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Important role of forest disturbances in the global biomass turnover and carbon sinks

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1	Biomass stocks in half of global forests controlled by large
2	disturbances
3	
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25 Forest disturbances leading to replacement of whole tree stands are a cornerstone 26 of forest dynamics, with drivers including fire, wind-throw, biotic outbreaks and 27 harvest. The frequency of disturbances may change over the next century, 28 impacting the age, composition and biomass of forests. However, the variation in 29 disturbance return time, i.e. the mean interval between disturbance events, across 30 the world's forested biomes remains poorly characterised, hindering 31 quantification of their role in the global carbon cycle. Here we present the global 32 distribution of stand-replacing disturbance return time inferred from satellite-33 based observations of forest loss. Prescribing this distribution within a vegetation 34 model with a detailed representation of stand structure, we quantify the 35 importance of stand-replacing disturbances for biomass carbon turnover globally 36 over 2001-2014. Return time varied from less than 50 years in heavily-managed 37 temperate ecosystems to over 1000 years in tropical evergreen forests. Stand-38 replacing disturbances accounted for 12.3% (95% confidence interval, 11.4-39 13.7%) of annual biomass carbon turnover due to tree mortality globally, and in 40 44% of forested area biomass stocks are strongly sensitive to changes in 41 disturbance return time. Relatively small shifts in disturbance regimes in these 42 areas would substantially influence the forest carbon sink, that currently limits 43 climate change by offsetting emissions.

44

45 Main text

46

The amount of carbon stored in global forest biomass is similar to that in the
atmosphere¹, and, excluding the effects of land-use change, has been estimated to
have grown at *ca.* 1.5 PgC a⁻¹ over recent decades². This uptake has significantly

50 slowed the atmospheric growth rate of carbon dioxide and thus the rate of climate 51 change³. The accumulation of carbon in the stems of growing trees results from 52 the balance between the growth rates of vegetation and the average length of time 53 carbon remains in live biomass (hereafter, "biomass"), the turnover time, 54 calculated as the carbon stock divided by the flux of carbon loss through plant and 55 tissue death⁴. Quantification of this turnover time is crucial because it governs the 56 size of biomass stocks for a given growth rate and it is one of the most significant 57 uncertainties affecting projections of the terrestrial carbon cycle^{5,6}. Large-scale 58 estimates of carbon turnover times for whole ecosystems and for biomass have 59 been recently developed^{7,8}, but offer limited insight into the processes governing 60 biomass stocks because the turnover flux could only be approximated by 61 estimates of net primary productivity (NPP). This conflates turnover of soft 62 tissues, such as leaves and roots, with that of the woody carbon pools that 63 dominate biomass carbon stocks¹. To understand forest biomass turnover times 64 globally, large-scale tree mortality rates must be quantified.

65

66 Tree death is often the culmination of a prolonged period of physiological stress 67 related to a shortage of essential resources required for the synthesis of basic 68 metabolites, for instance, due to shading by other plants, low water availability, or 69 a shortage of soil nutrients such as N and P in plant-available forms^{9,10}. 70 Alternatively, disturbances such as wind-throw, fire, insect and disease outbreaks, 71 or anthropogenic activities such as wood harvest may constitute the cause of 72 death¹¹⁻¹⁶. Disturbances act on scales ranging from a single tree to whole forest 73 stands or landscapes. Here we investigate stand-replacing disturbances, defined 74 operationally as discrete events resulting in the death of all, or almost all, living

tree biomass at a scale of 0.1 ha or larger. Such events affect the average tree age as well as the stature, density and composition of forest stands^{11,17}, in turn impacting carbon storage¹⁸. There is evidence that the frequency of disturbances may be changing globally, with continued change likely in the future^{8,13-15,19,20}. Yet, to understand the consequences of future changes, it is first necessary to provide a baseline of current conditions. Such a baseline is lacking for stand-replacing disturbances across global forests.

82

83 **The frequency of stand-replacing disturbances.** We estimated the frequency of 84 stand-replacing disturbances across all global forests at 1° spatial resolution. 85 Drawing on a Landsat-based (2000-2014) forest-loss product²¹, we performed a 86 space-for-time substitution, calculating the disturbance rotation period, τ , defined 87 as the mean time period for the area disturbed to equal the area of the grid cell¹¹,

88
$$\tau_i = \frac{A_{T,i}}{\overline{A_{L,i}} - \overline{A_{C,i}}} \quad (1)$$

where *i* is a grid-cell index, A_T is total forest area in that 1° x 1° grid cell, $\overline{A_L}$ the 89 90 annual mean total forest loss over 2000-2014 calculated based on all Landsatpixels within the grid cell and $\overline{A_c}$ the annual mean forest loss due to conversion to 91 92 a non-forest land-use type (Methods). This observation-based τ is referred to as 93 τ_0 . While rare disturbances may be undersampled for individual grid cells, this is 94 less of an issue across a neighbourhood of many grid cells, and the global-scale 95 pattern is expected to be robust (Suppl. Note 1). We take τ as indicative of the 96 typical or average disturbance return time from all causes at any location in a grid 97 cell, reflecting causes such as wood harvest, as well as natural disturbances such as fires, large-scale wind-throws and biotic outbreaks. Whilst the drivers of each 98

99 of these disturbance agents differ markedly, their consequences for carbon 100 turnover in live biomass are assumed to be similar at large scales. Disturbances 101 associated with the conversion of forests to other land-uses were explicitly 102 excluded (Methods), allowing us to focus on dynamics within closed-canopy 103 forests.

104

105 Disturbance return time varies substantially across the global forest area (Fig. 1a). 106 The stand-replacing disturbances quantified here are more common in needleleaf 107 and mixed forests (median τ_0 of *ca*. 300 years) than they are in temperate or tropical broadleaf forests (median τ_0 of 830 to more than 1000 years), however 108 109 forest type emerges as a poor predictor of the spatial distribution of τ_0 (Fig. 1c). 110 In large areas of forest, stand-replacing disturbances are actually very rare events; 111 35% of forest area experienced stand-replacing disturbances with an average 112 frequency equivalent to less than once every thousand years. In these forests the 113 vast majority of tree mortality must thus be non-stand-replacing. The 95% 114 confidence intervals for τ_0 typically span a range less than one third of the 115 absolute value of τ_0 , except in areas of substantial land-use change (Fig. 1b, Suppl. 116 Fig. 7).

117

We compared our results against an inventory-based compilation of forest stand age (GFADv1.1²²). Despite the different scale and characteristics of inventory data we found qualitative consistency in tropical evergreen and boreal forests, as well as some regions under intensive forest management, but also suggestions of a substantial amount of disturbance in some temperate forests below the scale captured in the Landsat data and of legacies of past afforestation (Suppl. Note 2,

Suppl. Fig. 1). We also found consistency between our results and previous studies
of disturbance frequencies in the tropics^{17,23,24} and Canada²⁵ (Suppl. Note 3; Suppl.
Fig. 2), and biotic outbreak disturbances in the U.S.A.²⁶ (Suppl. Note 3; Suppl. Fig.
3).

128

129 Influence of stand-replacing disturbances on the carbon cycle. We apply the 130 gridded estimates of τ_0 within a dynamic global vegetation model (DGVM) with an explicit representation of forest stand structural development. τ was kept 131 132 constant in each grid cell for the entire model simulation, calculating the pseudo-133 equilibrium effect of τ_0 on forest dynamics. Stand-replacing disturbances are 134 simulated to dominate overall tree mortality, and associated carbon turnover, 135 across large areas of the mid-latitude and boreal forests, accounting for over 60% 136 in some locations, but are not the dominant cause of mortality in most tropical 137 forests (Fig. 2a,b).

138

139 The total turnover of biomass carbon as a result of stand-replacing disturbances 140 at pseudo-equilibrium in our simulations is 1.00 (95% confidence interval, 0.91-1.11) PgC a⁻¹, equivalent to 4.4 (4.0-4.9) % of total biomass carbon turnover in 141 142 closed-canopy forests (i.e. including soft-tissue turnover) (Fig. 2b,c). These 143 numbers are supported by an empirical cross-check based on satellite-derived 144 NPP and biomass estimates which combined satellite LIDAR and radar 145 observations with ground-based reference plots (red dots in Fig. 2b,c; Methods). 146 The fraction of biomass carbon turnover due to mortality must be taken with 147 caution, however, until biomass turnover rates from other forms of mortality can

be fully constrained. Likewise the total turnover flux is dependent on accurate
calculation of global biomass stocks, which remain uncertain¹.

150

151 The tropical broadleaved evergreen forest type provides the largest contribution 152 to global biomass carbon turnover from stand-replacing disturbances, followed by 153 needleleaved evergreen forest (Fig. 2c). Although stand-replacing disturbances 154 are infrequent in tropical evergreen forest, the disturbance-related flux is significant, as the amount of biomass in these forests is very high compared to 155 156 elsewhere^{27–30}. Conversely, low τ will tend to suppress biomass stocks, limiting the turnover flux generated in each disturbance event. Our estimates of biomass 157 158 carbon losses for tropical evergreen forest will tend towards the upper limit of 159 uncertainty because τ_0 was capped at 1000 years for reasons of sampling 160 (Methods); disturbance return times could in fact be even longer in some parts of 161 the tropics³¹. However, as demonstrated below, the sensitivity of biomass to very 162 high τ is low. Different disturbance agents cannot be distinguished in our data, but 163 carbon emissions from wildfire taken from the GFED dataset³² summed over the 164 same global forest area give a mean of 0.12 Pg C a⁻¹ over 2000-2014 (Methods), 165 suggesting that fires are globally a relatively minor driver of stand-replacing 166 disturbances in closed-canopy forests.

167

168 **Sensitivity of forest biomass to disturbance return time.** We ascertain the 169 influence on ecosystem properties of possible changes in τ , or errors in its 170 determination, for a randomly-selected grid cell from each of the tropical, 171 temperate and boreal zones. For each grid cell we run 100 individual simulations 172 varying τ sequentially within a plausible range of 10 to 1000 years. The resultant

173 range in pseudo-equilibrium carbon stocks reflects variation in τ alone, 174 independent of other environmental conditions or vegetation attributes (in 175 particular that the resilience of vegetation to disturbance does not change with τ). 176 The resulting curve of biomass carbon versus τ shows two distinct regimes (Fig. 177 3a); a regime of strong sensitivity of biomass to τ when τ is low, and a weak 178 sensitivity regime when τ is high. These regimes result from shifts in the primary 179 cause of dominant tree death. With low τ the majority of trees die from stand-180 replacing disturbance before they get old enough to die from another cause. Thus, 181 τ emerges as the primary limit on simulated stand biomass across almost all 182 stands. In contrast, when τ is large most canopy trees die from causes other than 183 stand-replacing disturbances. In this case, τ is not a primary limit on simulated 184 stand maximum biomass and changes in τ will only affect a subset of stands across 185 the landscape at any time.

186

187 To map the sensitivity of forests to changes in τ globally we propose a new metric 188 based on a fractional reduction of τ_0 . Plotting τ_0 against the difference in simulated 189 biomass between global simulations run with τ_0 and $0.5\tau_0$ reveals a similar curve 190 to the site-based simulations (Fig. 3b). We categorise global forests into two 191 classes of sensitivity to disturbances: where the biomass under $0.5\tau_0$ is less than 192 90% of that under τ_0 , the forest is classified as having strong sensitivity to changes 193 in τ (i.e. stand-replacing disturbance frequency is a strong control on biomass), 194 with other areas having weak sensitivity (i.e. other forms of mortality control 195 biomass). The τ_0 at which this sensitivity threshold, $\tau_{crit.90}$, is crossed varies with 196 forest type, with a global average of 444 (429-457) years. This implies a mean 197 recovery time of 222 years (i.e. $0.5\tau_{crit,90}$) to 90% of biomass stocks under τ_0 . This

198 is substantially longer than the 66 years reported for regain 90% of old-growth biomass in individual tropical forest stands³³ and follows because our simulations 199 200 take account of succession and also scale one stage further to the net landscape 201 change in biomass. I.e. individual stands may recover rapidly, but across the 202 landscape more stands are in a recently-disturbed state. Overall, 44 (39-49) % of 203 global forest falls into the strongly sensitive category, with 23 (20-27) % falling 204 into a very strong sensitivity category in which halving τ_0 leads to biomass 205 dropping below 80% of that under τ_0 ($\tau_{crit,80}$). Forests in weak-sensitivity τ 206 regimes are particularly located in tropical and temperate zones.(Fig. 3c).

207

208 The biomass content of weak-sensitivity forests would still be vulnerable to very 209 large reductions in τ , for instance through a catastrophic shift to an entirely new 210 disturbance regime³⁴ or introduction of a new disturbance type³⁵, but is robust to 211 moderate changes in τ . This is demonstrated in additional simulations using τ_0 212 adjusted up or down by a factor of up to 4; weak-sensitivity forests show large 213 biomass changes only with substantial reductions in τ , whereas high-sensitivity 214 forests show a steep relationship with τ (Fig. 3d). These results were robust to 215 assumptions on the type of disturbance (shaded areas in Fig. 3d), although we 216 note that selective effects of disturbance type on species composition^{35,36}, and thus 217 potentially on biomass and turnover³⁷, could only be treated crudely within the 218 plant functional type classes used for global simulation and there may be non-219 linear shifts not accounted for in the model³⁸. The time taken for forest biomass to 220 approach a new pseudo-equilibrium state will depend on the new value of τ to 221 which the ecosystem is subjected. In general, changes in τ will only be fully reflected in carbon fluxes over the next century if that new τ is of the order of 100 years or less.

224

225 Unlike the influence of τ on biomass (Fig. 3d), its influence on soil carbon stocks is 226 strongly sensitive to the rate of decompositon of the resulting litter and soil 227 organic matter, and also depends on biomass removals, for instance in conjunction 228 with wood harvest (Fig. 3e). When disturbed biomass is transferred to the litter, 229 disturbance only has notable negative consequences when τ becomes very low, 230 reducing the fraction of longer-lived woody biomass entering the litter. However, 231 harvest removals or burning of biomass substantially reduce the input rate of 232 carbon to the soil, leading to a strong positive relationship between soil carbon 233 density and τ . This strong sensitivity of soil carbon storage to the type, as well as 234 the frequency, of disturbance, underlines the need for improved discrimination of 235 different disturbance types at the global scale³⁹. Response times for soil will lag 236 those for vegetation, and be influenced by the form of necromass left after 237 disturbance⁴⁰, another area of high process uncertainty. Summing over both 238 vegetation and soil, a widespread shift in disturbance regimes equivalent to a 239 halving of τ_0 across all closed-canopy forests would ultimately release 47-80 Pg C, 240 depending on the form of that disturbance, while an increase in time between 241 disturbances could promote carbon uptake (Suppl. Fig. 10).

242

243 **Concluding remarks.** The results of this study allow us to partition one important 244 component of overall biomass turnover rates in global forests^{1,8}. Although stand-245 replacing disturbances constitute a relatively small portion of the overall global 246 biomass turnover flux, small changes in τ would exert a strong influence on

247 biomass stocks in almost half of the world's forests. DGVMs and land-surface 248 models currently incorporating explicit representations of forest demography⁴¹ 249 must properly account for stand-replacing disturbances to avoid biases in net 250 carbon uptake or erroneous calibration of processes to account for these biases. 251 Our study highlights the importance of accounting for variability in forest 252 disturbance regimes, yet constitutes only a first step; 88% of global carbon 253 turnover due to tree mortality is not explained by stand-replacing disturbances. It 254 thus remains crucial to constrain other causes of mortality, including disturbances 255 below stand-scale^{23,42}, drought⁴³, and demography. High-resolution data from 256 satellites, along with forest inventories, will be key in this regard.

257

258 Our results provide a snapshot of a global stand-replacing disturbance pattern 259 that may be undergoing rapid change¹³⁻¹⁵. Drivers of such change, whether 260 climate, management or otherwise are uncertain and likely highly region-261 specific^{13-16,19,20}. Future work must consider how carbon emissions through 262 changes in τ are likely to interact with other aspects of environmental change, 263 such as the fertilising effects of rising CO₂ concentrations, which may reduce 264 vulnerability to disturbance⁴⁴, as well as seeking to close the feedback loops 265 between disturbances, climate and vegetation properties. Changing disturbances 266 could both augment and offset carbon loading of the atmosphere caused by 267 anthropogenic carbon emissions³; better understanding the role of forest 268 disturbances in the carbon cycle is therefore highly relevant to the assessment of 269 emissions reductions necessary to meet climate targets.

270

271 **References**

272 Erb, K. et al. Unexpectedly large impact of forest management and grazing 1. 273 on global vegetation biomass. *Nature* **553**, 73–76 (2018). 274 Pan, Y. et al. A large and persistent carbon sink in the world's forests. 2. 275 Science 333, 988–93 (2011). 276 3. Quéré, C. Le et al. Global Carbon Budget 2017. Earth Syst. Sci. Data 10, 277 405-448 (2018). 278 4. Sierra, C. A., Müller, M., Metzler, H., Manzoni, S. & Trumbore, S. E. The 279 muddle of ages, turnover, transit, and residence times in the carbon cycle. 280 *Glob. Chang. Biol.* 23, 1763–1773 (2017). Friend, A. D. et al. Carbon residence time dominates uncertainty in 281 5. 282 terrestrial vegetation responses to future climate and atmospheric CO2. 283 Proc. Natl. Acad. Sci. U. S. A. 111, 3280-3285 (2014). 284 Ahlström, A., Xia, J., Arneth, A., Luo, Y. & Smith, B. Importance of 6. 285 vegetation dynamics for future terrestrial carbon cycling. Environ. Res. 286 Lett. 10, 054019 (2015). 287 7. Carvalhais, N. et al. Global covariation of carbon turnover times with 288 climate in terrestrial ecosystems. *Nature* **514**, 213–217 (2014). 289 8. Erb, K.-H. et al. Biomass turnover time in terrestrial ecosystems halved by 290 land use. Nat. Geosci. 9, 674–678 (2016). 291 9. Waring, R. H. Characteristics of Trees Predisposed to Die. *BioSicence* 37, 292 569-574 (1987). 293 10. McDowell, N. G. et al. The interdependence of mechanisms underlying 294 climate-driven vegetation mortality. Trends Ecol. Evol. 26, 523–532 295 (2011).

296	11.	Pickett, S. T. A. & White, P. S. The ecology of natural disturbances and patch
297		dynamics. (Academic Press Inc, Orlando, 1985).
298	12.	Frolking, S. et al. Forest disturbance and recovery: A general review in the
299		context of spaceborne remote sensing of impacts on aboveground
300		biomass and canopy structure. J. Geophys. Res. 114, G00E02 (2009).
301	13.	Kurz, W., Stinson, G., Rampley, G., Dymond, C. & Neilson, E. Risk of natural
302		disturbances makes future contribution of Canada's forests to the global
303		carbon cycle highly uncertain. Proc. Natl. Acad. Sci. U. S. A. 105, 1551–
304		1555 (2008).
305	14.	Seidl, R., Schelhaas, MJ., Rammer, W. & Verkerk, P. J. Increasing forest
306		disturbances in Europe and their impact on carbon storage. Nat. Clim.
307		<i>Chang.</i> 4 , 806–810 (2014).
308	15.	Flannigan, M., Stocks, B., Turetsky, M. & Wotton, M. Impacts of climate
309		change on fire activity and fire management in the circumboreal forest.
310		<i>Glob. Chang. Biol.</i> 15, 549–560 (2009).
311	16.	Hurtt, G. C. et al. Harmonization of land-use scenarios for the period
312		1500–2100: 600 years of global gridded annual land-use transitions,
313		wood harvest, and resulting secondary lands. <i>Clim. Change</i> 109 , 117–161
314		(2011).
315	17.	Cole, L. E. S., Bhagwat, S. A. & Willis, K. J. Recovery and resilience of
316		tropical forests after disturbance. <i>Nat. Commun.</i> 5, 1–7 (2014).
317	18.	Pregitzer, K. S. & Euskirchen, E. S. Carbon cycling and storage in world
318		forests: biome patterns related to forest age. <i>Glob. Chang. Biol.</i> 10, 2052–
319		2077 (2004).

- 320 19. Seidl, R. et al. Forest disturbances under climate change. *Nat. Clim. Chang.*321 **7**, 395–402 (2017).
- Reyer, C. P. O. et al. Are forest disturbances amplifying or canceling out
 climate change-induced productivity changes in European forests? *Environ. Res. Lett.* 12, 034027 (2017).
- 325 21. Hansen, M. C. et al. High-resolution global maps of 21st-century forest
 326 cover change. *Science* 342, 850–853 (2013).
- 327 22. Poulter, B. et al. *The global forest age dataset and its uncertainties*
- 328 *(GFADv1.1)*. (2019). doi.pangaea.de/10.1594/PANGAEA.897392
- 329 23. Espírito-Santo, F. D. B. et al. Size and frequency of natural forest
 330 disturbances and the Amazon forest carbon balance. *Nat. Commun.* 5,
- 331 3434 (2014).
- 332 24. Chambers, J. Q. et al. The steady-state mosaic of disturbance and
- 333 succession across an old-growth Central Amazon forest landscape. *Proc.*
- 334 *Natl. Acad. Sci. U. S. A.* **110**, 3949–3964 (2013).
- 335 25. White, J. C., Wulder, M. A., Hermosilla, T., Coops, N. C. & Hobart, G. W. A
- nationwide annual characterization of 25 years of forest disturbance and
- recovery for Canada using Landsat time series. *Remote Sens. Environ.* 194,
 303–321 (2017).
- 339 26. Kautz, M., Meddens, A. J. H., Hall, R. J. & Arneth, A. Biotic disturbances in
 340 Northern Hemisphere forests a synthesis of recent data, uncertainties
- and implications for forest monitoring and modelling. *Glob. Ecol. Biogeogr.*
- **26,** 533–552 (2017).
- Avitabile, V. et al. An integrated pan-tropical biomass map using multiple
 reference datasets. *Glob. Chang. Biol.* 22, 1406–1420 (2016).

- 345 28. Santoro, M. et al. Remote Sensing of Environment Forest growing stock
 346 volume of the northern hemisphere: Spatially explicit estimates for 2010
 347 derived from Envisat ASAR. *Remote Sens. Environ.* 168, 316–334 (2015).
- 348 29. Avitabile, V. et al. Comparative analysis and fusion for improved global
- biomass mapping. in *Global Vegetation Monitoring and Modeling 3 7 February 2014*, Avignon (France) (2014).
- 351 30. Thurner, M. et al. Carbon stock and density of northern boreal and
 352 temperate forests. *Glob. Ecol. Biogeogr.* 23, 297–310 (2014).
- 353 31. Espírito-Santo, F. D. B. et al. Storm intensity and old-growth forest
- disturbances in the Amazon region. *Geophys. Res. Lett.* **37**, L11403 (2010).
- 355 32. van der Werf, G. R. et al. Global fire emissions estimates during 1997-
- 356 2016. Earth Syst. Sci. Data **9**, 697–720 (2017).
- 357 33. Poorter, L. et al. Biomass resilience of Neotropical secondary forests.
 358 *Nature* 530, 211–214 (2016).
- 359 34. Scheffer, M., Carpenter, S., Foley, J. A., Folke, C. & Walker, B. Catastrophic
 360 shifts in ecosystems. *Nature* 413, 591–6 (2001).
- 361 35. Johnstone, J. F. et al. Changing disturbance regimes, ecological memory,
- and forest resilience. *Front. Ecol. Environ.* **14**, 369–378 (2016).
- 363 36. Marra, D. M. et al. Large-scale wind disturbances promote tree diversity in
- 364 a Central Amazon forest. *PLoS One* **9**, e103711 (2014).
- 365 37. Marra, D. M. et al. Predicting biomass of hyperdiverse and structurally
- 366 complex central Amazonian forests a virtual approach using extensive
- 367 field data. *Biogeosciences* **13**, 1553–1570 (2016).
- 368 38. Marra, D. M. et al. Windthrows control biomass patterns and functional
- 369 composition of Amazon forests. *Glob. Chang. Biol.* **24,** 5867–5881 (2018).

370	39.	McDowell, N. G. et al. Global satellite monitoring of climate-induced				
371		vegetation disturbances. Trends Plant Sci. 20, 114–123 (2015).				
372	40.	Renninger, H. J., Carlo, N., Clark, K. L. & Schäfer, K. V. R. Modeling				
373		respiration from snags and coarse woody debris before and after an				
374		invasive gypsy moth disturbance. J. Geophys. Res. Biogeosciences 119,				
375		630-644 (2014).				
376	41.	Fisher, R. A. et al. Vegetation demographics in Earth System Models: A				
377		review of progress and priorities. <i>Glob. Chang. Biol.</i> 24, 35–54 (2018).				
378	42.	Marvin, D. C. & Asner, G. P. Branchfall dominates annual carbon flux				
379		across lowland Amazonian forests. <i>Environ. Res. Lett.</i> 11, 094027 (2016).				
380	43.	Allen, C. D., Breshears, D. D. & McDowell, N. G. On underestimation of				
381		global vulnerability to tree mortality and forest die-off from hotter				
382		drought in the Anthropocene. <i>Ecosphere</i> 6 , 129 (2015).				
383	44.	Dolan, K. A. et al. Disturbance Distance: quantifying forests' vulnerability				
384		to disturbance under current and future conditions. Environ. Res. Lett. 12,				
385		114015 (2017).				
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405

406 Author contributions

407

408 TP conceived and designed the study with contributions from AA and BS. BP and
409 MK contributed data. TP carried out the model simulations. TP led the analysis
410 and wrote the paper with contributions from all authors.

411

- 412 **Competing financial interests**
- 413

414 The authors declare no competing financial interests.

416 Figure 1. Forest disturbance rotation periods. a, τ_0 calculated over 2000-2014. 417 **b**, Uncertainty in τ_0 , displayed as the difference between the 95% confidence 418 intervals divided by the central estimate. Uncertainty values of zero reflect 95% 419 confidence that τ_0 is over 1000 years. **c**, Boxplot of τ_0 grouped by forest type (see 420 Methods for codes). Circles show the median value, black triangles the 95% 421 confidence limits of the median, thick lines the interquartile range and whiskers 422 extend to a maximum of 1.5 times the interquartile range. Numbers indicate the 423 number of grid cells for each forest type.

424

Figure 2. Carbon turnover fluxes from closed-canopy forest for 2001-2014. a, 425 426 Fraction of carbon turnover fluxes resulting from vegetation mortality due to 427 stand-replacing disturbances (colour scale capped at 50%), calculated using τ_0 to 428 drive LPJ-GUESS. Breakdown by forest type of: **b**, fraction of carbon turnover 429 fluxes resulting from vegetation mortality (whole bars) and from stand-replacing 430 disturbances (darker shading); **c**, total turnover flux of vegetation carbon due to 431 stand-replacing disturbance. Error bars show the range of simulations driven by 432 the 95% confidence intervals of τ_0 . Red dots show results from an observationally-433 based cross-check method (Methods). Forest types as in Fig. 1.

434

Figure 3. Sensitivity of biomass to changes in τ . **a**, Simulated biomass versus τ for three random forested locations. Dots show individual simulations and lines a fitted exponential function. **b**, Sensitivity to τ_0 of difference in simulated biomass between simulations with τ_0 and $0.5\tau_0$. **c**, Sensitivity of biomass carbon stocks to changes in τ . Shading indicates the sensitivity regime. **d**, **e**, Effect of multiplicative perturbation in τ on vegetation and soil carbon density averaged across the

- 441 different sensitivity classes. Shaded areas show range of sensitivity simulations
- testing assumptions on the type of disturbance assumed (solid lines for standard
- 443 simulation) (Methods).
- 444







446 Methods

447

448 **Calculation of** τ_0 **.** τ_0 was calculated as defined in Eq. 1. We first created a forest 449 mask by aggregating year 2000 forest canopy cover data at 0.00025° (*ca.* 30 m) 450 resolution²¹ to 0.01° resolution. Grid cells with at least 50% canopy cover at 0.01° 451 resolution were assigned as closed-canopy forest. Further aggregation then provided the fractional coverage of closed-canopy forest at 1° resolution (A_T). 452 453 Across each 1° grid cell we then summed up the total area of 0.00025° pixels which 454 underwent forest loss during 2000-2014 and were located within the 0.01° grid 455 cells assigned as closed-canopy forest. A grid cell could only be counted as lost 456 once during the period. Dividing this sum by the length of the 14 year observation period provided $\overline{A_L}$. A threshold of 25% forest cover at the 1° grid cell level was 457 458 used throughout this study in order to provide sufficient statistical power for 459 calculation of τ_0 . The total forested area meeting these conditions is 2.71×10^7 460 km². Fig. 1c was calculated from the gridded τ_0 estimate using the "boxplot" 461 function of Matlab® 2014b.

462

This above definition provides a calculation of τ as a function of forest area. An alternative definition of τ would be to define it relative to canopy area. In this case A_T would be the total canopy area within the 0.01° grid cells designated as closed canopy forest, making use of the fractional canopy cover metric provided by Hansen *et al.*²¹, and $\overline{A_L}$ would be the sum of pixels undergoing forest loss multiplied by the fractional canopy cover of those pixels before disturbance. Using this canopy-area definition slightly reduces our estimates of τ_0 in most locations

470 (Suppl. Fig. 4), but the forest-area definition is preferred as it recognises that
471 whilst disturbances reduce canopy cover, they do not reduce the area of forest
472 unless associated with a land-use change.

473

Forest losses due to land-use change, $\overline{A_c}$, were calculated for the period 2000-474 2014 using the ESA CCI landcover product v2.0.7 (accessed 29th June 2017). ESA 475 476 CCI landcover classes were simplified into forested (classes 50, 60, 61, 62, 70, 71, 477 72, 80, 81, 82, 90, 100, 160, 170) and non-forested (classes 10, 11, 20, 30, 110, 478 130, 190) classes, the latter corresponding to cropland, grassland and urban land 479 uses. Then the area of 0.0028° pixels which were classified as forested in 2000 but 480 non-forested in 2014 was calculated. The forest loss due to land-use change calculated from this dataset shows excellent consistency with the total forest loss 481 dataset based on Hansen et al.²¹, with only very few locations where the loss due 482 483 to land-use change is reported to be larger than the total (Suppl. Fig. 5).

484

485 Uncertainties in τ_0 due to the sample sizes in the forest loss data were estimated through bootstrapping. In each 1° grid cell 1000 samples of $\overline{A_L}$ were created by 486 resampling with replacement the 0.01° grid cells designated as closed-canopy 487 forest. Uncertainties in $\overline{A_c}$ result from classification accuracy and scaling 488 differences between the Hansen et al.²¹ and ESA CCI datasets. Producer's accuracy 489 for the forest and non-forest classification in ESA CCI v2.0.7 is 92% and 78% 490 491 respectively, whilst the corresponding user's accuracy is 78% and 85%⁴⁵. 492 However, because we count the whole area of the pixel when an ESA CCI pixel 493 changes from forest to non-forest, and the CCI pixel area is *ca.* 100 times that of 494 Landsat, a scaling inaccuracy is induced, whereby the fraction of forest conversion

495 within the grid cell may be enough to cause a land-cover classification switch, but 496 substantially less than complete deforestation of the ESA pixel. To conservatively 497 account for classification and scaling errors we thus assume a 95% confidence 498 interval of +- 50% in the forest conversion area values. For each 1° grid cell, 1000 samples of $\overline{A_c}$ were taken from a normal distribution defined by this confidence 499 interval. We crossed these 1000 samples of $\overline{A_c}$ with those from $\overline{A_L}$ to create a 500 matrix of 1 x 10^6 estimates of the denominator in Eq. 1. The 2.5th and 97.5th 501 502 percentiles of this matrix were then used to estimate the 95% confidence limits of 503 τ_0 . This resampling of the forest loss areas within the 1° pixel addresses the 504 uncertainty induced when the forest area in the pixel is relatively small, in which 505 case confidence in the fidelity of the space-for-time swap would be reduced. It also 506 accounts for classification errors if those errors are not correlated across the grid 507 cell. Hansen *et al.*²¹ report a tendency to underestimate forest loss by *ca.* 4% in the 508 tropics and overestimate it by *ca.* 6% in the temperate and boreal regions. These classification biases are not captured in our uncertainty estimate, nor are 509 potential biases from $\overline{A_T}$ for which global quantification was not available. Based 510 511 on the available information, these biases are expected to be small and focused in 512 regions where the uncertainty is already assessed as being large (Suppl. Note 4; 513 Suppl. Fig. 8). Note that the capping of τ_0 at 1000 years often leads to very low 514 uncertainty for these grid cells, i.e. there is very high certainty that $\tau_0 > 1000$ years. 515 Calculated τ_0 is robust to subsampling of the 14 year observational period, 516 especially when the data series exceeds 10 years (Suppl. Fig. 6).

517

518 The resolution of ESA CCI landcover means it will have limited sensitivity to very519 small-scale land-use conversions, such as have been recently reported in the

520 Amazon⁴⁶. However, given that our τ_0 values in the tropical evergreen forests are 521 very high, even in absence of the land-use correction (Suppl. Fig. 7) we expect the 522 influence on our results to be minimal.

523

The use of a 1° aggregation resolution represents a compromise between spatial detail and sufficient area to make an effective space-for-time substitution. Following the simplifying assumption that disturbance events are equally likely in all locations in the grid cell, the maximum τ that we can expect to reliably quantify, τ_m , for a given disturbance size, *D*, can be calculated as:

529
$$\tau_m = \frac{A_T \times t}{D}, (2)$$

where *t* is the total number of years sampled. The largest disturbance events are 530 531 generally fires, especially in the Canadian boreal region, for which the typical large 532 fire size is 6000 ha⁴⁷. Assuming a grid-cell area of 628 000 ha (60° latitude), a forest coverage of 25% of grid-cell area (i.e. $A_T = 157\ 000$ ha) and a 14-year 533 sampling period, τ_m is 350 years at this scale. For smaller disturbances much 534 535 larger values of τ can be expected to be reliably captured. Substantial 536 undersampling of large rare events at 1° resolution would be expected to induce 537 scatter in our results, but Fig. 1 shows spatial coherence in variation of τ , 538 suggesting any such under-sampling to have minimal effects. τ_0 was capped at 539 1000 years to avoid spuriously large values in grid cells with very infrequent 540 disturbance. The influence of this capping on simulated forest biomass is very 541 small (e.g. Fig. 3b).

542

543 Forest type classification. Forest types were classified based on ESA CCI
544 landcover v2.0.7. The mapping of landcover classes to the forest types used in this

545 analysis in shown in Suppl. Table 2. A map of these forest types is shown in Suppl. 546 Fig. 11. There is a small fraction of forest area that is not assigned to any of these 547 major forest classes, but is included in the global totals. Note that open canopy 548 forests (<50% canopy cover at 0.01° scale, see above) are not included in any of 549 the calculations herein. Forest type codes are: Tropical broadleaved evergreen 550 (TrBE), tropical broadleaved deciduous (TrBD), temperate broadleaved 551 evergreen (TeBE), temperate broadleaved deciduous (TeBD), needleleaved evergreen (NE), needleleaved deciduous (ND), broadleaved-needleleaved mixed 552 553 forest (MX).

554

555 Forest age dataset. For cross comparison of spatial patterns in our results, we used the Global Forest Age Dataset (GFAD v1.1)²², a forest stand age dataset 556 557 developed as part of the EU FP7 GEOCARBON project. It provides a distribution of 558 stand age and associated uncertainties in 10-year age bins up to an age of 140 559 years from a base year of 2010 on a 0.5° grid. The salient features are summarised 560 here and described in more detail in Pugh et al.⁴⁸. It combines datasets of forest 561 age distributions from the following forest inventories: United States Forest 562 Inventory and Analysis (v 5.1, state summaries, representative for the 2000s), 563 IIASA Russian Forests and Forestry Database (late 2000s), Canadian Forest 564 Inventory (CanFI, state summaries, 2001-2006), EFISCEN (Europe, 32 countries, 565 2000s), 6th National Forest Inventory (China, 1999-2003), and the national forest 566 inventories of Kazakhstan (2000s), New Zealand (2000s), Mongolia (2000s) and 567 Japan (2005). GFAD estimates forest age in tropical regions, where widespread 568 inventories are not available, by applying plant-functional-type-specific biomass-569 age curves⁴⁹ to a large-scale forest biomass dataset⁵⁰.

571 **Forest modelling.** The LPJ-GUESS DGVM v4.0⁵¹ was used to calculate the effects 572 of τ on forest structure, dynamics and carbon cycling. LPJ-GUESS explicitly 573 simulates forest stand development and canopy structure divided among age cohorts of trees co-occurring in patches representative of a wider landscape. Leaf 574 575 area to sapwood area ratio and maximum crown area for tropical evergreen tree 576 types were set to 10 000 and 130 m² respectively, in accordance with estimates 577 for tropical forests^{52,53}. Mortality and establishment are stochastic, with replicate 578 1000 m² patches simulated to capture the distribution of stands of different time-579 since-last-disturbance across each grid cell. Stand-replacing disturbances are 580 simulated by clearing all trees in a patch and transferring their biomass stocks to 581 litter or out of the ecosystem (see below). We introduced a spatially-varying 582 stochastic disturbance frequency with an annual probability defined by $1/\tau$. In 583 order to allow LPJ-GUESS to simulate the closed-canopy forest area unrestricted 584 by the 25% cover threshold used to calculate τ_0 , the nearest-neighbour rule was 585 used to assign τ values to grid cells with less than 25% forest coverage. All forest-586 type-level and global numbers are presented based on a 5% minimum forest 587 coverage mask at the grid-cell level to avoid overextrapolation of τ_0 to regions 588 with very low forest cover. The map in Fig. 2 is presented with a 25% closed-589 canopy forest cover map for consistency with Fig. 1. Inclusion of τ_0 in LPJ-GUESS 590 improves the simulation of biomass compared to the disturbance settings in the 591 standard version of the model (Suppl. Fig. 12).

592

In the standard simulation setting, all cleared biomass is transferred to the litterpools. For sensitivity simulations underlying ranges in Fig. 3d,e and Suppl. Fig. 10

595 two further setting types were employed to test the effect of the fate of disturbed 596 material. In the harvest sensitivity simulations fine root and leaf biomass, along 597 with 34% of woody biomass, are transferred to the litter, with the remaining 598 woody biomass being removed from the ecosystem, emulating product extraction. 599 The fire sensitivity simulations employ the interactive fire sub-model^{51,54} with a 600 local probability of fire occurrence (burnt area fraction) set to $1/\tau$, resulting in 601 most biomass carbon and some litter carbon being transferred to the atmosphere. 602 Stochastic processes use the same seed to ensure replication between simulations. 603 Simulations covered 1901-2014 using climate, atmospheric CO₂ mixing ratio and 604 N deposition as described in Le Quéré et al.⁵⁵. All model outputs shown are means 605 for 2001-2014. The standard simulations with τ_0 and $0.5\tau_0$ used 100 replicate 606 patches per grid cell. Simulations testing additional multiplicative perturbations 607 of τ (0.25, 2, 4) and using the confidence intervals of τ_0 used 10 replicate patches. 608 Differences at forest-type level were negligible between simulations with 10 and 609 100 patches.

610

611 Simulations used to create Fig. 3a used the standard model setup described above, 612 but the model was only run for the specified three grid cells. 100 simulations were 613 carried out for each grid cell using levels of τ from 10 years to 1000 years. A 614 second-order exponential equation of the form B = $ae^{b\tau} + ce^{d\tau}$ was fitted to these 615 simulations using the "fit" function of Matlab® 2014b.

616

617 **Sensitivity metric.** The metric is based upon differencing biomass between the 618 τ_0 and $0.5\tau_0$ simulations. The choice of a halving of τ_0 for the sensitivity metric 619 was informed by recent disturbance trends in Europe¹⁴, and is also similar to

620 changes in background mortality rates in the western U.S.A.⁵⁶. It thus represents 621 a reasonable sensitivity test. The sensitivity threshold τ_{crit} (Suppl. Table 1) was 622 estimated by first plotting against τ_0 the difference between biomass simulated 623 with $\tau = \tau_0$ and that simulated with $\tau = 0.5\tau_0$, (Fig. 3b, Suppl. Fig. 13). A second-624 order exponential function was fitted to the data as for Fig. 3a. These fits were 625 carried out both globally and for individual forest types. 95% confidence intervals 626 for the fits were calculated using 1000 bootstrapped samples of the modelled grid cells. $\tau_{crit,90}$ and $\tau_{crit,80}$ were taken as the intersection of the fitted line with a 627 628 difference of -10% and -20% biomass respectively (Suppl. Fig. 13), with 629 confidence intervals for $\tau_{\rm crit}$ estimated using the confidence intervals of the fitted 630 lines . Scatter in the results is caused by the stochastic nature of the LPJ-GUESS 631 model, as well as variation in climate across the domain. The 90% biomass 632 threshold is consistent with recent work on the recovery of forest biomass³³ and 633 with the character of the curve in Fig. 3b.

634

635 The area of forest in each sensitivity regime (Fig. 3c) was created by comparing τ_0 636 for each grid cell with the τ_{crit} for the forest type to which that grid cell was 637 assigned. Uncertainty in the areas of the regimes (Main text) was calculated based 638 on the 95% confidence intervals of τ_0 . For forest grid cells not classified by one of 639 the seven forest types, not enough data points existed to make a reliable fit to 640 calculate τ_{crit} . Therefore the global mean τ_{crit} was used to determine the sensitivity regime. Fig. 3d,e shows the difference in biomass and soil carbon density between 641 642 model sensitivity simulations with different multiplicative factors of τ (see above) 643 averaged across the area of forest allocated to each sensitivity class. Variation in

response across the vulnerability classes is much less than that between them(Suppl. Fig. 9).

646

647 Empirical cross-checks. For cross-checking of biomass carbon turnover flux due 648 to disturbance (F_d) we used the GEOCARBON global biomass dataset^{27–29}, in which 649 biomass values are based on linking satellite-based LIDAR and radar observations 650 with ground-based forest plot data. We replaced values for northern forests with 651 those of Thurner et al.³⁰ due to the latter's more sophisticated approach to linking 652 satellite-based radar observations with above- and below-ground biomass in 653 these regions. Below-ground biomass for the GEOCARBON dataset was estimated 654 following Saatchi et al.⁵⁰ and a biomass to carbon conversion factor of 0.5 was 655 assumed. We then multiplied the carbon content of this observationally-based 656 total biomass dataset by $1/\tau_0$ to calculate F_d . Cross-checking the fraction of total 657 turnover due to disturbance (T_{frac}) involved making the assumption that NPP and 658 turnover fluxes are not drastically out of equilibrium, and therefore NPP must be broadly equal to the turnover flux of biomass carbon in the multi-annual mean. 659 660 Annual mean NPP over the period 2001-2010 was calculated from Zhao and Running⁵⁷. T_{frac} was then approximated as F_d /NPP. Fire emissions from the GFED 661 662 dataset³² were calculated by summing the boreal, temperate and tropical forest 663 wildfire emissions, excluding the savannah category, which does not fit our 664 definition of closed-canopy forest. The mask of at least 5% forest cover per grid 665 cell was applied to all these cross-check calculations as above.

666

667 Data availability

669	Calcul	ations o	of τ _o , dat	a from th	e model s	mulations	and th	ne fore	st mask us	ed are
670	availa	ble	from	https://c	lataguru.lu	.se/app#P	ughDis	turban	<u>ce2019</u>	(dois:
671	10.182	161/dist	turbance_	_tauo.2019	905,	10.18161,	/distur	bance_	lpj-guess.2	01905,
672	10.182	161/dist	turbance_	forestmas	sk.201905)	. GFAD	v1.1	was	obtained	from
673	PANG	AEA ²² , a	and the G	lobal Fore	est Change	2000-2014	4 v1.2 t	forest l	loss produc	ct from
674	<u>https:</u>	//earthe	enginepar	tners.app	spot.com/	science-20	13-glot	<u>bal-</u>		
675	forest/download v1.2.html. The ESA CCI Landcover v2.0.7 was obtained from						d from			
676	http://maps.elie.ucl.ac.be/CCI/viewer/.									
677										
678	Code	availal	bility							
679										
680	Matla	b cod	e for t	he data	analysis	herein	is av	ailable	from C	GitHub,
681	https://github.com/pughtam/GlobalDist. Source code for LPJ-GUESS v4.0 can be						can be			
682	obtained on request through Lund University, see web.nateko.lu.se/lpj-guess.						ess.			
683										
684	Methods References									
685										
686	45.	ESA. I	Land Cove	er CCI Pro	duct User	Guide Ver	sion 2.	0. (ESA	Α,	
687		2017)). http://1	naps.elie.	ucl.ac.be/	CCI/viewe	er/dow	nload/	ESACCI-L	C-Ph2-
688		PUGv2	2_2.0.pdf							
689	46.	Kalam	nandeen,	M. et al. P	ervasive F	ise of Sma	all-scal	e Defoi	restation i	n
690		Amaz	onia. <i>Sci.</i>	<i>Rep.</i> 8, 16	00 (2018)).				
691	47.	de Gro	oot, W. J.	et al. A co	mparison	of Canadia	an and	Russia	n boreal fo	orest
692		fire re	egimes. Fo	or. Ecol. M	anage. 29	4, 23–34 ((2013)			

693	48.	Pugh, T. A. M. et al. Role of forest regrowth in global carbon sink
694		dynamics. Proc. Natl. Acad. Sci. U. S. A. 116, 4382–4387 (2019).
695	49.	Marin-Spiotta, E., Cusack, D. F., Ostertag, R. & Silver, W. L. Trends in above
696		and belowground carbon with forest regrowth after agricultural
697		abandonment in the neotropics. in Post-agricultural succession in the
698		Neotropics (ed. Myster, R. W.) (Springer, New York, 2008).
699	50.	Saatchi, S. S. et al. Benchmark map of forest carbon stocks in tropical
700		regions across three continents. Proc. Natl. Acad. Sci. 108, 9899–9904
701		(2011).
702	51.	Smith, B. et al. Implications of incorporating N cycling and N limitations
703		on primary production in an individual-based dynamic vegetation model.
704		Biogeosciences 11, 2027–2054 (2014).
705	52.	Herwitz, S., Slye, R., Erwitz, S. T. R. H. & Lye, R. O. E. S. Long-term
706		survivorship and crown area dynamics of tropical rain forest canopy
707		trees. <i>Ecology</i> 81, 585–597 (2000).
708	53.	Calvo-Alvarado, J. C., McDowell, N. G. & Waring, R. H. Allometric
709		relationships predicting foliar biomass and leaf area:sapwood area ratio
710		from tree height in five Costa Rican rain forest species. <i>Tree Physiol.</i> 28,
711		1601–1608 (2008).
712	54.	Thonicke, K., Venevsky, S., Sitch, S. & Cramer, W. The role of fire
713		disturbance for global vegetation dynamics: coupling fire into a Dynamic
714		Global Vegetation Model. <i>Glob. Ecol. Biogeogr.</i> 10, 661–677 (2001).
715	55.	Le Quéré, C. et al. Global Carbon Budget 2016. <i>Earth Syst. Sci. Data</i> 8, 605–
716		649 (2016).

- 717 56. van Mantgem, P. J. et al. Widespread increase of tree mortality rates in the
- 718 western United States. *Science* **323**, 521–524 (2009).
- 719 57. Zhao, M. & Running, S. W. Drought-Induced Reduction in Global
- 720 Terrestrial Net Primary Production from 2000 Through 2009. *Science*
- 721 **329,** 940–944 (2010).