MGR, MJBZ, MJG, MJO, MLG, MM, MV, MWJ, NGF, NGM, PJH, PJM, RCC, RV,
589	SML, SS, TEF, TEH, TEK, UH, WRLA, and WTP contributed to the discussion of results.
590	ADC, AG, AH, AOG, AS, BEE, CDA, CXU, DAW, DDB, DJB, DJL, DTT, EAP, FIP, FR,
591	GBG, HB, HDA, HH, JDM, JMK, JML, JMV, KR, LDLA, LGP, LTD, MGR, MJBZ, MJG,
592	MJO, MLG, MM, MV, MWJ, NGF, NGM, PJM, RCC, RH, REP, RV, SML, SS, TEH, TEK,
593	TJB, UH, and WRLA wrote the manuscript.
594	
595	Competing financial interests. The authors declare they have no competing financial interests.
596	
597	Figure Legends
598	
599	Figure 1. Physiological responses at, or prior to, mortality from drought for multiple tree
600	species. Percent loss of hydraulic conductivity (PLC) for ambient moisture, control, or surviving
601	trees and concurrently at mortality from drought is shown for both angiosperm and gymnosperm
602	species (A). PLC was either measured directly (red) for control (open symbols) and dying
603	(closed symbols) trees or estimated from either water potential with a hydraulic vulnerability
604	curve (green) for control (open) and dying (closed) trees, or modeled from hydraulic
605	conductance (orange) for control (open), and dying (closed) trees. An "NA" indicates that
606	control PLC data were not available. In all panels for cases where individual data were
607	available, boxes indicate the 25% and 75% quartiles, whiskers indicate the extent of data, and

608 black bars indicate the mean. For cases where only means and a measure of variability were

609 available, means are indicated with squares and error bars are one standard error. For each case 610 in A where control and dying tree data were available, PLC was significantly higher at mortality 611 than for controls concurrently (p < 0.05, Student's t-test). A potential threshold for hydraulic 612 failure is indicated by a line at 60%. Non-structural carbohydrate concentration (NSC) at 613 mortality, normalized as the percent deviation from concurrent measurements of ambient, 614 control, or unaffected trees in each study for each plant tissue, is shown for deciduous and 615 evergreen non-tropical angiosperm (B), evergreen tropical angiosperm (C), and evergreen 616 gymnosperm (D) species. Significant differences for each tree tissue between drought trees at 617 mortality (black bar or square) and ambient, control, or surviving trees (0% line) are indicated 618 with an asterisk (p < 0.05, ANOVA). Note that the absolute values in NSC concentration used in 619 statistical analysis varied for each tissue in each case, such that distances between the mean and 620 zero in B-D are not a consistent indicator of statistical significance among cases or for tissues 621 within a case. An "M" indicates data from a study on mature trees; all other data are from studies 622 of seedlings, saplings, and small trees (Supplementary Tables 1, 2). Numbers after species 623 names in all panels designate original studies (Supplementary Table 1). Sample size for all data 624 analyzed for Figure 1 are shown in Supplementary Table 4.

625

**Figure 2.** The relationship between the tree hydraulic traits related to xylem embolism resistance and normalized non-structural carbohydrates (NSC) in aboveground woody tissue at, or prior to, mortality from drought, expressed as a deviation from concurrent measurements of surviving control trees, for angiosperm (blue circles; A, B) and gymnosperm (red triangles; C, D) species. Tree hydraulic traits related to embolism resistance are the water potential at 50% loss of hydraulic conductivity ( $\Psi_{50}$ ; A, C) and point of xylem embolism entry ( $\Psi_e$ ; B, D). Xylem embolism resistance increases to the right. NSC data shown are means for aboveground woody
tissue (bole, branch, stem, or twig), normalized as a percent of ambient moisture, control, or
surviving trees in each case. Significant linear regressions were found for gymnosperms (C, D)
but not angiosperms (A, B). Values for *Callitris rhomboidea* (upper right in C, D) were
identified as potential outliers, but both relationships remain statistically significant (p < 0.01,</li>
linear regression) for the remaining data with the removal of these points (Supplementary
Methods).

639

640 Figure 3. Physiological responses associated with hydraulic failure and carbon starvation, as 641 defined by PLC and NSC deviation from control in 13 cases (study × species combinations) for 642 which both data were available. Among these cases, trees either died with high PLC and low 643 NSCs (8/13 cases), or with only high PLC (5/13 cases). NSC data are means for all sampled 644 tissues available for each case and normalized as a percent of difference from concurrent 645 measurements of control trees. PLC data are those shown in Figure 1A. NSC and PLC at 646 mortality for angiosperm (blue circles) and gymnosperm (red triangles) species are shown 647 relative to hypothesized drought mortality mechanisms. Numbers near points designate original 648 studies (Supplementary Table 1). Error bars are one standard error.





**Xylem Embolism Resistance** 

