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1 **Carbon accumulation rate peaks at 1000m elevation in tropical planted and**
2 **regrowth forests**

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60 **Keywords:** Tropical forests; Biomass carbon accumulation; Planted forests; Natural

61 growth forests; Elevational pattern

62 **Abstract**

63 The Planted and Regrowth forests (TPRFs) are one most low-cost component for
64 recovering biomass-stored carbon in the tropics. Nevertheless, challenges persist in
65 pinpointing which elevational ranges exhibit the largest carbon accumulation rate
66 (γ_{rapid}), due to the highly inconsistent previous assessments. This prevents the
67 selection of optimal locations for implementing large-scale reforestation in the tropics.
68 Here we proposed a refined approach that used a carbon accumulation threshold (< 80%
69 of the maximum value) to quantify γ_{rapid} in TPRFs at various elevations. We find that
70 γ_{rapid} increases with elevations from 300 m to 1000 m and declines at
71 elevations >1000 m. TPRFs at elevation ~1000 m exhibit three times more γ_{rapid} than
72 lowland TPRFs. This optimal elevation, highly dependent of background temperatures,
73 varies slightly but significantly across different mountains. These findings provide
74 guidelines for policymakers to determine the optimal elevations from regional to
75 continental scales when implementing reforestation initiatives in the tropics.

76

77 **Introduction**

78 Tropical forests account for approximately 40-50% of global forest carbon sink¹,
79 ². Nevertheless, tropical forests are under threat from ongoing deforestation, with high
80 risks of flipping into a net carbon source^{3,4}. Thus, it is essential to regenerate tree cover
81 pan-tropically, either in areas where forests have historically existed (termed
82 ‘reforestation’) or where they have never existed before (termed ‘afforestation’)⁵.
83 Forest cover can either return naturally following land abandonment (natural regrowth
84 forests) or intentionally (planted forests). These have been recognized as one of the
85 most low-cost approaches for recovering biomass-stored carbon in the tropics⁶⁻⁸.

86 Nonetheless, a comprehensive analysis on the optimal locations for implementing
87 afforestation and reforestation is still lacking⁹. One key aspect to be considered when
88 implementing afforestation and reforestation for climate mitigation is the carbon
89 accumulation rate^{10,11}, a factor varying significantly with elevation. Recent studies have
90 attempted to use *in situ* data¹²⁻¹⁶ and high-resolution European Space Agency Climate

91 Change Initiative (ESA-CCI) aboveground biomass (AGB) datasets to quantify the
92 carbon accumulation rates in forests^{10, 17-18}. Among them, some studies highlighted the
93 existence of elevational patterns of biomass carbon in tropical forests, although these
94 patterns vary across studies¹⁹⁻²⁸. For instance, some found that tropical biomass carbon
95 declined¹⁹⁻²² or increased^{23, 24} monotonically with elevation, while others indicated U-
96 shaped²⁵⁻²⁷ or inverted U-shaped²⁸ elevational patterns. Overall, we still lack a large-
97 scale understanding of how the carbon accumulation rate varies with elevation in the
98 tropics^{29, 30}. This is partly due to the fact that trees at different elevations were often at
99 diverse growth stages, with various carbon accumulation rates related to age, i.e., higher
100 in young than in mature forests^{31, 32}. Current studies commonly used a stand age
101 window to determine the temporal position along a Chapman–Richards curve¹⁰ for
102 calculating the carbon accumulation rate⁹. However, the rates of carbon accumulation,
103 as estimated from different stand ages, may lead to contradictory assessments,
104 complicating the determination of their elevational patterns on large scales³³. It is thus
105 necessary to develop approaches that enable comparing forest biomass carbon
106 accumulation rates at the same growth stage across various elevations.

107 Here we conducted a pan-tropical analysis of the growth trajectories in the
108 Tropical Planted and Natural regrowth forests (TPRFs) (**Figure 1a**). We proposed a
109 refined approach that used a carbon accumulation threshold (< 80% of the maximum
110 value), to quantify the biomass carbon accumulation rate (γ_{rapid}) of TPRFs regarding
111 their rapid growth phase prior to reaching maturity³⁴, which is the most important stage
112 during tree’s lifespan for accumulating biomass carbon. Our analysis reveals a robust
113 and consistently increasing trend in γ_{rapid} at elevations from 300 m to 1000 m, but a
114 subsequently declining trend when elevations > 1000 m. This optimal elevation also
115 varies across mountains with different background temperatures. Our findings can help
116 guiding reforestation initiatives at the optimal elevations from regional to continental
117 scales in the tropics.

118

119 **Results**

120 **Overview of experimental procedures**

121 Instead of using fixed stand ages, here we use the time at which a certain biomass
122 carbon threshold is reached, to estimate γ_{rapid} across varying elevations. As such,
123 the rapid growth age is defined as those before the year when the biomass carbon
124 reaches 80% of its peak value predicted by the Chapman–Richards model³⁵,
125 specifically the median biomass carbon of old-growth forests with stand age ≥ 100
126 years within each elevation bin (**Figures S1 and S2**). It coincides with the year at
127 which the growth rate experiences a slowdown breakpoint in the relationship with
128 stand age as stand age increases (see **Experimental Procedure** for details, **Figure**
129 **S3**). Since different datasets have different strengths and weaknesses; *in-situ* data
130 provide high precision but are limited by their spatial and temporal consistency, while
131 satellite-based data typically offer comprehensive coverage but often contain noise in
132 specific areas. On the other hand, model-based data are global in scope but generally
133 have low resolution and lack consistent validation; thus, we integrated four
134 independent forest biomass and carbon datasets to conduct a constraint assessment of
135 elevation-driven variations in γ_{rapid} : (i) **Dataset 1**: *in situ* observations of the total
136 forest biomass (aboveground and belowground), sourced from published literature
137 compilation⁹ and the Smithsonian Institution’s Global Forest Carbon database³⁶
138 (hereafter γ_{rapid}^{GFC}); (ii) **Dataset 2**: the total forest biomass carbon derived from the
139 satellite-based 100 m resolution single-year product of ESA-CCI AGB¹⁷ ($\gamma_{rapid}^{ESA-CCI}$);
140 (iii) **Dataset 3**: a 0.1° resolution time-series dataset of carbon stock in total live
141 woody biomass, generated through machine-learning (ML)³⁷ (γ_{rapid}^{ML}); and (iv)
142 **Dataset 4**: a $\sim 0.07^\circ$ resolution time-series dataset of net ecosystem productivity
143 (NEP) simulations obtained from the BEPS (Boreal Ecosystem Productivity
144 Simulator)³⁸ model (γ_{rapid}^{BEPS}). We also examined the elevational gradient of carbon
145 accumulation rates by comparing the rates with the growth rate of tree height
146 (γ_{rapid}^{GEDI}), which was obtained independently from 30 m resolution spaceborne LiDAR
147 observations in 2019 by the Global Ecosystem Dynamics Investigation (GEDI)³⁹

148 **(Dataset 5)**. Corresponding stand age data and methods used to quantify γ_{rapid} for
149 each carbon dataset were introduced in **Table 1**. Here, we focused exclusively on
150 forests with over 80% of the area covered by newly planted or naturally regrown
151 trees, situated at low-to-mid elevations up to 2000 m above mean sea level (a.m.s.l.)
152 as most afforestation and reforestation are implemented below this elevation⁴⁰.

153

154 *Continental-scale elevational pattern of γ_{rapid}*

155 The average growth rate of TPRFs before 80% of the peak biomass, γ_{rapid} ,
156 estimated from the four biomass carbon datasets (**Table 1, Figure 1a**) all shows no
157 significant elevation-dependence at lowlands (elevation < 300m), whereas there is a
158 strong effect of elevation above 300m (**Figures 1b-1e**). Generally, mountainous γ_{rapid}^{GFC}
159 increases substantially (slope = $0.036 \pm 0.0005 \text{ MgC} \cdot \text{ha}^{-1} \text{yr}^{-1} \text{ m}^{-1}$; $P < 0.05$) from
160 $1.41 \pm 0.26 \text{ MgC} \cdot \text{ha}^{-1} \text{yr}^{-1}$ at an elevation of $290 \pm 50 \text{ m}$ ($P < 0.001$) to $4.65 \pm 0.71 \text{ MgC} \cdot \text{ha}^{-1}$
161 yr^{-1} at $1090 \pm 50 \text{ m}$ ($P < 0.001$) and then declines as elevation exceeds approximately
162 $1090 \pm 50 \text{ m}$ (slope = $-0.0072 \pm 0.0059 \text{ MgC} \cdot \text{ha}^{-1} \text{yr}^{-1} \text{ m}^{-1}$, $P=0.35$). TPRFs at elevations
163 of 1000 m demonstrate the highest γ_{rapid}^{GFC} , accumulating approximately 3.0 times more
164 biomass carbon per year than that of their lowland counterparts (**Figure 1b**). Notably,
165 this “inverted U” elevational trend is consistently observed in $\gamma_{rapid}^{ESA-CCI}$ (**Figure 1c**),
166 in which $\gamma_{rapid}^{ESA-CCI}$ is $2.19 \pm 0.16 \text{ MgC} \cdot \text{ha}^{-1} \text{yr}^{-1}$ at $290 \pm 50 \text{ m}$ ($P < 0.001$), 3.94 ± 0.41
167 $\text{MgC} \cdot \text{ha}^{-1} \text{yr}^{-1}$ at $1090 \pm 50 \text{ m}$ ($P < 0.001$) and falls to $2.19 \pm 0.17 \text{ MgC} \cdot \text{ha}^{-1} \text{yr}^{-1}$ at
168 $1970 \pm 50 \text{ m}$ ($P < 0.001$). Similar patterns are also evident in γ_{rapid}^{ML} (**Figure 1d**) and
169 γ_{rapid}^{BEPS} (**Figure 1e**), although their average magnitudes and corresponding slopes of the
170 increasing elevational trends are much smaller due to the coarse spatial resolutions of
171 the datasets. The elevation dependence is also supported by the growth rate of tree
172 height (γ_{rapid}^{GED1}) calculated from an independent, widely-used spaceborne LiDAR tree
173 height dataset (**Figure 1f**).

174 We then compared the elevational pattern of γ_{rapid} between the natural regrowth

175 and planted forests, between the broadleaved and needle-leaved forests, between the
176 forests located in south- and north facing slopes of mountains, and among different tree
177 genera⁴⁴ (**Figure 2**). While the overall elevational patterns in γ_{rapid} are still valid,
178 γ_{rapid} of planted forests increases more rapidly with elevation than that of natural
179 regrowth forests in areas below 1000 m; conversely, in areas above 1000 m, natural
180 regrowth forests show more pronounced negative elevational gradients in their growth
181 rate than planted forests (**Figures 2a, 2d and 2g**). The elevational patterns of γ_{rapid}
182 also differ between the broadleaved and needle-leaved forests. Generally, the
183 broadleaved forests have a higher value of mean γ_{rapid} compared with needle-leaved
184 forests, while the elevational gradients of γ_{rapid} are more pronounced in needle-
185 leaved forests than in broadleaved forests (**Figures 2b, 2e and 2h**). We also found
186 consistent elevational patterns of γ_{rapid} from 300 m to 1000 m in some specific tree
187 genera (**Figure S4**). However, their elevational patterns above 1000 m were unclear
188 due to the limited availability of biomass carbon data above 1000 m. Additionally, there
189 are no significant differences in the elevational patterns of γ_{rapid} between south- and
190 north facing slopes (**Figures 2g-2i**). This finding aligns with the results observed by
191 Maass et al. (2002)⁴⁵, Mendez-Toribio (2016)⁴⁶, and Madhumali et al. (2023)⁴⁷, which
192 reported insignificant differences in tree height between south- and north- facing slopes
193 in tropical mountains.

194 Notably, the above-mentioned elevational patterns of γ_{rapid} cannot be correctly
195 represented by approaches that necessitate assuming a specific stand age to estimate the
196 carbon accumulation rate, which theoretically varies with the assumed stand age
197 (**Figure 3a**). This is due to the fact that, in the real world, at elevations below 1000 m,
198 the sensitivities of carbon accumulation to elevation are mostly positive and increase
199 with stand age in a range from 1 to approximately 20-25 years (black points, **Figure**
200 **3c**). Conversely, as stands age is beyond 30 years, the sensitivities decline and become
201 negative. In contrast, at elevations above 1000 m, the sensitivities of carbon
202 accumulation to elevation are mostly negative and decrease with stand age rapidly after
203 the age of 20 years (black triangle, **Figure 3c**). This implies that TPRFs at elevations

204 of 1000m exhibit the fastest growth between 20 and 25 years, while TPRFs at elevations
205 < 300m or > 1000m achieve peak growth rates beyond 30 years (**Figure 3c**). In addition,
206 we also found that planted forests achieves their peak growth rates earlier (about 20
207 years) than natural regrowth forests (about 25 years) (**Figure S5**).

208 We then explored various stand age windows (i.e., from zero to 20, 24, 28 and 32
209 years of age) for calculating the biomass average accumulation rate along the
210 Chapman–Richards curve to assess the reliability of stand-age-based approaches
211 (**Figure 3d**), in comparison with using different peak biomass thresholds (i.e., from
212 zero to the time of reaching 70%, 80% and 90% of the maximum value). We found that
213 the biomass carbon accumulation rate exhibits a positive relationship with elevation
214 below 1000 m when selecting stand age thresholds of less than 20 and 25 years (red
215 and yellow fitted lines, **Figure 3d**) and becomes independent of elevation when 28
216 years are used as the threshold (green fitted line, **Figure 3d**). However, shifting into
217 negative trends occurs when opting for a threshold stand age of 32 years (dark blue
218 fitted line, **Figure 3d**). These results suggest that using a fixed stand age threshold to
219 compare the carbon accumulation rate may lead to various elevational patterns that are
220 difficult to be interpreted.

221 In contrast, γ_{rapid} estimated using different stand age at which a biomass
222 threshold is reached (**Figure 3b**), identified by 70%, 80% and 90% of the maximum
223 biomass on the Chapman–Richards curve, show consistent elevational patterns (**Figure**
224 **3f**). We further verified that stand ages (age_{rapid}) with rapid accumulation rate
225 identified using our biomass carbon thresholds are comparable with the observed stand
226 age at which a breakpoint is found in the relationship between the carbon accumulation
227 rate and stand age (**Figure S3**). Results indicate that the estimated age_{rapid} using 80%
228 carbon accumulation window coincide the best with those estimated from observed data
229 (**Figure 3e**). Overall, γ_{rapid} is insensitive to various moving windows lengths and
230 enables a reasonable comparison across elevations, ensuring the robustness of our
231 findings.

232

233 *Climate drivers of elevational gradients in γ_{rapid}*

234 The carbon accumulation rate is determined by two gross carbon fluxes: gross
235 primary production (GPP) and total respiration (TER)^{50, 51}. Over a decadal time span,
236 nearly all augmented carbon goes into stand biomass, while the dynamics of soil carbon
237 are much slower^{52, 53}. Consequently, we used BEPS-simulated NEP (GPP minus TER,
238 **Dataset 4**)³⁸, which exhibits a more comparable magnitude with the carbon
239 accumulation rate than net primary production (NPP) (**Figure S7**), as a proxy to explore
240 the intra-annual variation of their elevation dependence on a monthly time scale.

241 Our analyses indicate that at elevations < 1000 m, GPP and TER both show positive
242 correlations with elevation, but their intra-annual sensitivities differ greatly (**Figure 4a**).
243 Low-land forests experience stronger high-temperature stress (> 24°C) from April to
244 September due to the negative elevational gradients in temperature (**Figure 4c**),
245 resulting in a sharper decrease in GPP than in TER (**Figure 4g**). This discrepancy causes
246 greater negative sensitivities of GPP to elevation compared with TER (**Figure 4i**),
247 thereby leading to positive elevational gradients in NEP from April to September
248 (**Figure 4a**). Furthermore, the decrease in VPD with elevation (**Figure S8a**), emerges
249 as the second crucial factor in co-driving the increasing trend in GPP with elevation.
250 Large VPD serves as a valuable proxy for atmospheric dryness and often imposes
251 significant constraints on GPP in tropical forests⁵⁴⁻⁵⁸, thus highlighting the impact of
252 elevation on NEP during periods of high-temperature stress (**Figure 4i**). During the
253 high-precipitation months from April to September (**Figure 4e**), elevation-associated
254 variations in precipitation exert a nearly equivalent inhibitory effect on both GPP and
255 TER (**Figures 4h and 4i**). Consequently, the asynchronous elevational sensitivities of
256 GPP and TER to temperature primarily contribute to the positive response of NEP
257 (orange curve) to elevation (< 1000 m).

258 At elevations above 1000 m, forests grow in moderate temperature conditions
259 (average monthly MAT < 24°C) (**Figure 4d**). In this condition, NEP shows insignificant
260 elevational gradients from April to November (**Figure 4b**), due to the consistent
261 elevational sensitivities of GPP and TER to temperature (< 24°C) (**Figure 4g**). During

262 the low-precipitation months from December to March, elevation-associated variations
263 in precipitation (**Figure 4f**) exerts divergent impacts on GPP and TER (**Figure 4h**).
264 This discrepancy results in a negative elevational pattern of GPP and a positive
265 elevational pattern of TER (**Figures 4b and 4j**), thereby leading to negative elevational
266 gradients in NEP from December to March (**Figure 4b**). The asynchronous elevational
267 sensitivities of GPP and TER to precipitation primarily contribute to the negative
268 response of NEP at elevations > 1000 m.

269 Overall, the pronounced elevational patterns in MAT, VPD and precipitation
270 primarily drive the seasonally-dependent but divergent sensitivities of GPP and TER to
271 elevation. These impacts contribute to the increasing elevational trend of γ_{rapid}^{BEPS} at
272 elevations < 1000m and the decreasing elevational trend of γ_{rapid}^{BEPS} at elevations > 1000
273 m. In contrast, soil moisture (SM) (**Figures S8c-S8d**) and total photosynthetically
274 active radiation (PAR) (**Figures S8e-S8f**) exhibit lower sensitivity to elevation and play
275 less important roles (**Figures 4i-4j**).

276

277 *Variations at individual mountains*

278 We further examined the elevation patterns of γ_{rapid}^{BEPS} and corresponding driving
279 mechanisms across four mountainous areas: the Sierra Madre del Sur mountain in North
280 America (SMS), the Ethiopian Highlands mountain in Africa (EH), the Serra do
281 Espinhaço mountain in South America (SE), and the Eastern Ghats mountain in Asia
282 (EG) (**Figure 5a**). γ_{rapid}^{BEPS} in individual mountains shows an increase with elevation in
283 low-to-mid elevation, followed by a decline as elevation continues to increase (**Figures**
284 **5b-5e**), consistent with the results for entire tropical mountain regions (**Figure 1**).
285 Mechanism analyses also support that the positive elevational gradient of γ_{rapid} is
286 mainly attributed to the asymmetrical response of GPP and TER to temperatures
287 (**Figures S9 and 5i-5l**); while the negative elevational gradient of γ_{rapid} is due to the
288 divergent response of GPP and TER to precipitation (**Figures S9 and 5m-5p**).

289 However, the optimal elevations where show the γ_{rapid}^{BEPS} peak varies across

290 different mountains. Among the four studied mountains, the Eastern Ghats mountain
291 experiences the highest mean temperature (average monthly TMF = 26.61°C) (**Figure**
292 **5f**) and thus has a highest optimal elevation (1390 m), corresponding to the places with
293 greatest γ_{rapid}^{BEPS} (**Figure 5e**). This results in the smallest elevational sensitivity in
294 γ_{rapid}^{BEPS} (**Figure 5h**). Conversely, the Serra do Espinhaço mountain has the lowest mean
295 temperature (average monthly TMF = 23.67°C) (**Figure 5f**) and shows the lowest
296 optimal elevation (900 m) (**Figure 5d**). This leads to the sharpest elevational trend in
297 γ_{rapid}^{BEPS} (**Figure 5h**). Overall, the optimal elevations with highest γ_{rapid}^{BEPS} in the four
298 mountains are ranked as follows: Eastern Ghats (1390 m) > Ethiopian Highlands (1200
299 m) > Sierra Madre del Sur mountain (1000 m) > Serra do Espinhaço (900 m) (**Figure**
300 **5g**); and in the opposite, the elevational sensitivities of γ_{rapid}^{BEPS} are ranked as: Serra
301 do Espinhaço > Sierra Madre del Sur mountain > Ethiopian Highlands > Eastern Ghats
302 (**Figure 5h**).

303

304 **Discussion**

305 Afforestation and reforestation stand out as pivotal land-based actions for
306 mitigating climate change, especially in the context of diminishing net gains from CO₂
307 fertilization and increasingly negative impacts on tree growth from climate warming⁵⁹.
308 Tropical mountain forests exhibit a large potential for carbon accumulation^{29, 60}.
309 However, the expansion of a new agricultural frontier has caused significant tree cover
310 loss in tropical mountain forests during the 21st century^{61, 62}. Thus, tropical mountains
311 have emerged as a region with significant potential for implementing reforestation and
312 afforestation efforts for future climate mitigation⁴⁰.

313 Challenges persist in pinpointing optimal elevations for afforestation and
314 reforestation in the tropics⁵⁹. Previous field studies generally presented inconsistent
315 elevational patterns of carbon accumulations rate^{19, 20, 22-26, 30, 63}. Studies, mainly
316 focusing on the net carbon dynamics of mature forests, observed a monotonically
317 decreasing trend of biomass carbon with elevation at a large elevational range (0-5000

318 m)^{22, 25, 30}. Other studies that accounted for both young and mature forests found that
319 biomass carbon accumulation rate show an inverted U-shaped²⁸ curve along increasing
320 elevation with a transition at approximately 1600 m^{20, 63}. Stand age differences across
321 elevations likely introduce considerable uncertainties when comparing carbon
322 accumulation rates across various elevations.

323 Notably, such stand-age-associated discrepancy across elevations raises an
324 important issue — how to choose an appropriate time period along a Chapman–
325 Richards curve for quantifying the carbon accumulation rate at different elevations⁹.
326 For instance, young secondary forests (< 40 years old) can accumulate 11 to 20 times
327 more biomass carbon than the mature forests^{31, 32}. Studies using the same stand age
328 window for different elevations to estimate the carbon accumulation rate⁹ may bring
329 uncertainties when comparing the carbon accumulation rate at various elevations¹⁰, as
330 the stand age varies greatly at different elevations in the real world⁶⁴. This provides
331 explanations for previous studies that have observed diverse elevational patterns of the
332 carbon accumulation rate^{9, 25, 30}. Thus, previous studies mostly did not accounted for
333 the uncertainties introduced by different stand age and likely compared the carbon
334 accumulation rate of forests at different growth stages⁶³. This can be reflected by the
335 analysis regarding the relationship between biomass carbon accumulation rate and
336 elevations, using 24, 28, and 32 years of age as thresholds, which showed positive,
337 insignificant, and ultimately negative trends, respectively (**Figure 3d**).

338 To reduce such uncertainties from stand age, we proposed a refined approach that
339 used a carbon accumulation threshold (< 80% of the maximum value) rather than the
340 stand age window to define the analysis time period. This novel approach enables
341 comparing the carbon accumulation rates during the rapid growth periods in both low-
342 and mid-elevation TPRFs (**Figure 3b**), revealing a robust, consistent and positive
343 elevational gradient in γ_{rapid} within elevation below 1000 m and conversely a
344 negative elevational gradient in γ_{rapid} within elevation above 1000 m. (**Figures 1 and**
345 **3f**). This discovery helps to reconcile the diverging elevational trends that were
346 observed in previous studies^{60, 63, 65} and provides a benchmark for comparing the

347 biomass carbon accumulation rate (γ_{rapid}) of forests across elevations.

348 We further found that seasonal variations in temperature and atmospheric dryness
349 play key roles in the positive elevational gradient of γ_{rapid} at elevations < 1000 m.
350 This is probably attributed to the decreases of high-temperature stress⁶⁶ and
351 atmospheric dryness constraint⁵⁴ with elevation (**Figures 4c and S8a**). This finding is
352 consistent with previous analyses, which found the greatest threat of high-temperature
353 to the biomass carbon accumulation rate in lowland TPRFs^{67, 68}. In contrast, TPRFs
354 above 1000m often live in less hot temperatures⁶⁹ and thus γ_{rapid} exhibits less impact
355 from high-temperature stress. Conversely, precipitation becomes the most important
356 climatic limiter, as elevations above 1000 m generally have small rainfall²⁵. The
357 constraint on GPP from low precipitation, which also causes an increase in TER, is
358 typically severe from December to March, resulting in a negative elevational pattern of
359 γ_{rapid} at elevations above 1000m. Therefore, elevations of around 1000 m likely
360 encounter less high temperature and water stress (**Figures 4c-4f**), making these regions
361 optimal for potential reforestation efforts aimed at climate mitigation.

362 Notably, the elevational pattern of γ_{rapid} may be further exacerbated by ongoing
363 climate warming^{70, 71}, as future extreme high temperatures and atmospheric drying
364 could impose more substantial limitations on carbon accumulation rates in the
365 lowlands^{72, 73}. With the declining strength of carbon sinks in lowland tropical forests⁶⁸,
366 the importance of montane systems for carbon management may be increasing in the
367 future⁷⁴. Climate change may also bring various impacts on the elevational gradients in
368 carbon accumulation between the planted and natural growth forests, and between the
369 broadleaved and needle-leaved forests, as well as in different mountains, due to their
370 diverse climatic sensitivities^{13, 14, 18}. Worthy of noting is that disturbances from human
371 activities^{10, 18} also decrease with elevation (**Figure S10b**) and may also contribute to
372 the increasing elevational trend of γ_{rapid} in TPRFs (**Figures S10c and S10d**).

373 In summary, using multiple data streams, as well as modelling and mapping
374 approaches, our analysis reveals a robust and consistently increasing trend in γ_{rapid} at
375 elevations between 300 m and 1000 m, but a subsequently declining trend when

376 elevations > 1000 m. Thus, 1000 m shows to be the optimal elevation for accumulating
377 biomass carbon in TPRFs, while this threshold varies slightly across different
378 mountains. Our findings underscores the importance of incorporating elevation as a
379 global factor when estimating biomass carbon sinks, and when considering suitable
380 areas for reforestation and afforestation, addressing both scientific understanding and
381 policy considerations.

382

383 **EXPERIMENTAL PROCEDURES**

384 **Methods**

385 *General summary*

386 In order to conduct a constraint assessment of elevation-driven variations in the
387 biomass carbon accumulation rate in the rapid growth stages (γ_{rapid}), we firstly
388 collected five independent forest biomass, carbon flux, and height datasets. (i) **Dataset**
389 **1**: *in situ* observations of total forest biomass (aboveground and belowground) from
390 published literature compilation⁹ and the Smithsonian Institution's Global Forest
391 Carbon database (ForC)³⁶; (ii) **Dataset 2**: total forest aboveground biomass carbon
392 derived from the satellite-based 100 m resolution single-year product of European
393 Space Agency Climate Change Initiative (ESA CCI) aboveground biomass (AGB)¹⁷;
394 (iii) **Dataset 3**: a 0.1° resolution time-series dataset of carbon stock in total live woody
395 biomass, generated through machine-learning (ML)³⁷; (iv) **Dataset 4**: a 0.072727°
396 resolution time-series of net ecosystem productivity (NEP) simulations obtained from
397 the BEPS (Biosphere-atmosphere Exchange Process Simulator)³⁸ model; and (v)
398 **Dataset 5**: a 30 m resolution tree height dataset from spaceborne LiDAR observations
399 in 2019 by the Global Ecosystem Dynamics Investigation (GEDI)³⁹.

400 While **Dataset 1** includes information on stand ages, **Datasets 2- 5** do not provide
401 this information. Consequently, we needed to supplement **Datasets 2- 5** with stand age
402 data from other sources before proceeding with the subsequent analysis. For ESA-CCI
403 AGB data (**Datasets 2**) and GEDI tree height data (**Datasets 5**), we utilized the 30 m
404 resolution satellite-based tropical moist forest cover change dataset (TMF)⁴¹ to identify

405 natural regrowth (TRF) pixels where a conversion from deforested land to forest
406 occurred, termed “Secondary Forest” in the TMF dataset (see more detail later in
407 methods). The consecutive years during which the land remained forest-covered were
408 then used to estimate the stand age in years¹⁸ for these TRF pixels. We also employed
409 a satellite-based global plantation years dataset⁴² with a resolution of 30 m to identify
410 planted forest (TPF) pixels and their corresponding planting years. For the 0.1°
411 resolution time-series machine-learning live woody biomass data provided by Xu et al.
412 (2021)³⁷ (**Dataset 3**) and the 0.072727° resolution time-series BEPS-modeled NEP data
413 (**Dataset 4**), we identified TPRF pixels where non-forests converted to forested lands
414 based on the 0.05° resolution MODIS MCD12C1 landcover products⁴³. And then, we
415 supplemented the elevation data for these five independent datasets with Shuttle Radar
416 Topography Mission (SRTM) DEM data⁷⁵ extracted from global digital elevation model
417 (DEM) maps.

418 After supplementing the information on stand age and elevation, we analyzed the
419 γ_{rapid} of TPRFs across the elevation gradient based on these five biomass-related
420 datasets. It is essential to highlight that we applied a space-for-time analogy with the
421 Chapman–Richards curve¹⁰ (**Equation 1**) to estimate γ_{rapid}^{GFC} , $\gamma_{rapid}^{ESA-CCI}$ and γ_{rapid}^{GEDI} at
422 an elevation bin of 100 m and a moving window step of 80 m. Previous studies
423 commonly used a stand age window to determine the temporal position along a
424 Chapman–Richards curve for calculating the carbon accumulation rate⁹. Nevertheless,
425 the rates of carbon accumulation, as estimated from different stand age windows, vary
426 largely (**Figure 3a**). Here, we use a carbon accumulation window approach to estimate
427 the biomass carbon accumulation rate of TPRFs during their rapid growth phase (γ_{rapid}),
428 across varying elevations (**Figure 3b**). The rapid growth stand age is defined as the
429 time before the maturity year when the biomass carbon reaches 80% of its peak value
430 predicted by the Chapman–Richards model, specifically the median biomass carbon of
431 old-growth forests with stand age ≥ 100 years within each elevation bin. The grid cells
432 corresponding to old-growth forests (≥ 100 years of age) were identified based on stand
433 age information sourced from a global database of forest carbon provided by Anderson-

434 Teixeira et al³⁶. Subsequently, the γ_{rapid} is calculated as the slope of the linear
435 regression fit between live biomass carbon and forest stand age of TPRFs before
436 reaching the defined maturity age (inset in **Figures 1b-1c, 1f; and S1-S2**). It implies
437 that this approach enables comparisons of the biomass carbon accumulation rate in
438 TPRFs during their rapid growth stage, reducing uncertainties arising from differences
439 in stand age across various elevations.

$$440 \quad Y_t = A(1 - e^{-kt})^c \pm \varepsilon; A, k \text{ and } c > 0 \quad (1)$$

441 Where Y_t refers to the biomass carbon at age t ; A is the median biomass carbon of
442 old-growth forests with stand age ≥ 100 years within each elevation bin; k is a growth
443 rate coefficient of Y_t as a function of age; c is a coefficient that determines the shape
444 of the growth curve; and ε is an error term.

445 While the γ_{rapid}^{GFC} , $\gamma_{rapid}^{ESA-CCI}$ and γ_{rapid}^{GEDI} are estimated using the space-for-time
446 method, the γ_{rapid}^{ML} and γ_{rapid}^{BEPS} are obtained using time series methods. Specifically,
447 γ_{rapid}^{ML} is estimated as the slope of the fitted linear regression fit between annual live
448 biomass carbon and stand age over the entire time series period for each TPRF (inset
449 in **Figure 1d**), since their stand ages typically span less than 20 years. On the other
450 hand, γ_{rapid}^{BEPS} is calculated as the mean NEP for each TPRF over the same period
451 (**Figure 1e**).

452 In addition to examining the entire tropical mountain zones, we analyzed the
453 changes in γ_{rapid} with elevation across four specific mountains: the Sierra Madre del
454 Sur in North America (SMS), the Ethiopian Highlands in Africa (EH), the Serra do
455 Espinhaço in South America (SE), and the Eastern Ghats in Asia (EG), using time-
456 series BEPS-modeled net ecosystem productivity (NEP) data (**Figures 5b-e**).
457 Furthermore, we conducted a comprehensive assessment to explore the driving
458 mechanisms behind the elevational gradients in γ_{rapid} , both across the entire tropical
459 regions (**Figure 4**) and among the individual mountains (**Figure 5**).

460 The overall technical roadmap is shown in **Figure S11** and detailed estimation
461 steps for each dataset are described in the subsequent sections.

462

463 *Estimation of the biomass carbon accumulation rate based on in situ forest biomass*
464 *data (Dataset 1) using the space-for-time analogy*

465 We used a global forest biomass carbon dataset that provided field measurements
466 of plant biomass carbon (including both above- and below-ground biomass carbon) (in
467 MgC ha⁻¹), stand age (in years), elevation (in meters), pre-disturbed land cover
468 information and plant establishment method (e.g. natural regrowth or planted)^{9, 36}. In
469 cases where records lacked elevation information, we supplemented the elevation data
470 from Shuttle Radar Topography Mission (SRTM) DEM data⁷⁵. Finally, the dataset
471 comprised 2,693 aboveground carbon estimates within tropical regions, originating
472 from 518 distinct sites and encompassing data from 164 studies (**Figure 1a**).

473 Due to the limited availability of time-series observations for each site (mostly
474 spanning over one or two years)⁹, we employed a space-for-time analogy to estimate
475 the carbon accumulation rate. In contrast to time-series analysis, which quantifies
476 changes in plant carbon over time at each site, the space-for-time analogy estimates an
477 average carbon accumulation rate across all sites within a specific elevation range.
478 **Figure 1b** was produced at an elevation bin of 100 m and a moving window step of 80
479 m. We conducted additional analyses with various combinations of elevation bins and
480 moving window steps to assess the robustness of the results (**Figure S12**).

481 In addition, using the plant establishment and pre-disturbed land cover information
482 recorded in this database, along with 500 m resolution MODIS MCD12Q1 v061 land
483 cover data⁴³ and 90 m resolution DEM data⁷⁵, we further conducted a comparative
484 analysis of the elevational patterns of γ_{rapid}^{GFC} between natural regrowth and planted
485 forests (**Figure 2a**), between broadleaved and needle-leaved forests (**Figure 2b**),
486 between south- and north-facing slopes of mountains (**Figure 2c**), and different land
487 use types before afforestation, reforestation or both (**Figure S13**).

488

489 *Estimation of the biomass carbon accumulation rate based on the 100 m resolution*
490 *total biomass carbon derived from the ESA-CCI aboveground biomass (AGB)*

491 ***product (Dataset 2) using the space-for-time analogy***

492 We first obtained the aboveground biomass (AGB) records for 2018 from the 100
493 m resolution ESA-CCI product¹⁷, which used elevation data to reduce errors arising
494 from the differences in radar backscatter between slopes facing the radar and those
495 facing away. We then used the root-to-shoot ratio developed from field measurements
496 to estimate the below-ground biomass (BGB) from the ESA-CCI AGB data^{37, 76}.
497 Subsequently, biomass carbon was estimated from the sum of AGB and BGB by
498 multiplying by a constant coefficient (0.49)^{76, 77}. It is important to acknowledge
499 potential uncertainties in these estimates, considering that AGB and BGB may exhibit
500 distinct elevational patterns globally^{75, 78-81}.

501 To estimate the $\gamma_{rapid}^{ESA-CCI}$ for natural regrowth forests (TRFs), we utilized the 30
502 m resolution satellite-based tropical moist forest cover change dataset (TMF)⁴¹ to
503 identify TRF pixels. This dataset tracked the extent and alterations of tropical moist
504 forests over the past three decades⁴¹ and characterized the degraded forests and
505 secondary forests at a spatial resolution of 30 m and a yearly temporal resolution,
506 generated from Landsat data spanning from 1982 to 2019. Degraded forests, in this
507 dataset, were defined as tree-covered pixels for which disturbances were visible for a
508 short time period (between 3 months and 2.5 years maximum), whereas secondary
509 forests were defined as pixels with natural regrowth vegetation after an absence of tree
510 cover for more than 2.5 years¹⁸. In this study, we only selected the secondary forests
511 for TRF analysis. Furthermore, we also removed the oil palm plantations from the TMF
512 secondary forests following the methodology of Heinrich et al.¹⁸. Stand age for TRFs
513 was estimated as duration since the most recent disturbance event for any recovering
514 forest pixel in tropical moist forests, based on the annual number of detected
515 disturbances in the TMF dataset. Subsequently, we superimposed the 30m resolution
516 stand age map onto the 100 m resolution total biomass carbon map derived from ESA-
517 CCI AGB production, calculating the average stand age for each ESA-CCI grid cell.
518 Only those 100 m resolution ESA-CCI grid cells, where stand age information was
519 presented in more than 10 sub-grid pixels with 30 m resolution, were included in our

520 analysis. Finally, we employed a space-for-time analogy with the Chapman–Richards
521 curve to estimate $\gamma_{rapid}^{ESA-CCI}$ for natural regrowth forests at an elevation bin of 100 m
522 and a moving window step of 80 m (**Figure 2d**).

523 Similarly, to estimate the $\gamma_{rapid}^{ESA-CCI}$ for planted forests (TPFs), we first utilized a
524 30 m resolution global map of plantation planting years (GPY) spanning from 1982 to
525 2020 to identify TPF pixels. The GPY dataset includes two tree categories: tree crops
526 and planted forests. We retained only the 100 m resolution ESA-CCI grid cells that
527 were overlapped with more than 10 sub-grid pixels identified as “planted forest” at 30
528 m resolution. Subsequently, we employed a space-for-time analogy with the Chapman–
529 Richards curve to estimate $\gamma_{rapid}^{ESA-CCI}$ for TPFs at an elevation bin of 100 m and a
530 moving window step of 80 m (**Figure 2d**).

531 It is worth noting that the $\gamma_{rapid}^{ESA-CCI}$ in **Figure 1c** is the result of using both TPFs
532 and TRFs data. For TPFs, we used the species information recorded in the GPY dataset
533 to classify them into broadleaved and needle-leaved forests. For TRFs, we classified
534 them into broadleaved and needle-leaved forests based on the 500 m resolution MODIS
535 MCD12Q1 v061 land cover data⁴³. Subsequently, we analyzed the elevational patterns
536 of $\gamma_{rapid}^{ESA-CCI}$ for broadleaved and needle-leaved forests, respectively (**Figure 2e**), and
537 between south- and north-facing slopes of mountains (**Figure 2f**).

538 Furthermore, we conducted an extensive analysis at the tree genus level by
539 integrating a comprehensive *in situ* mega-database of tropical African vascular plant
540 distributions compiled from 13 datasets⁴⁴. In this analysis, we selectively considered
541 tree species that met specific criteria: they had to have at least four sites in the 100m
542 bin within a single elevation bin (100 m) and spanned at least three elevation bins
543 between elevations of 300 to 1000 m, as there is no sufficient data for analysis above
544 1000 m. Moreover, we excluded data points exhibiting anomalously high biomass
545 carbon values ($> 200 \text{ MgC}\cdot\text{ha}^{-1}$) for trees aged 1 to 3 years or demonstrating an
546 exceptionally high biomass carbon accumulation rate ($> 15 \text{ MgC}\cdot\text{ha}^{-1}\text{yr}^{-1}$). Adhering to
547 these stringent selection criteria, our analysis finally included 4 species with a total of

548 215 records (**Figure S4**).

549

550 *Estimation of the biomass carbon accumulation rate based on a time-series 0.1°*
551 *resolution live biomass dataset from 2000 to 2019 (Dataset 3)*

552 A global vegetation live biomass dataset was generated by integrating ground
553 inventory data and remote sensing observations, including airborne laser scanning
554 (ALS) data and the satellite LiDAR inventory of global vegetation height structure
555 information. This dataset was developed using a self-improving ML algorithm³⁷. It
556 provides global time-series (2000–2019) and annual live biomass data at 0.1° resolution,
557 covering terrestrial ecosystems worldwide. Extensive validation efforts have been
558 conducted based on over 100,000 *in situ* observations³⁷. This dataset used elevation
559 data by incorporating SRTM data into machine learning techniques to capture
560 topographical features, thereby enhancing the accuracy of carbon stock estimates across
561 different elevations³⁷. Importantly, this dataset has found widespread application in
562 studies pertaining to forest carbon dynamics⁸²⁻⁸⁵.

563 In this study, the biomass carbon accumulation rate was estimated using time-series
564 methods, as delineated in the following procedural steps.

565 First, we identified the TPRFs within the latitudinal range of 23.5°S–23.5°N based
566 on MODIS MCD12C1 landcover products (version v6.1)⁴³
567 (<https://lpdaac.usgs.gov/products/mcd12c1v006/>), which are available from 2001 to
568 2019 and exhibit comparable spatial resolutions (0.05°) with the 0.1° resolution ML-
569 derived live biomass data. Stand ages were then determined using the following
570 approach⁸⁶⁻⁸⁸: (1) Firstly, tropical regions were classified into two categories: forest and
571 non-forest lands. Forest pixels include five land cover types: evergreen needle-leaved
572 forests, evergreen broadleaved forests, deciduous needle-leaved forests, deciduous
573 broadleaved forests, and mixed forests; (2) Then, the commencement of growth periods
574 was designated as the year when a transition occurred from non-forest to forest;
575 Conversely, termination of growth periods was marked when a transition occurred from
576 forest to non-forest; and (3) the stand age was calculated as the temporal length between

577 the commencement and termination of growth periods. Given the accessibility of
578 MODIS landcover maps from 2001 to 2019, the estimated stand ages in our study
579 encompass only immature stages, ranging from 1 year to 18 years. It is pertinent to note
580 that, in reality, forests often commence regrowth before being visually identified in
581 remote sensing imagery, potentially resulting in an underestimation of TPRF stand ages.

582 Second, to align the stand age map with the coarse spatial resolution (0.1°) of
583 gridded forest biomass carbon data, we employed a 2×2 search window (size: 2×2 0.05°
584 pixels) to track land cover changes. Only pixels meeting the specific criteria were
585 included in the analysis: (1) all four pixels within the 2×2 search window were
586 classified as non-forest lands in the year 2001; (2) more than half of the 0.05° pixels
587 within the 2×2 search window transitioned from non-forest to forest lands during the
588 same period; and (3) the duration of the growth period should exceed 8 years, ensuring
589 the reliability of carbon accumulation rate calculations. Applying these criteria, a total
590 of 1,754 $0.1^\circ \times 0.1^\circ$ grids were selected for analysis.

591 Finally, for each selected 0.1° grid cell, scatter diagrams were plotted to illustrate
592 the relationship between stand age and annual live biomass carbon. The OLS linear
593 regression model was employed to establish a linear regression fit, with its slope
594 designated as the biomass carbon accumulation rate (inset in **Figure 1d**). To mitigate
595 potential uncertainties arising from data anomalies, pixels exhibiting unreasonably high
596 carbon accumulation rates (i.e., a linear regression slope $> 15 \text{ MgC} \cdot \text{ha}^{-1} \cdot \text{yr}^{-1}$) were
597 excluded following the screening approach used in recent studies⁹. Ultimately, a total
598 of 1,512 TPRF grid cells were included in the final analysis (**Figure 1a**).

599

600 *Exploring the elevation pattern of NEP based on model-simulated time-series carbon* 601 *flux datasets spanning from 2000 to 2019 (Dataset 4)*

602 It was observed that, on a decadal time span, nearly all augmented carbon goes
603 into stand biomass during the initial period following afforestation, while the dynamics
604 of soil carbon are much slower^{52, 53}. NEP can therefore be used as a proxy for the
605 biomass accumulation rate. Here, we used the time-series NEP dataset simulated by the

606 Boreal Ecosystem Productivity Simulator (BEPS) model³⁸, providing daily carbon
607 fluxes at a resolution of 0.072727° spanning from 1981 to 2019. We resampled the
608 time-series BEPS NEP datasets into 0.1° using the nearest-neighbor method and
609 overlapped them with the 0.1° TPRF grid cells identified for 0.1° resolution ML -
610 derived live biomass data. We then calculated the mean NEP during the growth period
611 for each TPRF grid cell derived from MODIS MCD12C1 landcover products (**Figure**
612 **1e**).

613 Furthermore, we estimated the proportion of TRFs pixels (30 m resolution)
614 identified using TMF data; and estimated the proportion of TPFs pixels (30 m
615 resolution) identified based on GPT data within each 0.1° TPRF grid cell. If the
616 proportion of TRF pixels was greater than that of TPF pixels, we classified the 0.1°
617 TPRF grid cell as a natural regrowth forest; otherwise, it was classified as a planted
618 forest. Similarly, we classified the 0.1° TPRF grid cells into broadleaved and needle-
619 leaved forests, as well as north-facing and south-facing forests, based on the land cover
620 data from MCD12C1 v061⁴³ and high resolution DEM data⁷⁵. Subsequently, we
621 compared the elevational patterns of γ_{rapid} in various areas and different forest types
622 (**Figures 2g-2i**).

623

624 ***Estimation of the tree height growth rate based on GEDI tree canopy height data***

625 Tree height data were extracted from a global map of forest canopy height with 30
626 m resolution for the year 2019³⁹. This map was produced by integrating LiDAR-derived
627 canopy height metrics, specifically from the GEDI level 2 product, with Landsat multi-
628 temporal surface reflectance data. This dataset underwent calibration against elevation
629 during its generation, utilizing the Shuttle Radar Topography Mission (SRTM)⁷⁵
630 elevation data in its regression model. This calibration improves the accuracy of tree
631 height measurements, particularly in hilly or mountainous regions³⁹.

632 To further mitigate uncertainties in the LiDAR tree height data, a comparative
633 analysis was conducted with another widely used dataset on tree height⁸⁹. The selection
634 process involved retaining only those pixels where the two tree height datasets

635 exhibited a high level of consistency, with tree height difference $< \pm 5$ m (**Figure S14**).

636 Subsequently, we employed a space-for-time analogy to calculate the tree height
637 growth rate (**Figure 1f**) within a specific elevation interval of 100 m using a moving
638 window step of 80 m. This approach closely mirrors that employed for calculating the
639 carbon accumulation rate from the ESA-CCI-derived biomass carbon data.

640

641 *Exploring the impact of the differences in elevation patterns of monthly GPP and*
642 *TER on the elevation patterns of NEP.*

643 Given that the increasing elevational patterns of carbon accumulation rates were
644 predominantly observed between March and September (**Figure 4a**), our investigation
645 thus focused on identifying potential environmental drivers at the monthly time scale.
646 Currently, there is a lack of available data pertaining to monthly plant carbon
647 accumulation rates in TPRFs. Our analyses uncovered a strong correlation between the
648 net ecosystem productivity (NEP) and the rate of plant carbon accumulation (**Figure**
649 **S7a**, $P < 0.001$, slope=1.14, $R^2=0.5$), closely aligning with the 1:1 diagonal line. We
650 therefore used NEP as a proxy of carbon accumulation rate to explore its potential
651 environmental drivers in TPRFs. NEP, in this context, is defined as the difference
652 between the amount of organic carbon fixed by photosynthesis in an ecosystem (gross
653 primary production, GPP) and total ecosystem respiration (the sum of autotrophic and
654 heterotrophic respiration, TER).

655 Worthy of note is that, although NPP is strongly linearly correlated with the carbon
656 accumulation rate, it exhibits a much larger magnitude compared with the carbon
657 accumulation rate (**Figure S7b**, $P < 0.001$, slope = 0.21, $R^2 = 0.4$). This is because
658 biomass carbon accumulation is one of the four components of NPP, while the other
659 three components, foliage turnover, fine root turnover, and mortality, can occupy 50%
660 to 80% of NPP⁹⁰. For young forests, mortality is usually very low. In tropical forests,
661 the foliage and fine root turnovers to the soil would decompose within a few years,
662 resulting in little change in the soil organic matter^{52, 53}. Therefore, NEP performs as a
663 better proxy for the biomass carbon accumulation rate (only stemwood and coarse root

664 biomass accumulates with time) compared with NPP.

665 For our investigation, we utilized a global carbon flux product created by Chen et
666 al.³⁸ using the BEPS model. It provides the global daily GPP, TER and NEP at a spatial
667 resolution of 0.07272727° (~10 km) from 1981 to 2019. The BEPS model is a process
668 based diagnostic model driven by remotely sensed vegetation parameters, including
669 biophysical variables such as Leaf Area Index (LAI), climate data (temperature and
670 precipitation), nitrogen deposition, and atmospheric CO₂ concentrations. It initializes
671 carbon pools based on historical net primary production (NPP) data from 1901. The
672 model simulates carbon dynamics by stratifying biomass carbon into four pools (leaf,
673 stem, coarse root, and fine root) and soil carbon into nine pools. Key processes include
674 heterotrophic and autotrophic respiration, as well as net ecosystem production (NEP),
675 which is derived from gross primary production (GPP) minus total respiration. In
676 comparison with 15 prognostic models used by Global Carbon Project (GCP), BEPS is
677 among the best in terms of Pearson's coefficient (R^2) and root mean square error
678 (RMSE) between simulated and the observation-based annual global residual land sink
679 (RLS)³⁸.

680 Although the BEPS model has a resolution of approximately 0.07272727°, it can
681 still effectively capture differences along elevation gradients. As shown in **Figure 15**,
682 in the Eje Volcanico Transversal Mountain range in North America, along the latitude
683 of 18.5°N and longitude ranging from 104°W to 96°W, there are 83 grid cells at a 10
684 km x 10 km resolution distributed below an elevation of 2000 m. This implies that the
685 10 km resolution data has sufficient capability to capture the elevation gradients below
686 2000 m and corresponding variations of ecological factors associated with these
687 elevations.

688 Although many studies have demonstrated the satisfactory performance of the
689 BEPS model in simulating the global carbon sink⁹¹, some uncertainties may still
690 remain⁹². For instance, BEPS model used MODIS LAI as an essential input data to
691 simulate NEP, while MODIS LAI is found to be less accurate in mountainous areas
692 compared to flatlands⁹³. This may bring uncertainties in assessing the elevational

693 pattern of NEP⁹⁴. Additionally, BEPS model mainly relies on the maximum
694 carboxylation rate (V_{cmax})⁹⁵ to simulate GPP, while other photosynthesis-related
695 parameters that may vary with elevation are mostly overlooked⁹⁶. Although results are
696 verified by multiple biomass- and height- related datasets, assessments are needed in
697 the future based on accurately simulating the GPP, TER, and NEP.

698

699 ***Investigation of climatic drivers influencing the elevational patterns in GPP and TER***
700 ***using a multiple linear regression model***

701 Prior research has suggested that precipitation, air temperature, sunlight, vapor
702 atmospheric dryness, and soil moisture are potential key factors in influencing the rate
703 of carbon accumulation in tropical forests. In this study, we delved into the impacts of
704 five key drivers, i.e. the mean air temperature (MAT), the vapor pressure deficit (VPD),
705 precipitation (PRE), soil moisture (SM), and total photosynthetically active radiation
706 (PAR) on the rate of carbon accumulation in restored forests (**Figures 4 and S8**). For
707 our analyses, we used the time-series mean air temperature, VPD and precipitation from
708 the TerraClimate global gridded meteorological and water balance variables dataset⁹⁷.
709 SM data was from the RSSSM global surface soil moisture dataset⁹⁸. Total PAR data
710 were from 0.05° resolution MODIS-derived global land products of total
711 photosynthetically active radiation from 2000 to 2019⁹⁹. Given the distinct spatial
712 resolutions of these products (**Table S1**), all variables were resampled to a 0.1° spatial
713 resolution and monthly for the analyses. It is worth noting that air temperature, VPD,
714 and soil moisture are represented as daily means on a monthly scale, while radiation
715 and precipitation are presented as daily totals on a monthly scale.

716 Due to the significant differences in the elevation gradients of GPP and TER
717 between months experiencing high-temperature stress and those without high-
718 temperature stress, we then separately quantified the contributions of the elevation
719 gradient of climatic drivers, specifically MAT, VPD (only for GPP), PRE, SM, and total
720 PAR (only for GPP), to the elevational gradient of GPP or TER for the months under
721 high-temperature stress and those without such stress, respectively. This quantification

722 was achieved by decomposing the elevation gradient of GPP ($\frac{dGPP}{delevation}$) and TER
 723 ($\frac{dTER}{delevation}$) for each warm month into the additive contributions of five or four
 724 components $X \left(\frac{dY}{delevation} \right)^X$, which was represented as the product of the partial
 725 derivative against that variable X as $\frac{\partial Y}{\partial X}$ and the elevation gradient of X itself as
 726 $\frac{dX}{delevation}$ ⁸⁰, as shown in **Equation (1)**.

$$\begin{aligned}
 727 \quad \frac{dY}{delevation} &= \frac{\partial Y}{\partial MAT} \frac{dMAT}{delevation} + \frac{\partial Y}{\partial VPD} \frac{dVPD}{delevation} + \frac{\partial Y}{\partial PRE} \frac{dPRE}{delevation} + \frac{\partial Y}{\partial SM} \frac{dSM}{delevation} + \\
 728 \quad \frac{\partial Y}{\partial PAR} \frac{dPAR}{delevation} + \varepsilon &= \left(\frac{dY}{delevation} \right)^{MAT} + \left(\frac{dY}{delevation} \right)^{VPD} + \left(\frac{dY}{delevation} \right)^{PRE} + \\
 729 \quad \left(\frac{dY}{delevation} \right)^{SM} &+ \left(\frac{dY}{delevation} \right)^{PAR} + \varepsilon \quad (1)
 \end{aligned}$$

730 where $\frac{\partial Y}{\partial X}$ represents the sensitivity of Y (GPP or TER) to an explanatory variable X
 731 (MAT, VPD, PRE, SM, total PAR [only for GPP], respectively). These sensitivities
 732 were estimated as the regression coefficients of a multiple linear regression performed
 733 with GPP or TER against all listed explanatory variables. $\frac{dY}{delevation}$ (or $\frac{dX}{delevation}$)
 734 represents the sensitivity of Y or X to elevation (300-1000 m or 1000-2000 m) for each
 735 warm month. The sensitivity was calculated as the slope of the simple linear regression
 736 of mean Y (or X) values against the elevation.

737

738 ***Analysis of potential uncertainties arising from forest biomass carbon data, carbon***
 739 ***accumulation rate calculations, soil fertility, types of disturbances, and various sub-***
 740 ***regions***

741 The reliability of forest biomass carbon data is a pivotal factor influencing the
 742 calculation precision of the biomass carbon accumulation rate. Previous studies have
 743 suggested that the ESA-CCI data may underestimate AGB in regions characterized by
 744 high AGB density ($> 250 \text{ Mg ha}^{-1}$), particularly in low-elevation tropical forests¹⁷. To
 745 address this issue, we refined the bias in the ESA-CCI-derived total biomass data by
 746 integrating in situ observations, using the techniques proposed by Zhao et al.¹⁰⁰. An
 747 upward trend persists in the biomass carbon accumulation rate with elevation from 300

748 m to 1000 m based on the adjusted biomass carbon data (slope = $0.20 \pm 0.05 \text{ MgC} \cdot \text{ha}^{-1}$
749 $\text{yr}^{-1} \text{ m}^{-1}$; mean carbon accumulation rate = $2.94 \pm 0.28 \text{ MgC} \cdot \text{ha}^{-1} \text{ yr}^{-1}$, **Figure S16**). This
750 consistency aligns with our findings based on the original ESA-CCI-derived total
751 biomass carbon data (slope = $0.23 \pm 0.04 \text{ MgC} \cdot \text{ha}^{-1} \text{ yr}^{-1} \text{ m}^{-1}$; the mean rate = 2.85 ± 0.28
752 $\text{MgC} \cdot \text{ha}^{-1} \text{ yr}^{-1}$, **Figure 1e**).

753 To determine the biomass carbon accumulation rate of TPRFs during the rapid
754 growth phase across various elevations, this study initially focused on data points where
755 the biomass carbon was below 80% of that in old-growth forests for analysis.
756 Subsequently, we adjusted these thresholds to 70% and 90% to examine their potential
757 effects on the elevation dependence of the biomass carbon accumulation rate (**Figure**
758 **3f**). Results reveal very slight variations in the slopes of the linear regression curve
759 between biomass carbon accumulation rate and elevation (for altitudes between 300 m
760 and 1000 m: 70% threshold: slope = $0.11 \pm 0.05 \text{ MgC} \cdot \text{ha}^{-1} \text{ yr}^{-1} \text{ m}^{-1}$; 80% threshold: slope
761 = $0.23 \pm 0.04 \text{ MgC} \cdot \text{ha}^{-1} \text{ yr}^{-1} \text{ m}^{-1}$; 90% threshold: slope = $0.19 \pm 0.06 \text{ MgC} \cdot \text{ha}^{-1} \text{ yr}^{-1} \text{ m}^{-1}$). We
762 further verified the results of this method for detecting the rapid growth stand age
763 (age_{rapid}) with the observed age_{rapid} , which was determined as the stand age when
764 there was a break point change in the relationship of five-years carbon accumulation
765 rate and age. This time point was identified using segmented regression models¹⁰¹
766 (**Figure S3**). Results indicated that the estimated age_{rapid} using the 80% peak
767 biomass coincided well with those estimated from the derivative change of
768 accumulation rates (**Figure 3e**).

769 Human activities have the potential to influence the biomass carbon accumulation
770 rate in TPRFs¹⁸. Here, we both tested the intrinsic influences from previous land use
771 types before converting to TPRFs and external influences from surrounding non-forest
772 lands. The fraction of non-forest lands (urban and croplands) nearby the TPRFs
773 (neighboring 10×10 1km resolution pixels) decreases with elevation, contributing
774 slightly to the increasing elevational trend of the carbon accumulation rate in TPRFs
775 (**Figure S10**). In contrast, the elevational patterns in TPRFs that used to be shifting
776 cultivation and pasture both show marginally small variations (**Figure S13**). Worthy of

777 note is that, we did not analyze other land use types, such as fire, clear-cut harvest, and
778 mining, due to a lack of sufficient biomass carbon data across the studied elevation
779 range.

780 Moreover, organic carbon (<https://openlandmap.org>), total phosphorus¹⁰², sand and
781 clay concentrations (<https://openlandmap.org>) in soil all exhibited insignificant trends
782 ($P>0.05$) along with elevation (**Figure S17**), indicating a limited influence of soil
783 fertility on the elevation dependence of carbon accumulation rate.

784 All these additional analyses confirm the robustness of our findings regarding the
785 elevational patterns of biomass carbon accumulation rates in TPRFs.

786

787 **Resource availability**

788 *Lead contact*

789 For further information on the analysis, please contact the corresponding author,
790 Yongxian Su (yxsu@rcees.ac.cn).

791 *Materials availability*

792 This study has not generated any new, unique materials.

793 *Data and Code Availability*

794 All the original datasets used in this research are publicly available from their
795 sources: a global Forest Carbon database (ForC): <https://github.com/forc-db>; ESA-CCI
796 AGB map: <https://catalogue.ceda.ac.uk/uuid/af60720c1e404a9e9d2c145d2b2ead4e>;
797 0.1° global live biomass carbon: <https://zenodo.org/records/4161694>; Carbon flux
798 (GPP/TER/NPP/NEP) simulations obtained from the BEPS model:
799 <https://datadryad.org/stash/landing/show?id=doi%3A10.5061%2Fdryad.q573n5tgb>;
800 Jung et al.'s FLUXCOM data: <https://fluxnet.org/data/fluxnet2015-dataset/>; Tree
801 height dataset: <https://glad.umd.edu/dataset/GLCLUC2020>;
802 <https://www.nature.com/articles/s41559-023-02206-6>; JRC-TMF dataset
803 (<https://forobs.jrc.ec.europa.eu/TMF/download/>); MCD12C1 v061 land cover:
804 <https://lpdaac.usgs.gov/products/mcd12c1v061/>; TerraClimate MAT, VPD, PRE:
805 <https://www.climatologylab.org/terraclimate.html>; Global remote-sensing-based

806 surface soil moisture (RSSM): <https://doi.pangaea.de/10.1594/PANGAEA.912597>;
807 BESS total PAR: <https://www.environment.snu.ac.kr/bess-rad>; Plantation year dataset:
808 [https://figshare.com/articles/dataset/A_global_map_of_planting_years_of_plantations](https://figshare.com/articles/dataset/A_global_map_of_planting_years_of_plantations/19070084/1)
809 [/19070084/1](https://figshare.com/articles/dataset/A_global_map_of_planting_years_of_plantations/19070084/1); Global reforestation potential map: <https://zenodo.org/records/883444>;
810 Global soil total phosphorus concentration dataset:
811 <https://doi.org/10.6084/m9.figshare.14583375.v9>; Global soil organic/sandy/clay
812 carbon dataset: <https://zenodo.org/records/2525663>;
813 <https://doi.org/10.5281/zenodo.2525662>; <https://doi.org/10.5281/zenodo.2525553>;
814 Tree species dataset: <https://gdauby.github.io/rainbio/index.html>

815 The code used for this analysis is available in a Zenodo repository at
816 <https://doi.org/10.5281/zenodo.13922571> (ref.103).

817

818 **CONTRIBUTIONS**

819 Y.S. and X.C. designed the study and wrote the initial manuscript. Y.S., C.Z., W.
820 Y. and X.L. collected the data and performed the analysis. P.C., S.C.P, O.L.P., J.S., A.C.,
821 J.M.C., J.L., J.C., C.E.D., F.T., Y.Q.L., Y.L., Z.Y., D.H., S.T. and Y.Z. contributed to
822 discuss the scientific question and revise the manuscript. All authors reviewed, revised
823 and approved the manuscript.

824

825 **COMPETING INTEREST STATEMENT**

826 We thank Dr. Ranga Myneni for his valuable suggestions to improve the
827 manuscript. The authors declare no competing interests.

828

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832

833

TABLE

834

Table 1 Independent biomass, carbon, and height datasets and corresponding methods to quantify the γ_{rapid} (see **Experimental Procedure** for details).

835

NO.	Biomass carbon proxies	Biomass carbon data sources	Stand age data sources	Stand age estimation method	γ_{rapid} estimation method	Acronyms
Dataset 1	forest biomass (<i>in situ</i> , global)	Smithsonian Institution's Global Forest Carbon (ForC) database ³⁶ and other in-situ datasets ⁹		Original stand age records	γ_{rapid} was determined as the slope of linear regression fit between observed forest biomass carbon densities and the stand age of TPRFs where biomass carbon was less than 80% of old-growth forests (stand age ≥ 100 years) simulated in the Chapman–Richards curve using the space-for-time method.	γ_{rapid}^{GFC}
Dataset 2	Aboveground forest biomass (100m, global, 2018)	ESA-CCI ¹⁷	Cover change map of tropical moisture forest (TMF) (30m, Tropical, 1982-2019) ⁴¹ and global plantation years dataset (GPY) (30m, global, 1982-2020) ⁴²	Stand age of natural regrowth forests was determined based on TMF cover change data. Stand age of planted forests was calculated based on GPY dataset.		$\gamma_{rapid}^{ESA-CCI}$
Dataset 5	Forest canopy height dataset (30m, global, 2019)	GEDI LiDAR dataset ³⁹				γ_{rapid}^{GEDI}
Dataset 3	Total live woody biomass (0.1°, global, 2000-2019)	Machine-learning (ML)-derived terrestrial live biomass dataset ³⁷	MODIS MCD12C1 landcover products (0.05°, global, 2001-2019) ⁴³	Stand age was calculated based on the time series MODIS MCD12C1 land cover dataset.		γ_{rapid}^{ML}
Dataset 4	Net ecosystem productivity (0.07273°, global, 1981-2019)	BEPS model ³⁸				γ_{rapid}^{BEPS}

836

837

FIGURE CAPTIONS

838 **Figure 1. Average carbon accumulation rates (γ_{rapid}) in TPRFs by elevation up to**
839 **2000 m above mean sea level (a.m.s.l.). a, *In-situ* sites of field measurements and grid**
840 **cells of raster-based data used in this study. Purple triangles represent *in situ* sites of**
841 **forest biomass data used for γ_{rapid}^{GFC} estimation. The dots in different green colors**
842 **represent the $100\text{ m} \times 100\text{ m}$ grid cells used for $\gamma_{rapid}^{ESA-CCI}$ or γ_{rapid}^{GEDI} estimation. Black**
843 **dots represent the $0.1^\circ \times 0.1^\circ$ grid cells used for γ_{rapid}^{ML} or γ_{rapid}^{BEPS} estimation. **b-f,****
844 **Elevation pattern of annual γ_{rapid}^{GFC} (b), $\gamma_{rapid}^{ESA-CCI}$ (c), γ_{rapid}^{ML} (d), γ_{rapid}^{BEPS} (e) and**
845 **γ_{rapid}^{GEDI} (f), respectively. The dashed lines represent the linear fit between γ_{rapid} and**
846 **elevation, with shading representing the 95% confidence interval. Significant**
847 **relationships ($P \leq 0.05$) are denoted in blue, and insignificant ones ($P > 0.05$) in grey. In**
848 **panels b, c, and f, each histogram represents the slope of the corresponding ordinary**
849 **least squares (OLS) regression line (orange dashed lines in the inset plot)⁹ between**
850 **biomass carbon and age during the rapid growth stage of forest (γ_{rapid}) before reaching**
851 **maturity (black dashed vertical lines in the inset plot, representing 80% of its maximum**
852 **biomass carbon and grey dashed lines in the inset plots represent the fitted Chapman–**
853 **Richards curve) within each elevation bin ($100 \pm 50\text{ m}$ in an 80 m step) using the space-**
854 **for-time analogy. Color gradients of histogram graphs indicate the R^2 between**
855 **simulated and observed γ_{rapid} in each elevation bin. Error bars indicate one standard**
856 **error. In panel d, black dots represent the slope of the corresponding linear regression**
857 **curve (orange dashed lines in inset plot) between model-simulated plant carbon**
858 **densities and stand age, using the time-series data from each TPRF. In panel e, each**
859 **black triangle represents the mean NEP of each targeted TPRF, with error bars**
860 **indicating one standard error. Numbers at the top of panels d and e represent the slope**
861 **of linear regression between γ_{rapid} and elevation, with significance indicated in the**
862 **legend as $\bullet P < 0.5$, $* P < 0.1$, $** P < 0.01$, and $*** P < 0.001$. Notably, the assessment of the**
863 **relationship between γ_{rapid} and elevation is only up to 1300 m based on in-situ data**

864 (panel **a**), while analyses using other datasets (panels **b-f**) are up to 2000 m.
865 **Figure 2. Elevational patterns of average carbon accumulation rates (γ_{rapid})**
866 **derived from three datasets across various forest characteristics at elevations up**
867 **to 2000 m above mean sea level (a.m.s.l.). a, d, g,** The elevational patterns of
868 γ_{rapid}^{GFC} (**a**), $\gamma_{rapid}^{ESA-CCI}$ (**d**) and γ_{rapid}^{BEPS} (**g**) in natural regrowth compared to planted
869 forests. The same as for **a, d, g**, but in broadleaved compared to needle-leaved forests.
870 **c, f, i,** The same as for **a, d, g**, but in north-facing compared to south-facing slopes of
871 mountains. In panels **a-f**, each dot represents the mean γ_{rapid} for each elevation bin
872 (100 ± 50 m in an 80 m step), estimated based on the Chapman–Richards curve, with
873 the error bars indicating one standard error in each elevational bin. In **g-i**, each dot
874 represents the mean NEP for each targeted forest grid cell, with the error bars indicating
875 one standard error. The lines in panels **a-i** represent the linear fit between γ_{rapid} and
876 elevation, with shading indicating the 95% confidence interval. Significant
877 relationships ($P\leq 0.05$) are denoted by solid lines, and insignificant ones ($P>0.05$) are
878 represented by dashed lines. Numbers at the top of panels represent the slope of linear
879 regression between γ_{rapid} and elevation, with significance indicated in the legend as
880 • $P<0.5$, * $P<0.1$, ** $P<0.01$, and *** $P<0.001$.

881 **Figure 3. The influence of stand age on the elevational pattern of γ_{rapid} .** **a,**
882 Illustrations of the fixed stand age approach for calculating the carbon accumulation
883 rate. The slopes of three blue dashed curves represent the carbon accumulation rates
884 during $T_1 - T_2$, $T_2 - T_3$ and $T_1 - T_3$ time periods, respectively. **b,** Illustrations of the carbon
885 accumulation window approach for calculating γ_{rapid} . The slopes of red solid and red
886 dashed curves represent the γ_{rapid} at low- (i.e., < 500 m) and mid-elevations (i.e., 500
887 -1500 m^{48, 49}), respectively. The selected window is intensified when carbon
888 accumulation reaches 80% of the median biomass carbon of old-growth forests (stand
889 age > 100 years). The black solid and dashed curves represent the Chapman–Richards
890 curves for TPRFs at low- and mid-elevations, respectively. **c,** Sensitivity of total
891 biomass carbon derived from ESA-CCI data to elevation (i.e., 300-1000 m and 1000-

2000 m) across different stand ages. Sensitivity is defined as the slope of the linear regression curve illustrating the relationship between biomass carbon and elevation within each one-year stand age bin (**Figure S6**). **d**, Elevational patterns of biomass carbon accumulate rates using diverse stand age windows (i.e., 20, 24, 28, and 32 years of age) to locate the analysis period along the Chapman–Richards curve, as used in previous studies⁹. **e**, Comparisons between the stand ages (age_{rapid}) with rapid accumulation rate identified using 80% peak biomass thresholds and observed age_{rapid} , which was determined as the stand age when there was an abrupt change with the relationship with stand age as stand age increases. **f**, Elevational patterns of biomass carbon accumulate rates using different peak biomass thresholds (i.e., from zero to the time of reaching 70%, 80% and 90% of the maximum value). In panels **d** and **f**, each dot represents the slope of the corresponding ordinary least squares (OLS) regression curve between ESA-CCI-derived total biomass carbon and the stand age within given age windows (**d**) or using different peak biomass thresholds (**f**) at a given elevation bin (100 ± 50 m in an 80 m step). The colored curves depict the linear regressions between γ_{rapid} and elevation, ranging from 300 m to 1000 m and 1000 m to 2000 m, respectively.

Figure 4. Impact of climatic factors on the elevation patterns of seasonal carbon fluxes. **a-b**, Linear regressions between each month's GPP (red), TER (blue), NEP (orange) simulated by the BEPS model (**Dataset 4**)³⁸ and elevation for ranges of 300-1000 m (**a**) and 1000-2000 m (**b**), respectively. **c-f**, Linear regressions between each month's mean air temperature (MAT) (**c-d**) or precipitation (PRE) (**e-f**) and elevation for ranges of 300-1000 m (**c, e**) and 1000-2000 m (**d, f**), respectively. Shadings indicate 95% confidence intervals. Significant relationships ($P \leq 0.05$) are shown in solid lines, and non-significant relationships ($P > 0.05$) in dashed lines. **g-h**, Changes in monthly GPP and TER with MAT (**g**) or PRE (**h**), respectively. Each dot denotes the median value of GPP (red) (or TER [blue]) within a 1°C MAT bin (or 20mm PRE bin), respectively. Error bars depict one standard deviation. The dashed curves are fitted using local polynomial regression based on the 'loess' function in the R 'stats' package

921 with default settings. The solid curves in panel **g** represent the linear fitting regressions
922 between the GPP, TER, and MAT for MAT higher than 24°C, while in panel **h** represent
923 the linear fitting regressions between the GPP, TER, and PRE for PRE lower than 150
924 mm. In panels **c-d** and **g**, the brown background indicates that MAT is $\geq 24^\circ\text{C}$. In panel
925 **h**, the light blue background indicates that PRE < 150 mm. **i-j**, Absolute contributions
926 of the elevational trends in climate factors to the elevational variations in GPP and TER
927 determined by the multiple linear regression model for months with MAT $\geq 24^\circ\text{C}$ (**i**)
928 and $< 24^\circ\text{C}$ (**j**), respectively. Error bars represent one standard error. The asterisks
929 indicate the P values: * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$.

930 **Figure 5. Elevational patterns of γ_{rapid}^{BEPS} and the underlying mechanisms across**
931 **four mountains. a**, Locations of four tropical mountains: the Serra do Espinhaço
932 mountain in South America (SMS), the Ethiopian Highlands mountain in Africa (EH),
933 the Serra do Espinhaço mountain in South America (SE), and the Eastern Ghats
934 mountain in Asia (EG). **b-e**, Elevational pattern of γ_{rapid}^{BEPS} in four mountains. Blue
935 dashed curves and red solid lines represent the smoothed trend fitted by a generalized
936 additive model (GAM) and the linear fit at both sides of each threshold, respectively. **f**,
937 Elevational pattern of mean air temperature and precipitation in four mountains. **g**, The
938 optimal elevation, i.e., the places with the highest values of γ_{rapid}^{BEPS} in four mountains.
939 **h**, Elevational sensitivity of γ_{rapid}^{BEPS} below (green histogram) and above (grey
940 histogram) the optimal elevation in four mountains. **i-l**, Absolute contributions of the
941 elevational trends in climate factors to the elevational variations in GPP and TER
942 determined by the multiple linear regression model for months with MAT $\geq 24^\circ\text{C}$ in
943 SMS (**i**), EH (**j**), SE (**k**), and EG (**l**), respectively. **m-p**, Absolute contributions of the
944 elevational trends in climate factors to the elevational variations in GPP and TER for
945 months with MAT $< 24^\circ\text{C}$ in corresponding four mountains. Error bars represent one
946 standard error.

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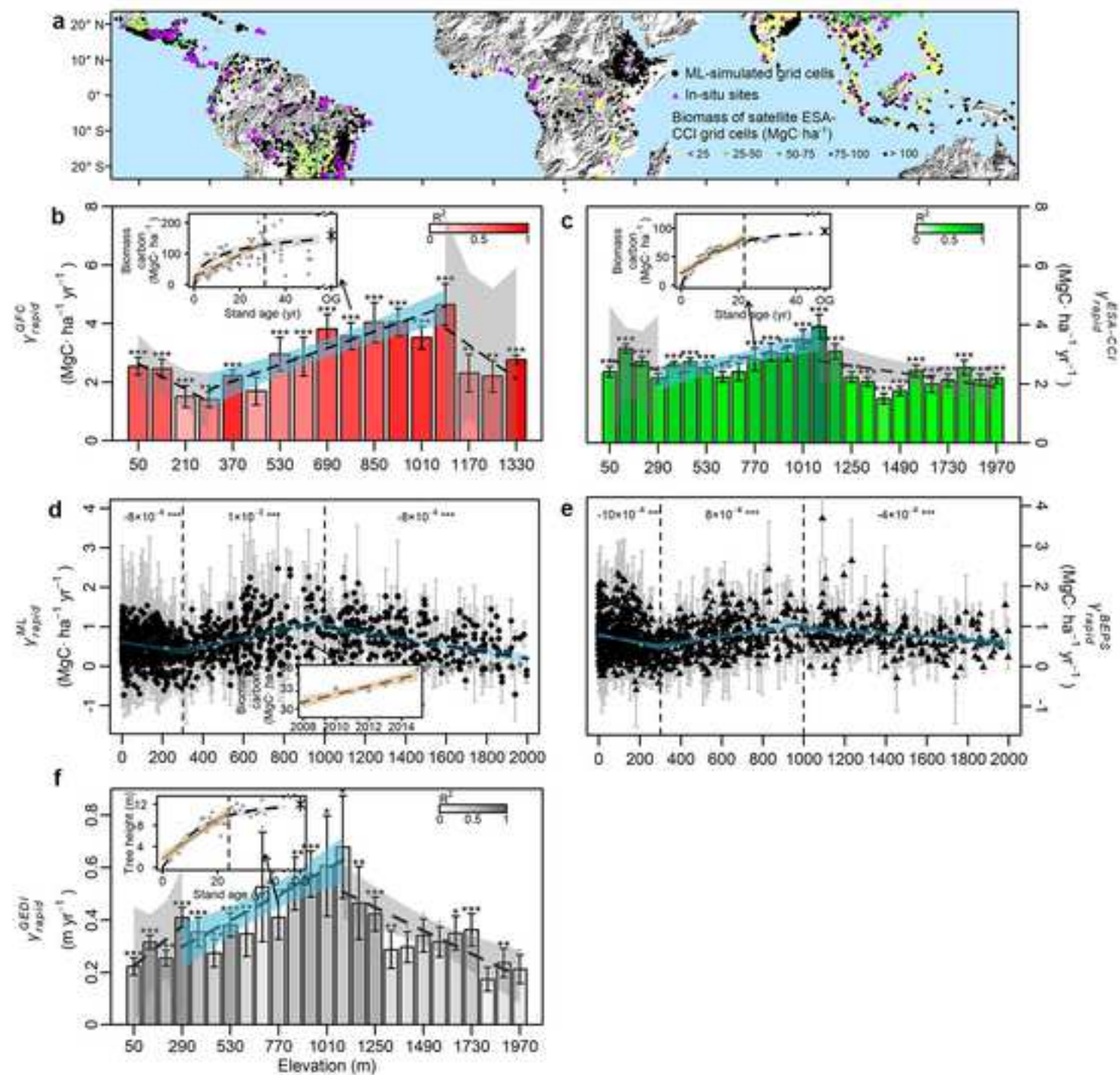
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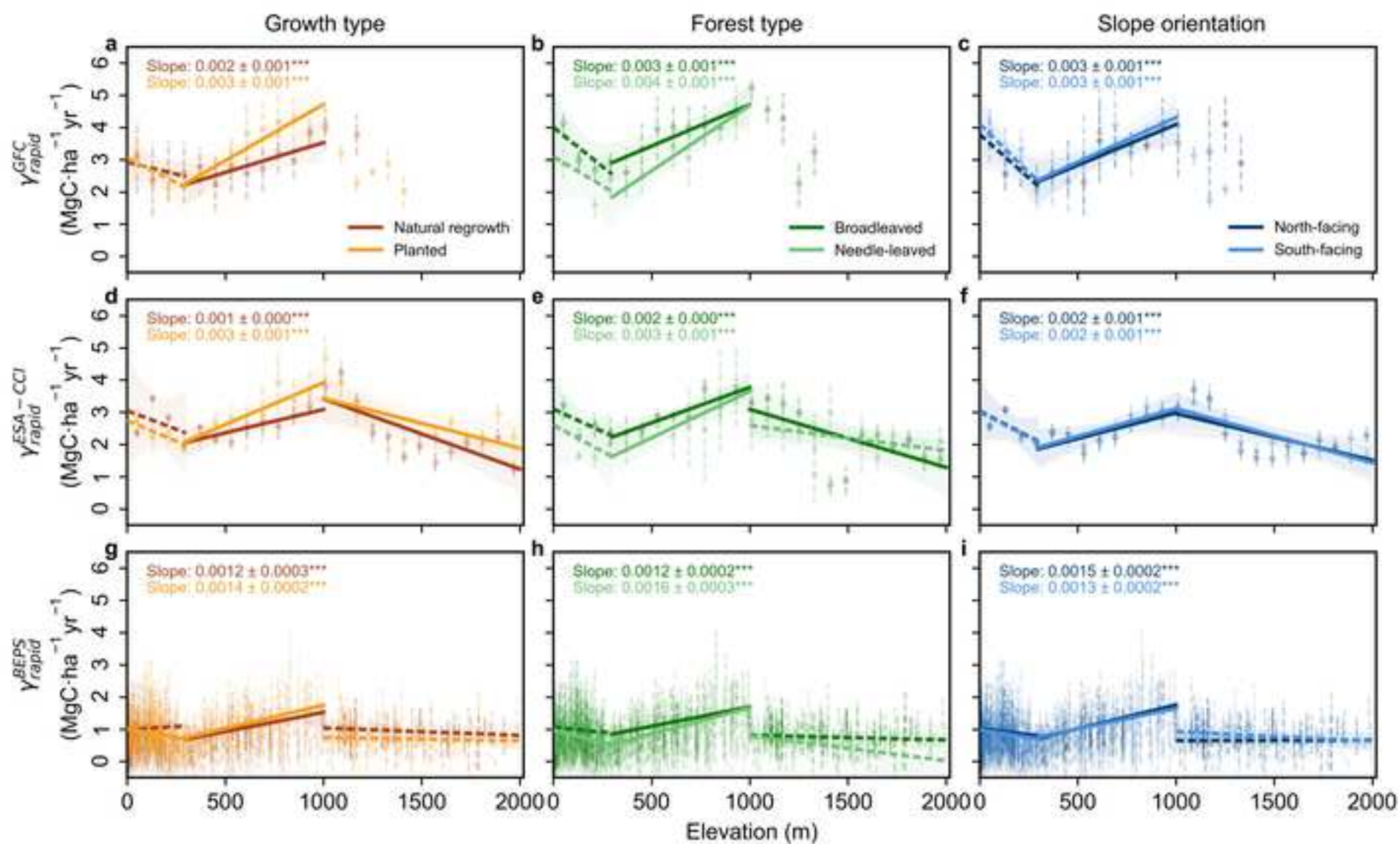
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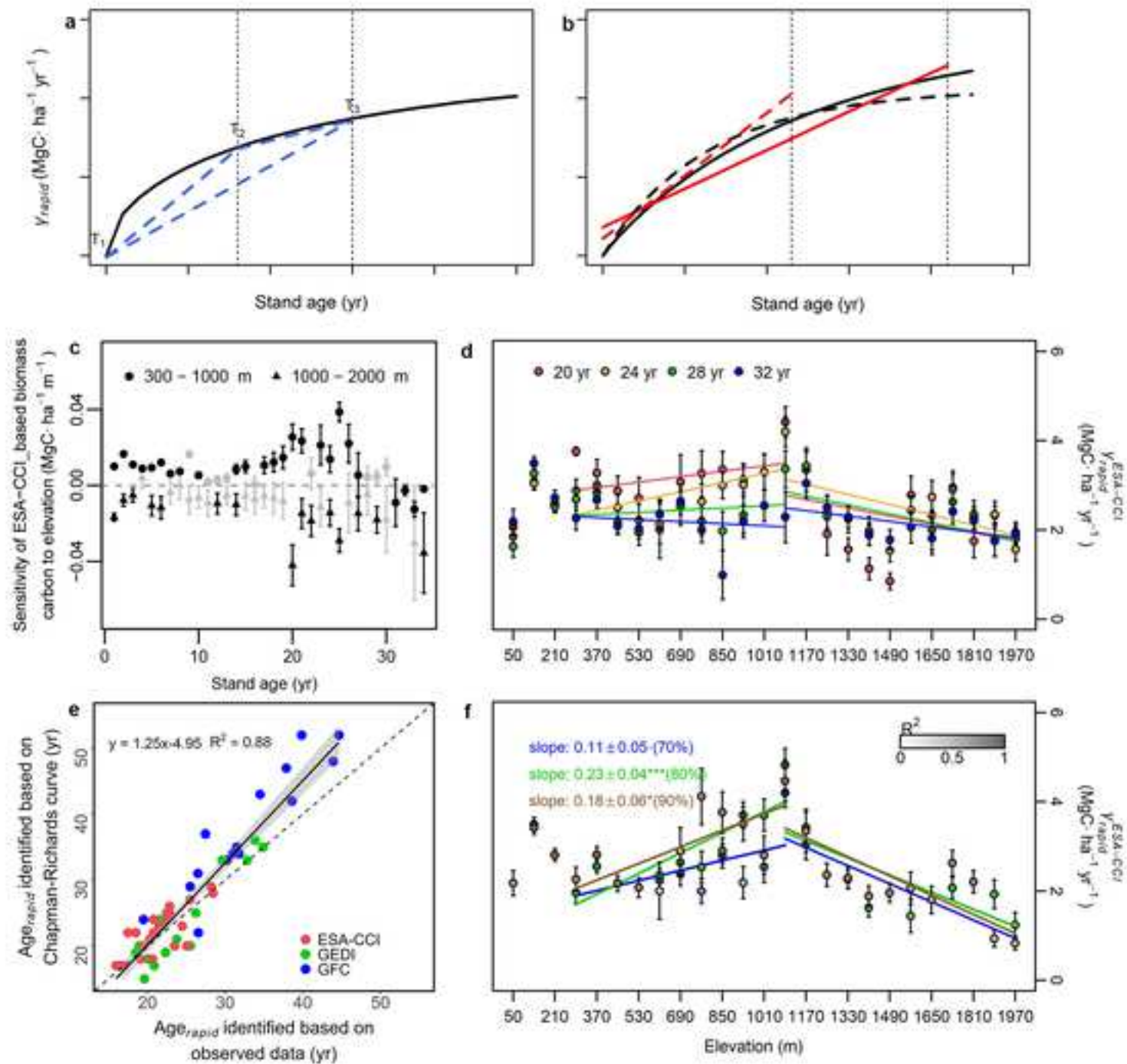
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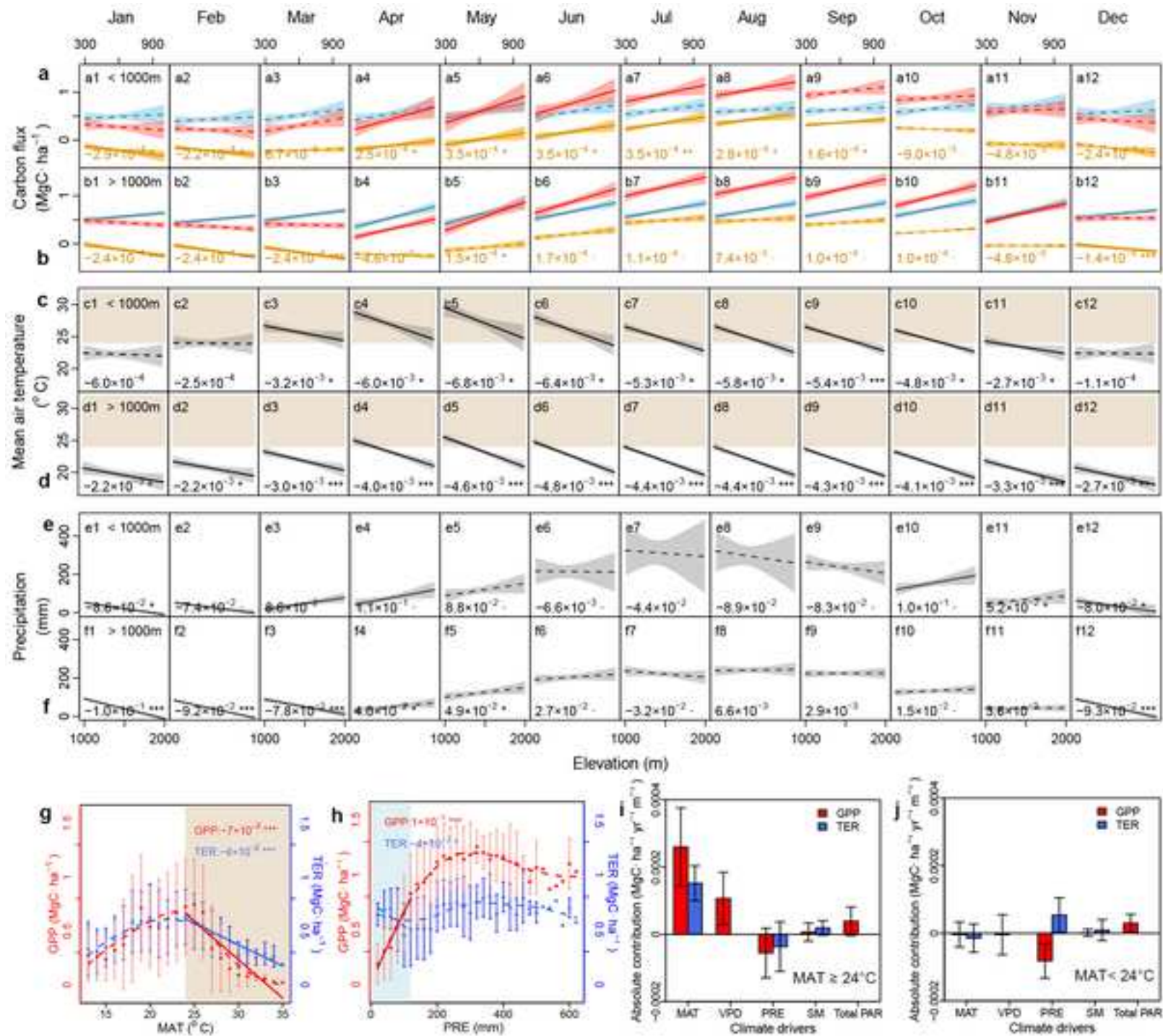
1 **Table 1** Independent biomass, carbon, and height datasets and corresponding methods to quantify the γ_{rapid} (see **Experimental Procedure** for
2 details).

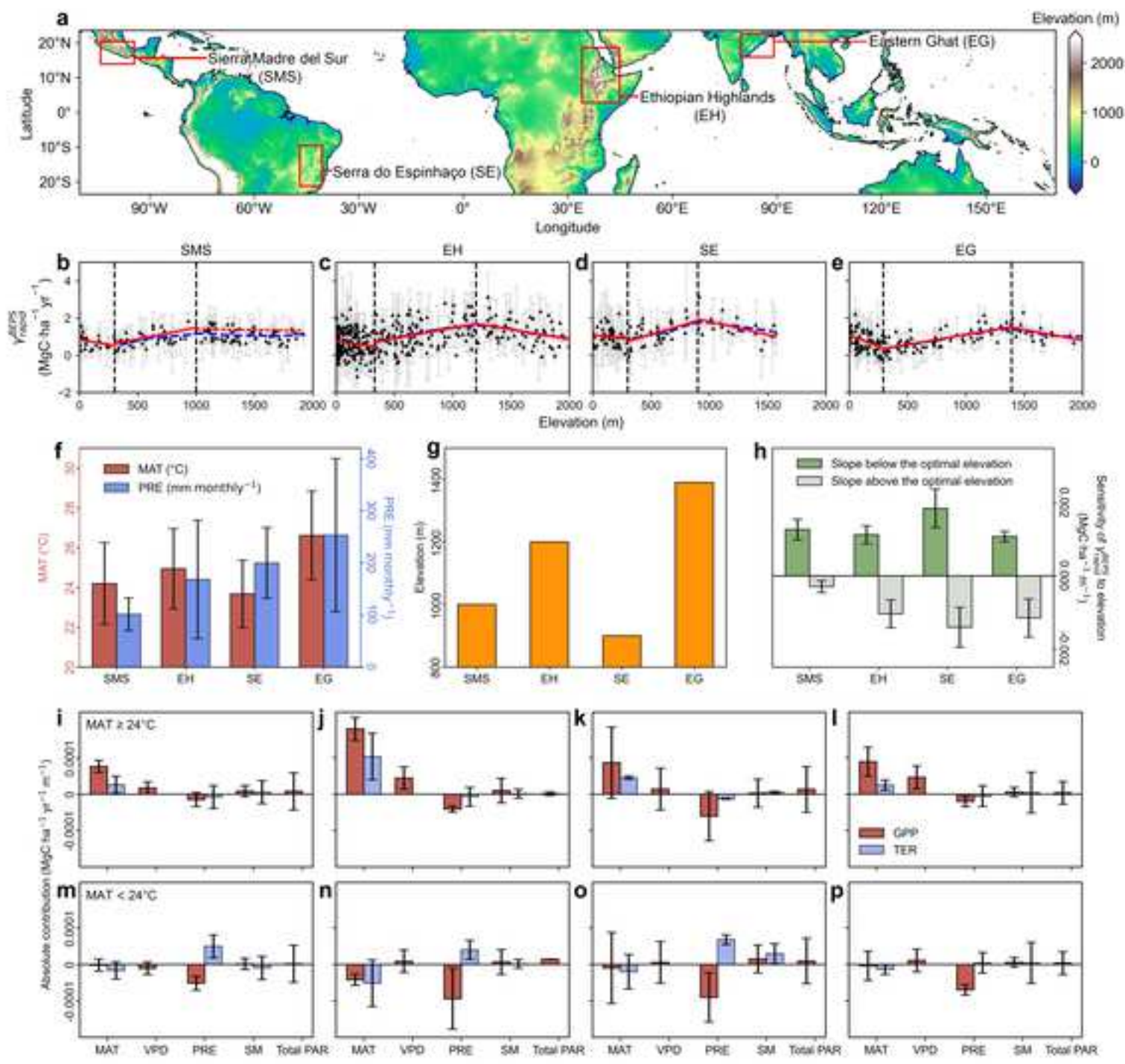
NO.	Biomass carbon proxies	Biomass carbon data sources	Stand age data sources	Stand age estimation method	γ_{rapid} estimation method	Acronyms
Dataset 1	forest biomass (<i>in situ</i> , global)	Smithsonian Institution's Global Forest Carbon (ForC) database ³⁶ and other in-situ datasets ⁹		Original stand age records	γ_{rapid} was determined as the slope of linear regression fit between observed forest biomass carbon densities and the stand age of TPRFs where biomass carbon was less than 80% of old-growth forests (stand age ≥ 100 years) simulated in the Chapman–Richards curve using the space-for-time method.	γ_{rapid}^{GFC}
Dataset 2	Aboveground forest biomass (100m, global, 2018)	ESA-CCI ¹⁷	Cover change map of tropical moisture forest (TMF) (30m, Tropical, 1982-2019) ⁴¹ and global plantation years dataset (GPY) (30m, global, 1982-2020) ⁴²	Stand age of natural regrowth forests was determined based on TMF cover change data. Stand age of planted forests was calculated based on GPY dataset.		$\gamma_{rapid}^{ESA-CCI}$
Dataset 5	Forest canopy height dataset (30m, global, 2019)	GEDI LiDAR dataset ³⁹				γ_{rapid}^{GEDI}
Dataset 3	Total live woody biomass (0.1°, global, 2000-2019)	Machine-learning (ML)-derived terrestrial live biomass dataset ³⁷	MODIS MCD12C1 landcover products (0.05°, global, 2001-2019) ⁴³	Stand age was calculated based on the time series MODIS MCD12C1 land cover dataset.		γ_{rapid}^{ML}
Dataset 4	Net ecosystem productivity (0.07273°, global, 1981-2019)	BEPS model ³⁸				γ_{rapid}^{BEPS}





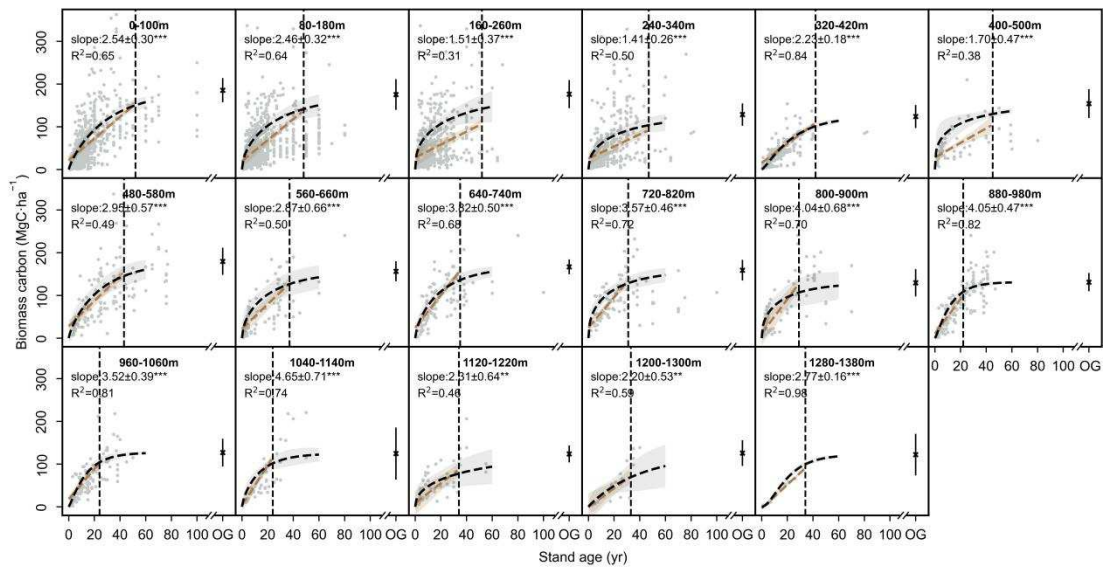






1

Supplementary information



2

3 **Figure S1. γ_{rapid}^{GFC} for each elevation bin (100 ± 50 m) in 80 m step.** Grey points

4 represent *in situ* tree biomass carbon. The Grey dashed curves represent the smoothed

5 trend fitted by Chapman Richards growth model, which was used to find the mature

6 age threshold when the accumulated carbon reached 80% of median carbon of old-

7 growth forest (the black crosses). Error bar indicates one standard error. Brown dashed

8 lines represent the linear regression curves between *in situ* tree biomass carbon and

9 stand age of all sites of TPRFs during the rapid growth stage of trees before approaching

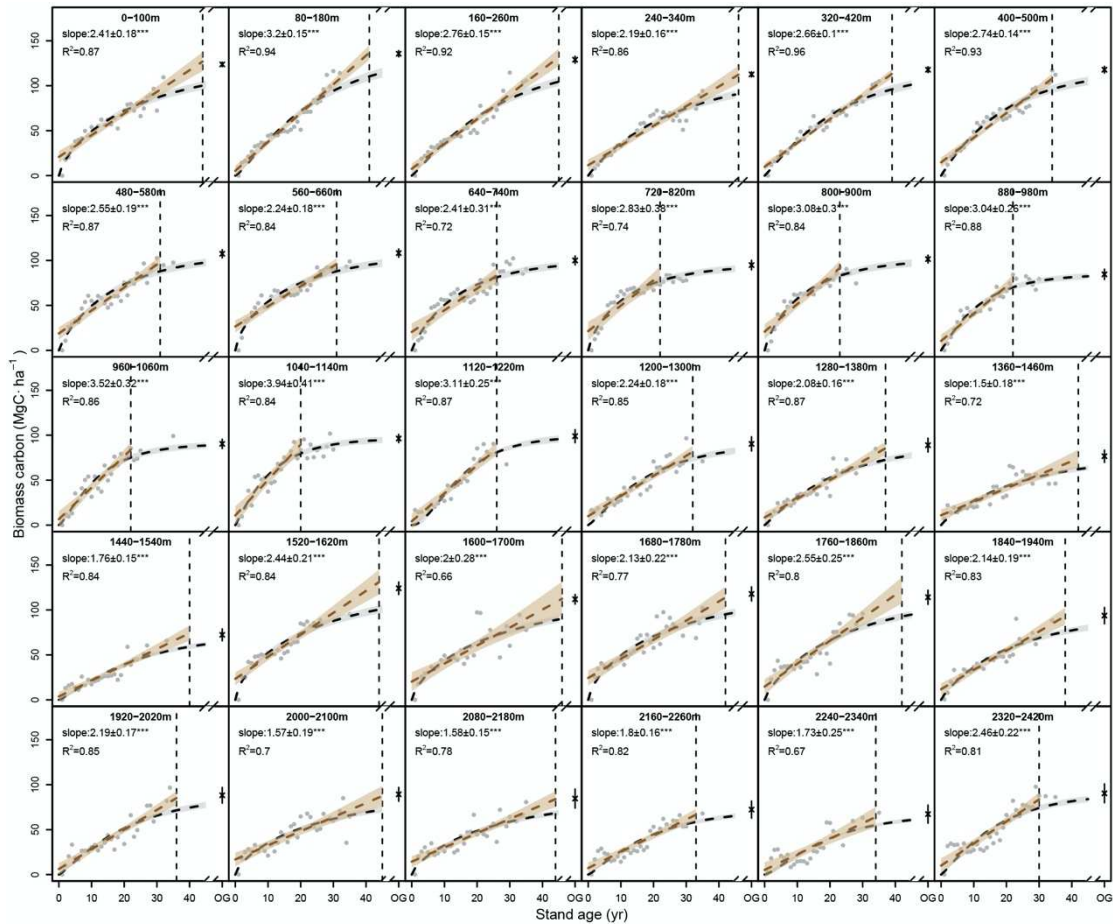
10 maturity within the given elevation bin. Shading represents the 95% confidence interval.

11 Significance of linear regression is indicated in the legend as: • $P < 0.5$, * $P < 0.1$,

12 ** $P < 0.01$, and *** $P < 0.001$.

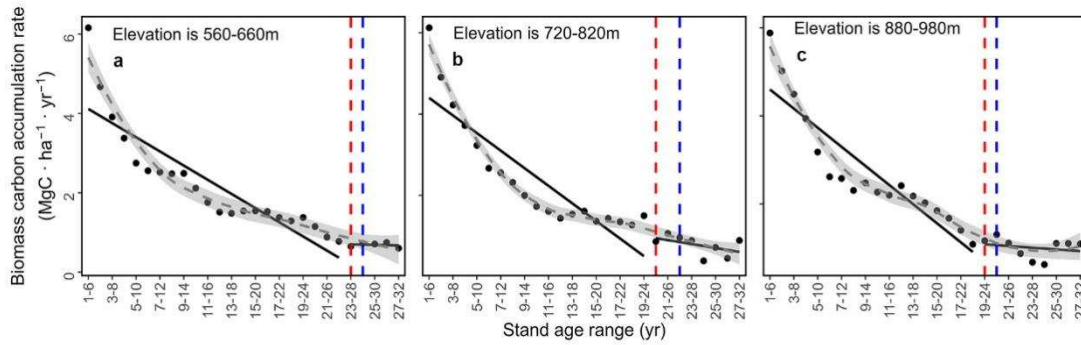
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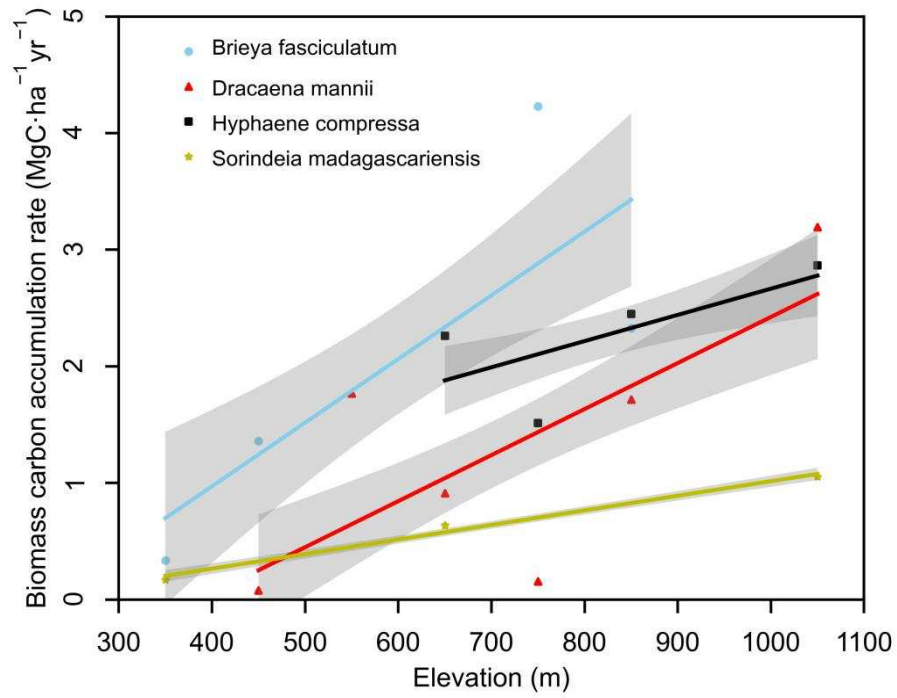
16 **Figure S2.** $\gamma_{rapid}^{ESA-CCI}$ for each elevation bin (100 ± 50 m) in 80 m step. Grey points
 17 denote the median biomass carbon value calculated for each stand age bin (± 1 year).
 18 Grey dashed curves represent the smoothed trend fitted by Chapman Richards growth
 19 model, which was used to find the mature age threshold when the accumulated carbon
 20 reached 80% of median carbon of old-growth forest (the black crosses). Error bar
 21 indicates one standard error. Brown dashed lines represent the linear regression curves
 22 between ESA-CCI-derived total biomass carbon and stand age of all sites of TPRFs
 23 during the rapid growth stage of trees before approaching maturity within the given
 24 elevation bin. Shading represents the 95% confidence interval. Significance of linear
 25 regression is indicated in the legend as: • $P < 0.5$, * $P < 0.1$, ** $P < 0.01$, and *** $P < 0.001$.
 26



27

28 **Figure S3. Illustrations for determining the stand age (age_{rapid}) of the rapid**
 29 **growth phase in TPRFs. age_{rapid} was defined as the stand age when there was a**
 30 **break point of the relationship with stand age as stand age increases. Panels a-c are**
 31 **examples of age_{rapid} thresholds identified for three altitude ranges: 560-660 m (a),**
 32 **720-820 m (b), and 880-980 m (c), respectively, based on ESA-CCI data. The black**
 33 **dots represent the slopes of ordinary least squares regression applied to the**
 34 **relationship between biomass carbon and stand ages for each five-year interval,**
 35 **defining $\gamma_{rapid}^{ESA-CCI}$ for each five years bin. Grey dashed curves and black solid lines**
 36 **represent the smoothed trend fitted by a generalized additive model (GAM) and the**
 37 **linear fits at both sides of each threshold, respectively. The age_{rapid} with an abrupt**
 38 **change between the relationship with stand age as stand age increases was identified**
 39 **using the segmented package in R. Red vertical dashed line represents the observed**
 40 **age_{rapid} identified using this method. Blue vertical dashed line represents the**
 41 **simulated age_{rapid} identified by 80% of the maximum value on the Chapman-**
 42 **Richards curve.**

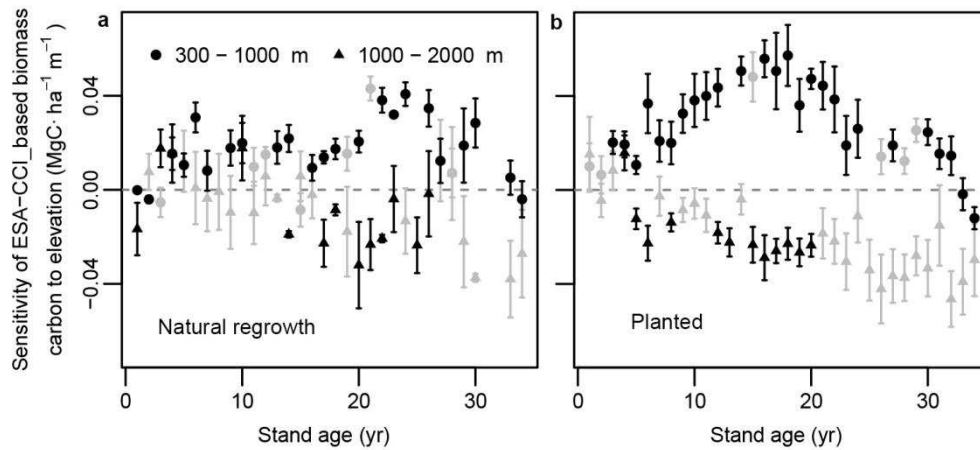
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44

45 **Figure S4. The elevation patterns of $\gamma_{rapid}^{ESA-CCI}$ for four tree species based on a**
 46 **comprehensive mega-database of tropical African vascular plants distributions²**
 47 **(Methods).**

48

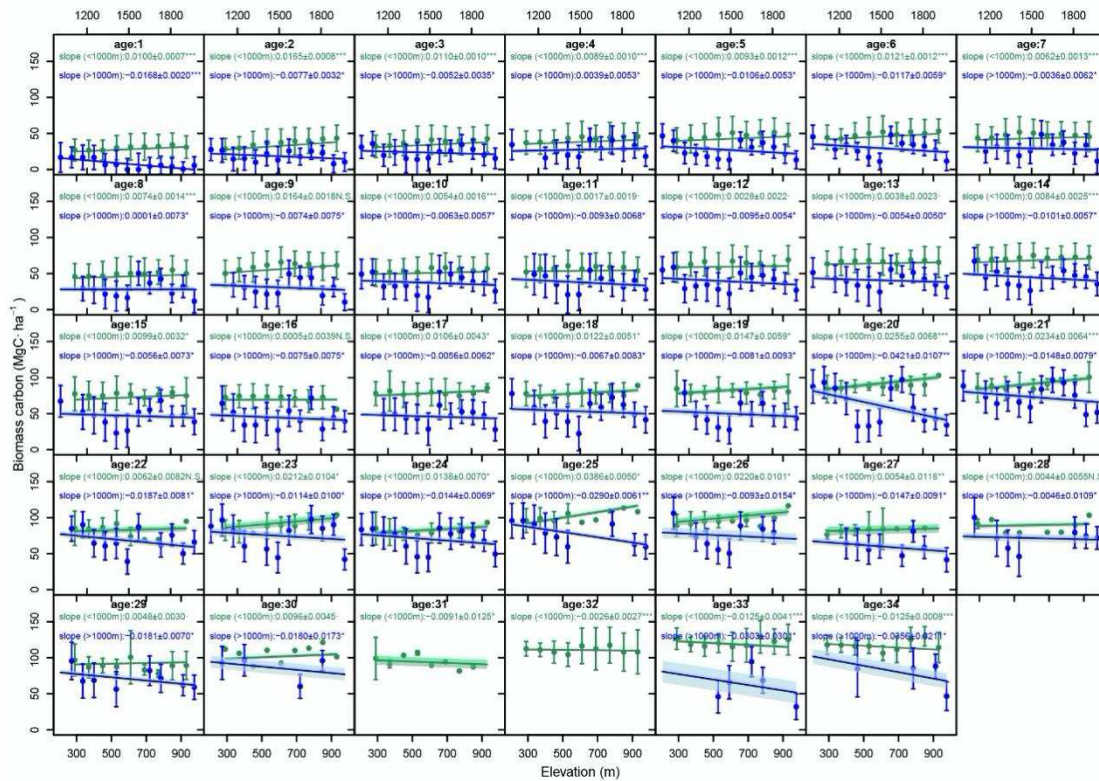


49

50 **Figure S5. Sensitivity of total biomass carbon derived from ESA-CCI data to**
 51 **elevation (i.e., 300-100 m and 1000-2000 m) across different stand ages for natural**
 52 **regrowth (a) and planted forest (b), respectively.** Sensitivity is defined as the slope
 53 of the linear regression curve illustrating the relationship between biomass carbon and
 54 elevation within each one year stand age bin. Significant relationships ($P \leq 0.05$) are
 55 shown in black samples, and non-significant relationships ($P > 0.05$) in grey samples.

56

57

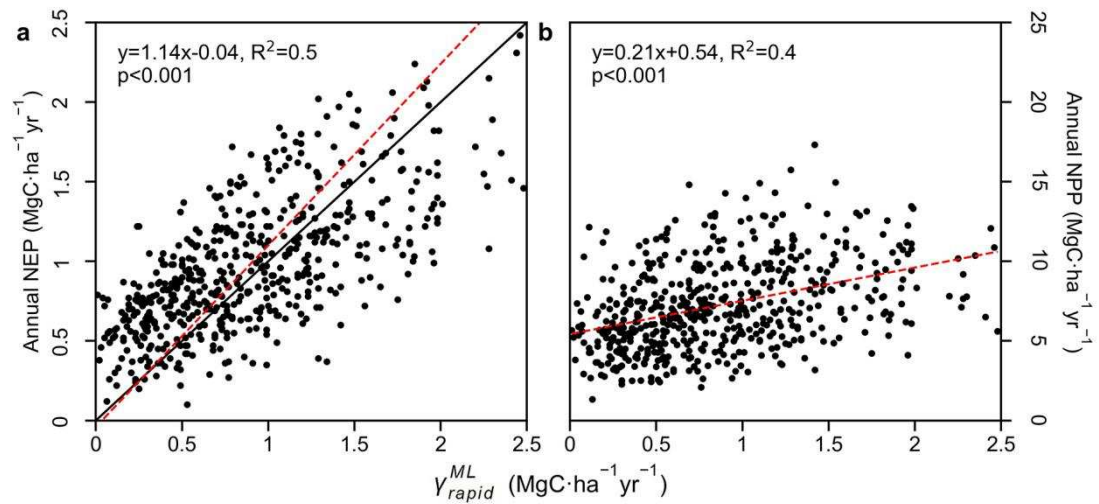


58

59 **Figure S6. The elevational patterns of ESA-CCI-derived biomass carbon in TPRFs**
 60 **with different stand age classes.** The dots represent the mean biomass carbon of all
 61 TPRF grid cells at each 100 m elevation bin; while error bars represent the
 62 corresponding one standard deviation. Green and blue curves represent the linear fitting
 63 regressions between the biomass carbon accumulation and elevation for elevation from
 64 300 to 1000 m and elevation from 1000 to 2000 m, respectively. Shading represents the
 65 95% confidence interval. Significance of linear regression is indicated in the legend as:
 66 • $P<0.5$, * $P<0.1$, ** $P<0.01$, and *** $P<0.001$.

67

68



69

70 **Figure S7. Correlation between annual BEPS-simulated NEP, NPP and γ_{rapid}^{ML} in**

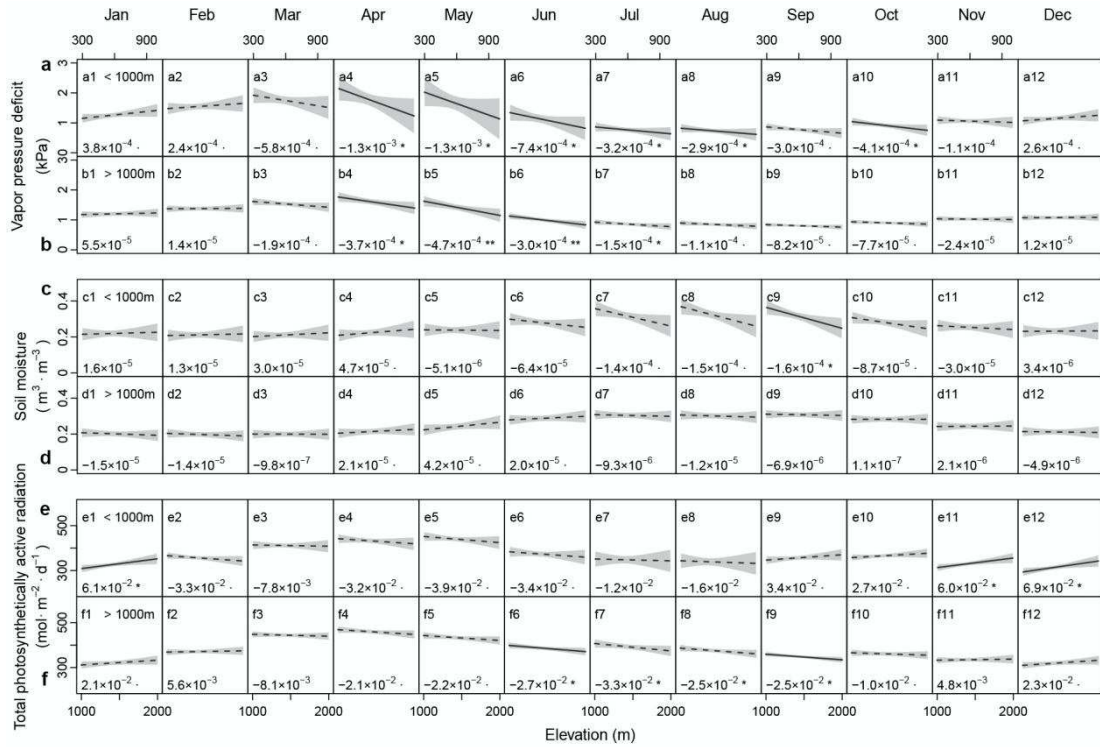
71 **TPRFs. a,** Relationship between BEPS-simulated NEP and γ_{rapid}^{ML} . The magnitudes of

72 NEP are similar as those of γ_{rapid}^{ML} . **b,** Relationship between NPP and γ_{rapid}^{ML} . The

73 magnitudes of NPP are much larger than those of carbon γ_{rapid}^{ML} . The black dashed line

74 is the 1:1 line.

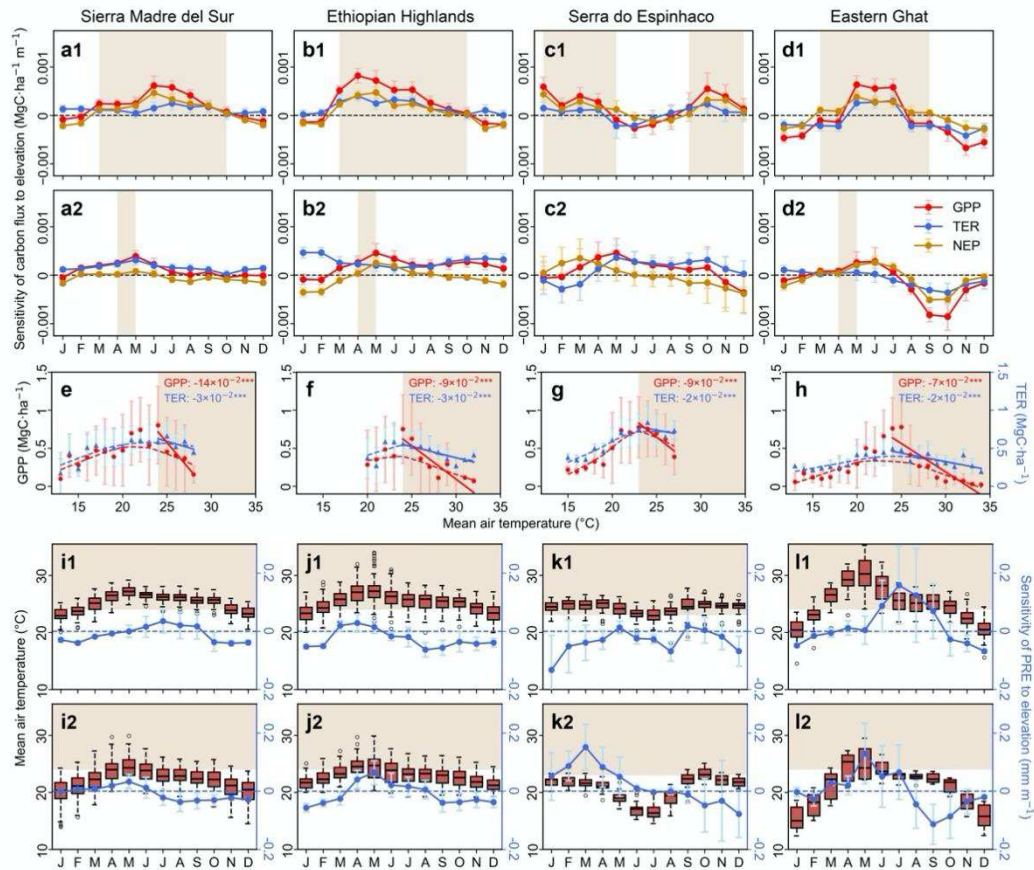
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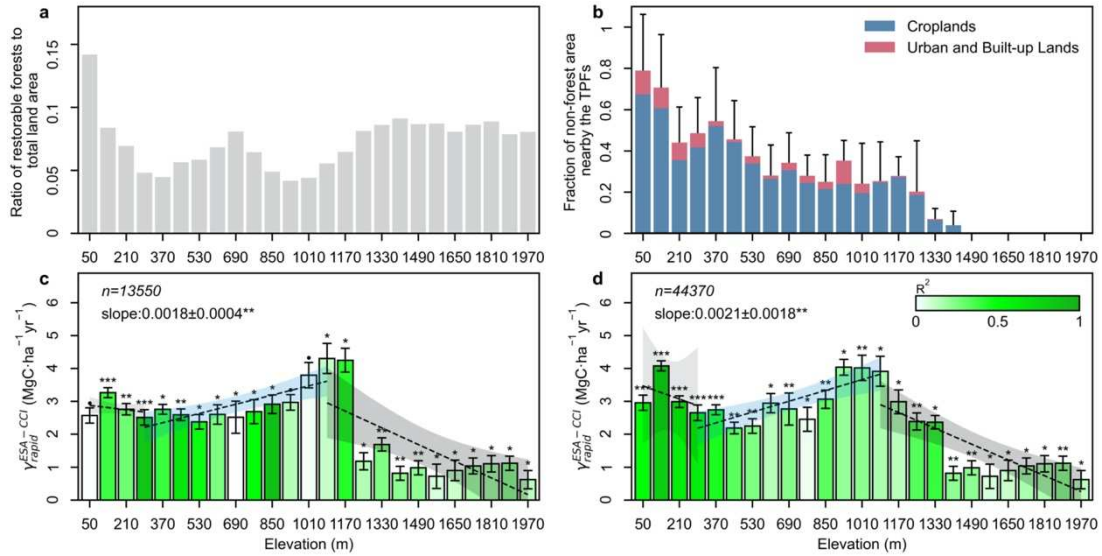
77 **Figure S8. Elevational patterns of seasonal climate factors in TPRFs.**

78 Linear regressions between monthly VPD (a-b), soil moisture (SM) (c-d) or total
 79 photosynthetically active radiation (Total PAR) (e-f) and elevation for range of 300-
 80 1000 m (a, c, e) and 1000-2000 m (b, d, f), respectively. Shadings indicate 95%
 81 confidence intervals. Significant relationships ($P \leq 0.05$) are shown in solid lines, and
 82 non-significant relationships ($P > 0.05$) in dashed lines. Significance of linear regression
 83 is indicated in the legend as: • $P < 0.5$, * $P < 0.1$, ** $P < 0.01$, and *** $P < 0.001$.



84

85 **Figure S9. Sensitivity of carbon fluxes to elevation and the climatic driving**
 86 **mechanisms in four individual mountains. a-d**, Seasonality of elevational sensitivity
 87 of GPP (red), TER (blue), NEP (orange) simulated by the BEPS model (**Dataset 4**)³⁰
 88 for elevations between 300 and 1000 m (**a1-d1**) and for elevations between 1000 and
 89 2000 m (**a2-d2**) in four individual mountains. Error bars represent one standard error.
 90 **e-h**, Changes in monthly GPP and TER with MAT in four individual mountains. Each
 91 dot denotes the median value of GPP (red) (or TER [blue]) within a 1°C MAT bin and
 92 error bars depict one standard deviation. The asterisks indicate the *P* values: **P*<0.05,
 93 ***P* < 0.01, and ****P* < 0.001. **i-l**, Seasonality of MAT and elevational sensitivity of
 94 TER for elevations between 300 and 1000 m (**i1-l1**) and for elevations between 1000
 95 and 2000 m (**i2-l2**) in four individual mountains. The upper, center, and bottomed lines
 96 in the brown boxplot indicate the first, median, and third quartiles of monthly MAT.
 97 The blue points indicate the slope of linear regressions between PRE and elevation.
 98 Error bars represent one standard error.



99

100 **Figure S10. Land use characteristics and potential influences on the elevational**

101 **pattern of $\gamma_{rapid}^{ESA-CCI}$ in TPRFs. a,** The ratio of restorable forests to total land area for

102 different elevation bin. The reforestation potential data was extracted from a global

103 reforestation potential map proposed by Griscom et al.⁵, while the total land area was

104 calculated from MODIS MCD12C1 landcover products⁶. **b,** The fraction of non-forest

105 lands (urban and croplands) nearby the TPRFs (neighbouring 10×10 1 km resolution

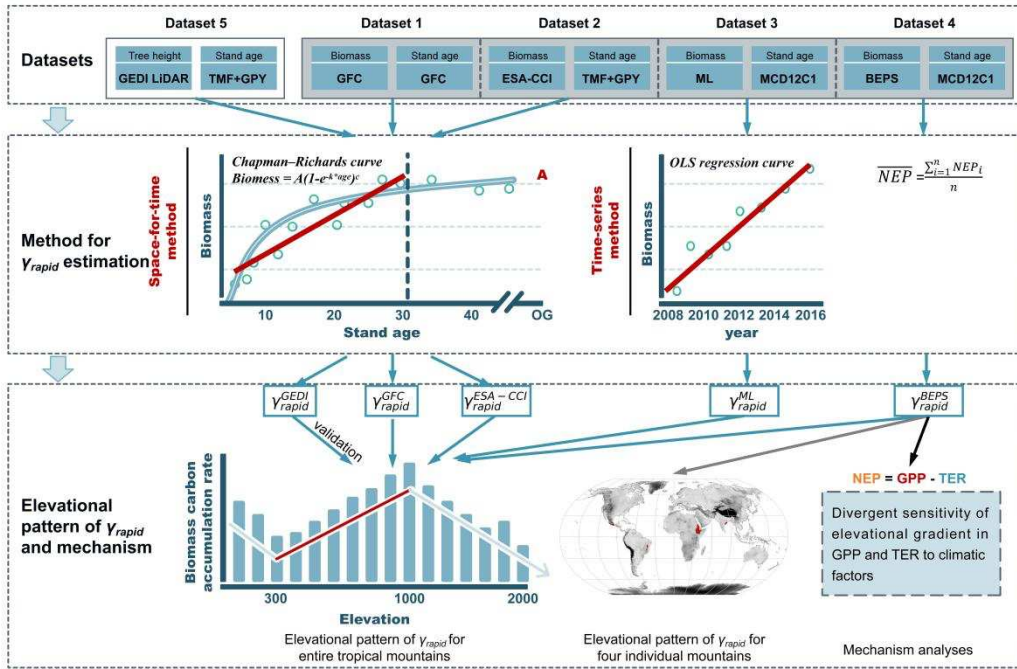
106 pixels), calculated based on MODIS MCD12C1 landcover products⁶. **c,** The elevation

107 patterns of $\gamma_{rapid}^{ESA-CCI}$ in TPRFs where their nearby fractions of non-forest lands are >

108 0.5%. **d,** The elevation patterns of $\gamma_{rapid}^{ESA-CCI}$ in TPRFs where their nearby fractions of

109 non-forest lands are < 0.5%.

110



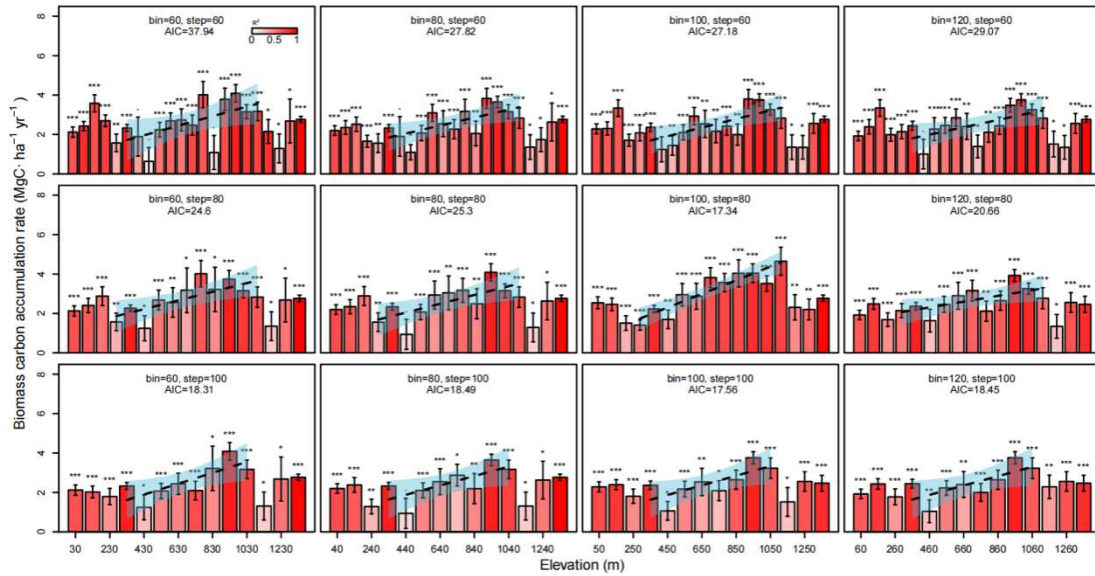
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112 **Figure S11. The flowchart illustrating the steps for analyzing the elevational**

113 **pattern of γ_{rapid} and its underlying mechanisms.**

114

115



116

117 **Figure S12. The elevation patterns of γ_{rapid}^{GFC} in TPRFs using various**

118 **combinations of elevation bin and step settings.** Each histogram represents the slope

119 of the corresponding ordinary least squares regression (OLS) curve between *in situ* tree

120 biomass of all sites of TPRFs during the rapid growth stage of trees before approaching

121 maturity (80% of the maximum biomass carbon) within the given elevation bin and

122 moving step, using the space-for-time analogy method. R^2 of OLS regression is shown

123 in a light-dark color gradient. Significance of OLS regression is indicated in the legend

124 as: • $P < 0.5$, * $P < 0.1$, ** $P < 0.01$, and *** $P < 0.001$. The error bars indicate one standard

125 error of the estimated carbon accumulation rates. The dotted lines with shading

126 represent the linear fitting curves between plant carbon accumulation rates and

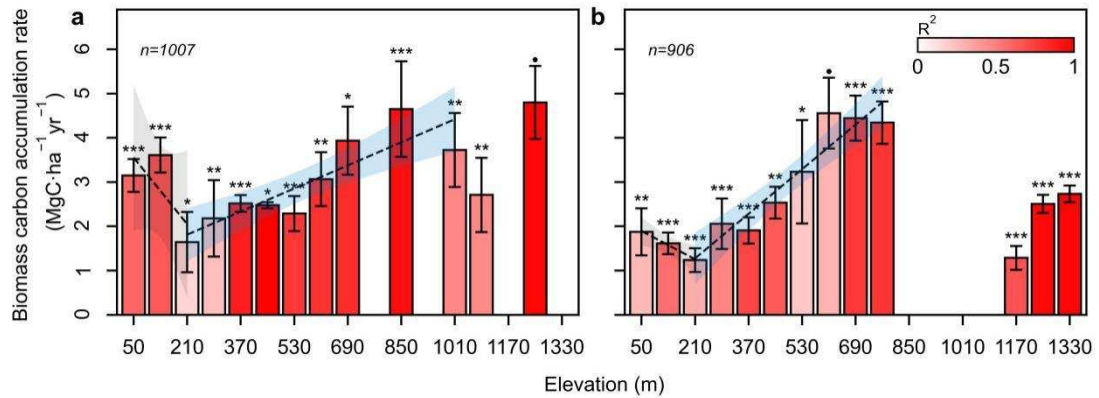
127 elevation with 95% confidence interval. Significant relationships ($P \leq 0.05$) are shown

128 in blue shading color while insignificant ones ($P > 0.05$) are displayed in grey shading.

129

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132

133 **Figure S13. The elevation patterns of γ_{rapid}^{GFC} in TPRFs which used to be different**

134 **land use types before afforestation. a,** The elevation patterns of γ_{rapid}^{GFC} in TPRFs

135 where were pasture lands before afforestation. **b,** The elevation γ_{rapid}^{GFC} in TPRFs

136 where were shifting cultivation before afforestation. Each histogram represents the

137 slope of the corresponding ordinary least squares regression (OLS) curve between *in*

138 *situ* tree biomass of all sites of TPFs during the rapid growth stage of trees before

139 approaching maturity (80% of the maximum biomass carbon) within each elevation bin

140 ($100 \pm 50m$ in 80m step) using the space-for-time analogy method. R^2 of OLS regression

141 is shown in a light-dark color gradient. Significance of OLS regression is indicated in

142 the legend as: • $P < 0.5$, * $P < 0.1$, ** $P < 0.01$, and *** $P < 0.001$. The error bars indicate one

143 standard error of the estimated carbon accumulation rates. The dashed lines with

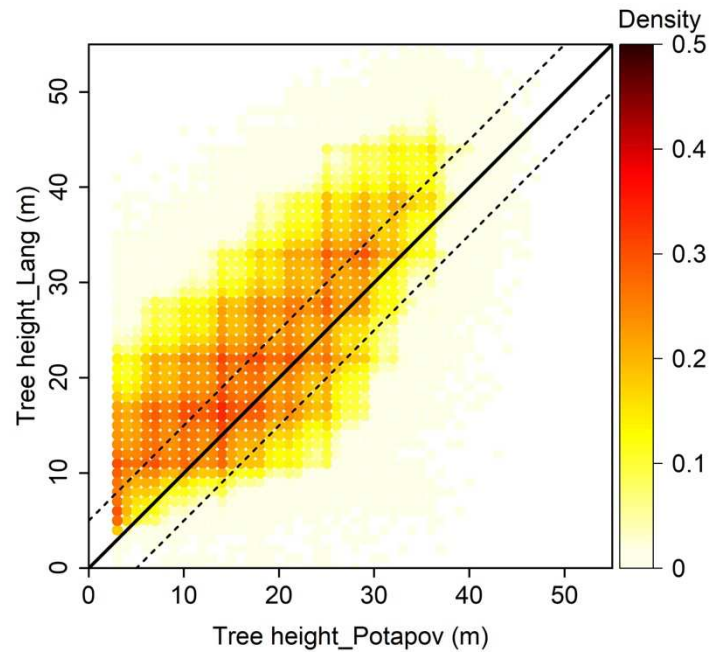
144 shading represent the linear fitting curves between plant carbon accumulation rates and

145 elevation with 95% confidence interval. Significant relationships ($P \leq 0.05$) are shown

146 in blue shading color while insignificant ones ($P > 0.05$) are displayed in grey shading.

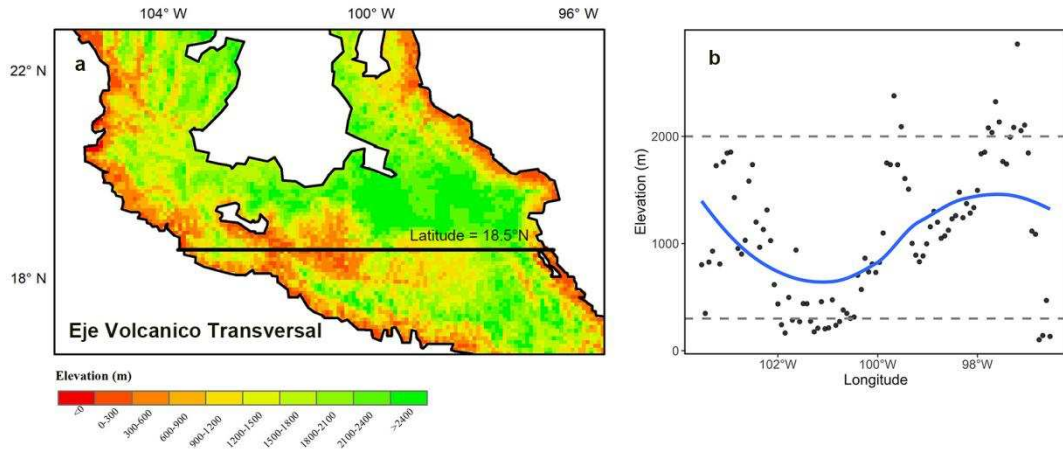
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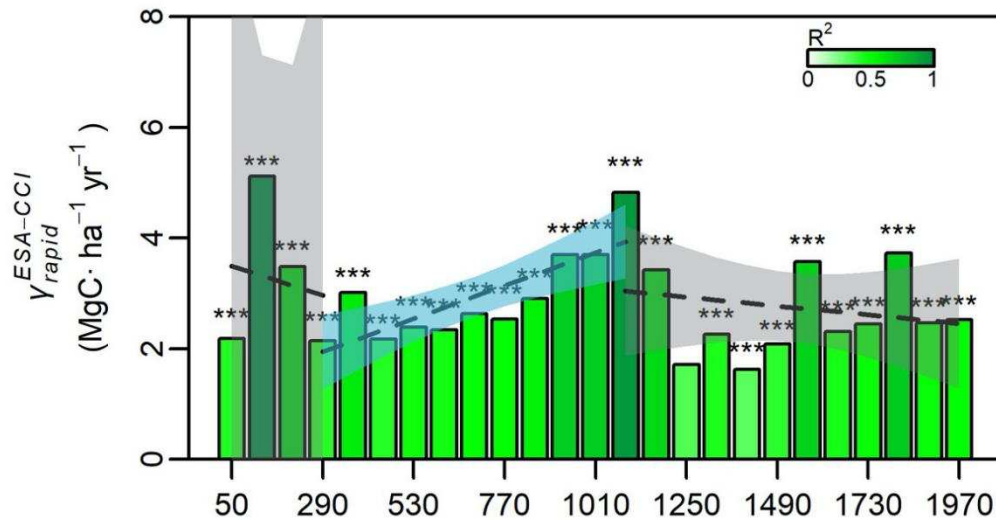
150 **Figure S14. Selection of robust tree height data for estimating the tree height**
 151 **growth rate.** We only selected those pixels where Potapov et al. (2021)'s³ and Lang et
 152 al. (2023)'s⁷ tree height data are highly consistent (± 5 m) (**Methods**).
 153



154

155 **Figure S15. Topography variations (10 km resolution) of the Eje Volcanico**
 156 **Transversal mountain range in North America. a, DEM map at a 10 km resolution.**
 157 **b, Elevation data along the black line (Latitude:18.5°N; Longitude: 104°W ~ 96°W) on**
 158 **the DEM map extracted from this 10 km resolution DEM for Eje Volcanico Transversal**
 159 **mountain range.**

160

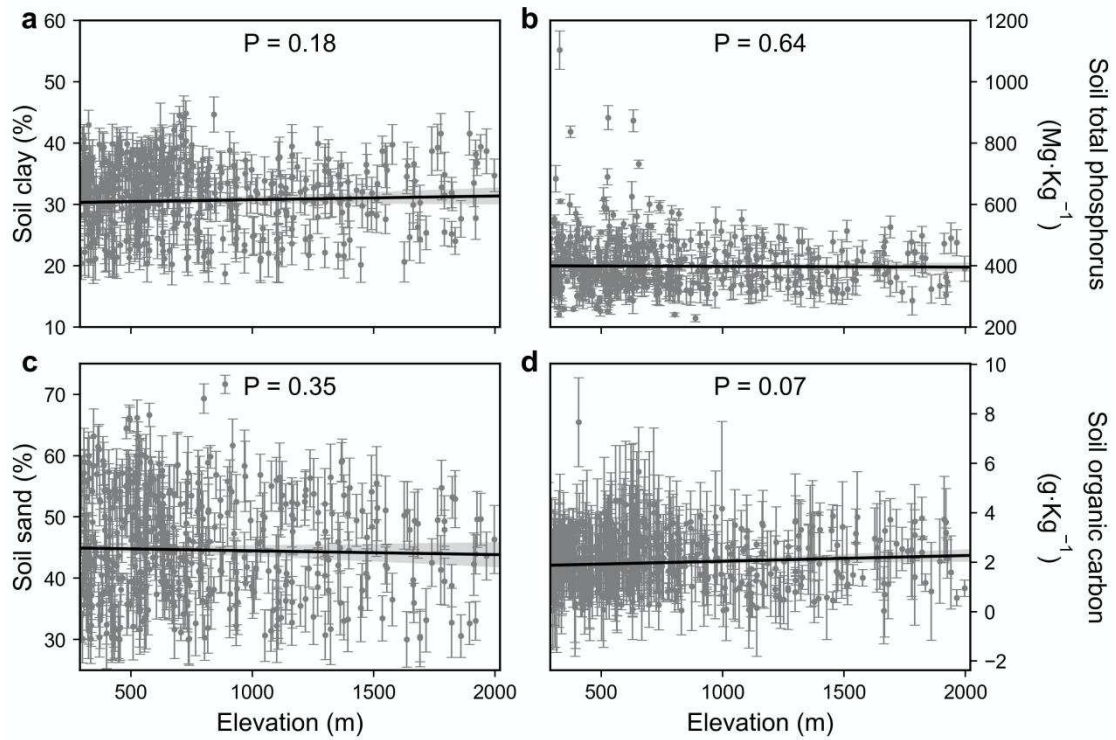


161

162 **Figure S16. The elevation patterns of biomass carbon accumulation rates in TPFs**
 163 **based on adjusted ESA-CCI data.** Each histogram represents the slope of
 164 corresponding ordinary least squares regression (OLS) curve between satellite-based
 165 tree biomass carbon and stand age of all pixels of TPFs within the given elevation bin
 166 (100 ± 50 m in 80 m step), using the space-for-time analogy method (**Methods**). R^2 of
 167 OLS regression are shown in a light-dark color gradient. The error bars indicate one
 168 standard error. Significance of OLS regression is indicated in the legend as: • $P < 0.5$,
 169 * $P < 0.1$, ** $P < 0.01$, and *** $P < 0.001$. The error bars indicate one standard error of the
 170 estimated carbon accumulation rates. The dashed lines with shading represent the linear
 171 fitting curves between plant carbon accumulation rates and elevation with 95%
 172 confidence interval. Significant relationships ($P \leq 0.05$) are shown in blue shading color
 173 while insignificant ones ($P > 0.05$) are displayed in grey shading.

174

175



176

177 **Figure S17. The elevation patterns of soil fertility in TPRFs.** The elevation patterns
 178 of soil organic carbon⁸, total phosphorus⁹, sand¹⁰ and clay¹¹ concentrations, respectively.
 179 Each dot represents the mean values of corresponding soil nutrient concentration of all
 180 soil layers at each TPRF site. Error bars represent one standard error. The solid lines
 181 represent the linear fitting curves between soil fertility and elevation. Shading
 182 represents the 95% confidence interval.

183

184

185 **Table S1.** Information on the data used in this study.

Name	Parameters	Spatial resolution	Temporal resolution	Reference	Applications
Field observation sites	Plant carbon, stand age and disturbance type	Multiple	Multiple	Cook-Patton et al., 2020 ⁴ , Anderson-Teixeira et al., 2018 ¹²	To calculate carbon accumulation rate
ESA-CCI forest above-ground biomass product	AGB	100 m	Yearly	Santoro & Cartus, 2021 ¹³	To extract forest AGB
Global vegetation live biomass	AGB	0.1°	Yearly	Xu et al., 2021 ¹⁴	To extract forest AGB
MCD12C1 v061	Land Cover	0.05°	Yearly	Friedl & Sulla-Menashe, 2022 ⁶	To identify forest regrowth period
Terra Climate	MAT, VPD, PRE and SM	1/24°, ~4- km	Monthly	Abatzoglou et al., 2018 ¹⁵	To extract environmental variables
BESS PAR	Total PAR	5km	Daily	Ryu et al., 2018 ¹⁶	To extract total PAR variable
Global sensing-based surface soil moisture (RSSSM) dataset	Surface soil moisture	0.1°, ~10 km	~10 days	Chen et al., 2021 ¹⁷	To extract soil moisture variable
Global carbon flux	GPP, TER,	0.07272727	Daily	Chen et al.,	To extract

(GPP/NPP/NEP) simulation product	NEP and NPP	° ~10 km			2019 ¹⁸	carbon flux variables
Tropical moist forests	forest cover change	30 m	Yearly	Vancutsem et al., 2021 ¹⁹		To calculate forest age
FLUXCOM data	NEP	0.5°	Yearly	Jung et al., 2017 ¹		To extract NEP
Tree height	tree height	10m	_	Lang et al., 2023 ⁷		To extract tree height
Tree height	tree height	30m	-	Potapov et al., 2021 ³		To extract tree height
Global reforestation potential map	reforestation potential	1km		Griscom et al., 2017 ⁵		To extract reforestation potential area
Global soil phosphorus concentration dataset	total phosphorus	0.05°	-	He et al., 2021 ⁹		To extract soil fertility variables
Global soil organic carbon dataset	soil organic carbon	250m	-	Hengl & Wheeler, 2018 ⁸		To extract soil fertility variables
Global soil sandy content dataset	soil sandy concentration	250m	-	Hengl et al., 2018 ¹⁰		To extract soil fertility variables
Global soil clay content dataset	soil clay concentration	250m	-	Hengl et al., 2018 ¹¹		To extract soil fertility variables
RAINBIO database	mega tree species	Multiple	Multiple	Dauby et al., 2016 ²		To extract tree species information

SRTM DEM	elevation	30m	-	Van Zyl et al., 2001 ²⁰	To extract elevation
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