

1 **Coordination of physiological traits involved**
2 **in drought-induced mortality of woody**
3 **plants**

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17 **Coordination of physiological traits involved in**
18 **drought-induced mortality of woody plants**

19

20 **Abstract**

- 21 • Accurate modelling of drought-induced mortality is challenging. A steady-state model is
22 presented integrating xylem and phloem transport, leaf-level gas exchange and plant
23 carbohydrate consumption during drought development.
- 24 • A Bayesian analysis of parameter uncertainty based on expert knowledge and literature
25 review is carried out. The model is tested by combining six data compilations covering
26 170 species using information on sensitivities of xylem conductivity, stomatal
27 conductance and leaf turgor to water potential.
- 28 • The possible modes of plant failure at steady-state are identified (i.e., carbon starvation,
29 hydraulic failure and phloem transport failure). Carbon starvation occurs primarily in the
30 parameter space of isohydric stomatal control, whereas hydraulic failure is prevalent in
31 the space of xylem susceptibility to embolism. Relative to carbon starvation, phloem
32 transport failure occurs under conditions of low sensitivity of photosynthesis and high
33 sensitivity of growth to plant water status, respectively.
- 34 • These three failure modes are possible extremes along two axes of physiological
35 vulnerabilities, one characterized by the balance of water supply and demand and the
36 other by the balance between carbohydrate sources and sinks. Because the expression of
37 physiological vulnerabilities is coordinated, we argue that different failure modes should
38 occur with roughly equal likelihood, consistent with predictions using optimality theory.

39

40 Key-words: phloem transport, xylem embolism, phloem viscosity, photosynthetic down-
41 regulation, water stress, drought-induced mortality, source-sink relationships, optimality.

42

43 Introduction

44

45 Mortality of plants as a consequence of drought events has become a major focus of attention
46 recently as a result of reports highlighting severe mortality episodes around the globe (e.g., Allen
47 et al., 2010; Peng et al., 2011). Current process-based models do not adequately represent local
48 and regional mortality, because they have been constructed primarily to represent the fluxes of
49 carbon, water and nutrients and because they have not been calibrated against datasets of severe
50 drought episodes (Powell et al., 2013; McDowell et al., 2013). Consequently, doubts exist as to
51 their capacity to predict shifts in vegetation composition a consequence of increased drought
52 frequency and intensity (e.g., Anderegg et al., 2012, Adams et al., 2013; Xu et al., 2013; Meir et
53 al., 2014).

54 One central element of uncertainty is given by the lack of detailed understanding of the
55 environmental, ecological and physiological processes leading to mortality (McDowell et al.,
56 2013). The existing datasets which have documented plant mortality paying sufficient attention
57 to some of the underlying physiology (e.g., Adams et al., 2009; Fisher et al., 2010; Anderegg et
58 al., 2012; Hartmann et al., 2013; Mitchell et al., 2013; Poyatos et al., 2013) differ in the emphasis
59 given to different aspects of the mortality process. Partly as a consequence, different
60 interpretations of the main processes affecting mortality have emerged (McDowell et al., 2008;
61 Adams et al., 2009; Sala, 2009; Sala et al., 2010; McDowell & Sevanto, 2010). In addition, biotic
62 interactions can interact significantly with the physiological status of the plants to increase the
63 chances of drought-related mortality (Dobbertin & Rigling 2006; Wermelinger et al., 2008;
64 Galiano et al., 2011; Heiniger et al., 2011; McDowell, 2011; Zweifel et al., 2012; Krams et al.,
65 2012; McDowell et al., 2013).

66 Being able to avoid death is arguably the most important attribute that living organisms must
67 possess to reach reproductive age and transmit their genes to future generations. A tenable
68 assumption is that, over evolutionary times, plants have adopted strategies that minimise their
69 chances of failing quickly in response to multiple abiotic hazards such as drought (e.g., Anderegg
70 et al., 2013). From this perspective, the threats of mortality caused by failure of the hydraulic
71 transport systems (xylem or phloem) and of starvation caused by lack of carbon can be viewed as
72 possible extremes across a continuum of physiological vulnerabilities (Meir et al., 2014). From an
73 evolutionary perspective, the expression of functional traits might be optimally coordinated to
74 minimise the chance that any one source of mortality risk prevails. If that was not the case,
75 plants would arguably be over-built with respect to the risk posed by individual hazards.

76 The present work has three main objectives. Firstly, we present a steady-state model that
77 incorporates many of the processes involved in drought-induced mortality, with an emphasis on
78 the interaction between water and carbon fluxes. In the framework proposed by McDowell et al
79 (2008), the central distinction is between length and intensity of drought events, mediated by the
80 degree of isohydric regulation of water potential. Here, we expand that analysis. Secondly, we
81 explore the biological parameter space of the model, which constrains the range of water- and
82 carbon-related processes leading to physiological failure and mortality. Finally, we employ
83 empirical data to test the optimality idea set out above, that mortality risks should be equally
84 likely across species, thanks to the coordination of the relevant functional traits. A steady-state
85 model has distinct advantages compared to time-dependent approaches, because assumptions
86 about poorly known processes (such as thresholds and regulatory dynamics of carbohydrate
87 pools) are avoided and because the number of parameters is small enough that fitting to
88 empirical datasets with quantified uncertainty is possible (Meir et al., 2014). It suits our objective
89 to determine the trait set involved in mortality, rather than predicting the time courses to death.

90

91 **Description**

92 **Model structure**

93 The steady-state model develops a previously published coupled xylem and phloem transport
94 model (Hölttä et al. 2009a). Definitions, symbols, units and choice of values for all the
95 parameters employed in the model are given in Tables 1 and 2 (for the parameters whose values
96 were changed and those that were kept fixed, respectively). A diagrammatic representation of
97 model structure is given in Fig.1, with the represented processes individually numbered. In the
98 two parallel transport systems of the xylem and the phloem, axial hydraulic conductances of all
99 vertical elements are calculated from cross-sectional areas and hydraulic conductivities.
100 Following Minunno et al. (2013), we determined the number of finite elements required to
101 resolve the system's nonlinear responses. We progressively shortened the number of elements of
102 the catena from 100 to 10. For all state variables, the difference in outputs between catenas with
103 100 and 40 elements was very small. The difference in outputs between catenas with 40 and 10
104 elements was less than 5%. The final simulations were carried out with 100 elements.

105 The xylem water pressure at each element is calculated from the water pressure of the element
106 underneath it (for the bottommost element of the catena, this is the soil water potential, Ψ_{soil} in
107 Fig.1) minus the effects of gravity and the viscous pressure losses caused by xylem sap flux (F_{xyl}
108 in Fig.1). For each vertical phloem element, equations of radial water exchange with the xylem
109 (F_{radial} in Fig.1), phloem axial sap flow (F_{ph} in Fig.1), water conservation and solute conservation

110 are written (Hölttä et al., 2009a). The boundary condition at the bottommost element of the
 111 phloem (the ‘sink’) is such that the sugar unloading rate at the sink maintains a "target" turgor
 112 pressure (U_{100} in Fig.1). Different values of this target turgor pressure were employed, with their
 113 range given in Fig.S1A. The viscosity of the phloem sap is a function of its sucrose concentration
 114 at each element using an equation describing this dependency accurately up to osmotic potentials
 115 of about -8 MPa (cf., Morison, 2002 and the green dashed double-arrowed link in Fig.1 linking
 116 phloem sucrose osmotic potential c_{ph} with phloem conductance K_{ph}). This is an essential feature
 117 of the Hölttä et al. (2009a) model resulting from sucrose being the only solute transported and
 118 also the cause for the viscosity increases. It predicts a point of potential vulnerability for the
 119 phloem if the system fails to transport all the products of photosynthesis. A very dilute solution
 120 minimizes viscosity but requires large volume fluxes, while a very concentrated solution
 121 minimizes volume fluxes but increases viscosity. Jenssen et al (2013) showed that this problem
 122 leads to an optimal solute concentration that is broadly consistent with the concentrations
 123 normally measured in plants under well-watered conditions (cf., Lang (1978) and Hölttä et al.,
 124 (2009a), for similar arguments). In practice, it translates into a vulnerability curve for the phloem
 125 as a function of phloem osmotic potential (Fig.S1B), equivalent to the one for the xylem as a
 126 function of xylem water potential. The two main parameters affecting the shape of this phloem
 127 vulnerability curve are maximum phloem hydraulic conductance and the type of transported
 128 osmoticum.

129 Simulations are driven only by soil water potential (MPa), while transpiration and photosynthesis
 130 do not depend on other environmental variables. For each value of soil water potential, a steady-
 131 state solution is first found for the xylem water potential profile, PLC and stomatal conductance,
 132 by iterating equations (1) to (5) below plus Darcy’s law, until water potential of the uppermost
 133 element varies by less than 0.001MPa. The procedure is repeated to find steady-state values of
 134 phloem transport rates, photosynthesis and respiration, following an approach similar to the one
 135 presented in Hölttä et al. (2009a). The model normally converges very quickly and 500,000 runs
 136 take a few hours on a desktop computer.

137

138 **Xylem vulnerability to cavitation**

139 Xylem conductance is assumed to decrease with decreasing water potential according to
 140 (Pammenter & Willigen 1998)

$$141 \quad k_x = k_{0,x} (1 - PLC_i) \quad (1)$$

142 where

$$143 \quad PLC = \frac{1}{(1 + \exp(A_x(\Psi - B_x)))} \quad (2)$$

144 In Equation (2), Ψ is 'xylem' water potential at any point in the catena and the parameter B_x can
 145 be interpreted as the water potential at which xylem conductance reaches 50% of its maximum
 146 value (referred to as P50, Pammenter & Willigen 1998). Parameter A_x represents instead the
 147 slope of the relationship, i.e., the gradient of change in PLC with changes in water potential. It
 148 has been shown (Cochard 2006, Choat et al., 2012) that these two parameters are related, i.e.,
 149 plants with vulnerable xylem (i.e., high B_x) also have steep vulnerability curves (high A_x) and vice
 150 versa. Two examples of curves drawn with extreme values of A_x and B_x taken from the sampled
 151 distribution are given in Fig.S1C. Beyond its sensitivity to xylem water potential, plant
 152 conductance in Eqn. (1) also depends strongly on the value of maximum hydraulic conductance
 153 $K_{0,x}$.

154

155 **Transpiration rate and stomatal conductance**

156 Transpiration rate is represented as (e.g., Jarvis & McNaughton 1986)

$$157 \quad T = g_s T_0 \quad (3)$$

158 Maximum transpiration rate T_0 is given a fixed value of $2.25 \cdot 10^{-6} \text{ m}^3 \text{ s}^{-1}$ (i.e., $50 \text{ mmol m}^{-2} \text{ s}^{-1}$ for a
 159 25 m^2 tree) and the parameter space of suitable hydraulic values is varied by changing maximum
 160 plant hydraulic conductance $K_{0,x}$. Fig.S1D shows how the two most extreme values of the
 161 parameter $K_{0,x}$ coupled with T_0 affect plant water potentials, following Darcy's law.

162 While water flux affects xylem Ψ , leaf Ψ affects stomatal conductance g_s , reducing the chances of
 163 extremely low water potentials. Similarly to the case for xylem hydraulic conductance, stomatal
 164 conductance declines with plant water potential following a sigmoidal curve:

$$165 \quad g_s = g_{s,0} (1 - PLC_{g_s}) \quad (4)$$

166 and

$$167 \quad PLC_{g_s} = \frac{1}{(1 + \exp(A_{g_s}(\Psi - B_{g_s})))} \quad (5)$$

168 where $g_{s,0}$ is set to 1.00 and g_s is constrained to vary in the range $0 \leq g_s \leq 1$ in Eqns. (3) and (4). In
 169 Equation (5) above, Ψ is 'leaf' (the top element of the catena) water potential and the parameter
 170 B_{g_s} can also be interpreted as the leaf water potential at which stomatal conductance reaches 50%
 171 of its maximum value (cf., Tuzet et al. (2003) for a representation of the relationship between

172 photosynthesis and stomatal conductance that responds to leaf Ψ). Two extreme examples of
 173 the relationship employed here are given in Fig.S1E.

174

175 **Photosynthesis rate**

176 Photosynthesis rate is modelled as (Mäkelä et al. 1996)

$$177 \quad P = P_{\max} \frac{g_s}{g_s + \gamma} f_{ns} \quad (6)$$

178 where P_{\max} is a parameter setting the maximum photosynthesis rate, g_s is the stomatal
 179 conductance (in relative units from 0 to 1), γ is a parameter describing the saturation of
 180 photosynthesis with respect to stomatal opening, and f_{ns} is a factor accounting for the down-
 181 regulation of photosynthesis as a function of the osmotic pressure at the source (not included in
 182 Mäkelä et al., 1996). Albeit empirical (cf., von Caemmerer & Farquhar 1981), equation (6)
 183 incorporates the effects of stomatal aperture and of sink regulation of photosynthesis (e.g., Paul
 184 & Foyer 2001). In a preliminary analysis, we let parameter γ vary, but found that its effect on
 185 output variables was very small. We therefore kept it fixed in all analyses at a value of 0.2.

186 Because photosynthesis occurs at the top of the catena of phloem transport cells (the ‘source’)
 187 and sucrose is assumed to be loaded directly into the phloem, f_{ns} is dependent on the sucrose
 188 osmotic pressure of the first phloem cell c_1 at the top of the catena (where c_1 is in MPa). The
 189 effect of progressive concentration of the products of photosynthesis is therefore represented as:

$$190 \quad f_{ns} = \frac{c_{\max} - c_1}{c_{\max}} = 1 - \frac{c_1}{c_{\max}} \quad (7)$$

191 where c_{\max} is the parameter giving the maximum osmotic pressure of the phloem (MPa). If $c_1=0$,
 192 there is no down-regulation ($f_{ns}=1$); if $c_1=c_{\max}$, photosynthesis is depressed to zero ($f_{ns}=0$) to
 193 avoid further phloem loading. Fig.S1F gives a representation of this relationship using the two
 194 extreme values of P_{\max} and c_{\max} employed. Because of our steady-state assumption,
 195 photosynthesis rates and phloem loading rates are equal, unless the plant fails. In additional
 196 model runs, we compared this model against a representation of the down-regulation of
 197 photosynthesis by leaf water potential using the following formulation for f_{ns} :

$$198 \quad f_{ns} = \exp(P_{\text{mod}} \Psi) \quad (8)$$

199 where P_{mod} ($P_{\text{mod}} > 0$) is the parameter setting the direct sensitivity of P_{\max} to leaf water potential.

200

201 **Growth and respiration**

202 Growth and growth respiration are not explicitly separated from maintenance respiration, but we
 203 assume that the substrate is partially consumed during its transit along the phloem catena (e.g.,

204 Dewar, 1993; Cannell & Thornley, 2000). This approach is similar to the idea that ‘source’
 205 photosynthesis and ‘sink’ respiration are co-limiting processes, resulting in a near-homeostasis of
 206 phloem solute osmotic potential profiles as drought develops (i.e., Thornley & Johnson 1990;
 207 Minchin et al., 1993; Minchin & Thorpe 1996; Farrar 1996; Bancal & Soltani, 2002; Bijlsma &
 208 Lambers 2000).

209 The respiration rate R from each element i along the catena of phloem transport is assumed to
 210 be constant for values of the osmotic pressure c_i between 0 and c_{\min} . Above c_{\min} (i.e., for more
 211 positive osmotic pressures than c_{\min}), R increases linearly as a function of c_i . Therefore:

$$212 \quad R_i = R_0 = 0.10 \frac{P_{\max}}{N}$$

$$213 \quad R_i = R_0 \frac{c_i}{c_{\min}} \quad (9)$$

214 For $c_i \leq c_{\min}$ and $c_i > c_{\min}$, respectively (Fig.S1G). R_0 is the base respiration rate at minimum levels
 215 of substrate availability. The central value of R_0 was set at 10% of P_{\max} divided by N , the number
 216 of elements of the catena. Uncertainty in this parameter was introduced by Monte Carlo
 217 sampling of the parameter space (cf., later on). In the baseline scenario, we assumed no direct
 218 dependency of respiration on plant water status. In additional simulations (cf., Supplementary
 219 Materials, Section C), a direct dependency of respiration on water potential was introduced using
 an additional parameter, as done above for P_{\max} , i.e.:

$$220 \quad R_i = R_0 \exp(R_{\text{mod}} \Psi) \quad (10)$$

221 where R_{mod} ($R_{\text{mod}} > 0$) is the parameter setting the direct sensitivity of R_0 to water potential.

222
 223 In the baseline simulations, increased phloem concentrations during drought always lead to
 224 increased respiratory losses (cf., Eqn.9 above). The dependency on water potential of Eqn. (10),
 225 either alone or in combination with Eqn. (9), allows for the moderating effects of low plant
 226 water status on plant respiration when phloem concentrations are high.

227 Because we assumed a constant sink turgor pressure (see above), each solution for steady-state
 228 photosynthesis and respiration resulted in a certain amount of carbohydrates not being employed
 229 for respiration and being unloaded at the sink. We refer to this fraction as F_{resid} , the residual flux
 230 of transported carbohydrates. This metric is useful as an indicator of carbohydrate availability or
 231 potential carbohydrate storage, as it represents the fraction produced in the leaves, transported
 through the phloem, not respired by the catena and unloaded at the sink.

232

233 **Definition of modes of failure**

234 Some of the processes represented in the model contain negative feedback loops that tend to
 235 stabilize plant performance and avoid run-away failure (red arrows in Fig.1). Two of the
 236 processes (drought-induced cavitation in the xylem and viscosity-induced reductions of
 237 conductance in the phloem, green arrows in Fig.1) are destabilising feedback loops that can lead
 238 to failure. We classified the possible modes of plant failure at steady state as:

239 1. Hydraulic failure (HF). A combination of parameters was assumed to lead to HF when
 240 the calculated rate of xylem hydraulic conductance fell to zero as a result of complete
 241 xylem cavitation, i.e.,

$$242 \quad K_x = 0 \quad (11)$$

243 2. Carbon starvation (CS). A combination of parameters was assumed to lead to CS when
 244 the calculated steady-state rate of photosynthesis was lower than the steady-state rate of
 245 respiration by the catena, i.e.,

$$246 \quad P < R_{\text{tot}} \quad (12)$$

247 Because respiration was calculated for each element of the model separately, R_{tot}
 248 represents the sum of the N respiratory terms. By definition, $F_{\text{resid}} = 0$ when $P \leq R_{\text{tot}}$.

249 3. Phloem transport failure (PF). A combination of parameters was assumed to lead to PF
 250 when the rate of photosynthesis was greater than the rate at which carbohydrates could
 251 be transported out of the leaf as a result of excess phloem viscosity, i.e.,

$$252 \quad F_{\text{ph}} < P \quad (13)$$

253

254 The definitions of such modes of failure need to be interpreted in the narrow sense that is
 255 consistent with the use of a steady state model, as opposed to the broader definitions applicable
 256 to the field. For example, the definition of CS above should be relaxed to the broader negative
 257 carbon balance under prolonged non-steady state conditions, because a negative carbon balance
 258 during a short time period does not necessarily lead to failure. Similarly, the narrow criterion of
 259 PF for steady state conditions should be relaxed to the broader lack of equilibrium between
 260 photosynthesis and phloem transport (and therefore changing storage pools) under non-steady
 261 state conditions.

262

263 **Exploration of parameter space**

264 Of the 17 model parameters, eleven have the potential to affect the likelihood and the mode of
 265 plant failure. The behaviour of 11 of these parameters (13 including P_{mod} and R_{mod}) was examined
 266 by carrying out a prior parameter uncertainty quantification (van Oijen et al., 2013) to determine
 267 the sensitivity of model outputs to uncertainty in the global parameter space, as opposed to

268 changes in individual parameters (i.e., Beven and Binley, 1992). We defined the prior parameter
269 space based on literature estimates. We examined compilations that summarised hydraulic traits
270 for different biomes and plant functional types (cf., Notes S1). For each compilation, we
271 extracted the range of the main hydraulic parameters to set the limits of our prior distributions.
272 Values of maximum photosynthetic rates were constrained based on values from the
273 GLOPNET database (Wright et al., 2005).

274 We used log-normal distributions for our sampled parameter space (Table 1 and Hölttä et al.,
275 2009a), with 95% of the values within limits obtained by multiplying and dividing the central
276 estimate by 10. Parameters were generally sampled using univariate log-normals. Multivariate
277 lognormals were sampled using the function `mvnorm` in the library MASS (Venables & Ripley
278 2002) in R 3.0.2 (R Development Core Team 2013) for the parameters related to xylem
279 vulnerability curves and for those related to the response of stomatal conductance to water
280 potential. For the first set of parameters (i.e., A_x and B_x and K_x), the covariances ensured that A_x
281 and B_x were positively and curvilinearly related (Cochard 2006; Choat et al., 2012) and that high
282 values of B_x (i.e., values of P50 close to zero) corresponded to high values of xylem K . For the
283 second set of parameters, the covariance ensured that A_{gs} and B_{gs} were similarly positively but
284 loosely related (Manzoni et al., 2013; 2014).

285 Sampling was repeated 500,000 times. For each of the 500,000 parameter combinations, a
286 drought sequence was imposed on the model plant, starting from a soil water potential of -
287 0.005 MPa and continuing in steps of 0.005 MPa. At each step, the model calculated the steady-
288 state values of all state variables and checked whether the three conditions defining the modes of
289 failure (Eqn. 11, 12 and 13) were encountered. If steady-state values could be found for all state
290 variables and none of those conditions were satisfied (i.e., if $P=F_{ph}>R_{tot}$ and $K_{xyl}>0$), the soil
291 water potential was lowered. This process continued until a value of soil water potential was
292 reached at which one of the conditions above was satisfied. At this point, failure was deemed to
293 have been reached as CS, HF, or PF.

294 The 500,000 combinations of initial parameter values, output variables and classified modes of
295 failure were screened to eliminate runs that were clearly outside the range of realistic values
296 ('non-behavioural simulations'; Beven and Binley, 1992). This was accomplished by selecting
297 limits to two variables, i.e., leaf water potential and water use efficiency. Runs were given a
298 probability of 1 only if: a) steady-state values of 'leaf' water potentials Ψ at a soil water potential
299 of -0.005 MPa were within the range $-3.0 < \Psi < -0.2$ MPa, and b) the internal water use efficiency
300 (i.e., the ratio of assimilation divided by stomatal conductance) did not decrease between the soil
301 water potential of -0.005 MPa and the critical soil water potential at failure. Alternatively, runs

302 were given a probability of 0. Condition a) ensured a loose coupling between transpiration rate T
303 and xylem hydraulic conductance, forcing realistic values of water potentials. Condition b)
304 ensured that those parameter combinations resulting in reductions in internal water use
305 efficiency during a drought (caused by, e.g., a combination of stomatal conductance being very
306 insensitive to leaf water potential and photosynthesis rate being very sensitive to calculated
307 sucrose concentrations or leaf water potential) were excluded.

308

309 **Model sensitivity analyses**

310 To determine the sensitivity of model outputs to input parameters, we conducted a canonical
311 correlation analysis (CCA, Hair et al., 1998). CCA is a multivariate technique allowing the study
312 of the relationships among sets of correlated multiple dependent (model outputs) and
313 independent variables (model parameters, cf., Notes S1 and Table S2). In addition, we
314 determined the sensitivity of the frequency distributions of the three failure modes to the model
315 boundary conditions and carried out additional simulations varying model parameters that were
316 kept fixed for all the other runs (i.e., phloem radial hydraulic conductance, tree height, degree
317 and direction of correlations between stomatal and xylem parameters). Finally, we compared
318 these results with those obtained after introducing a direct dependency of basal respiration rate
319 and/or maximum photosynthetic rate on plant water potential.

320

321 **Empirical data analysis**

322 To analyse model behaviour, we used studies that reported values of the sensitivity of xylem
323 conductivity to Ψ , of stomatal conductance to leaf Ψ and of leaf turgor to Ψ (Choat et al., 2012;
324 Bartlett et al. 2012; Manzoni et al., 2013; Nardini & Luglio 2014; Klein 2014; Manzoni et al.,
325 2014). Six additional species came from Vilagrosa et al. (2014). The P50 values given by Choat et
326 al. (2012), Vilagrosa et al. (2014), Klein (2014) and Manzoni et al. (2013) were directly equated
327 with B_x . Manzoni et al. (2013) and Klein (2014) directly reported B_{gs} , using stomatal conductance
328 and sap flux data against leaf Ψ . A significant overlap in the species coverage of these two
329 datasets was found, even though absolute values of B_{gs} were frequently different between them.
330 The Manzoni et al. (2014) dataset is an expanded version of the Manzoni et al. (2013) version.
331 Bartlett et al. (2012), Nardini & Luglio (2014) and Vilagrosa et al. (2014) reported Ψ_{tp} (water
332 potential at turgor loss point, i.e., the Ψ at which leaves, on average, lose turgor). Ψ_{tp} is an index
333 of plant resistance to water stress and does not directly control the dependency of stomatal
334 conductance to water potential. Estimates of B_{gs} obtained from the relationship between sap flux
335 data and water potentials have similar limitations. Values of Ψ_{tp} were only assumed proportional

336 to B_{gs} and the assumption of proportionality between Ψ_{tp} to B_{gs} was tested in three ways. Firstly,
337 we let the proportionality coefficient between Ψ_{tp} and B_{gs} vary between 0.3 and 1.0 and we
338 checked whether changes in these coefficients affected our conclusion on the distribution of
339 species values in model parameter space (cf., Notes S2, Tables S4-S5 and Figures S2-S3).
340 Secondly, we checked databases for species with pairs of values of B_{gs} and Ψ_{tp} . We found 14
341 species, giving a correlation coefficient of 0.57 ($P < 0.05$), confirming that a relationship between
342 the two estimates can be postulated. Thirdly, to avoid systematic biases, we employed additional
343 categorical variables ('dataset' and 'method'), to test the effects of the individual datasets and of
344 the two methods employed to calculate B_{gs} . We crossed these seven data-sets for common
345 species, checked nomenclature, standardised definitions for biome and eliminated duplications
346 for individual species by value averaging. Plants were separated into the groups of angiosperms
347 and gymnosperms. Coupled values of B_x and B_{gs} were found for 243 independent observations
348 and 170 species across all compilations. The relationship between B_x and B_{gs} was tested using a
349 general linear model in R 3.0.2 (R Core Development Team, 2013), using 'dataset', 'biome' and
350 'plant group' as additional categorical factors.

351

352 **Results**

353 **Sensitivity analyses and distributions of simulations by failure modes**

354 The boundary conditions selecting the 'behavioural' simulations screened out a significant
355 number of parameter combinations (92% in the baseline case). Of the simulations that were
356 retained under the baseline case, 25% resulted in HF, 71% in CS and only 4% in PF. These
357 proportions varied greatly (cf., Table S3) depending on the imposed boundary conditions,
358 especially tree height (varied between 1m and 100m) and radial hydraulic conductance (varied
359 between $2 \cdot 10^{-13}$ and $2 \cdot 10^{-9}$). The parameter that most affected the frequency distributions of the
360 failure modes was the dependency of plant respiration on water potential. Including this
361 additional parameter (varied from $0.1 \cdot 10^{-6}$ to $1 \cdot 10^{-6}$) increased the proportion of HF (from
362 generally <20 to >30%) and PF (from ~5 to >10%) at the expense of CS (from >75 to <60%).

363 Model output variables showed sensitivity to a range of parameters for the first five canonical
364 variates (cf., Table S2 in Notes S1). Two parameters with opposing effects (i.e., xylem K and the
365 slope of the stomatal response to Ψ) affected almost the entire set of output variables. Plant
366 Failure mode was primarily related to xylem K, the slope of the stomatal response to Ψ and both
367 xylem and stomatal P50.

368

369 **Distribution of parameters**

370 The general distribution of the input parameters by mode of failure is given in Fig.2. The last
 371 columns (in red) are the reference empirical distributions for those parameters for which data
 372 were available from the meta-analyses. In general, the distributions obtained for the three failure
 373 modes (in black) encompassed the distributions from the empirical compilations (in red). Xylem
 374 conductance K_x showed a significant difference ($P < 0.001$) in the parameter distribution between
 375 the three modes of failure, with higher values for PF than CS. An even more accentuated
 376 difference was found for B_x (xylem P50) and A_x , with much higher values found for mode HF,
 377 followed by CS and PF ($P < 0.001$). Conversely, B_{gs} (stomatal P50) and A_{gs} showed higher values
 378 for CS ($P < 0.001$), with no difference between HF and PF. PF was characterized by a
 379 combination of parameter distributions, i.e., relatively high K_x , low B_x and A_x (both $P < 0.001$),
 380 large c_{max} ($P < 0.001$) and relatively higher c_{min} and sink turgor U . The distributions of the input
 381 parameters by mode of failure did not vary by varying the boundary parameters in the sensitivity
 382 analysis (data not shown).

383 The distribution of the main output variables at failure showed (Fig. 3) that soil and leaf Ψ varied
 384 across modes of failure, with significantly more negative values for PF ($P < 0.001$). This was
 385 associated with higher source turgor pressures ($P < 0.001$), more negative osmotic potentials (in
 386 turquoise, $P < 0.001$, as expected for PF by viscosity) and larger turgor drops from leaves to sink
 387 ($P < 0.001$). Parameter combinations that resulted in HF showed 100% loss of xylem conductance
 388 (in turquoise), almost complete stomatal closure and no photosynthesis. Relative to PF, CS was
 389 characterised by lower photosynthetic rates at failure (but not by higher cumulative respiration)
 390 and lower cumulative residual fraction of transported carbohydrates at the sink (in turquoise, as
 391 expected for this mode of failure). The range of soil water potentials at failure did not differ
 392 between HF and CS. The distributions of the output variables at failure varied only marginally by
 393 varying the boundary parameters (data not shown).

394

395 **Controls on modes of failure**

396 A plot of xylem P50 versus stomatal P50 separated HF versus CS (Fig.4). HF was characterized
 397 by points distributed at the top of the space delimited by B_x (xylem P50), whereas CS was
 398 characterized by points distributed on the side of the parameter space characterized by high
 399 values of B_{gs} (stomatal P50). Around a diagonal space from top right to bottom left (i.e., from
 400 sensitive stomata plus vulnerable xylem to insensitive stomata plus resistant xylem), a relatively
 401 wide region of overlap between the two modes of failure was found. Fewer points were found in
 402 the left bottom corner of the parameter space. For each of the two modes of failure, plant water

403 potential at failure depended on parameter combinations. Low (negative values) of xylem P50
404 resulted in low critical leaf water potentials for the case of HF. Similarly, for the case of CS, low
405 values of stomatal P50 resulted in low critical leaf water potentials.

406 When the 170 species from the meta-analytical compilations were plotted on the B_x - B_{gs} space (as
407 in Fig.4), the vast majority of the points fell within a region covering the bivariate 99% ranges of
408 these two modes of failure around the main diagonal line (Fig.5). A significant positive
409 relationship was found between xylem P50 and stomatal P50 across all datasets (all $P < 0.001$,
410 depending on the assumed relationship between stomatal turgor loss point and B_{gs} , cf., Table 3
411 and Tables S4-S5 in Notes S2), with a significant negative intercept for the gymnosperms
412 ($P < 0.001$), indicating a lower P50 (between about -1.1 and -1.9 MPa) for a fixed stomatal P50.
413 Highly significant effects were also found for 'dataset' (with significant differences for the
414 Vilagrosa dataset, $P < 0.001$) and 'biome' (with significant differences for the dry sclerophyllous
415 biome, $P < 0.001$). The overall model including stomatal P50, the three categorical variables and
416 their interactions explained between 59 and 60% of the variance (Table 3, S4 and S5). Despite
417 changes in the distribution and linear fits in Fig.5 depending on the assumption made for the
418 conversion between leaf stomatal P50 and Ψ_{tp} , the bulk of the data points remained in the area
419 of joint overlap between the two bivariate distributions of 99% of the simulations for HF and CS
420 (Figures S2 and S3 in Notes S2).

421 CS and PF differed for parameter combinations regulating plant carbon source-sink balance.
422 Because multiple parameters affected the photosynthetic and respiratory responses, composite
423 response parameters were calculated for each, following the response curves given in Eqns.9-15.
424 Relative to PF, CS was characterised by parameter combinations leading to a weak regulation of
425 respiration ($-R_0/(U^*c_{min})$, i.e., base respiration; degree of respiration down-regulation by osmotic
426 pressure -or water potential- and phloem turgor pressure, cf., Fig. 2) and a strong regulation of
427 photosynthesis ($P_{max} * B_{gs}/c_{max}$, i.e., maximum photosynthesis, sensitivity of stomatal closure to
428 water potential and photosynthetic down-regulation by osmotic pressure -or water potential-) in
429 response to water stress (Fig.6a). Conversely for PF, the combination of parameters regulating
430 carbon fixation, phloem transport and respiration during drought led to a less sensitive
431 regulation of carbon losses and to a more sensitive regulation of the sinks (Fig. 6b). This resulted
432 in combinations leading to PF being situated above the line of carbon supply/demand and those
433 leading to CS being situated below or on it (Fig. 6b).

434

435 Discussion

436 Model structure and major assumptions

437 The model incorporates many of the interactions among the processes of carbohydrate fixation
438 and transport and water transport and transpiration. By way of comparison, the Sperry et al
439 (1998) model includes a very detailed representation of the linkage between gaseous and liquid
440 water transport processes in the soil and the plant, but the processes linked to C fixation and
441 transport are not represented (cf., Mackay et al., 2012, for an advanced combination of water-
442 and carbon-related processes). Conversely, models by Cannell & Thornley (2000) and Dewar
443 (1993) represent C fixation and allocation using concepts related to source and sink strength, but
444 the biophysical representation of xylem and phloem transport is missing. Finally, the model by
445 De Schepper & Steppe (2010) is close to the approach presented here, but its focus is in
446 simulating short-term (minutes to hours) dynamics. The fundamental feature of this model is to
447 include both stabilizing and de-stabilizing processes for xylem and phloem. In the case of
448 phloem transport, the effect of viscosity on conductance is the main de-stabilizing process (cf.,
449 Hölttä et al, 2009), viscosity being a strong nonlinear function of sucrose osmotic concentration
450 (Morison, 2002).

451 We used a Bayesian approach based on literature information and expert knowledge to analyse
452 parameter and model output distributions. In our case, limits to parameter distributions were set
453 using global compilations of parameter values. In addition, screening criteria were set to create
454 boundaries for the parameter space ('behavioural' values). In Bayesian parlour, we constrained
455 partially informative priors by logical criteria based on expert knowledge. Criterion a) is well
456 supported in the literature (Mencuccini 2003; Martínez-Vilalta et al., 2014). Criterion b) is also
457 regarded as a universal observation.

458

459 **Co-ordination among modes of failure along water supply-demand axis**

460 A plot of xylem versus stomatal P50 discriminated between HF and CS (McDowell et al., 2008).
461 The distribution of these two modes of failure is delimited by a diagonal space going from
462 combinations of sensitive stomata plus vulnerable xylem to combinations of insensitive stomata
463 and resistant xylem. Inside this diagonal space, both types of failure occurred. The significance of
464 this diagonal space can be understood as follows. Firstly, the variability in xylem conductance
465 and stomatal conductance in relation to water and carbon fluxes depends on parameters that are,
466 at least to some degree, correlated with one another (e.g., A_x , B_x with K_x and A_{gs} with B_{gs}). This
467 reduces the dimensionality of the problem. Indeed our sensitivity analysis (cf., Table S2) showed
468 that failure mode was affected by a number of parameter combinations reflected in the
469 covariances mentioned above. Secondly, one would expect that plants evolved strategies to
470 minimise the relative risks caused by different mortality hazards. Traits that would cause plants

471 to be situated entirely within the space of only one dominant hazard type would likely be
472 evolutionary unstable. It is possible that different optimal solutions evolved such that different
473 sets of functional traits lead to roughly equal chances of mortality by different processes. For
474 example, levels of xylem PLC were higher than 90% for some of the simulations of CS (Fig.3),
475 while total cumulative F_{resid} were also comparatively higher for simulations of HF (Fig.3). One
476 may expect *a priori* that mortality be brought about by a coincidence of several different
477 processes. Recent experiments directly testing mechanisms of mortality show that a single
478 species can die by different causes depending on the circumstances (Sevanto et al., 2014). It is
479 interesting that the vast majority of the species for which empirical data were available were
480 contained within this diagonal space (Fig.5). This dataset of 170 species covered all major
481 biomes, climate conditions and plant types (Table S1). The significant terms for ‘dataset’ found
482 in the relationship between B_x and B_{gs} in the meta-analytical compilation suggests that caution is
483 needed when different datasets are combined. However, when tested, we did not find a
484 significant effect of the method employed to estimate B_{gs} (i.e., either from sap flow/conductance
485 measurements or from Ψ_{tp}) based on three different tests. This finding supports the use of Ψ_{tp}
486 as an indicator also of stomatal behaviour across species.

487

488 **Co-ordination among modes of failure along carbon supply-demand axis**

489 A plot of photosynthetic versus respiratory parameters discriminated combinations leading to CS
490 from PF. Interestingly many combinations could lead to both modes of failure. PF was
491 associated with an altered balance between carbohydrate sources (less sensitively regulated in
492 relation to drought) and sinks (more sensitively regulated) (Figure 6). This altered balance
493 produced larger residual carbohydrate fluxes and led to wider carbon safety margins (*sensu*
494 Mitchell et al., 2014). These results suggest that plants in which growth continues at low water
495 potentials may be more likely to suffer CS. *Vice versa*, plants may risk PF when consumption of
496 carbohydrates responds sensitively while stomatal and photosynthetic rates remain high during
497 drought. Interestingly, the proportion of combinations resulting in PF was strongly increased
498 when a direct sink limitation by plant water status was introduced (Table S2).

499 Evidence for the response of photosynthetic non-stomatal parameters to drought was recently
500 reviewed by Zhou et al (2013). While growth is very sensitive to turgor reductions, the response
501 of respiration to drought is more rarely documented. Duan et al. (2013, 2014) and Ayub et al.
502 (2011) found that leaf dark respiration declined only at the end of severe droughts while Metcalfe
503 et al. (2010) reported increases in stem respiration during drought in a tropical rainforest. The

504 response of growth to drought is almost never documented (cf., Mitchell et al., 2014 for an
505 exception).

506 The diagonal 1:1 line of Fig.6b is the line of source/sink balance. CS and PF can both be avoided
507 provided a plant can co-regulate source and sink activity with equal sensitivity during drought.
508 This appears to be possible for some, but not all, parameter combinations (cf., regions of overlap
509 between the two failure modes in Fig.6a). Combinations leading to HF were found well below
510 the 1:1 source/sink balance line of Fig.6b, i.e., in the same region as CS (data not shown). This is
511 because hydraulic regulations of stomatal conductance during drought led to stomatal closure
512 and lower photosynthesis, but not necessarily lower respiration rates. In our model, we assumed
513 that photosynthesis and respiration, but not F_{resid} , were actively controlled by plant water status.
514 In other words, the assumptions in our model are equivalent to the assumption that allocation to
515 carbohydrate storage is a residual term.

516 It is important to note that CS was affected also by phloem properties, albeit indirectly, via the
517 effects of changed phloem turgor, phloem osmotic potentials and phloem conductance. This is
518 supported by the results of the sensitivity analyses of Table S2 and S3. The osmotic and turgor
519 variables at failure (leaf osmotic pressure at failure, leaf turgor pressure at failure) were affected
520 by a combination of xylem, phloem and gas exchange parameters (Table S2).

521 Our steady state model constrains the solutions to a space where turgor is kept constant, but
522 phloem transport may also temporarily fail under dynamic conditions by reaching turgor loss for
523 limited but crucial time periods (e.g., McDowell et al. 2013, Sevanto et al. 2014). For example,
524 under drought, low photosynthesis may result in sucrose concentrations barely capable of
525 maintaining a positive turgor pressure.

526

527 **Non-steady-state behaviour and time scales to mortality**

528 How much would the conclusions drawn on the basis of Figs.5-6 change, had we incorporated
529 non-steady state conditions? It is likely that additional failure modes exist that can only be
530 identified under non-steady state conditions. However, the characterization of these additional
531 modes is prevented by our lack of mechanistic understanding of the underlying processes. In
532 addition, non-steady-state models tend to be parameter-rich and their calibration within known
533 uncertainty margins is difficult. Considering these limitations, a steady-state approach seems a
534 reasonable first approximation. In the context of the variables studied here, the behaviour of a
535 xylem hydraulic capacitor may primarily affect the magnitude of the declines in xylem water
536 potentials, slowing down xylem cavitation and HF. For example, Meinzer et al (2003) showed
537 that diurnal changes in plant water potential and sap flow can be moderated significantly as a

538 result of the presence of hydraulic capacitors and cavitation of xylem conduits may have
539 temporary moderating effects (cf., Hölttä et al., 2009b). Alternatively, a leaf capacitor may
540 primarily slow down the declines of water potential, thereby reducing stomatal closure and CS.
541 Dynamic carbohydrate storage under high photosynthetic rates may lower phloem loading and
542 prevent excessive solute concentrations (and viscosity) in the phloem but at the same time,
543 carbohydrate release may prevent dangerously low levels of sugar concentrations and loss of
544 turgor under conditions of long and intense respiratory losses. Empirical data are currently
545 unavailable to help tease out these possibilities.

546 Incorporating processes resulting in non-steady state conditions may be useful under significant
547 hydraulic disequilibrium between soil and plant. Several causes of hydraulic disequilibrium have
548 been reported (i.e., transient accumulation of solutes, lack of over-night equilibration in plant
549 hydration, continued night-time transpiration; cf., Donovan et al., 2003). Expanding this model
550 to include processes occurring during longer time periods would allow probing the significance
551 of progressive leaf shedding, changing rooting depth and root/shoot ratios, xylem growth and
552 refilling and cavitation fatigue.

553

554 **Conclusions**

555 The interpretation of mortality given here, of a process occurring along two independent axes
556 representing the dimensions of water supply/demand and carbon supply/demand differs
557 significantly from McDowell et al. (2008), where the primary axis driving mortality was the
558 degree of isohydric/anisohydric regulation of water potential. Stomatal behaviour turns out to be
559 just one component of a strategy that minimizes the risk of three different modes of mortality. A
560 plot of stomatal versus xylem P50 separated out the possible parameter combinations leading to
561 HF from those leading to CS. Conversely, PF could be separated from CS by parameter
562 combinations regulating phloem transport, respiration and photosynthesis. PF occurred
563 especially when growth was assumed to respond sensitively to plant water status while stomatal
564 regulation and photosynthetic down-regulation were limited. Maintaining phloem turgor via
565 regulation of osmotic pressure, and the link between solute concentration and viscosity were
566 crucial in understanding the relative sensitivity of growth and gas exchange to drought. With
567 regard to model validation, this exercise showed that only about half of the parameters currently
568 in the model could be constrained empirically. Some of the remaining parameters (i.e., sink
569 turgor pressure) can be constrained using analogous leaf or root turgor measurements
570 (Mencuccini M., Minunno F., Salmon Y, Poyatos R, Hölttä T, Martínez-Vilalta J., unpublished),
571 however empirical calibration remains difficult for others (e.g., phloem-related parameters).

572

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582

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763

764 **Legends of items in Supporting Information**

765 Table S1. Compilation of datasets of plant hydraulic traits by biome or plant functional types.

766 Notes S1. Sensitivity analysis of model outputs in relation to inputs.

767 Notes S2. Sensitivity analysis of definition of leaf turgor loss point as point of stomatal P50.

768 Figure S1. Illustration of the theoretical relationships used in the model.

769 Table S3. Sensitivity analysis of frequency distribution of failure modes.

770 Table 1. The 11 parameters employed to explore the sensitivity of model structure to the various
 771 modes of plant failure to drought. Each parameter is defined, the symbol and the units are given,
 772 as well as the central value employed in the simulations and the range of values sampled.

773

Parameter	Symbol	Units	50% percentile	2.5 and 97.5% percentiles of the distribution sampled
Maximum xylem hydraulic conductance	K_x	$\text{m}^3 \text{MPa}^{-1} \text{s}^{-1}$	2.42×10^{-6}	2.21×10^{-7} 3.68×10^{-5}
Maximum phloem hydraulic conductance	K_{ph}	$\text{m}^3 \text{MPa}^{-1} \text{s}^{-1}$	1.69×10^{-4}	1.86×10^{-5} 1.58×10^{-3}
Water potential Ψ causing 50% loss of K_{xyl}	B_x	MPa	-3.69	-14.38 -1.21
Water potential Ψ causing 50% loss of g_s	B_{g_s}	MPa	-0.79	-12.21 -0.11
Maximum photosynthesis	P_{max}	mol s^{-1}	6.62×10^{-5}	6.72×10^{-6} 6.02×10^{-4}
Leaf osmotic pressure at which P goes to zero	c_{max}	MPa	9.24	2.18 60.19
Base respiration rate	R_0	mol s^{-1}	3.36×10^{-6}	1.61×10^{-7} 6.05×10^{-5}
Osmotic pressure above which R begins to increase as a function of c	c_{min}	MPa	1.46	0.22 10.78
Slope of the xylem vulnerability curve	A_x	% PLC MPa^{-1}	2.25	0.11 3.43
Slope of relationship between stomatal conductance and water potential Ψ	A_{g_s}	% closure MPa^{-1}	8.09	1.34 30.52
Turgor pressure at the bottom of the phloem	U	MPa	0.63	0.09 3.43

774

775

776 Table 2. The six parameters of the model which were kept fixed in the simulations carried out to
 777 explore the sensitivity of model structure to the various modes of plant failure to drought. Each
 778 parameter is defined, the symbol and the units are given, as well as the fixed value employed in
 779 the simulations.

Parameter	Symbol	Units	Central value
Tree height	b	m	10
Phloem cross-sectional area	A_p	m ²	1.2*10 ⁻⁴
Xylem cross-sectional area	A_x	m ²	2*10 ⁻³
Xylem-phloem radial conductance	K_{rad}	m ³ Pa ⁻¹ s ⁻¹	2*10 ⁻¹¹
Transpiration rate at full stomatal opening	T_0	m ³ s ⁻¹	2.25*10 ⁻⁶
Slope of the photosynthetic response curve to stomatal conductance	γ	-	0.2

780

781

782 Table 3. Results of the general linear model employed to explain xylem P50 as a function of
 783 stomatal P50, dataset, biome and plant group. For the datasets based on estimates of turgor loss
 784 point (TLP), stomatal P50 was defined here as 70% of TLP (See text for further explanation and
 785 Tables S4/S5 for tests using different assumptions) (n=170, $R^2_{adj} = 0.60$). ***, $P < 0.001$.

786

	Degrees freedom	Sum Squares	Mean Square	F value	Prob (>F)
Stomatal P50	1	114.482	114.48	80.79	9.24 e-16***
dataset	5	214.34	42.87	30.25	< 2.2 e-16***
Biome	6	38.146	6.36	4.49	3.26 e-04***
Plant.group	1	51.027	51.03	36.01	1.38 e-08***
Biome * Plant.group	4	53.692	13.42	9.47	7.28 e-07***
Residuals	152	215.384	1.42		

787

788

789 **Figure Legends**

790 Fig 1. Diagrammatic representation of model structure. The two central open tubes indicate
 791 xylem and phloem transport (brown and green, respectively). P , photosynthesis; g_s , stomatal
 792 conductance; R , respiration, F_{ph} , F_{radial} and F_{xyl} , phloem, radial and xylem transport rates; c_{ph} ,
 793 phloem osmotic pressure; U_{100} , turgor pressure in unloading element; K_{ph} and K_{xyl} , phloem and
 794 xylem conductance; Ψ_{soil} and Ψ_{xyl} , soil and xylem water potential, respectively. Numbers 1 to 100
 795 inside the green (phloem) tube for R refer to the corresponding finite elements of the numerical
 796 model. Each of the blow-up circles represents one or more processes or feedbacks that are
 797 incorporated in the model. The progressive numbers from 1 to 14 inside the grey circles refer to
 798 the 14 processes represented in the model and discussed in the text.

799 Figure 2. Boxplot distribution of the 11 parameters varied in the model as a function of the three
 800 modes of failure (HF = hydraulic failure; CS = carbon starvation; PF = phloem transport
 801 failure). Boxplots provide mean and interquartile ranges for each parameter and each mode of
 802 failure. The first three boxes on the left in black give the modelled distributions, the last box on
 803 the right in red gives the distributions from the empirical data compilations, for those parameters
 804 for which empirical data were available. Symbols follow Table 1.

805 Figure 3. Boxplot distribution of 12 variables calculated at the soil water potentials at which the
 806 plants failed, as a function of the three modes of failure (HF = hydraulic failure; CS = carbon
 807 starvation; PF = phloem transport failure). Boxplots provide mean and interquartile ranges for
 808 each variable and each mode of failure. For each of the three modes of failure, the variable most
 809 closely associated with that mode is shown in turquoise in the respective plot (i.e., leaf osmotic
 810 potential for PF; xylem PLC for HF; residual flux (or cumulative stores) for CS). Note that the
 811 last three variables are plotted on log scale.

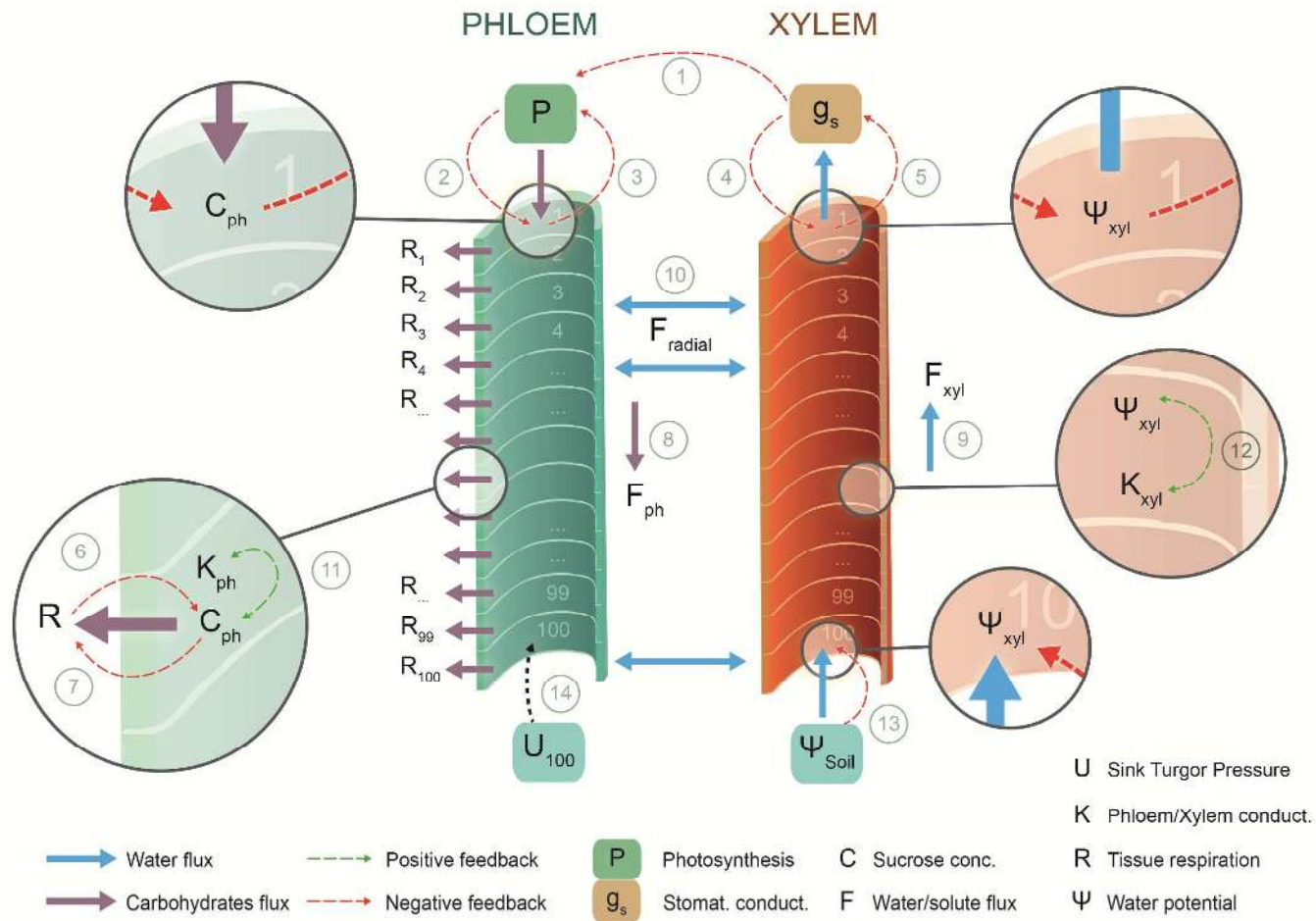
812 Figure 4. Distribution of the model simulations in the space defined by the xylem vulnerability to
 813 cavitation (P50) and the stomatal sensitivity to water potentials (stomatal P50) for the three main
 814 modes of failure (hydraulic failure, carbon starvation, phloem transport failure). For each panel,
 815 the color scheme follows the leaf water potentials at failure (with warmer colors indicating more
 816 negative values), following the legend in the first panel.

817 Figure 5. Distribution of the model simulations in the space defined by xylem P50 and stomatal
 818 P50 as per Figure 4. The two main modes of failure (hydraulic failure and carbon starvation) are
 819 highlighted with grey and pink points, respectively. Red contour lines indicate 99% relative
 820 densities of points for each distribution (e.g., less than 1% of the grey points is located outside

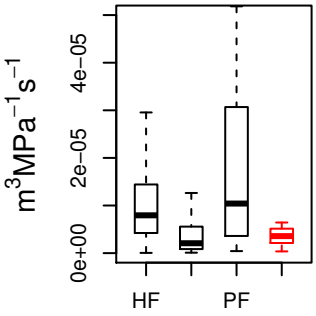
821 the corresponding thick red 1% contour). Red contour distributions are given separately for the
822 grey points (hydraulic failure) and the pink points (carbon starvation). The area of joint
823 occurrence of the two failure modes is therefore indicated by the intersection of the two 1%
824 contour red lines. Turquoise, green, black, red, pink and blue colours indicate boreal, tropical
825 evergreen, tropical seasonal, temperate evergreen, temperate deciduous and dry sclerophyllous
826 biomes, respectively. For each colour, circles indicate angiosperms and squares indicate
827 gymnosperms.

828 Figure 6. A) Distribution of the model simulations in the space defined by the combination of
829 parameters controlling respiration versus those controlling photosynthesis. The pink points
830 indicate the simulations resulting in CS, the black points those resulting in phloem transport
831 failure. Red and black contour lines indicate the respective 99% relative densities of points for
832 each distribution, as per Figure 5. The composite parameter controlling respiration is calculated
833 as $(-R_0)/(U^*c_{min})$. The composite parameter controlling photosynthesis is calculated as
834 $P_{max} * B_{gs}/c_{max}$. B). The values of photosynthesis and respiration at failure are given for the runs
835 resulting in CS (pink points) and phloem transport failure (black points). The blue line gives the
836 1:1 line of source-sink balance.

837

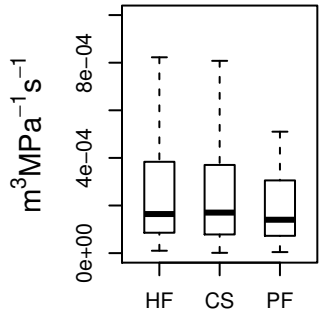


K_x



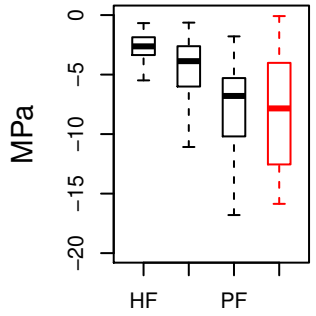
modes of failure

K_{ph} New Phytologist



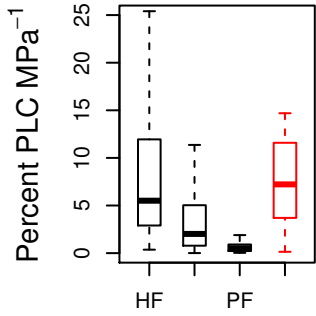
modes of failure

B_x



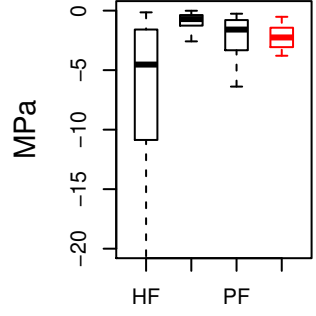
modes of failure

A_x



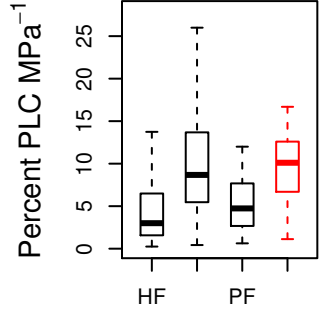
modes of failure

B_{gs}



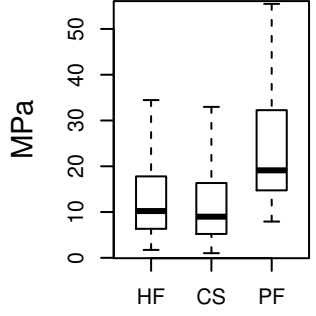
modes of failure

A_{gs}



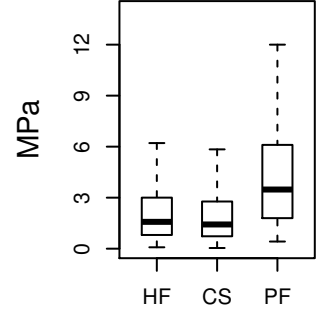
modes of failure

C_{max}



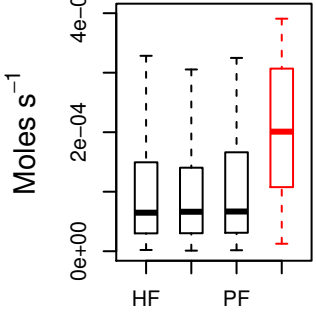
modes of failure

C_{min}



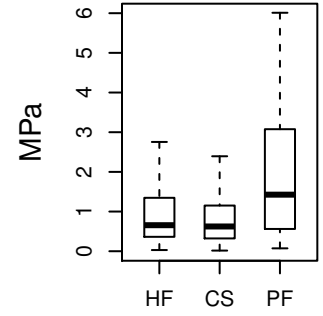
modes of failure

P_{max}



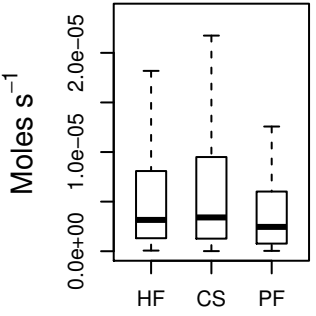
modes of failure

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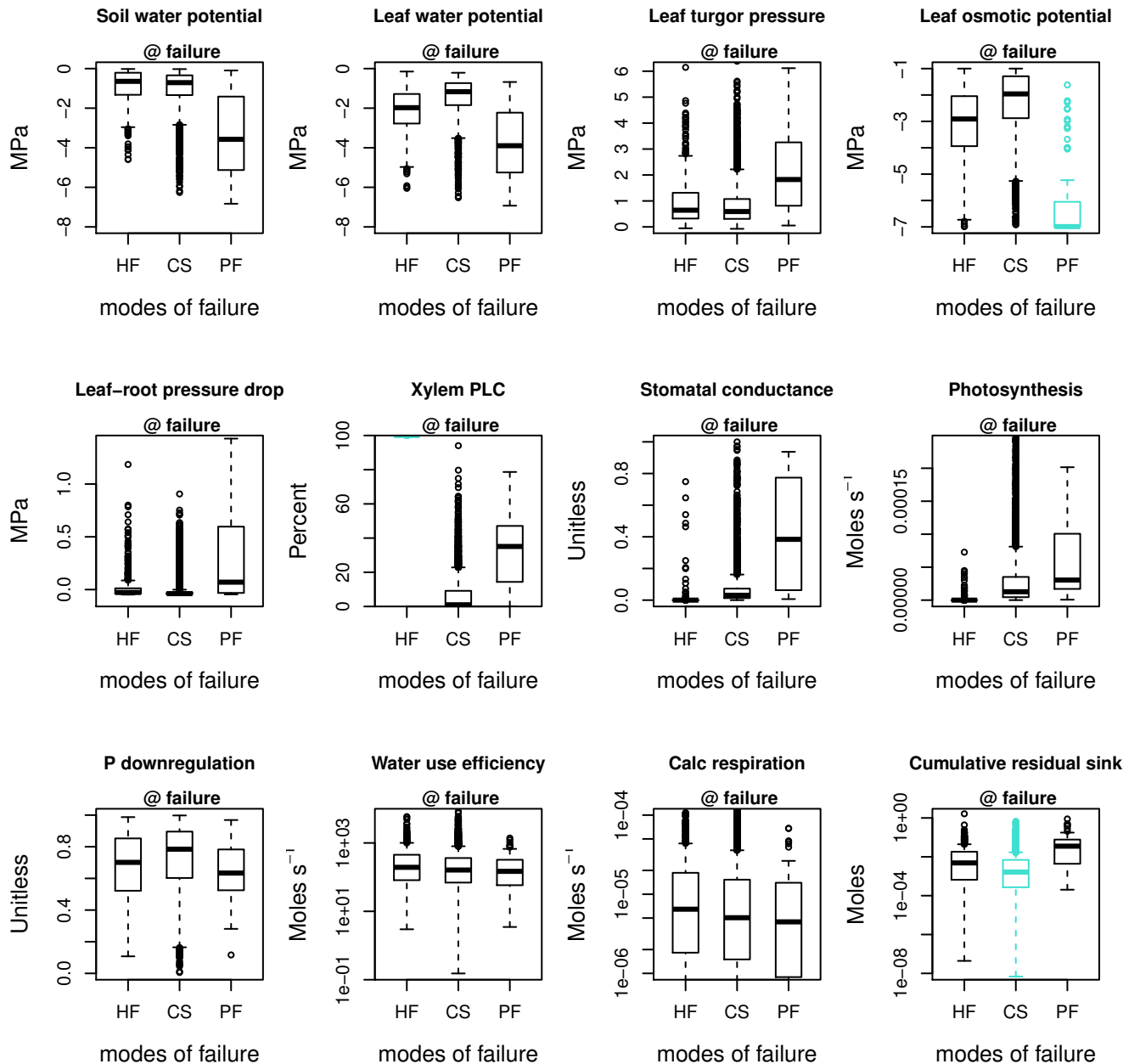


modes of failure

R_0



modes of failure



Coloured by levels of Leaf water potential at failure

