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1 **Death from drought in tropical forests is triggered by hydraulics not carbon starvation**

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18 Drought threatens tropical rainforests over seasonal to decadal timescales¹⁻⁴, but the drivers
19 of tree mortality following drought remain poorly understood^{5,6}. It has been suggested that
20 reduced availability of non-structural carbohydrates (NSC) critically increases mortality risk
21 through insufficient carbon supply to metabolism ('carbon starvation')^{7,8}. However little is
22 known about how NSC stores are affected by drought, especially over the long term, and
23 whether they are more important than hydraulic processes in determining drought-induced
24 mortality. Using data from the world's longest-running experimental drought study in tropical

25 rainforest (in the Brazilian Amazon), we test whether carbon starvation or deterioration of the
26 water-conducting pathways from soil to leaf trigger tree mortality. Biomass loss from
27 mortality in the experimentally-droughted forest increased substantially after >10 years of
28 reduced soil moisture availability. The mortality signal was dominated by the death of large
29 trees, which were at a much greater risk of hydraulic deterioration than smaller trees.
30 However, we find no evidence that the droughted trees suffered carbon starvation, as their
31 NSC concentrations were similar to those of un-droughted trees, and growth rates did not
32 decline in either living or dying individuals. Our results indicate that hydraulics, rather than
33 carbon starvation, triggers tree death from drought in tropical rainforest.

34

35 Drought-response observations from both field-scale experiments and natural droughts have
36 demonstrated increased mortality over the short-term (1-3 years), with notably higher
37 vulnerability for some taxa, and for larger trees^{6,9,10}. After several years of drought,
38 recovering growth rates in smaller trees, dbh (diameter at breast height) <40 cm, and reduced
39 mortality have been recorded at different locations^{6,11,12}. However, the long-term (>10 yr)
40 sensitivity of tropical forests to predicted prolonged and repeated water deficit¹⁻³ and the
41 physiological mechanisms influencing this are poorly understood. Through-fall exclusion
42 (TFE) studies, that create soil moisture deficit by the exclusion of a fraction of incoming
43 rainfall, provide the only current means to assess the long-term response in mechanistic
44 detail^{5,13}.

45 Trees experiencing drought stress are thought to die from direct physiological failure and/or
46 from injury and biotic attack associated with a decline in physiological vigour¹⁴. A global
47 effort to identify the relevant physiological mechanisms triggering death and thus to improve
48 predictions of forest tree mortality has focussed on the twin possibilities of: (i) failure to

49 supply sufficient carbon substrate to metabolism following drought-related reductions in
50 photosynthesis and increased use of NSC, theoretically leading to carbon starvation; and (ii)
51 deterioration of the water-conducting xylem tissue, causing a rapid or gradual failure of key
52 dependent processes (e.g., gas exchange, photosynthesis, phloem transport), and potentially
53 leading to tissue desiccation^{14,15}, ultimately leading to mortality. Despite recent intensive
54 research, it is unclear how important these two mechanisms are in different biomes and how,
55 or whether, to model them¹⁶.

56 Since 2002 a 50% TFE treatment has been implemented at a 1 ha-scale drought experiment in
57 old-growth forest at Caxiuanã National Forest Reserve, Pará State, Brazil^{6,12}, to simulate
58 possible rainfall reductions predicted to occur in parts of Amazonia by 2100¹. Mortality
59 surveys, recruitment and growth rates of all trees ≥ 10 cm dbh, have been monitored through
60 the experimental period (see Methods). Recently, seasonal data on NSC concentrations were
61 measured on leaves, branches and stems of 41 trees (20 trees on the control, 21 trees on the
62 TFE) of the most common genera in the experiment (Extended Data Table 1). Xylem
63 vulnerability curves were also performed on the branches of these trees (see Methods). Here,
64 we synthesise these data to test whether long-term soil moisture deficit alters NSC storage
65 and use in tropical rainforest trees, and if this, or hydraulic processes, are most strongly
66 associated with increased mortality rates.

67 By 2014, following 13 years of the TFE treatment, cumulative biomass loss through mortality
68 was $41.0 \pm 2.7\%$ relative to pre-treatment values (Fig. 1a), and the rate of loss had increased
69 substantially since the previous reported value of $17.2 \pm 0.8\%$, after 7 years of TFE⁶.
70 Accelerating biomass loss and failure to recover substantially, or to reach a new
71 equilibrium¹³, has led to a committed flux to the atmosphere from decomposing necromass of
72 101.9 ± 19.1 Mg C ha⁻¹ (Fig. 1a). This biomass loss has been driven by elevated mortality in

73 the largest trees (Fig. 1b), as previously observed over shorter timescales⁶, and has created a
74 canopy that has had a persistently lower average leaf area index during 2010-2014
75 (12.0±1.2% lower; Extended data Fig. 1).

76 Remarkably, individual tree growth rates for the four years prior to death showed no
77 significant reduction in either the TFE or control plots (Fig. 2a), indicating that growth is
78 prioritised to the point of death irrespective of the soil moisture deficit treatment. From 2008,
79 tree growth in every wet season (January-June) on the TFE treatment relative to the control
80 was significantly elevated ($P<0.05$) in the small and medium trees (up to 4.6±0.2 times
81 higher in small trees, and 2.9±0.2 times higher in medium trees), and maintained in the
82 largest trees (10-20 cm, 20-40 cm and >40 cm dbh, respectively; Fig. 2b-d). Elevated wet
83 season growth occurred despite 0.1-0.9 MPa reduction in average soil water potential (Ψ_s) at
84 depths of 0-4 m on the TFE and a loss of seasonality in Ψ_s (Extended Data Fig. 2). Increased
85 growth in the small trees occurred from 2008 onwards, following earlier substantial mortality
86 of large trees (Fig. 1), which generated canopy gaps. Increased light availability to smaller
87 trees, and presumably reduced below-ground competition for water and nutrients, allowed
88 competitive release of trees on the TFE⁸, and elevated growth rates. Competitive release on
89 the TFE implies that, following 13 years of drought-stress, photosynthetic production is
90 sufficient not only to maintain growth in the largest trees (Fig. 2d), but to increase growth in
91 trees <40 cm dbh (Fig. 2b-c). This response would not be possible if the majority of trees
92 were severely carbon limited, unless very considerable long-term (or renewed) carbon
93 resources were being drawn upon.

94 Prioritisation of growth under drought in the TFE is consistent with recent observations
95 following short-term drought in Amazonia⁷. However, the maintenance of NSC
96 concentrations in the TFE treatment suggests that the prioritisation of growth during drought

107 does not occur at the expense of depleted carbon stores, as previously hypothesised⁷. Neither
108 the concentrations of soluble sugar (carbon immediately available to metabolism) nor starch
109 (stored carbon which can be converted to sugars) were significantly depleted in stem, leaf and
110 branch tissue from the TFE, relative to control (Fig. 3). The seasonal changes in both sugar
111 and starch concentrations, which varied by 50-90%, were much larger than any differences
112 associated with the TFE treatment (Fig. 3). Despite 13 years of severely reduced soil moisture
113 availability, the seasonal cycle and use of NSCs was unaltered, implying that the sampled
114 trees did not draw significantly upon their NSC reserves to buffer against the long-term
115 effects of soil moisture deficit. Large changes in carbon allocation from roots and leaves to
116 maintain stem growth during drought¹⁷ have not been reported on the TFE¹². Similarly no
117 drought-induced reductions in photosynthetic capacity occurred on the TFE¹⁸, although how
118 total canopy productivity is affected remains uncertain. Considering this and additional
119 evidence of no increase in herbivore attack on the TFE (Extended Data Fig. 3), our results
120 suggest progressive carbon starvation and biotic foliar consumption are not important drivers
121 of the mortality patterns observed in the TFE forest following extended severe soil moisture
deficit (>10 years).

113 Deterioration of the water transport system in the xylem tissues following drought can also
114 lead to death^{17,19}. The vulnerability of the xylem to drought is described by a vulnerability
115 curve²⁰, which relates water potential in xylem conduits to loss of hydraulic conductivity
116 because of occlusions by gas emboli. The water potential at which 50% loss of xylem
117 conductivity occurs (P_{50} , MPa) is a commonly used index of embolism resistance²⁰. We
118 determined xylem P_{50} for the trees on the control and TFE plots, with tree dbh ranging from
119 15 to 48 cm. A highly significant decrease in P_{50} with dbh was found across TFE and control
120 (Extended Data table 3, $P < 0.01$). As dbh increased from 15 to 48 cm there was a 1.3 ± 0.2
121 MPa reduction in the P_{50} value, with significant genus-to-genus differences (Fig.4). Leaf

122 water potential (Ψ_1) could only be measured during limited sampling campaigns (2-3 days)
123 that were characterised by low vapour pressure deficit (VPD, 54-59% of peak dry season
124 values) and unseasonal rainfall in the preceding days. Differences between treatment and
125 control Ψ_1 were not detected. Mean midday Ψ_1 recorded across all the trees together with the
126 vulnerability curves determined for each genus were used to predict the percentage loss of
127 xylem conductivity (PLC) with dbh. Values of PLC at mean Ψ_1 increased with dbh, with the
128 largest diameter trees predicted to have reductions in conductive capacity of about 80% in
129 some genera, indicating significant vulnerability to hydraulic deterioration (inset of Fig. 4).

130 Given no evidence of carbon starvation and similar Ψ_1 across plots in the sample dates, why
131 did many more trees die in TFE than control? The lack of treatment differences in Ψ_1
132 contrasts starkly with the long-term records of lower Ψ_s (Extended Data Fig. 2). The lack of
133 difference in midday Ψ_1 could have been caused by sampling constraints or by isohydric
134 behaviour. We found evidence of non-isohydric behaviour in our diurnal Ψ_1 measurements
135 (Extended Data Fig. 4), with overall strong linear declines in Ψ_1 observed with increasing
136 VPD on the control ($R^2=0.18$, $P<0.01$) and in particular on the TFE ($R^2=0.33$, $P<0.01$).
137 Consequently limited sampling is the most likely cause of equal Ψ_1 between the two plots,
138 with TFE trees likely having more negative Ψ_1 and lower hydraulic conductance during VPD
139 maxima in the dry season. Reduced carbon uptake because of stomatal closure in some TFE
140 trees is possible²¹, but is unlikely to have caused carbon starvation considering growth rates
141 were maintained or elevated on the TFE (Fig. 2) and radial growth should decline prior to
142 photosynthesis in drought conditions²². Even with an isohydric response, trees on the TFE
143 would still be likely to suffer greater hydraulic deterioration caused by greater PLC in the
144 roots and main stem. Strongly reduced Ψ_s on the TFE (Extended Data Fig. 2) and significant
145 hydraulic vulnerability of the tall trees are consistent with the hypothesis of hydraulic

146 deterioration as the most likely trigger of greater mortality, particularly in the largest trees, as
147 observed. Why the xylem tissue of larger trees is more vulnerable to embolism deserves
148 further study. Taller trees are predisposed to greater hydraulic stress, from elevated
149 atmospheric demand and longer hydraulic path lengths²³. As the canopies are exposed to
150 rainfall in the TFE, smaller trees could avoid hydraulic deterioration through leaf water
151 uptake²⁴⁻²⁶, but this may not be sufficient to save the largest trees, which we hypothesise are
152 forced to maintain their high growth rates until death to continually replace dysfunctional
153 xylem.

154 Following decadal-scale soil moisture depletion, our results suggest tropical rainforests will
155 experience accelerating biomass loss and a likely transition to a lower statured, lower
156 biomass forest state, due to substantially elevated mortality of the largest trees. This mortality
157 is most likely triggered by hydraulic processes, which lead to hydraulic deterioration and
158 subsequent, potentially rapid, limitations in carbon uptake²¹, instead of being caused directly
159 by gradual carbon starvation. Under natural drought these forests may be under greater risk
160 than from experimental drought, as severe soil moisture deficit is combined with low
161 humidity and high air temperature increasing hydraulic demand. Improved prediction of the
162 sensitivity of tropical tree mortality to drought should therefore focus on improved model
163 simulation of plant hydraulics and modelling environmental controls on growth^{27,22}. Decadal-
164 scale ecological data such as these are rare, but they are invaluable for testing and improving
165 predictions from vegetation models over timescales that are relevant to climate change²⁸.
166 They also underpin the long-term environmental policy needed to manage the natural capital
167 that is embedded in tropical rainforests.

168

169 **Author contributions**

170 LR, PM, ALDC and MM designed and implemented the research. PM conceived and led
171 the experiment and this study. LR led recent measurements; all authors contributed to
172 data collection, led by ALDC. LR analysed the data with MM, PM, OB and AMP. LR
173 wrote the paper with PM and MM, with contributions from all authors.

174

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271 **Figure Legends**

272 **Fig. 1| Changes in biomass and mortality rates.** a. Biomass on the control and TFE plot
273 from 2002-2014 ($\text{Mg C ha}^{-1} \text{ yr}^{-1}$). Error bars show the s.e.m. calculated from 12 estimates of
274 biomass, accounting for uncertainty in wood density and allometric equations (see Methods).
275 b. Mortality rate ($\% \text{ stems yr}^{-1}$) calculated separately for trees of 10-20 cm dbh, 20-40 cm
276 dbh and >40 cm dbh on the control (black) and TFE (grey). The genus at date of death for
277 each tree used in the mortality rate calculations is shown in Extended data Table 2.

278 **Fig. 2| Tree growth of living and dead trees.** a. Average annual stem increment in the four
279 years prior to death for all trees which died on the control ($n=35$) and TFE ($n=60$) from 2005
280 onwards. Growth rates are normalised by the growth in year 0, the year in which the tree
281 died. b-d. Stem increment for small (a. 10-20 cm dbh), medium (b. 20-40 cm dbh) and large
282 (c. >40 cm dbh) trees on the control and TFE plot from 2005 to 2014. Values for b-d are
283 normalised by the maximum increment on the control plot. Error bars show the s.e.m.

284 **Fig. 3| Leaf, branch and stem NSC concentrations.** Percentage values for soluble sugars
285 (a.) and starch (b.) in dried biomass of leaves, branches and stems on the control and TFE for
286 late dry season (November 2013), mid wet season (March 2014) and in the wet-to-dry
287 transition (June 2014) for trees on the control ($n=20$) and TFE ($n=21$). Error bars show s.e.m.
288 and * indicates a significant difference at $P < 0.05$ using the Wilcoxon test. June 2014 has
289 significantly elevated sugar values on the TFE plot, however the absolute values for sugar
290 concentration are very low, and the absolute differences are very small.

291 **Figure 4| Xylem vulnerability to embolism and predicted loss of xylem hydraulic**
292 **conductivity as a function of tree diameter (dbh).** A general linear model was employed to
293 test for the effects of genus and tree dbh on the estimates of xylem P_{50} obtained for each
294 species ($n=37$ trees in total). The dashed lines give the predicted regressions for each of the

295 six genera, whereas the bold black line gives the overall mean regression line across all
296 species (The adjusted R^2 in the figure refers to the overall model, while the P value refers to
297 the significance of tree dbh). The insert shows the changes in predicted losses of hydraulic
298 conductivity (PLC) as a function of tree size. In the main graph, each genus is represented
299 with a different symbol and/or colour, as detailed in the legend. In the inset, the bold lines
300 represent the average prediction for the most vulnerable genus (red: *Pouteria*) and for the
301 most resistant one (blue: *Licania*) to drought induced embolism, with the shaded areas (pink,
302 light blue) giving the respective 95% confidence intervals.

303

304 **Extended data Figure 1| Leaf area index change:** leaf area index (LAI; m^2 leaf area m^{-2}
305 ground area) for the period of 2001-2014 on the control (black, solid) and TFE (grey, dashed)
306 plots. Error bars show the s.e.m. associated with LAI calculation (see Methods).

307 **Extended Data Fig. 2| Seasonal soil water potential:** average soil water potential (- MPa)
308 in the control and TFE during dry season (July-December) and wet season (January-June),
309 calculated from monthly average volumetric soil moisture content data, collected from 2008-
310 2014, using sensors installed 0, 0.5, 1, 2.5 and 4 m below the surface and the necessary van
311 Genuchten parameters previously calculated from soil hydraulics measurements at this site
312 (see Methods). They are averaged from hourly data collected from 2008-2011 and error bars
313 show s.e.m.

314 **Extended data Figure 3| Leaf herbivory comparison:** average percentage loss of leaf area
315 from herbivore attack calculated from 9770 leaves collected in litter-traps on the control and
316 TFE plot from 2010-2014. Error bars show s.e.m. A separate analysis of herbivore attack on
317 13,694 top-canopy living leaves from branches of the 41 trees used for the P_{50} analysis

318 support these results, also showing no significant differences in percentage herbivory
319 between the control and the TFE (data not shown).

320 **Extended Data Figure 4| Diurnal patterns of Ψ_1** : diurnal Ψ_1 measured every 2 hours from
321 6.00 until 18.00 in dry season on trees accessible from the walk up tower. Each box shows
322 the diurnal Ψ_1 against diurnal air vapour pressure deficit (VPD) from one of seven trees
323 accessible on the control (C), or one of four trees accessible on the TFE. Note a majority of
324 trees demonstrate some a negative relationship with VPD. Combined separately for each plot,
325 a significant negative linear relationship is observed between Ψ_1 and VPD on the control
326 ($R^2=0.18$, $P=0.002$) and even more strongly on the TFE ($R^2=0.33$, $P=0.001$).

327

328

329

330 **Methods**

331 *Site:* The through-fall exclusion (TFE) experiment is located in the Caxiuanã National Forest
332 Reserve in the eastern Amazon (1°43'S, 51°27'W), ~400 km west of the nearest city, Belém,
333 State of Pará, Brazil. The experiment is located in *terra firme* forest, on yellow oxisol soils
334 which are 75-83 % sand, 12-19 % clay and 6-10 % silt²⁹. The site is 15 m above sea level, has
335 a mean annual rainfall between 2000-2500 mm and a pronounced dry season between June
336 and November².

337 The experimental site has two 1 ha plots: the TFE, over which plastic panels and gutters have
338 been placed at a height of 1-2 m, and which exclude 50% of the incident rainfall; and a
339 corresponding control plot, <50 m from the TFE, on which there has been no manipulation of
340 incident rainfall. The TFE was trenched to between 1-2 m to remove the effect of through-
341 flow of soil water; and to control for any temporary damage to roots from the trenching, the
342 control plot was also trenched to the same depth. The TFE treatment has been installed and
343 running continuously since January 2002 to the present, except for a 1 week period in
344 November 2002 (full removal), a month period during the dry season in November 2014
345 (sequential removal of all panel) and during 2004 (30% removal).

346 *Soil Moisture Data:* In both the control and TFE plots there are soil access pits in which
347 volumetric soil water content sensors (CS616, Campbell Scientific, Logan, USA) located at
348 depths of 0, 0.5, 1, 2.5 and 4 m, monitor soil moisture every hour (*cf.* Fisher et al.³⁰, for full
349 methodology). New data presented here are from March 2008 - December 2014, averaged
350 into monthly values. Due to equipment failure, some soil moisture data are missing for
351 August and December 2013, for 2008 and 2010 on the control plot, and for November and
352 December 2013 on the TFE plot. Volumetric soil water content was converted in to soil water

353 potential (Ψ_s) using the necessary van Genuchten parameters previously calculated by Fisher
354 et al.³¹ based on soil hydraulics measurements at this site.

355 *Biomass data:* Trees ≥ 10 cm diameter at breast height (dbh) on both the control and the TFE
356 plots were tagged and identified to species level in September 2000. Diameters were
357 measured on these trees at 1.3 m, unless buttress roots were present, in which case the
358 measurement was made above the buttressing. Individuals on the plots were re-censused at
359 varying intervals from January 2001, until November 2014. During each census the trees
360 were also assessed to be either dead or alive. A tree was considered dead if leaflessness was
361 accompanied by a persistent zero or negative stem increment, and/or the tree had snapped or
362 fallen to the ground. Recruitment of new trees into the ≥ 10 cm dbh size class was enumerated
363 in 2005, 2009 and 2014.

364 Trees in the 10 x 10 m subplots adjacent to the trenches were excluded from our biomass and
365 growth rate analysis to eliminate possible effects of changes in mortality resulting from root
366 damage². Consequently, 369 trees on the control and 358 trees on the TFE, each in 0.64 ha
367 were analysed and the biomass scaled to 1 ha. Following da Costa et al. (2010)⁶ trees were
368 grouped into small (10-20 cm dbh) and medium (20-40 cm dbh) and large (>40 cm dbh) size
369 classes.

370 Biomass was calculated using the Chave et al.³² equation which uses diameter, wood density
371 and environmental predictors. A mean and standard deviation of wood density for each tree
372 was calculated from data in the global wood density database^{33,34} and from Patiño et al.³⁵.
373 Multiple estimates of wood density at the species, genus and family level were used to
374 calculate standard deviations on our wood density estimates. Of all the trees on both plots, 68
375 % had values for wood density at species level, 18 % at genus level and 3 % at family level;
376 11 % of trees were not identified and were given a plot level average wood density with an

377 associated standard deviation. A standard error on our biomass estimations was calculated,
378 which accounted for the error associated with wood density estimation and variations in
379 commonly used allometric equations. Twelve calculations were used to calculate the error on
380 our biomass values; these 12 biomass estimates were from combinations of four biomass
381 equations: Chave et al.³² and Chave et al.³⁶ both with and without height and three wood
382 density estimates: mean wood density and mean wood density \pm one standard deviation. The
383 allometric equations selected represent one of the most commonly used biomass equations for
384 Amazonia³⁶ (with and without height as a predictor variable) and the most recent and most
385 comprehensive biomass equations for our study area³² (with and without height as a predictor
386 variable). Measurements of height were not available and were calculated from an equation
387 developed specifically for the region of the Amazon in which our plots are located³⁷.
388 Mortality rate was calculated according to da Costa et al. (2010)⁶ and separately for trees of
389 10-20 cm dbh, 20-40 cm dbh and >40 cm dbh on the TFE and control plot.

390 *Seasonal growth data:* Dendrometers were installed at just above or below the point of dbh
391 measurement on all trees ≥ 10 cm dbh³⁸. Circumference measurements from the dendrometer
392 bands were made monthly to tri-monthly from January 2005 to November 2014, with the
393 exception of a six month gap from July 2007 and January 2013. Seasonal shrinkage was
394 calculated by taking the average growth rate of 19 trees on the control and TFE plots which
395 experienced no overall growth, but demonstrated a seasonal pattern of shrinkage and
396 expansion. Tropical trees can experience net diameter shrinkage in the dry season, due to a
397 lack of growth, accompanied by reduced water content of stem tissues, with subsequent
398 swelling when tissues are rehydrated in the wet season³⁹. An average pattern of shrinkage and
399 expansion was therefore subtracted from all trees to ensure that dry season growth was not
400 under-estimated and wet season growth over-estimated. Dendrometer increments were
401 filtered to remove the growth spikes from measurement errors following Rowland et al.

402 (2014)³⁹. Subsequent to this procedure, gap-filling was done using linear interpolation³⁹; on
403 the control and TFE plot 16 % and 17 % of the data were gap-filled respectively. Growth
404 rates per day calculated from the dendrometer measurements were averaged into three
405 monthly periods to give continuous tri-monthly growth rates from 2005-2014. When trees in
406 the 10 x 10 m subplots adjacent to the trenches, and trees with poor quality dendrometer
407 measurements throughout the 10 year study period for growth were excluded, 316 and 310
408 trees on the control and TFE plot remained, respectively. All growth and mortality analyses
409 were done using the R statistical package (version 3.1.2). Tests for significance were
410 performed using Wilcoxon signed-rank test.

411 *Non Structural Carbohydrate (NSC) analysis.* Using samples cut by a tree climber from fully
412 sunlit branches, three leaves and a branch sample of ~ 8-10 mm diameter were taken from 20
413 trees on the control plot and 21 trees on the TFE plot in November 2013, March 2014 and
414 June 2015. The numbers and species of the trees selected for analysis are shown in Extended
415 Data Table 1. The selected trees were all >10 cm dbh and represented the most common
416 genera existing on both plots²; samples were not taken from the external 10 x 10 m subplots
417 to avoid any impacts of trenching. Samples of tree stem tissue from the same trees were taken
418 with a 5 mm increment borer at breast height once in November 2013; this sampling was not
419 repeated to avoid excessive damage incurred by repeated boring. We followed the enzymatic
420 method proposed by Sevanto et al. (2014)⁴⁰ to analyze the NSC content. Here, NSC is
421 defined as free, low molecular weight sugars (glucose, fructose and sucrose) and starch.
422 Immediately after collection, samples were microwaved to stop enzymatic activity. After that,
423 samples were oven-dried at 70°C for 24–48 h and ground to fine powder. We prepared
424 approximately 12 mg of plant material with 1.6 mL of distilled water for the analysis. We
425 used amyloglucosidase from *Aspergillus niger* (Sigma-Aldrich) to digest total NSC to
426 glucose, and invertase, glucose hexokinase kits (GHK) and phosphorus glucose (Sigma-

427 Aldrich) to quantify the low molecular weight sugars. The concentration of free glucose was
428 determined photometrically in a 96-well microplate spectrophotometer (BioTek, Epoch).
429 NSC values are expressed as percent of dry matter. For further method details, see Sevanto et
430 al. (2014)⁴⁰.

431 *Xylem vulnerability to cavitation (Percentage Loss of Conductivity; PLC) and leaf water*
432 *potential measurements*. Samples were cut from fully sunlit branches following the same
433 protocol as above for NSC during late dry season (November 2013), and using the same
434 sample trees as used for the NSC analysis (Extended Data Table 1), with the exception of
435 branches of the genus *Manilkara*. To maintain a balance of number of genera and trees
436 sampled, four additional trees were therefore sampled from the genus *Inga* in the TFE and
437 control plots for both sets of analyses. One to three 1.0-1.5 m long branches per tree were cut
438 and left to rehydrate overnight under a black plastic bag in a bucket of water. Maximum
439 vessel length was determined for one branch out of a set of 5-6 branches for each species by
440 injecting low pressure air at the branch base under water and progressively re-cutting the
441 stem until bubbles emerged (maximum conduit length varied between 25 and 50 cm across
442 samples)⁴¹. Axial slits were made in the branch segments selected for PLC analysis to
443 increase the efficacy of the air injection. These partially debarked segments were mounted on
444 a 4.6 cm long air injection apparatus. Water, filtered to 0.2 μ , flowed gravimetrically through
445 the sample. After 10 minutes of equilibration at low pressure (10 kPa), the sample was
446 pressurised for 20 min. The pressure was increased in steps of 0.3 to 0.5 MPa, with each step
447 followed by 10 min relaxation and flow measured at a constant background 100 kPa air
448 pressure, until a residual flow lower than 5 % of the initial flow was found. Five to 10
449 measurements of water flux were taken at the distal end. The interval for each conductivity
450 measurement ranged from 2 to 10 minutes depending on stem length, conductivity and

451 pressure head employed (normally 3 kPa). Segment lengths, cross-sectional diameters and
452 leaf areas of the leaves subtended by the measured segment were determined.

453 We employed a two-parameter Weibull function to model the changes in percent loss of
454 xylem hydraulic conductivity as a function of xylem pressure⁴². The two parameters
455 represented P_{50} and slope of the conductivity-pressure curve. We estimated P_{50} and slope for
456 all trees using tree as a random factor in a mixed-model analysis (nlme library⁴³) in R
457 (Version 3.02, R Core Team). We let P_{50} vary for each tree while keeping slope constant
458 across trees to achieve convergence. We then employed these conditional estimates of P_{50} in
459 a general linear model to test for the effects of plot, genus and dbh (Extended Data Table 3a).

460 We then confirmed the above results by running a second mixed-effect model, in which we
461 accounted for random tree-by-tree variation and variance driven by phylogeny, by nesting
462 individual trees within genera as the random component of the model and incorporating plot
463 and dbh as fixed effects (Extended Data Table 3b). We tested whether tree diameter affected
464 P_{50} by comparing the performance of the full model with the same model without the effect
465 of dbh on P_{50} using a likelihood ratio test, and by conducting simultaneous hypothesis tests at
466 the 95% significance level (library multcomp in R). For both analyses, distributional
467 assumptions were tested by looking at plots of residuals for fixed and, if the case, random
468 effects.

469 Leaf water potential Ψ_1 was measured during two campaigns (one during the dry and one
470 during the wet season, i.e., October 2013 and May 2014, respectively). Two to four leaves
471 were measured for each branch of each tree, following the same sampling protocol above for
472 NSC and hydraulic measurements. During each campaign, Ψ_1 measurements were conducted
473 at pre-dawn (prior to 07.00 hrs) and at midday (between 11.30 hrs and 1300 hr). Midday Ψ_1
474 measurements of both campaigns were used in conjunction with the P_{50} and slope values

475 determined for each genus to estimate percent loss of hydraulic conductivity (PLC), assuming
476 xylem water potential to be equal to measured leaf water potential²⁰. We acknowledge that
477 this may overestimate PLC; however in counterpoint, only one seasonal campaign could be
478 conducted and so the minimum of leaf Ψ_1 for that year was thus likely to have been
479 underestimated because fuller sampling through the dry season was impractical. In addition,
480 earlier (2003) studies in the same experiment⁴⁴ reported minimum values of stem xylem
481 water potentials of around -1.6 MPa, across control and TFE plots, suggesting that xylem
482 values substantially more negative than those assumed here can be experienced.

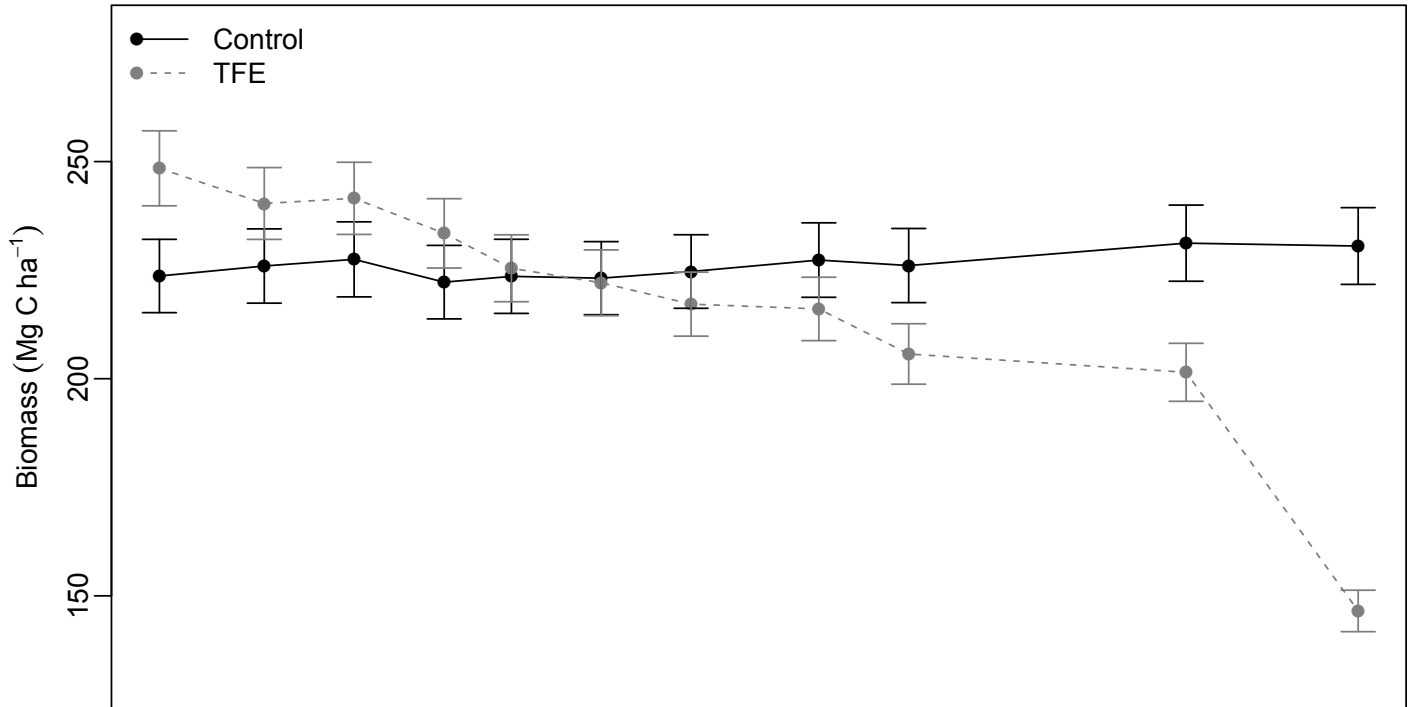
483 *Herbivory:* All leaf material from 25 x 1 m² litter-traps on the control and TFE was collected
484 13 times from 2010-2014 at 3-6 month intervals, with one eight month interval in 2011. Each
485 collection of leaf material represented two weeks of litter-fall in the forest. The 9770 leaves
486 collected from all 25 per plot litter-traps over the study period were scanned and the images
487 were analysed according to Metcalfe et al.⁴⁵ to calculate the percent leaf area lost to herbivore
488 attack on the control and TFE.

489 *LAI measurements:* LAI values from 2001 to 2007 are taken from Figure 1f in Metcalfe et al.
490 (2010)³⁸. For 2009-2014 LAI was measured at the same 25 permanent points³⁸ on a grid
491 throughout the control and TFE plot. Measurements were made using hemispherical photos
492 taken every 3-6 months from 2009-2014. Photos were taken before sunrise (~6.00 am) from a
493 height of 1.5 m on the control plot and 2 m (above the TFE structure) on the TFE plot. The 25
494 photos per plot were analysed together using the CAN_EYE software (INRA, Avignon,
495 France). Standard errors were calculated using three different estimates of LAI given by the
496 CAN_EYE software.

497 **Extra References for Methods**

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