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Pervasive shifts in forest dynamics in a changing world

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1 Pervasive shifts in forest dynamics in a changing world

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51 Enhanced abstract.

52 **Background:** Forest dynamics arise from the interplay of chronic drivers and transient

53 disturbances with the demographic processes of recruitment, growth, and mortality. The

resulting trajectories of vegetation development drive the biomass and species composition of

55 terrestrial ecosystems. Forest dynamics are changing due to anthropogenic-driven exacerbation

- of chronic drivers, such as rising temperature and CO₂, and increasing transient disturbances,
- 57 including wildfire, drought, windthrow, biotic attack, and land-use change. There are
- 58 widespread observations of increasing tree mortality due to changing climate and land use,
- accompanied by observations of growth stimulation of younger forests due to CO₂ fertilization.
- These antagonistic processes are co-occurring globally, leaving the fate of future forests
- uncertain. We examine the implications of changing forest demography and its drivers as a
- critical uncertainty for both future forest management and forecasting impacts of global climateforcing.

64 Advances: We reviewed the literature of forest demographic responses to chronic drivers and

transient disturbances to generate hypotheses on future trajectories of these factors and their

subsequent impacts on vegetation dynamics, with a focus on forested ecosystems. We

- 67 complemented this review with analyses of global land-use change and disturbance datasets to
- 68 independently evaluate the implications of changing drivers and disturbances on global-scale tree
- 69 demographics. Ongoing changes in environmental drivers and disturbance regimes are
- 70 consistently increasing mortality and forcing forests towards shorter and younger stands,
- reducing potential carbon storage. Acclimation, adaptation, and migration may partially mitigate
- these effects. These increased forest impacts are due to natural disturbances (e.g. wildfire,
- drought, windthrow, insect/pathogen outbreaks) and land-use change, both of which are
- 74 predicted to increase in magnitude in the future. Tree growth, and potentially recruitment, may
- have increased globally in the 20th century based on atmospherically derived estimates of the terrestrial carbon sink and based on remote sensing data, but the growth of this carbon sink has
- slowed. Variability in growth stimulation due to CO_2 fertilization is evident globally, with
- observations and experiments suggesting that forests benefit from CO₂ primarily in early stages
- 79 of secondary succession. Furthermore, increased tree growth typically requires sufficient water
- and nutrients to take advantage of rising CO_2 . Collectively, the evidence reveals that it is highly
- 81 likely that tree mortality rates will continue to increase while recruitment and growth will
- respond to changing drivers in a spatially and temporally variable manner. The net impact will
- 83 be a reduction in forest canopy cover and biomass.

84 **Outlook:** Pervasive shifts in forest vegetation dynamics are already occurring and are likely to

- accelerate under future global changes, with consequences for biodiversity and climate forcing.
- 86 This conclusion is robust with respect to the abundant literature evidence and our global
- 87 assessment of historical demographic changes, but it also forms the basis for hypotheses
- regarding the patterns and processes underlying the shifts in forest dynamics. These hypotheses
- 89 will be directly testable using emerging terrestrial and satellite-based observation networks. The
- 90 existing evidence and new observations provide a critical test of Earth system models that
- 91 continue to improve in their ability to simulate forest dynamics and resulting climate forcing.

- 92 Ultimately, forest managers and natural resource policies must confront the consequences of
- 93 changing climate and disturbance regimes to ensure sustainable forests and accrue their
- 94 associated benefits.

- 96 **Abstract:** Forest dynamics arise from the interplay of environmental drivers and disturbances
- 97 with the demographic processes of recruitment, growth, and mortality, subsequently driving the
- biomass and species composition of terrestrial ecosystems. However, forest disturbances and
- subsequent recovery are shifting with global changes in climate and land use, altering forest
- 100 dynamics. Ongoing changes in environmental drivers, land use, and disturbance regimes are
- forcing forests towards younger, shorter stands. Rising CO₂, acclimation, adaptation, and
 migration can influence these impacts. Recent developments in Earth system models support
- 103 increasingly realistic simulations of vegetation dynamics. In parallel, emerging remote sensing
- 104 datasets promise qualitatively new and more abundant data on the underlying processes and
- 105 consequences for vegetation structure. When combined, these advances hold promise to improve
- the scientific understanding of changes in vegetation demographics and disturbances. Pervasive
- shifts in forest dynamics are already occurring and are likely to accelerate under future global
- 108 changes, with consequences for climate forcing.

110 Introduction

The interplay of vegetation demography—recruitment, growth, and mortality—with 111 environmental conditions and disturbances drives forest dynamics of biomass, function, and 112 113 species composition (Figure 1; see Box 1 for definitions). In old-growth forests that approximate steady-state demographics, the recruitment, growth, and mortality of trees are approximately 114 balanced; in contrast, rapid recruitment often follows widespread disturbance-induced mortality 115 116 (1). Vegetation dynamics may now be changing because the environmental context in which plant demography and disturbances interact is shifting with anthropogenic change (Figure 1). 117 The interaction between episodic forest disturbances such as wind-throw or wildfire, and 118 119 chronically changing drivers such as rising temperature, vapor pressure deficit (VPD), and CO₂, 120 together with land-use change (2), leads to both compounding and antagonistic impacts that alter demographic rates (3), with consequences for terrestrial biogeochemical cycles and climate (4,5). 121 Understanding the drivers of vegetation dynamics is thus critical for accurate prediction of global 122 123 terrestrial biogeochemistry under future conditions (6).

The impacts of global change on forest demographic rates may already be materializing. 124 125 In mature ecosystems, tree mortality rates have doubled throughout much of the Americas and in 126 Europe over the last four decades (7-9). Simultaneously, global carbon budgets indicate either a 127 growing or constant terrestrial carbon sink (10-12), which implies increased or constant 128 vegetation production rates (13,14). However, satellite evidence suggests that forests might be 129 switching from a CO₂ fertilization dominated period to a VPD dominated period (15). Terrestrial greening indices indicate a shift from a CO₂-driven increase in greenness in the late 20th century 130 131 to a VPD-driven decrease in the last decade (16). Thus, increasing mortality due to anthropogenic changes, along with potentially increasing or stable growth and recruitment due to 132

133 CO₂ fertilization (5), represent opposing processes that are co-occurring globally, leaving the fate134 of future forests uncertain.

135 Beyond changing vegetation dynamics within "intact" or relatively undisturbed forests, 136 episodic disturbances are tending to be larger, more severe and, in some regions, more frequent under global change(17-20). Similarly, the rates and types of land-use change (LUC) vary widely 137 138 (21) but have, on average, increased globally in the past few centuries (2,22,23). Thus, at the global scale, disturbances and LUC have likely amplified tree mortality beyond that suggested by 139 140 the doubling of background mortality rates in undisturbed forests (7-9). Current understanding of the net balance of tree losses (mortality) and gains (recruitment and growth) under a changing 141 142 environment characterized by more extreme drivers and disturbances is limited, preventing prediction of whether recruitment and growth can balance increased mortality rates in the future. 143

To evaluate whether environmental changes and increasing disturbances are causing 144 globally widespread shifts in vegetation demography, we reviewed global observations of 145 recruitment, growth, and mortality of forests and woodlands. Our expert-derived compilation of 146 the state-of-the-art knowledge on vegetation dynamics, their drivers, and disturbances, allowed 147 148 us to address four questions: i) Is there evidence for shifts in demography over recent decades? ii) What physiological and disturbance-mediated processes underlie these demographic shifts? 149 150 iii) What are the potential consequences of disturbance-mediated changes in demography for 151 climate forcing? iv) How can global predictions of future vegetation dynamics best be improved?

152 Evidence for changing drivers and disturbances and their impact on demography.

Determining the impacts of changing drivers on demography is difficult given the lack of global observation platforms. However, evidence abounds from individual published studies on the drivers and their impacts on plant communities, and new modeling and observational efforts

now enable a more complete picture of disturbances and forest demography (24-26). In this 156 section, we first examine if there are global trends in stand ages and test the sensitivity of the 157 158 stand-age distribution to changes in disturbance rate using global datasets on LUC (27) and non-LUC (25,28) disturbances. We subsequently draw upon the wealth of published studies on 159 changes in forest demographics and their drivers to investigate the potential changes leading to 160 161 global age-trends. Ultimately, the combination of our global estimates along with the large literature base allows us to generate testable hypotheses regarding trends and impacts of the 162 drivers of forest demographics. 163

164

165 Is disturbance changing forest demography at the global scale?

We re-analysed the Land-use Harmonization (LUHv2) dataset (28) with respect to forest 166 167 age, revealing that the area of young forest stands (here defined as <140 years old) resulting 168 directly from LUC (conversion of forest to non-forest) or wood harvest (reduction of biomass and age but retained as forest) has increased from 4.8 million km² in 1900 to 12.5 million km² in 169 170 2015 (or from 11.3% to 33.6% of forest area; Figure 2A). The results were insensitive to 171 assumptions regarding the link of disturbance likelihood to stand age (Figure 2A). These forest 172 stand age distributions exhibit different trajectories in different regions. Tropical forests have progressively lost old-growth area due to LUC over the 20th century (Figure 3a, black dashed 173 174 line). Wood harvest has increased from a minor driver of tropical forest age distribution in 1900 to a major one in 2015 (difference between solid and dashed lines). The split between 175 176 deforestation and shifting cultivation drivers is broadly consistent with a satellite-based analysis for the period 2001-2015 (29). Temperate and Mediterranean forest ages are strongly influenced 177 by wood harvest, which has made old-growth forests increasingly scarce in these regions. Boreal 178

forests saw little influence of *LUC* on stand age, but wood harvest has substantially shifted theage distribution towards young forests.

181 In reality, old-growth forest are scarcer than that due to LUC and wood harvest (Figures 182 2a and 3), due to other disturbances that have shifted landscapes from old- to young-dominated stands (14), such as wildfire (29), windthrows (30), and biotic outbreaks (31). To address these 183 184 additional disturbances, we integrated recent observation-based estimates of non-LUC disturbance for closed-canopy forests (25,28) with LUC from LUHv2, to obtain a first principles 185 estimate of the combined effect of human and natural disturbances on forest age structure (Figure 186 2B). A twofold increase in non-LUC disturbance rates over the period 2015-2050 would result in 187 a substantial increase in the fraction of young forests (Figure 2B, C). Thus, realistic shifts in 188 disturbance rates can have substantial impacts on the age structure of forests in the future. As 189 discussed below, such an increase in disturbance rate is consistent with the magnitude of changes 190 observed or predicted in individual ecosystems. 191

Notably, calculations based on the Global Forest Age Dataset (GFAD) v1.1 (14,32)
yielded 16.5 M km2 old-growth and 26.3 M km2 of young forest (32), which differs from that in
Figure (2B, C). This disparity is likely attributable to consideration of different forest types
(closed-canopy versus all forests) and to differences in definition of stand size and age used in
inventories versus used in satellite-based estimates

197 *Chronically changing drivers: Atmospheric CO*₂ has risen more than 125 ppm since the
198 industrial revolution (11), and is projected to rise an additional 50 to 200 ppm by 2100. Higher
199 CO₂ increases leaf-level water use efficiency, and rising CO₂ has positive but uncertain
200 feedbacks on plant demographic rates (Figure 4A-B). Maturation and seed production can be
201 accelerated under elevated CO₂ (33); however, seedling growth is not always stimulated by CO₂

(34). Recruitment response to rising CO₂ is variable (35,36). Forest inventory and tree-ring 202 203 studies show limited evidence for CO_2 fertilization of growth (37-43), potentially due to the 204 overwhelming influence of increasing drought and nutrient limitations (44). Ecosystem-scale CO₂ enrichment experiments in young forests suggest a 30% gain in decadal biomass increment 205 (45), but experiments in mature forests have found minimal growth stimulation (46,47). This is 206 207 consistent with evidence for an initially strong CO2-related growth stimulation in young forests that decreases with tree age and size³⁹ due perhaps to nutrient (7,48) and hydraulic path-length 208 209 limitations (49).

210 Limited studies suggest elevated CO₂ causes increased mortality or no change in mortality. Mortality rates of saplings during experimental drought were not mitigated by 211 elevated CO₂ (50,51) while accelerated self-thinning due to CO₂ fertilization-induced stand 212 density increases may lead to higher mortality (6,52,53) (Figure 4B). The latter process would 213 214 be consistent with increases in recruitment at large scales. Because tree mortality is dominated 215 by large size classes i.e. (54) (for details see section on size-related mortality below) faster growth via CO₂ fertilization may expose trees to size-related mortality risks earlier (7). Such 216 CO₂-induced increases in mortality may be global (55). Furthermore, faster growth is often 217 218 associated with lower wood density (56), rendering fast growing trees more susceptible to high 219 winds. Thus, future CO₂ fertilization could increase recruitment, growth, and mortality (Figure 220 4B), though there is significant uncertainty about these effects.

221

Chronically changing drivers: Temperature and vapor pressure deficit are rising globally and
will continue to rise into the future (57). Both temperature and *VPD* can have impacts on
demographic rates. Rising temperature forces an exponential rise in *VPD*, which prompts

stomatal closure and limits photosynthesis, leading to lower growth, higher mortality (58), and 225 226 reduced regeneration (59), and ultimately driving community shifts (60,61). These observations 227 are consistent with hydraulic theory, which suggests that as VPD rises, potential maximum tree height declines (62) (Figure 5). This results from the dependency of water transport limitations 228 on tree size (49) that are exacerbated by elevated VPD (Figure 5), making short stature 229 230 advantageous with rising VPD. Because most plants cannot reduce their size (beyond limited reductions in leaf area or crown dieback), forests respond through increased mortality of large 231 232 plants, which are replaced by smaller ones (62), as has been observed in many studies (26,54). 233 While rising air temperature may also increase respiratory carbon loss, leaving less carbon for growth (63), warming in wetter and cooler regions may actually stimulate reproductive output, 234 recruitment, and growth (64-66). Changes in temperature and VPD also can produce asynchrony 235 in floral and pollinator phenology (67) and can reduce cold stratification (68), both of which 236 reduce seed abundance (69), and negatively affect recruitment (70,71). Sapling mortality is 237 238 accelerated by elevated temperature (71,72), but recruitment has increased in moist areas (73). Thus, rising temperature and VPD may be beneficial in cooler or wetter areas, but most evidence 239 suggests negative impacts on plant demographic rates (Figure 4C, D; Figure 5). 240

241

Changing disturbance regimes: Droughts are anticipated to increase in frequency, duration,
and severity globally (Figure 4E-F) and are more stressful to plants due to increases in
temperature, *VPD*, and associated water loss (57). Drought can directly cause tree death or
indirectly lead to mortality through associated increases in insect or pathogen attack (51).
Hydraulic failure and carbon starvation remain the most likely, mutually inclusive, underlying
physiological mechanisms for drought-induced mortality (74) and both processes are likely to
increase tree susceptibility to biotic agents (75). Evidence suggests that drought-induced

mortality occurs more rapidly under warmer conditions (51.72). Consistent with these empirical
results, models suggest far greater mortality of temperate conifer trees in the future (76).
Reproductive output is often reduced by drought (but see (64)), which combined with drought
impacts on seedling survival, leads to reduced recruitment (77). However, growth was relatively
stable across a drought in Amazonia (78), while mortality increased. Thus, like rising
temperature and *VPD*, it appears that drought may increase mortality regardless of location,
while having variable impacts on recruitment and growth (Figure 4F).

256

257 Changing disturbance regimes: Land-use change and forest management have reduced vegetation stature and biomass, and altered species composition, with profound consequences for 258 forest dynamics (Figures 2A, 4G-H). Today's global vegetation biomass stocks may amount to 259 only ~50% of their potential due to LUC (79). Wood harvest and shifting cultivation are the 260 land-use activities primarily responsible for the conversion from primary to secondary vegetation 261 262 cover and associated demographic shifts (2). For systems that return to wild-vegetation or to managed forest after human clearing, demographic rates are typically accelerated. The increased 263 resource availability after forest removal facilitates establishment of early-successional species, 264 265 reduces species diversity (80,81), and causes a transition to younger, smaller plants (82). Postdeforestation recruitment is often prolific even in the absence of management (83). Globally, the 266 267 recovery of harvested forests and abandoned agricultural land, along with establishment of new 268 plantations, has resulted in younger forests (Figure 2A), with associated reductions in tree size and biomass (84). Such post-deforestation recruitment may be limited by elevated VPD or 269 270 drought, as is the case with recruitment following all natural disturbances. Overall the net effect

of historical *LUC* and wood harvest has resulted in a substantial loss of forest area, along with
altered demographic rates, leading to younger, shorter, less diverse ecosystems (Figure 4H).

Changing disturbance regimes: Wildfire is increasing in many forests worldwide (85) (Figure 274 4I), although human management of landscapes has led to wildfire suppression in some biomes 275 276 (86). Given sufficient fuel, burned area increases exponentially with aridity (87), and future fire frequencies may exceed those documented over the past 10,000 years (88). Increased fire 277 278 activity causes increased mortality and potentially higher recruitment and growth of either pre-279 existing or new species, but rates of recruitment and growth may be slowed under climate warming. Forests characterized by stand-replacing fire regimes are dominated by obligate 280 seeders, and typically have effective seedling recruitment (89). However, high-severity and high-281 frequency fires can reduce recruitment by reducing seed supply through the repeated and severe 282 loss of reproductively mature vegetation (90), and high frequency fires can cause recruitment 283 284 losses via direct mortality of the seedbank, seedlings, and saplings (91), which is worsened by elevated VPD (92). Woody species that can resprout following fire, including shrubs that 285 suppress tree regeneration (92), may be favored by increased fire frequency and severity. 286 287 Increased fire severity results in high tree mortality in forests historically adapted to low-severity fires, and subsequent recruitment and growth may be slow or absent, resulting in conversion of 288 289 forests to low-biomass ecosystems (93). Thus, wildfire can result in higher demographic rates, 290 though rising temperature and VPD can negatively impact recruitment and growth (Figure 4J). 291

292 *Changing disturbance regimes: Wind throw* from cyclonic storms represents the dominant
293 natural disturbance in coastal forests across the globe (94). Cyclonic storms are expected to

increase in frequency, wind velocities, and precipitation intensity (95) (Figure 4K), with more 294 295 extreme flooding that promotes tree instability. Wind throw also results from convective 296 thunderstorms and topographically mediated winds, and warming is expected to increase the frequency of atmospheric conditions conducive to severe thunderstorms (96). Canopy damage 297 and whole-tree mortality are the most immediate impacts of wind throw (97) (Figure 4L). Larger 298 299 trees dominate mortality from storms (98), and the loss of large canopy trees during wind disturbance favors growth of surviving trees (98,99) and advances regeneration, recruitment of 300 301 early successional species (100), or resprouting of trees broken by wind (101). Depending on the 302 resprouting or seeding capacity of surviving species, wind damage may slow or accelerate succession (102). We note that storms may also be associated with lightning, which may be a 303 significant cause of large-tree mortality (103). Thus, windstorms should result in changes in all 304 three demographic rates, though with large uncertainty at the global scale (Figure 4L). 305

306

307 Changing disturbance regimes: Biotic agents Biotic disturbances from insects, insect-pathogen complexes, and other biotic agents have been increasing in frequency, severity, and extent in 308 recent decades (17,104,105) (Figure 3M). Such trends reflect a changing climate (106), altered 309 310 land use (107), and introductions of non-indigenous insects and pathogens (108). Climate change is expected to further amplify biotic disturbances (109) in part through enhanced host 311 312 vulnerability (Figure 4M). However, shifts in frequency or dampening of disturbance regimes 313 could also emerge (110), leading to some uncertainty in outbreak dynamics under future conditions (Figure 4M). While insects and associated pathogens are globally widespread, lianas, 314 315 or vines that use other plants as host structures, are increasing in abundance and are thought to be 316 causing increasing mortality in the tropics (7,111).

Response of insects and pathogens to climate change is likely to increase plant mortality 317 (4), with variable impacts on growth and recruitment (Figure 4N). Tree mortality can result from 318 319 girdling of the phloem and xylem by bark beetles (75) and from repeated defoliation events that exhaust the capacity of trees to recover (112). Tree mortality during outbreaks is usually partial 320 at the stand-level because many biotic agents preferentially attack trees of specific size- or 321 322 health-classes, or are host-specific (16). Suppressed, smaller trees and non-host tree species may survive and grow rapidly when released from competition for resources (113,114). Thus, similar 323 324 to many other disturbances, mortality increases while recruitment and growth show variable 325 responses to biotic disturbances, including a dependency on post-disturbance temperature, VPD, and drought. 326

327

On size and age demographics. The combination of LUC, disturbances, and chronic drivers is 328 329 likely to have already shifted forests to younger and shorter stands, with these impacts increasing 330 under expected future changes in drivers and disturbances (Figure 2A-C). These results are consistent with our review of the literature (Figure 4). Large trees are the most susceptible to die 331 from LUC-caused forest fragmentation (115,116), drought (26), rising temperature or VPD 332 333 (54,62) (Figure 5), windthrow (117,118), biotic attacks (119), and lightning (103), with variable size-impacts of fire (120). The abundance of size-dependent mortality drivers and disturbances 334 335 should logically push stands towards younger/smaller distributions of trees and shorter statured 336 species assemblages (121).

There are exceptions to the pattern of climate drivers and disturbances reducing tree height and stand age. Non-stand-replacing fires that kill smaller trees and leave the larger, older trees will shift forests towards larger size distributions. Similarly, on occasions when droughts

preferentially kill younger but fast-growing trees, this would impact the subsequent size
distribution and rate of carbon accumulation. Rising CO₂ and increased precipitation in some
areas also counter the general decrease in size because they may lead to faster growth and hence
taller trees (122). Thus, the antagonistic drivers promoting larger trees (e.g. rising CO₂) vs
smaller trees (e.g. rising *VPD*, increasing disturbances) co-occur, but the general pattern of
decreasing size and younger ages reveals that processes driving down size and age (Figures 2-5)
are dominant globally.

347

348 Mitigation of demographic-disturbance impacts

349

350 The literature patterns suggest most drivers and disturbances will increase tree mortality now and in the future, with variable effects on recruitment and growth (Figure 4). The 351 352 uncertainty grows, however, when we consider multiple feedbacks that can mitigate the changes 353 in forest demography induced by chronically changing drivers and disturbance regimes. These 354 processes include acclimation, adaptation, migration, and compensatory mechanisms of resource use. With global change, forests will be influenced by a combination of phenotypic plasticity 355 (i.e. acclimation (123)), adaptation to novel biotic and abiotic stresses (124), and the ability to 356 migrate as conditions change (125). Failure to acclimate, adapt, or migrate, including due to 357 358 human infrastructure (126), could lead to recruitment and growth reductions and local extinctions. Plants have demonstrated acclimation of phenology, seed longevity, and metabolic 359 360 processes to single and/or multiple stressors (127-130). Acclimation and adaptation will likely 361 depend on an array of factors including genetic variation, fecundity, dispersal, population size, and environmental variability (123). Many tree species have migrated in response to past 362 363 climatic cycles but at rates slower than the current pace of climate change (131). Regarding

resource use, reductions in stand density as a result of increased mortality or reduced recruitment 364 should allow greater resource availability to surviving individuals, and therefore subsequently 365 366 higher growth and survival rates (132). Such stand-resource mechanisms can manifest at the landscape scale, as most disturbances are patchy (133), and the size, shape, and arrangement of 367 surviving forest patches can play a key role in recovery of the disturbed landscape (20). Taken 368 369 together, the mitigating factors can play a significant role in buffering the impacts of changing drivers on plant survival, but it remains unclear if these factors will enable trees to keep pace 370 371 with ongoing climate change (50,123). Ultimately, the uncertainty around future demographic 372 rates shown in Figure 3 is partially because of the influence of these mitigating factors.

373

375

374 Consequences for community assembly and for climate forcing

The widespread shift in vegetation dynamics begets questions regarding consequences for 376 community assembly and climate forcing. Hydraulic theory suggests that under rising VPD, 377 378 functional traits of high conductance, low stature, and low leaf area should best enable survival, all of which are characteristics of pioneer, shrub and weed species (62). Consistent with this, 379 diversity (e.g. species richness) temporarily increases post-disturbance for many systems, as 380 381 short-statured, opportunistic species invade (134). If forest communities shift towards trait assemblages better suited to the new disturbance regime, such shifts may confer some resistance 382 383 to future disturbances (134,135). Alternatively, if disturbance regimes shift faster than 384 recruitment, growth, and subsequent community assembly can respond, resistance to future disturbances will likely decline. 385

Climate forcing responds to changing vegetation dynamics in complex ways. Changes in
 forest disturbance regimes and vegetation dynamics can affect climate via biogeochemical,

hydrological, and land-surface energy budgets (136). Reductions in biomass result in a loss of 388 carbon to the atmosphere despite younger, shorter stands often having higher gross 389 390 photosynthesis; this is due to the loss of carbon through decomposition of necromass, which is a particularly large flux from mortality of older, larger trees such as those in old-growth forests 391 (137), and reduced landscape-mean carbon storage under an intensified disturbance regime 392 393 (138). The time required to re-achieve the same live carbon storage of an ecosystem after disturbance can be decades to centuries, particularly if the disturbance cycle is increased, thus the 394 395 net effect of the biomass loss is increased CO₂ to the atmosphere and hence greater climate 396 forcing. This impact may be mitigated by increased carbon uptake due to CO₂ fertilization (139) or enhanced recruitment. Calculations of the terrestrial carbon sink from atmospheric inversions 397 indicate that the sink grew over recent decades¹² due in part to increased leaf area (13), 398 consistent with increased recruitment and growth. However, evidence suggests that forests are 399 switching from a CO₂ fertilization dominated period to a VPD dominated period (15,16), despite 400 401 sustained high gross photosynthesis at the global scale (140). The increased mortality throughout much of the terrestrial biosphere (7-9) further minimizes potential carbon storage 402 through the enhanced biomass loss. Ultimately, the terrestrial contribution to climate forcing 403 404 through carbon uptake and release results from the antagonistic process of rising CO₂ and forest recovery from LUC, which enhance the carbon sink, and rising VPD and disturbances that reduce 405 the carbon sink. 406

407 Changing vegetation dynamics also influence regional and global surface energy budgets 408 and hydrological cycles. Disturbances frequently shift albedo of ecosystems from darker to 409 lighter, resulting in a decline in radiative forcing through less light absorption (141). The rate of 410 recruitment post disturbance influences the temporal period of this negative feedback (142). The

impact of changing vegetation dynamics on the water cycle is particularly complex. Evaporation
from canopies shifts as stands become taller because taller trees transpire less (per unit leaf area)
than smaller trees (49), but larger trees often have better rooting access to water sources and have
greater total leaf area. The net effect of disturbance is a transient decrease in evaporative loading
to the atmosphere along with albedo shifts, causing a feedback of decreasing precipitation
downwind (143,144). Ultimately, carbon storage is at least transiently reduced by disturbances,
with mixed impacts on the water and energy budgets.

418

419 The path to improved prediction

420 Changes in the global drivers of temperature, CO₂, VPD, and disturbances including LUC, drought, wildfire, windstorms, and insect outbreaks, should all force forests towards 421 shorter, younger, lower biomass ecosystems. This trend is supported by hydraulic theory (62) 422 423 (Figure 5) and by abundant empirical evidence demonstrating a consistent increase in mortality 424 across the global-spectrum of drivers and disturbances and variable but often declining recruitment and growth (Figure 4). While the bulk of evidence points to reduced plant stature 425 due to changing drivers, large uncertainty remains in the magnitude and slope of demographic 426 427 trajectories in the future (Figure 4). Given these trajectories, and the large uncertainties around them, what are the critical next steps to allow improved global prediction? Continued long-term 428 429 observations (ground and remotely sensed) are essential to reveal the patterns of demographic responses to drivers and disturbances; likewise, manipulative experiments are needed that alter 430 431 conditions such as CO₂ or drought to provide cause-and-effect understanding of the interactions among mechanisms of demographic responses. However, for global-scale prediction of 432

responses and climate consequences we need to mainstream insights from observations andexperiments into Earth system models (*ESM*s).

435 ESMs simulate the exchange of fluxes between the atmosphere, land, and ocean and 436 stores of carbon, water, and energy; the land-surface modules of *ESMs* simulate the vegetation. *ESMs* have made great progress in simulating land use, disturbances, and demography, including 437 438 representation of wildfire (145), drought-induced mortality (146), and cohort-age structured models that enable representation of succession and associated shifts in physiological traits (6). 439 440 The global Coupled Model Intercomparison Projet CMIP6 now includes a dedicated model intercomparison activity focused on the effects of changes of land-use on carbon and climate 441 442 (147). Advances in remote sensing and forest inventory integration are enhancing global datasets of forest structure (148) and age (32) that can be used in model initialization, data 443 assimilation benchmarking, and sensitivity analyses (Figure 2A-C). These advancements set the 444 stage for developments in *ESMs* such as the prediction of disturbances and demographic rate 445 446 responses under climate and LUC scenarios.

The newest generation of *ESMs* utilize size or age-structured approaches to explicitly 447 448 model demography in the Earth system (6), which should ultimately enable model-based representation of observed shifts in age structure (e.g. Figure 2). However, representation of 449 vegetation demographic rates remains relatively simplistic. Simulation of growth responses to 450 451 global change requires model refinement in light capture, belowground water and nutrient acquisition, and responses of respiration to temperature (6). Recruitment, including reproduction 452 and dispersal, is the most undeveloped demographic process in *ESM* simulations. Reproductive 453 454 allocation is invariant with plant functional type (*PFT*), and seed is assumed to mix evenly 455 throughout a grid cell (but see (149)). Environmental constraints to *PFT* establishment are

derived from prior distributions of major taxa, and while recruitment rates can be influenced by 456 light or space availability, they are not responsive to temperature, CO₂, or soil moisture 457 458 (150,151). Simplistic dispersal assumptions are typically either overly permissive or restrictive. Improvements in representing recruitment under global change are critical for improving 459 predictions of vegetation dynamics. These advancements will require data synthesis and new 460 461 data collection to support PFT-specific, environmentally sensitive parameterizations of regeneration processes, such as reproductive allocation; effective dispersal; seedling 462 establishment, survival and growth; and post-disturbance recovery strategies (e.g., serotiny and 463 resprouting). 464

Disturbance-induced mortality is better developed for landscape-scale models than for 465 ESMs. ESM modeling of disturbance-induced mortality exists for wildfire and drought 466 (145,146), although significant challenges remain to represent both reliably globally, while ESMs 467 are under-developed for wind and insect mortality. Currently only one ESM to our knowledge 468 469 represents canopy damage (152); this causes ESMs to potentially underestimate the impacts of drought and wind, as both disturbances cause lagged tree mortality associated with canopy loss 470 years after the inciting event (153,154). For insects, there have been prescriptive studies 471 472 examining the impact of insect outbreaks on land processes within ESMs, but no ESM has yet explicitly considered the interaction between plant defense and insect population dynamics for 473 prediction of large-scale insect-induced tree mortality. For predicting wildfire, models should be 474 sensitive to both fuels and climate interactions and represent spatial patterns of burn severity 475 476 because the burn mosaic strongly influences postfire vegetation dynamics (145). Nextgeneration demographic models are evolving to include explicit, mechanistic representations of 477 drought-associated mortality, including carbon starvation and hydraulic failure (155). The 478

evaluation of new hydraulics models (155) for prediction of mortality is an essential next step.
Ultimately, model formulations that include environmentally sensitive, *PFT*-specific processes
compatible with the cohort-based approach are likely to provide the best compromise between
process-detail and parsimony and therefore most likely to capture changes in large-scale forest
dynamics under future conditions.

484

485 **Summary** Forest vegetation dynamics (Figure 1) are already strongly influenced by global change (Figure 2) and will continue to be affected in the future (Figures 2-5) by changes in land 486 use, chronic drivers such as CO₂ and VPD, and increasing frequency and severity of transient 487 488 disturbances such as windthrow, wildfire, and insect outbreaks. Effects on forests are driven largely by consistent increases in tree mortality from these drivers, and variable responses of 489 recruitment and growth depending on stand-age, disturbance type, and geographic location 490 (Figure 4). The consequences of changing demographics suggest an increasing constraint in 491 492 terrestrial carbon storage due, at least, to the consistent increase in mortality. Any declines in 493 recruitment or growth, especially when disturbance-recovery cycles are disrupted, will exacerbate this carbon-cycle constraint. Shifts in other terrestrial radiative forcing terms such as 494 energy and water budgets are also likely. While well-supported by the literature, data, and 495 496 sensitivity analysis (Figure 2), the trends in Figure (4) represent hypotheses to be tested by the next-generation of observational platforms, both terrestrial and space-borne. Forest management 497 must ultimately confront the elevated mortality and uncertainty in recruitment and growth when 498 499 considering options for sustaining forest benefits to society into the future.

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920

922 **Box 1 Vegetation dynamics definitions**

- 923 We focus on three main plant demographic processes: recruitment, growth, and mortality.
- 924 Recruitment (including reproduction) results in the seedling and sapling composition of a plant
- 925 community following disturbance (156). Growth from sapling to mature plants results in
- 926 development of mature forests and includes competitive processes. Mortality is a key rate
- controlling carbon storage and species composition in a plant community and is a dominant
- 928 demographic rate during a pulse-disturbance (157,158).
- Abiotic drivers. Physical factors that cause changes in demography and that respond to global
 change or to disturbances, such as light, CO₂, soil moisture, humidity, temperature, etc.
- Biotic drivers. Biological factors that may drive changes in demography, such as pathogens,
 insects, herbivores, or competition with other individuals.
- 933 Chronic environmental change. Persistently changing drivers of demographic rates. These
- 934 drivers have a non-stable and directional trajectory, such as rising CO₂, temperature, and *VPD*.
- 935 Demographic rate. Any individual-, population-, or community-level parameter that affects the
 936 age- and/or size-structure of a population or community, including rates of recruitment, growth,
 937 and death.
- 938 Demographic driver. An abiotic or biotic factor that, when undergoing a change itself, also
 939 leads to a change(s) in one or more demographic rates.
- 940 **Disturbance.** The destruction of live plant biomass in a discrete event (159,160).
- 941 Disturbance regimes. Spatial and temporal characteristics of disturbances in a landscape over a
 942 long time period, including frequency, return interval, duration, intensity, severity, and size.
- 943 **Growth.** The rate of biomass production over time at the individual or ecosystem scale (i.e. Net 944 Primary Production grams C $m^{-2} yr^{-1}$).
- Land-use and land-cover change. Anthropogenic shifts in forms of cultivation or in vegetation
 cover such as due to forestry, or conversion of woodlands to crop ecosystems.
- 947 Mortality. Defined herein as the complete loss of a plants' ability to reproduce and ultimately948 loss of cellular metabolism.
- **Recruitment.** The rates of transition of plants from one size class into another (typically in units of individuals m⁻² yr⁻¹). Recruitment results from the birth and growth of individuals. Herein we consider recruitment from the stage of seed dispersal through seedling growth into the sapling stage.
- 953 Self-thinning. Reduction in the number of live plants within a stand, occurring via competition954 for resources.
- 955 Vegetation dynamics. The net outcome of the interplay between disturbances and vegetation956 demographic rates.
- 957

958 Figures



959

960 Figure 1. A conceptual diagram of the components of forest dynamics and disturbances

that drive them. In the far-left panel, a mature ecosystem is responsive primarily to localized 961 962 mortality, and the primary drivers of demography are chronically changing variables such as 963 CO₂, temperature, and VPD. The system is disturbed in the second panel, due to fire, insect outbreak, or another large-scale perturbation that removes much of the overstory trees, and 964 species adapted to rapid post-disturbance recruitment become established. In the third panel 965 966 recruitment and growth dominate demographic processes, with mortality increasing over time as competition leads to self-thinning. In the last panel, a mature ecosystem is dominated by species 967 968 that have replaced the original community in response to chronic environmental changes, leading 969 to a novel ecosystem.



Figure 2. (a) Human activities have increased the amount of young forest area (stands < 140 years old) over the 20th century as a result of both landuse change (LUC) and wood harvest (WH). Forest stand age distribution was reconstructed using forest-cover transitions from LUHv2, initialized using forest cover fractions in 1750 and incrementing forest cover each year, tracking the age of forest up to 140 years. Solid lines show the effect of LUC and WH, dashed lines LUC alone. Total forest area is based on *LUHv2*. The nominal minimum size of a stand is assumed to be ca. 0.1 ha. (b) Sensitivity of age distribution in closed-canopy (CC) forests to plausible changes in disturbance rate. Forest stand age distribution was reconstructed using forest cover transitions due to LUC from LUHv2 alongside non-LUC observation-based disturbance rates (25). In the baseline scenario (solid lines), non-LUC disturbance is assumed constant at observed 2001-2014 values throughout. In the Inc. dist scenario (dashed lines), disturbance rates are incremented linearly to 200% of the 2001-2014 values over the period 2015 to 2050 and held constant at that level thereafter. The underlying LUC scenario is GCAM RCP 3.4 that includes land-based mitigation for CO₂ emissions. Results are presented for closed-canopy forests only (25), which is why total forest area is lower between (a) and (b), as non-LUC disturbance rate information is not currently available for

open-canopy forests. The shaded areas in panels **a** and **b** indicate the effect of assuming that disturbances 1007 are five times more likely to affect the youngest forests versus old-growth, or vice versa, as opposed to an 1008 1009 even probability across ages (solid lines). The apparent large dampening of this assumption in **a** vs **b** is primarily due to the different y-scales. (c) Changes in the disturbance regime propagate through 1010 forest age structure at decadal time scales. Closed-canopy young (<140 years old) forest area is 1011 shown on the left-hand y-axis. Old-growth (>140 years old; OG) forest area is shown on the right axis 1012 (same units) and refers to the data points in the upper right-hand of the panel. Scripts used and additional 1013 1014 methods can be accessed at https://github.com/pughtam/AgeClassReconst rel.git.

1015

1017 Figure 3. Human activities have increased the amount of young forest area irrespective of biome.

1018 As for Figure 2a, but broken down by biome 161 .





Figure 4. Drivers, disturbances, and demographics are changing both historically and into
the future. A graphical summary of the literature evidence of changing drivers and disturbances
(left-hand column) and subsequent demographic rates (right-hand column). Shown are the
chronically changing drivers A) CO₂, C) *VPD* and temperature and the more transient
disturbances of E) drought (low precipitation), G) deforestation, I) wildfire, K) wind, and M)
insect outbreaks. Each driver or disturbances' corresponding demographic responses (shown as
carbon fluxes per area⁻¹ time⁻¹) are shown on the right-hand panels.



VPD (kPa)

1032 Figure 5. Rising VPD forces declines in potential plant stature. Predictions of plant height in 1033 response to rising VPD from the hydraulic corollary to Darcy's law. The equation is h = $A_s*k_s*(\Delta\Psi)/G*A_l*VPD$, where h is height, A_s is sapwood area, k_s is specific conductivity, $\Delta\Psi$ is 1034 the leaf to soil water potential gradient, G is stomatal conductance, and A_1 is leaf area⁵³. The 1035 different lines represent different levels of acclimation of A_s , k_s , $\Delta \Psi$, G, and A_l , all allowed to 1036 adjust simultaneously from 0 to 60% from their initial values. In the case of G it is assumed to 1037 1038 decrease due to rising atmospheric CO₂. Acclimation can help, but not completely mitigate, the impacts of rising VPD on plant size. 1039

1040