

# Multi-fold increase in rainforests tipping risk beyond 1.5-2°C warming

Chandrakant Singh<sup>1,2,3,\*</sup>, Ruud van der Ent<sup>4</sup>, Ingo Fetzer<sup>1,2,5</sup>, Lan Wang-Erlandsson<sup>1,2,5</sup>

<sup>1</sup>Stockholm Resilience Centre, Stockholm University, Stockholm, Sweden

<sup>2</sup>Bolin Centre for Climate Research, Stockholm University, Stockholm, Sweden

<sup>3</sup>Department of Space, Earth and Environment, Chalmers University of Technology, Gothenburg, Sweden

<sup>4</sup>Department of Water Management, Faculty of Civil Engineering and Geosciences, Delft University of Technology, Delft, The Netherlands

<sup>5</sup>Potsdam Institute for Climate Impact Research, Potsdam, Germany

\*Corresponding author; E-mail: [chandrakant.singh@su.se](mailto:chandrakant.singh@su.se), [chandrakant.singh@chalmers.se](mailto:chandrakant.singh@chalmers.se)

## ORCID

Chandrakant Singh: <http://orcid.org/0000-0001-9092-1855>

Ruud van der Ent: <https://orcid.org/0000-0001-5450-4333>

Ingo Fetzer: <http://orcid.org/0000-0001-7335-5679>

Lan Wang-Erlandsson: <http://orcid.org/0000-0002-7739-5069>

**Abstract.** Tropical rainforests rely on their root systems to access moisture stored in soil during wet periods for use during dry periods. When this root-zone soil moisture is inadequate to sustain a forest ecosystem, they transition to a savanna-like state, losing their native structure and functions. Yet the influence of climate change on ecosystem's root-zone soil moisture storage and their impact on rainforest ecosystems remain uncertain. This study assesses the future state of rainforests and the risk of forest-to-savanna transitions in South America and Africa under four shared socioeconomic pathways (SSP1-2.6, SSP2-4.5, SSP3-7.0, and SSP5-8.5). Using a mass-balance-based empirical understanding of root zone storage capacity ( $S_r$ ), defined as the maximum volume of root zone soil moisture per unit area accessible to vegetation's roots for transpiration, we project how rainforest ecosystems will respond to future climate changes. We find that under the end-of-the-21<sup>st</sup>-century climate, nearly one-third of the total forest area will be influenced by climate change. As the climate warms, forests will require a larger  $S_r$  than they do under the current climate to sustain their ecosystem structure and functions, making them more water-limited. Meanwhile, recovering to a less water-limited state gradually diminishes. Furthermore, warming beyond 1.5-2°C will significantly elevate the risk of a forest-savanna transition. In the Amazon, the forest area at risk of such a transition grows by about 1.7-5.8 times in size compared to their immediate lower warming scenario (e.g., SSP2-4.5 compared to SSP1-2.6). In contrast, the risk growth in the Congo is less substantial, ranging from 0.7-1.7 times. These insights underscore the urgent need to limit global surface temperature rise below the Paris Agreement to conserve rainforest ecosystems and associated ecosystem services.

## 37 **1 Introduction**

38 Tropical rainforests in the Amazon and Congo basins are critical to the Earth system since they store and  
39 sequester a large amount of carbon, host vast biodiversity, and regulate the global water cycle (Malhi et al.,  
40 2014). However, these forests are under severe pressure from climate change and land-use change (Davidson  
41 et al., 2012; Lewis et al., 2015; Malhi et al., 2008), which risk amplifying further warming and forest degradation  
42 (Lawrence et al., 2022). Climate change and land-use change lead to a decrease in precipitation, an increase in  
43 seasonality and atmospheric water demand (Malhi et al., 2014). This causes a deficit in soil moisture availability  
44 that inhibits plant growth (Singh et al., 2020; Wang-Erlandsson et al., 2022). Furthermore, climate-induced  
45 hydroclimatic changes, including the projected increases in drought frequency, severity, and duration (Dai,  
46 2011; Liu et al., 2018), present imminent threats to the capacity of rainforests to maintain their native ecological  
47 structure and functions (i.e., forest resilience) (Bauman et al., 2022; Grimm et al., 2013; Jones et al., 2009).

48 Under water-deficit conditions, rainforests adapt by investing in their root systems to gain better access  
49 to soil moisture necessary to maintain their structure and functions (Singh et al., 2020, 2022). At the same time,  
50 the availability of surplus moisture at shallow depths minimises the need for ecosystems to invest in extensive  
51 (deeper and lateral) root systems (Bruno et al., 2006). Furthermore, forest ecosystems adapt to climate change  
52 by optimising water distribution through mechanisms such as hydraulic redistribution (Liu et al., 2020; Oliveira  
53 et al., 2005), enhancing water-use efficiency by regulating stomatal conductance, and even shredding leaves  
54 (Wolfe et al., 2016) to minimise moisture loss (Barros et al., 2019; Brum et al., 2019; Lammertsma et al., 2011).  
55 Despite their critical role, the dynamic influence of climate change on vegetation's rooting structure and subsoil  
56 moisture is challenging to measure at the ecosystem scale (Fan et al., 2017). Thus, understanding how moisture  
57 from wet periods is stored, transmitted, and lost from soil, and how it is accessed by vegetation during dry  
58 periods, is critical to the ecohydrology and resilience of terrestrial ecosystems under climate change.

59 However, such ecohydrological dynamics remain challenging to incorporate in Earth System Models  
60 (ESMs) (Lenton, 2011; Maslin and Austin, 2012; Valdes, 2011) – complex mathematical representations of Earth  
61 system processes and interactions across different biospheres. This limits ESM's capacity to simulate tipping  
62 points as an emergent property of the system (i.e., properties that emerge due to multiple interactions between  
63 several system components, and are not the property of an individual component) (Hirota et al., 2021; Reyer  
64 et al., 2015). This constraint is mainly due to our poor understanding of complex mechanisms governing the  
65 ecosystem, which are not well represented in ESMs. This includes a limited understanding of vegetation-climate  
66 feedbacks (Boulton et al., 2013, 2017; Chai et al., 2021), subsoil moisture availability (Cheng et al., 2017),  
67 adaptation dynamics (Yuan et al., 2022), the response time of forest ecosystems to climate change  
68 perturbations, and assumptions about future (i.e., prescribed) land-use change (Hurtt et al., 2020) in the ESMs.  
69 Furthermore, in the Earth system, some interactions still remain largely unknown, thereby making the  
70 prediction of (abrupt) forest-to-savanna transition (referring to changes in the dense-canopy structure of

71 forests to one that mimics an open-canopy structure similar to savanna; hereafter referred to as forest-savanna  
72 transition) challenging (Drijfhout et al., 2015; Hall et al., 2019; Koch et al., 2021).

73 To understand the extent of rainforest tipping risks, there is a need to assess and contrast the forest  
74 resilience consequences of low-emission and current commitment trajectories with the more commonly used  
75 high-emission scenario (Jehn et al., 2022). However, the risk of forest-savanna transitions under various possible  
76 climate future scenarios is relatively under-investigated. As a result of the conflicting findings and scenario-  
77 dependent uncertainties, the Intergovernmental Panel on Climate Change (IPCC) has only low confidence about  
78 the possible tipping of the Amazon forest by the end of the 21<sup>st</sup> century (Canadell et al., 2021). However, with  
79 mounting empirical evidence on how climate change influences rainforest ecosystems (Boulton et al., 2022;  
80 Küçük et al., 2022; Singh et al., 2020, 2022), the research on rainforest resilience loss has accelerated  
81 considerably in the recent decade (Ahlström et al., 2017; Huntingford et al., 2013). Yet, forest resilience is often  
82 assessed based on changes in forest carbon stocks (Huntingford et al., 2013; Parry et al., 2022) or precipitation  
83 (Hirota et al., 2011; Staal et al., 2020; Zemp et al., 2017); and rarely on the subsoil moisture availability of the  
84 ecosystem (Singh et al., 2022).

85 This study aims to assess the state of rainforests and the risk of a forest-savanna transition under the end  
86 of the 21<sup>st</sup>-century climate based on an empirical understanding of ecosystems' root zone storage dynamics.  
87 For this, we use mass-balance derived root zone storage capacity ( $S_r$ ) – representing the maximum amount of  
88 soil moisture vegetation can access for transpiration (Gao et al., 2014; Singh et al., 2020; Wang-Erlandsson et  
89 al., 2016). Our use of  $S_r$  is grounded in its effectiveness in representing ecosystems' access to soil moisture and  
90 their ability to modify above-ground structures accordingly (de Boer-Euser et al., 2016; Singh et al., 2020;  
91 Stocker et al., 2023; Wang-Erlandsson et al., 2016). It should be noted that we refer to rainforest tipping as a  
92 forest-savanna transition 'risk' since the timing of such transitions depends on the stochastic fluctuations of  
93 other environmental factors, beyond just hydroclimate (e.g., fire, human influence, species composition) (Cole  
94 et al., 2014; Cooper et al., 2020; Higgins and Scheiter, 2012; Poorter et al., 2016). Therefore, to project if an  
95 ecosystem is a forest or has tipped to savanna in the future, we assume the hydroclimate projected by the end  
96 of the 21<sup>st</sup> century (i.e., 2086-2100) and ecosystem are in equilibrium. However, we do not account for the time  
97 required for ecosystems to reach their (long-term) equilibrium state, which previous studies suggest can take  
98 between 50-200 years after crossing the tipping point (Armstrong McKay et al., 2022).

99

## 100 **2 Methodology**

### 101 **2.1 Study Area**

102 This study focuses on forest ecosystems (i.e., excluding savanna/grassland and vegetation in human-influenced  
103 ecosystems) extending between 15°N–35°S for South America and Africa.

104

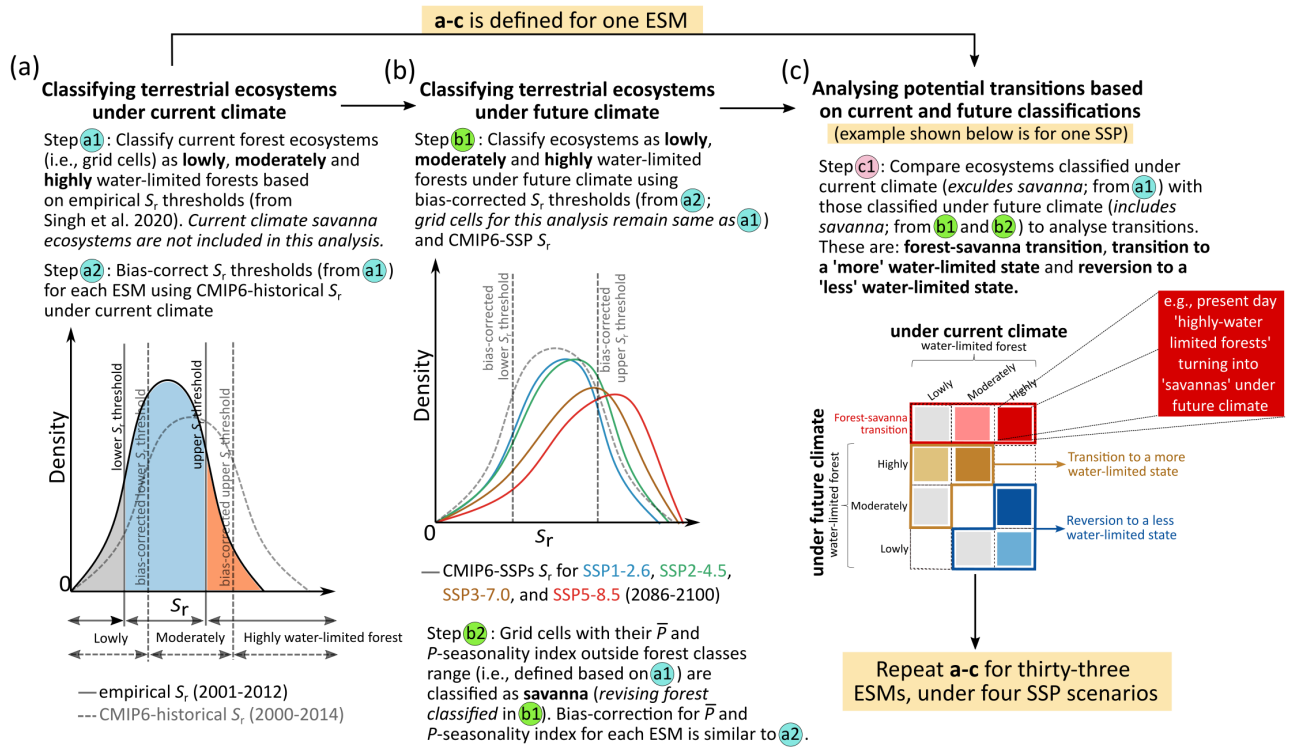
## 105 2.2 Data

106 This analysis uses both empirical and ESM-simulated datasets of precipitation and evaporation. Empirical  
107 datasets include remotely sensed and observation-corrected precipitation and evaporation time-series.  
108 Empirical precipitation estimates at daily timestep are obtained from the Climate Hazards Group InfraRed  
109 Precipitation with Station data (CHIRPS; 0.25° resolution) (Funk et al., 2015). Furthermore, empirical  
110 evaporation is derived using an equally-weighted ensemble of three different datasets – (i) Breathing Earth  
111 System Simulator (BESS; 0.5° resolution) (Jiang and Ryu, 2016) (ii) Penman-Monteith-Leuning (PML; 0.5°  
112 resolution) (Zhang et al., 2016) and (iii) FLUXCOM-RS (0.083° resolution) (Jung et al., 2019) – at monthly  
113 timestep. Here, evaporation represents the sum of all evaporated moisture from the soil, open water and  
114 vegetation, including interception and transpiration. We only selected evaporation datasets free from biome-  
115 dependent parameterisation (such as plant function types, stomatal conductance, and maximum root  
116 allocation depth) and soil layer depth (represents maximum depth of moisture uptake). Ultimately, all  
117 evaporation datasets are bilinearly interpolated to 0.25° resolution and downscaled to daily timestep using  
118 ERA5 evaporation (0.25° resolution) estimates (Hersbach et al., 2020). All empirical datasets are obtained for  
119 2001-2012.

120 We also obtained precipitation and evaporation estimates from 33 ESMs (from 22 different institutes),  
121 which includes CMIP6-historical and four SSP scenario simulations (SSP1-2.6 leads to approx. 1.3-2.4°C  
122 warming; SSP2-4.5 corresponds to 2.1-3.5°C warming and is closest to the current trajectory according to the  
123 nationally determined contributions (Anon, 2015); SSP3-7.0 around 2.8-4.6°C warming; and SSP5-8.5  
124 represents 3.3-5.7°C warming; °C warming represents an increase in mean global surface temperature change  
125 by the end of 21<sup>st</sup> century relative to 1850-1900 (IPCC, 2021) (Fig. 1; Table S1 and S2). The historical estimates  
126 are obtained at a monthly timestep for 2000-2014, and the estimates under different SSPs are obtained for  
127 2086-2100. Though obtained estimates from different ESMs are at different spatial resolutions, we bilinearly  
128 interpolated them to 0.25° for this analysis.

129 Finally, to minimise the influence of human activity and non-forest land cover on the natural water cycle,  
130 we utilised land-cover data to remove pixels with such features from our analysis. We began by removing  
131 human-influenced and non-forest land cover, such as savanna, grasslands, and water bodies, from Globcover,  
132 a global land-cover classification dataset by the European Space Agency (ESA) at 300m resolution (GlobCover  
133 land-use map, 2022). We then performed majority interpolation to convert the dataset to a 0.25° resolution  
134 and to mask grid cells with less than 50% forest cover. This step ensured that only grid cells with over 50% forest  
135 cover were classified as forests for further analysis.

136



137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

160

161

**Figure 1: Methodological framework for analysing the potential transitions in tropical terrestrial ecosystems using empirical and CMIP6-Earth System Models (ESMs) hydroclimate estimates.** (a) We use root zone storage capacity ( $S_r$ )-based classification thresholds (obtained from Singh et al., 2020) – calculated using empirical precipitation ( $P$ ) and evaporation ( $E$ ) estimates (Fig. S1; see Methodology section and Appendix A1) – to classify terrestrial ecosystems under the current climate. Savanna ecosystems under the current climate are excluded from this analysis. We bias-correct these  $S_r$  thresholds for all ESMs using the histogram equivalence method (Piani et al., 2010) (Table S1). (b) We then use these bias-corrected  $S_r$  thresholds to classify ecosystems under future climate conditions (Fig. S2 and S3). Furthermore, we use mean annual precipitation ( $\bar{P}$ ) and  $P$ -seasonality index range ( $S_r$ -based forest classes from a) – as a proxy for ecosystem state – to revise our classification under future climate (Appendix A3 and Fig. S4). (c) We then analyse the potential transitions by comparing ecosystems classified under the current climate (analysed in a) with those classified under future climate (analysed in b) individually for all ESMs (Fig. S5 and S6). The transition analysis assumes that the vegetation and hydroclimate are in equilibrium, and does not account for the time required for transitions to occur. A detailed description is provided in the Methodology section. An exemplification of this methodological framework is shown in Fig. S7.

### 2.3 Root zone storage capacity-based framework for projecting forest transitions

Vegetation uptakes soil moisture from its roots; thus, the availability of root zone moisture is a key element that mediates the interaction between vegetation and climate (Brooks et al., 2015; Küçük et al., 2022; Rosas et al., 2019; Wang-Erlandsson et al., 2022). However, measuring soil- (such as texture and porosity) and root-characteristics (such as vertical and lateral extent and soil moisture uptake profiles) that influence access to subsoil moisture are challenging to measure at ecosystem scales (Bruno et al., 2006). Furthermore, land-system models tend to oversimplify the transfer and storage of water in root-zone due to insufficient knowledge about soil-vegetation-climate interactions (Albasha et al., 2015; Hildebrandt et al., 2016; Wang et al., 2004). In such cases, the mass-balance approach-based  $S_r$  provides a tangible and comprehensive understanding of ecosystem

162 access to moisture stored in the soil (de Boer-Euser et al., 2016; Gao et al., 2014; McCormick et al., 2021; Stocker  
163 et al., 2023).

164

### 165 **2.3.1 Estimating mass-balance derived root zone storage capacity ( $S_r$ )**

166 Derived using the mass-balance approach,  $S_r$  represents the maximum amount of soil moisture accessed by  
167 vegetation for transpiration (Singh et al., 2020; Wang-Erlandsson et al., 2016). This methodology calculates the  
168 maximum extent of soil moisture within the reach of plant roots, assuming that ecosystems do not invest in  
169 expanding their root-zone storage beyond what is necessary to bridge the maximum (accumulated) water-  
170 deficit experienced by the vegetation during dry periods (i.e., periods in which evaporation is greater than  
171 rainfall, irrespective of the seasons). This maximum annual accumulated water deficit ( $D_{a,y}$ ) experienced by the  
172 ecosystem is calculated using daily precipitation and evaporation estimates (Appendix A1 and Fig. A1). Subsoil  
173 moisture beyond the reach of plant roots is primarily controlled by gravity-induced gradients (de Boer-Euser et  
174 al., 2016) and is not available for transpiration. The rationale is that any extensive investment (i.e., more than  
175 necessary) in root expansion would require carbon allocation and, thus, is inefficient from the perspective of  
176 the plants (Gao et al., 2014; Schenk, 2008). Since, this approach does not rely on prior information about  
177 vegetation, soil, or land cover-based, by using empirical (observation-based) datasets (Appendix A1 and Fig.  
178 A1), we capture the dynamics of actual soil moisture available for the ecosystems (Wang-Erlandsson et al.,  
179 2016). The detailed methodology for calculating  $S_r$  using precipitation and evaporation estimates is outlined in  
180 Appendix A1.

181 In this mass-balance approach,  $S_r$  only represents a hydrological buffer essential for maintaining the  
182 ecosystem's structure and functions (Gao et al., 2014; Wang-Erlandsson et al., 2016). However, other biotic and  
183 abiotic factors, such as root morphology, soil depth, and geological formations, can physically restrict  $S_r$  by  
184 limiting rooting depth, rooting structure, and the soil's water-holding capacity (Canadell et al., 1996; Jackson et  
185 al., 1996; Schenk and Jackson, 2002) (Appendix A2). Additionally, soil properties like porosity or field capacity  
186 could necessitate a deeper rooting strategy in different soil types (e.g., between sandy and clayey soil) to  
187 achieve a comparable level of  $S_r$  to sustain the ecosystem under future climate (Kukal and Irmak, 2023).  
188 However, this study assesses the impact of future climate change on the ecosystem's hydrological regime,  
189 focusing on the changes to the ecosystem's equilibrium state. Therefore, the direct influence of soil and root  
190 characteristics under future climate change on  $S_r$  (Appendix A2) and forest transitions falls outside our current  
191 scope.

192

### 193 2.3.2 Determining root zone storage capacity thresholds for forest transitions

194 A recent study by Singh et al. (2020) demonstrated that  $S_r$  effectively represents an ecosystem's above-ground  
195 state (i.e., whether it is a forest or savanna) and its level of water–stress, based on root-zone moisture  
196 availability. In this study, we refine their terminology from 'water-stressed state' to 'water-limited state' to more  
197 precisely describe the effects of hydroclimatic conditions on forest ecosystems, specifically in terms of inhibiting  
198 plant growth based on subsoil moisture availability and the potential of them approaching the threshold of  
199 forest-savanna transition. According to Singh et al. (2020), in response to water-limited conditions, forests  
200 adapt their rooting strategies and modify their above-ground forest cover. These adaptations aim to allocate  
201 carbon in the most efficient way possible to maximise the hydrological benefits available to the ecosystem.  
202 They classified terrestrial ecosystems into four distinct categories based on the relationship between tree cover  
203 density and root zone storage capacity ( $S_r$ ), illustrating the various drought coping strategies of ecosystems  
204 (detailed description provided in Singh et al., 2020):

- 205 i. **Lowly water-limited forest:** Dense forests (>70% tree cover) that receive ample rainfall (with daily  
206 precipitation exceeding evaporation year-round) results in a very low  $D_{a,y}$ . In such an environment, the  
207 top layer of the soil remains consistently damp, allowing for efficient soil moisture uptake through  
208 shallow roots (<1m;  $S_r$  and maximum rooting depth comparison in Singh et al., 2020), as vegetation  
209 typically utilises the shortest available pathway for moisture uptake (Bruno et al., 2006). Consequently,  
210 these forest ecosystems can sustain themselves with a low  $S_r$  (<100 mm).
- 211 ii. **Moderately water-limited forest:** Although these forests retain a dense structure (>65% tree cover),  
212 the increased precipitation seasonality (evaporation rates remain same as before) leads to a relatively  
213 higher  $D_{a,y}$ . This necessitates a greater investment in their rooting systems to access subsoil moisture  
214 for dry periods, with  $S_r$  for these ecosystems ranging between 100-400 mm in South America and 100-  
215 350 mm in Africa. Notably, this enhanced below-ground investment does not compromise the above-  
216 ground ecosystem structure.
- 217 iii. **Highly water-limited forest:** With further increase in precipitation seasonality (even negligible  
218 precipitation during dry seasons) and duration of dry period, forests need to maximize their  $S_r$   
219 (maximum rooting depths typically between 15-20m). Maintaining ecosystems under these conditions  
220 is costly from a subsoil investment perspective (Schenk, 2008), with regions in South America and Africa  
221 showing  $S_r$  values as high as 750 mm and 450 mm, respectively. Consequently, these values represent  
222 the upper limits beyond which forest ecosystems cannot further enhance their  $S_r$ .

223 Possible mechanisms suggest that these trees adapt by shedding leaves to minimise moisture  
224 loss (Wolfe et al., 2016). However, this adaptation can reduce photosynthetic activity, leading to  
225 declines in root growth, and heightening the risk of mortality from hydraulic failures due to the  
226 unavailability of soil moisture at accessible depths (Guswa, 2008). Furthermore, accumulation of dry

227 leaves also perpetuates forest fires, thinning the ecosystem even further (tree cover can drop as low as  
228 30%) (Nepstad et al., 1999). Although increased tree mortality reduces competition for water, enabling  
229 some trees to survive, the heightened risk of hydraulic failures and forest fires makes these ecosystems  
230 highly susceptible to transitioning to savanna.

231 iv. **Savanna-grassland regime** (hereafter referred to as **savanna**): These ecosystems, typically  
232 characterised by an open, grass-dominated structure (tree cover <40%), have both a lower water  
233 availability and demand (both precipitation and evaporation are lower than in forest ecosystems). Thus,  
234 requiring a lower hydrological buffer to sustain their structure and functions. For these ecosystems,  $S_r$   
235 values can be as low as 100 mm. Although tree species in this ecosystem can develop deep roots  
236 (extending up to 20m), the majority of the root biomass is concentrated in the shallow soil layers (top  
237 30–50 cm; shallow water uptake profile) (Nippert and Holdo, 2015; Schenk, 2008). This strategy allows  
238 for complete moisture uptake between trees and grass species. This also suggests that, for savanna,  
239 deeper roots don't always necessitate a high  $S_r$ .

240

241 The difference in  $S_r$  thresholds between both continents is due to the presence of water-use-efficient  
242 C4 grasses in Africa (Still et al., 2003), which reduces the competitiveness for moisture uptake between tree  
243 species and grasses – leading to a lesser need for extensive  $S_r$  in the African forest ecosystem (Singh et al., 2020).  
244 Furthermore, these adaptation dynamics align with the alternative stable state theory (i.e., forest's stabilising  
245 feedback under hydroclimatic changes and tipping risk beyond certain hydroclimatic extremes) (Hirota et al.,  
246 2011), which makes  $S_r$  more representative of the transient state of the ecosystem than precipitation (Singh et  
247 al., 2022). We, thus, use these mass-balance derived  $S_r$  thresholds to project rainforest transitions and tipping  
248 risk under future climate change. A detailed description of how previous studies have projected rainforest  
249 tipping (Table S3), and how  $S_r$ -based framework builds upon their shortcomings is mentioned in the  
250 Supplement.

251

### 252 **2.3.3 Projecting forest transitions under future climate change**

253 To project forest transitions under future climate, we have to first classify forests based on  $S_r$  thresholds under  
254 the current and future climate. Based on this classification, we analyse potential transition for each ESM and  
255 aggregate the results (Fig. 1). We start by classifying forests under current climate following the approach by  
256 Singh et al. (2020), which uses the (empirical) daily estimates of CHIRPS precipitation and ensemble evaporation  
257 (2001-2012) (Appendix A1 and Sect. 2.3.2) (Fig. 1a). Since we are only interested in forest transitions, the  
258 ecosystems classified as savanna under the current climate are excluded from this analysis.

259 Next, for classifying ecosystems under future climate scenarios, we follow the same mass-balance  
260 approach (Appendix A1). However, since precipitation and evaporation estimates from ESMs do not align with



261 empirical estimates (Baker et al., 2021; McFarlane, 2011), we employ a bias-correction method. Specifically, we  
262 use a histogram equivalence method (Piani et al., 2010) to adjust empirical  $S_r$  thresholds to comparable CMIP6  
263  $S_r$  thresholds for various ESMs (Table S1). This involves, first, calculating  $S_r$  using CMIP6-historical precipitation  
264 and evaporation estimates between 2000-2014 (Appendix A1 and Fig. S8). We then determine a percentile-  
265 equivalent  $S_r$  thresholds for each of the thirty-three CMIP6-ESMs under the current climate. For example, if an  
266 empirical  $S_r$  of 100 mm corresponds to the 10<sup>th</sup> percentile ( $n = 20\%$  of total pixels), we find the 10<sup>th</sup> percentile  
267 in the CMIP6-historically  $S_r$ , which may be higher or lower than 100 mm for each ESM (Fig. 1 and Table S1).  
268 These percentile-equivalent  $S_r$  thresholds are then used to classify ecosystems both under current (CMIP6-  
269 historical; 2000-2014) and future climate (CMIP6-SSPs; 2086-2100) (Fig. 1b). Classifying savanna under future  
270 climate requires an additional step as outlined in Appendix A3.

271 Ultimately, we evaluate potential transitions by comparing ecosystems classified under current climate  
272 conditions (*this excludes savanna*) with those under future climate conditions (*this includes savanna*) (Sect.  
273 2.3.2). These transitions are divided into three distinct categories (Fig. 1c and Fig. A2):

- 274 i. **Forest-savanna transition:** This refers to current climate forest ecosystems that risk transitioning to a  
275 savanna under future climate change.
- 276 ii. **Transition to a more water-limited state:** This includes ecosystems that shift to a higher water-limited  
277 state in the future. For example, if a forest currently classified as lowly water-limited transitions to  
278 either a moderately or highly water-limited state in the future, it would fall under this category.
- 279 iii. **Reversion to a less water-limited state:** This includes ecosystems that shift to a lower water-limited  
280 state in the future.

281  
282 To aggregate the results from all ESMs, grid cells with  $> 50\%$  convergence are referred to as ‘moderate-  
283 high model agreement’, 20-50% as ‘moderate model agreement’ and  $\leq 20\%$  as ‘low model agreement’. In the  
284 Results section, we primarily discuss estimates from scenarios  $>20\%$  and  $>50\%$  model convergence. While a  
285 threshold of  $>20\%$  may seem low given the total number of ESMs analysed, it is important to recognise the  
286 variable and often limited capabilities of these ESMs, particularly in simulating biophysical interaction and  
287 emerging properties due to our limited understanding of the Earth system (Lenton et al., 2019; Stevens and  
288 Bony, 2013). Opting for a majority-based consensus in ESMs could overlook critical tipping risks identified by a  
289 minority of models, which might provide insights as valid as those from more widely agreeing models (Arora et  
290 al., 2023; Reyer et al., 2015).

291

## 292 **2.4 Sensitivity analyses**

293 Our methodology operates under two key assumptions: (i) the empirically derived  $S_r$  thresholds remain valid in  
294 the future, and (ii) the hydroclimatic estimates projected by ESMs accurately represent the actual climate, even

295 though these models have prescribed land-cover (Hurtt et al., 2020). To address the uncertainties related to  
296 the first assumption, we conduct four sensitivity analyses to assess the robustness of our analysis: (a) assuming  
297 that the regions exceeding the 99<sup>th</sup> percentile  $S_r$  are prone to a forest-savanna transition, as high  $S_r$  investment  
298 could be unrealistic from the perspective of plants under future climate change, (b) evaluating forest transitions  
299 using three different evaporation datasets, (c) assessing forest transitions under 10- and 40-year drought return  
300 periods, and (d) adjusting the forest-savanna transition thresholds.

301 Regarding the second assumption, we explicitly apply this methodology across a wide range of available  
302 ESMs under four SSP scenarios to identify consistencies and discrepancies in the results. Additionally, the  
303 discrepancies between the prescribed land-use and the forest transitions derived from our methodology, as  
304 well as the implications of these assumptions are detailed in the Discussion section.

305

### 306 **3 Results**

307 We find that under future climate conditions (2086-2100), considering >50% models' agreement, about one-  
308 fourth of the forests in both South America and Africa are projected to transition (Fig. 2b-g). With >20% models'  
309 agreement, these transitions are projected to occur for about three-fourths of the forests for both continents.  
310 Considering a lower threshold for models' agreement causes double or triple counting of some transitions (Fig.  
311 2b-g). To minimise this in further analyses, we only consider >50% models' agreement for forests that transition  
312 to a more and less water-limited state. Furthermore, because (abrupt) forest-savanna transitions are under-  
313 represented in ESMs (Drijfhout et al., 2015; Lenton, 2011; Maslin and Austin, 2012; Valdes, 2011), we consider  
314 >20% models' agreement for them. Considering this, we not only reduce the overlap to <0.4% of the total forest  
315 area (Fig. S9), but we also maximise highlighting forest-savanna transition risk for both continents.

316 We find that the risk of forest-savanna transitions mainly occurs in the Guiana Shield of South America,  
317 and the southern and south-eastern regions of Africa (Fig. 3). Compared to Africa, forest-savanna transitions  
318 are more prominent in South America under warmer climates (i.e., higher SSPs; Fig. 2b and 3). Our analysis  
319 reveals that the extent of forest-savanna transitions in South America decreases from almost  $1.32 \times 10^6$  km<sup>2</sup>  
320 (16.3% of total forest area in South America) under the highest emission scenario to  $0.04 \times 10^6$  km<sup>2</sup> (0.5%) under  
321 the lowest emission scenario (Fig. 2b). Interestingly, for Africa, the extent of forest-savanna transition did not  
322 change much for different SSPs, i.e., (median)  $0.25 \times 10^6$  km<sup>2</sup> with a maximum deviation of  $\pm 0.11 \times 10^6$  km<sup>2</sup>  
323 (minimum and maximum extent of transition between 3-6.6% of total forest area in Africa) (Fig. 2c).

324 When comparing the changes in forest-savanna transition risk areas relative to their immediate lower  
325 warming scenarios, we find considerable increases for South America. The highest relative growth of  
326 approximately 5.75 times is observed between SSP1 and SSP2, with the forest area under risk increasing from  
327  $0.04 \times 10^6$  km<sup>2</sup> to  $0.23 \times 10^6$  km<sup>2</sup>, respectively. It increases by 3.48 times from SSP2 to SSP3 ( $0.23 \times 10^6$  km<sup>2</sup> to  
328  $0.80 \times 10^6$  km<sup>2</sup>), and by 1.65 times from SSP3 to SSP5 ( $0.80 \times 10^6$  km<sup>2</sup> to  $1.32 \times 10^6$  km<sup>2</sup>). For Africa, however,

329 the increases are more modest: the risk grows by 1.29 times from SSP1 to SSP2 ( $0.17 \times 10^6 \text{ km}^2$  to  $0.22 \times 10^6$   
330  $\text{km}^2$ ), by 1.63 times from SSP2 to SSP3 ( $0.22 \times 10^6 \text{ km}^2$  to  $0.36 \times 10^6 \text{ km}^2$ ), and is observed to decrease by 0.72  
331 times from SSP3 to SSP5 ( $0.36 \times 10^6 \text{ km}^2$  to  $0.26 \times 10^6 \text{ km}^2$ ).

332 By evaluating changes to their hydroclimate, we find that under warmer climates, forest-savanna  
333 transition regions in both continents are projected to experience a decrease in precipitation. Furthermore, we  
334 observe an increase in precipitation seasonality for South America, whereas Africa shows a decrease (Fig. S12).  
335 Here, an increase in precipitation seasonality (seasonal variability in precipitation over the year) creates water-  
336 limited conditions for the ecosystem. In contrast, a decrease in seasonality and precipitation in Africa  
337 corresponds to a lower moisture availability altogether. Nevertheless, for both these continents, this transition  
338 seems to occur for the previously highly water-limited forests under the current climate, followed by  
339 moderately, with the least contribution from lowly water-limited forests (Fig. 3). This highlights the looming risk  
340 on highly water-limited forests to experience a forest-savanna transition under warmer climates.

341 Forests that transition to a 'more' water-limited state in South America are spatially aggregated towards  
342 the border between Brazil, Colombia, and Peru – covering a considerable portion of the Central Amazon (Fig.  
343 3). Whereas for Africa, these forests exist in moderate to small patches towards the northern and southern  
344 extent of central Congo rainforests. We observe that these transitions account for most of the projected  
345 changes to forests' states across both continents (Fig. 2d,e), with the transition to just the 'highly water-limited  
346 forest' accounting for more than three-fourths of all such transitions (Fig. 3). We observe that South American  
347 forests gradually become increasingly water-limited under warmer climates, with maximum and minimum  
348 projected transition of  $1.89 \times 10^6 \text{ km}^2$  (23.4%) and  $1.61 \times 10^6 \text{ km}^2$  (19.9%) observed under the highest and lowest  
349 emission scenarios, respectively (Fig. 2d,e). Whereas for Africa, the change in the water-limited state of the  
350 forests under different SSP scenarios remains almost similar (i.e., median  $1.14 (\pm 0.06) \times 10^6 \text{ km}^2$ ; 19.6-22.2%).  
351 Analysis of their hydroclimatic changes reveals that water-limitation is induced by both a decrease in  
352 precipitation and an increase in seasonality in South America (Fig. S13). In contrast, water-limitation in Africa is  
353 driven solely by an increase in seasonality. We observe that these newly water-limited forests seem to have  
354 permeated to regions that were previously (under the current climate) dominated by lowly and moderately  
355 water-limited forests (Fig. 3). Here, this shift only signifies the changes to hydroclimatic conditions allowing  
356 forests to transition to a more water-limited state, rather than the changes to the floristic composition of  
357 terrestrial species from one location to another. Although such a shift under changing climate is not unlikely  
358 (Esquivel-Muelbert et al., 2019), they are not analysed in this study.

359 Forests that revert to a 'less' water-limited state in South America are primarily observed in the south-  
360 eastern Amazon, with small patches observed towards eastern Brazil and the western coast of Equatorial  
361 Guinea and Gabon (Fig. 3). For Africa, the reverted forests exist in patches in the northern and southern regions  
362 of the Congo rainforest. Furthermore, for South America, we observe a gradual decrease in these reversions

363 with an increase in warming. Here, we observe the lowest reversion of  $0.23 \times 10^6 \text{ km}^2$  (2.8%) under the highest  
364 emission scenario and the highest reversion of  $0.67 \times 10^6 \text{ km}^2$  (8.4%) under the lowest emission scenario (Fig.  
365 2f,g). For Africa, these trends remain almost similar under all SSPs (i.e., median  $0.18 (\pm 0.05) \times 10^6 \text{ km}^2$ ; 2.2-  
366 3.5%). Comparing these transitions with their hydroclimatic changes reveals an overall increase in precipitation  
367 (Fig. S14). Interestingly, we observe a much higher precipitation increase for South America under high-  
368 emission scenarios than those in lower-emission scenarios. However, we find that precipitation seasonality is  
369 also higher for these ecosystems under warmer climates (Fig. S14). This suggests that increased precipitation  
370 without changes to precipitation seasonality helps decrease the water-limitation of the ecosystem, compared  
371 to the ecosystems that experienced a simultaneous increase in both.

372 Our sensitivity analysis, detailed in Appendix B1, reveals a consistent pattern of forest transitions across  
373 various scenarios.

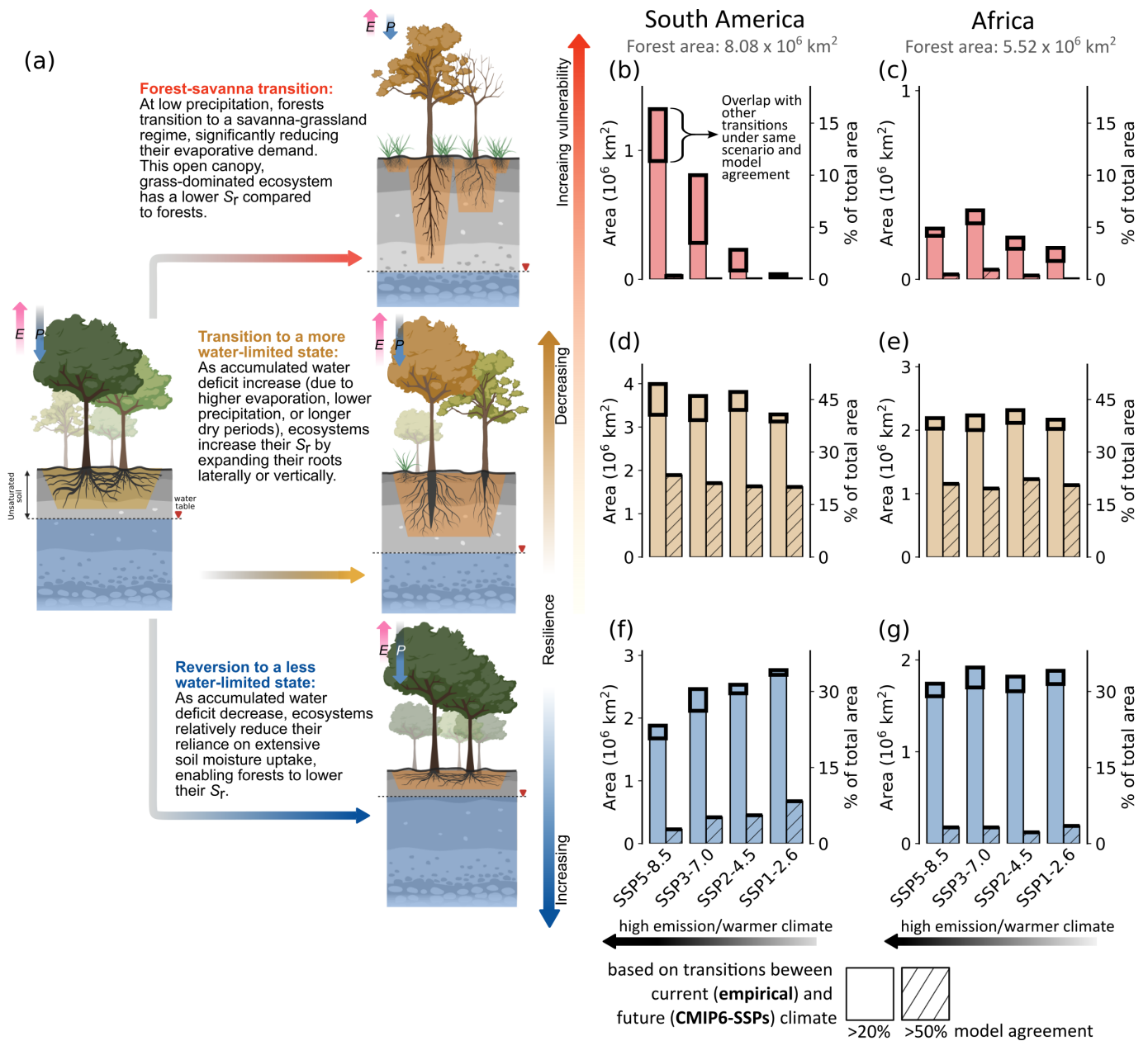
374

375

376

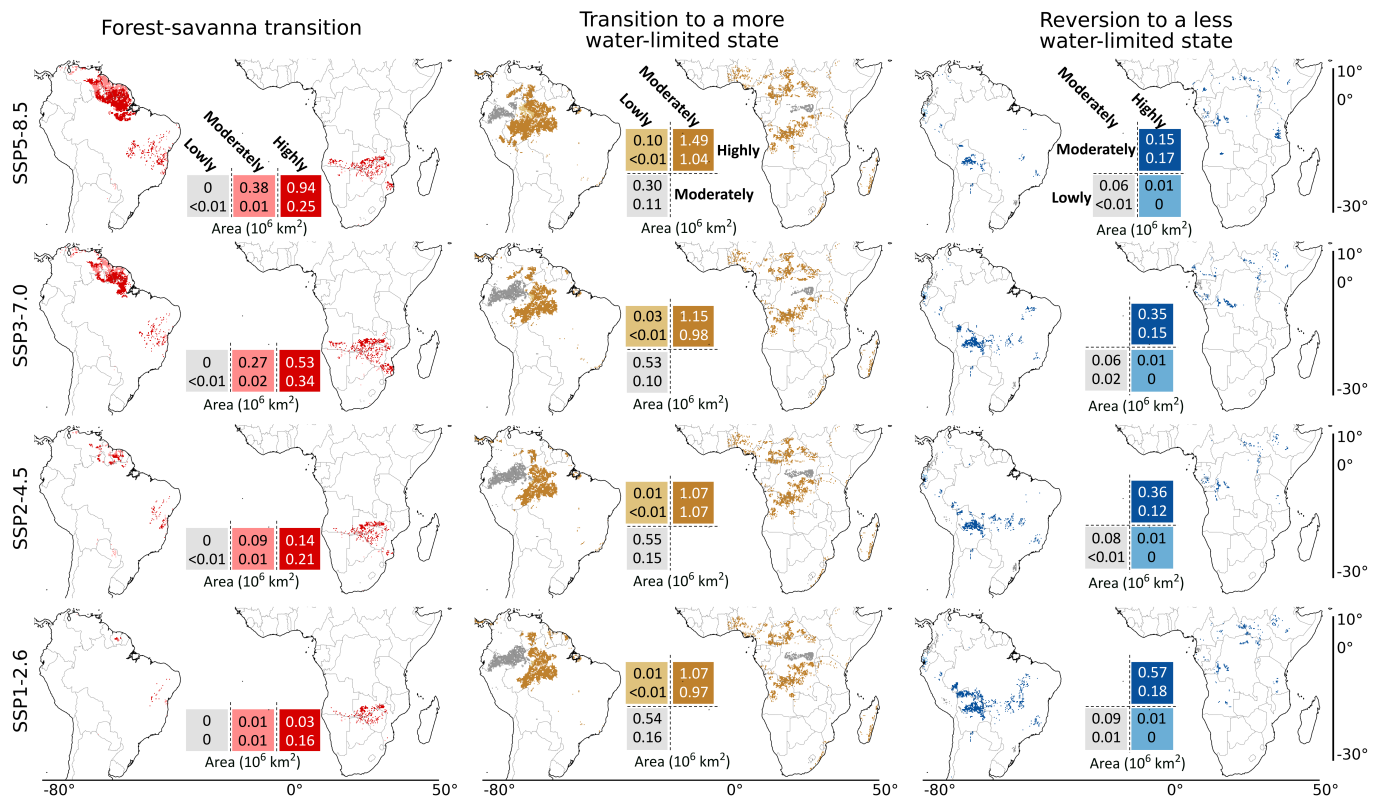
377

378

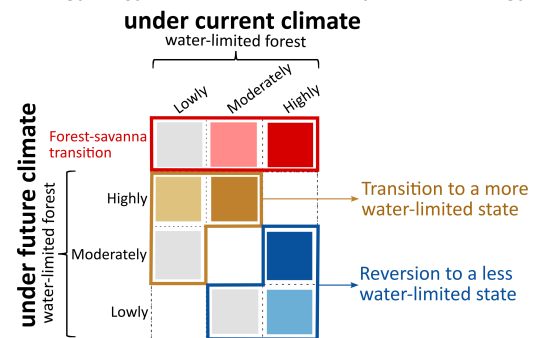


379  
 380 **Figure 2: Comparing the potential transitions under different SSP scenarios.** (a) The state of the ecosystem,  
 381 both above- and below-ground, (post-transition) under future climate, quantifying (b,c) forest-savanna  
 382 transition, (d,e) forests' that transition to a more water-limited state and (f,g) revert to a less water-limited  
 383 state for South America and Africa (present forest area mentioned on the top of (b,c)), respectively. For the  
 384 analysis above, transitions are calculated for grid cells with model agreement  $>20\%$  (plain bar plot) and  $>50\%$   
 385 (hatched bar plot). These quantifications show changes in the forest area based on ecosystem transitions under  
 386 empirical-current (2001-2012) and future (2086-2100) climate conditions. For all these transitions, we assume  
 387 that the hydroclimate and vegetation are in equilibrium. Analyses comparing ecosystem transitions based on  
 388 CMIP6-historical (2000-2014) and future (2086-2100) climate conditions are shown in Fig. S10 and S11. For each  
 389 transition, the total area of spatial overlap with other transitions under the same SSP scenario and model  
 390 agreement is highlighted with thick black bars. The  $P$  and  $E$  arrows in (a) describe the relative magnitude of  
 391 precipitation and evaporation fluxes. The illustration in (a) is adapted from Singh et al. (2020) and created with  
 392 [BioRender.com](https://www.biorender.com).

393



Total forest area for South America:  $8.08 \times 10^6 \text{ km}^2$   
 and Africa:  $5.52 \times 10^6 \text{ km}^2$



394

395 **Figure 3: Spatial extent of potential transitions with respect to their current state under different SSP**  
 396 **scenarios.** We analysed transitions, explicitly focusing on forest-savanna transition, transition to a more water-  
 397 limited state, and reversion to a less water-limited state, by comparing different ecosystem classes under  
 398 current (empirical; 2001-2012) and future (SSPs; 2086-2100) climate conditions (as defined in Fig. 2). All  
 399 transitions shown above are analysed for moderate-high (>50%) model agreement, except forest-savanna  
 400 transition, for which moderate (>20%) model agreement is considered. Values overlaying the legends  
 401 correspond to the total area of transition for South America (top values) and Africa (bottom values).

402

## 403 4 Discussion

### 404 4.1 Asynchronous resilience risks under future climate change

405 Our analysis reveals the spatial extent of potential ecosystem transitions in South America and Africa and their  
 406 vulnerability to future climate change (Fig. 2 and 4). For South America, we find a clear indication of a decrease  
 407 in forest resilience (i.e., an increase in water-limited forests) and an increase in forest-savanna transition risk  
 408 under warmer climates (Fig. 2b,d,f). In contrast, these trends are not symmetric for Africa, where transition risk

409 shows only slight variation across the different SSPs (Fig. 2c,e,g). Similar to the results of this study, previous  
410 studies on rainforest tipping have also suggested that exceeding 1.5-2°C will considerably increase the tipping  
411 risk (Flores et al., 2024; Jones et al., 2009; Parry et al., 2022), with the Guyana Shield in the Amazon being the  
412 most susceptible under future climate change (Cox et al., 2004; Staal et al., 2020) (Fig. 3 and Table S3). Previous  
413 studies also agree that, in contrast to the Amazon, the projected risk to Congo rainforests is not substantial  
414 (Higgins and Scheiter, 2012; Staal et al., 2020) (Fig. 2). Despite it being unclear to what extent the ESMs  
415 represent the correct carbon-water dynamics (Koch et al., 2021), our results show a further divergence between  
416 Amazon's and Congo's responses to different SSPs (Fig. 2 and Fig. S12-S14). This could either be caused simply  
417 by a different response to changes in precipitation patterns over the respective regions (Kooperman et al., 2018;  
418 Li et al., 2022) or a different response to increased CO<sub>2</sub> levels in the atmosphere (Brienen et al., 2015; Hubau et  
419 al., 2020; Trumbore et al., 2015).

420 Previous empirical studies have linked these divergent responses to evolutionary and biogeographical  
421 differences between the ecosystems, which resulted in distinct species pools that uniquely influence each  
422 ecosystem's adaptability and response to climate change (Fleischer et al., 2019; Hahm et al., 2019; Hubau et  
423 al., 2020; Slik et al., 2018). These studies found that forest ecosystems in the Amazon tend to be more dynamic  
424 – grow faster due to high CO<sub>2</sub> levels in the atmosphere – than those in the Congo rainforests. However, these  
425 fast-growing trees also die young due to them investing substantially less in their adaptive strategies against  
426 perturbations than (less dynamic) old-growth forests (Brienen et al., 2015; Körner, 2017; Rammig, 2020). This  
427 makes the Amazon rainforest especially sensitive to CO<sub>2</sub> emissions pathways, as the positive influence of CO<sub>2</sub>  
428 fertilisation-induced growth is counteracted by the negative impact of warming and droughts, thereby  
429 exacerbating the risk of forest mortality under high emission scenarios (Brienen et al., 2015; Hubau et al., 2020;  
430 Yang et al., 2018). In this case, the projected changes to the future hydroclimate could be an artefact of  
431 decreased transpiration and precipitation due to forest mortality, rendering the rainforests vulnerable to  
432 tipping. In contrast, terrestrial species in Congo rainforests appear more resilient, having adapted to severe  
433 droughts during glacial periods, which makes them better equipped to handle episodic water-induced  
434 perturbations than Amazon (Cole et al., 2014).

435 Nevertheless, with compounding influence from land-use and climate-induced hydroclimatic changes  
436 (Davidson et al., 2012), these rainforests risk tipping to a savanna state. Our results highlight that by keeping  
437 the mean global surface temperature below 1.5-2°C warming (which in this case is equivalent to SSP1-2.6  
438 relative to the pre-industrial), we minimise forest-savanna transition risk and maximise recovery – thereby  
439 improving the resilience of rainforest ecosystems (Fig. 2, 3 and 4).

## 440

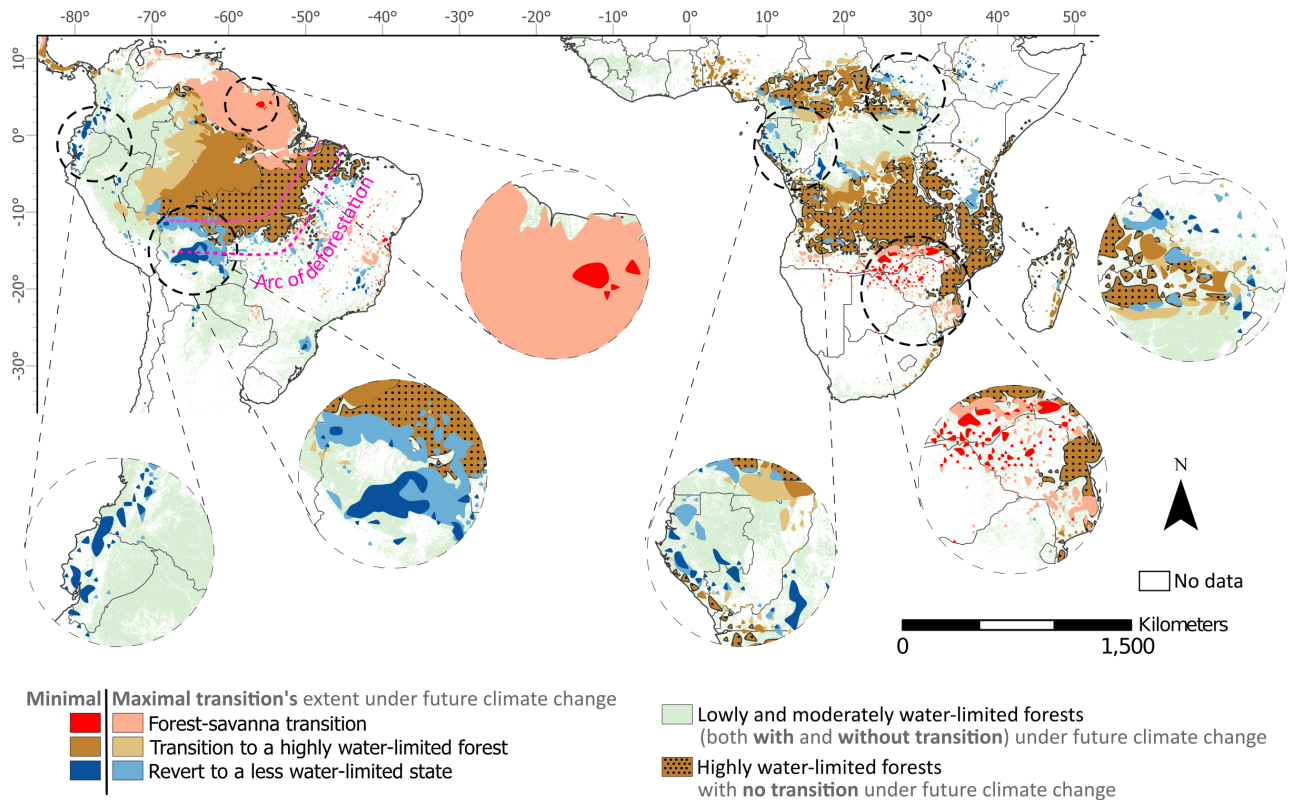
## 441 **4.2 Changes in atmospheric moisture flow drives forest-savanna transition**

442 Among all transitions, the most noticeable and catastrophic (since it is difficult to revert) is the forest-savanna  
443 transition projected in the Amazon's Guiana Shield of South America, and over the southern and south-eastern  
444 parts of Africa (Fig. 3 and 4). These transitions are associated with the shifting of the inter-tropical convergence  
445 zone (ITCZ) (Mamalakis et al., 2021), which decreases precipitation and increases precipitation seasonality over  
446 the continents. For South America, the creation of these low-pressure bands allows the trade winds to bring in  
447 considerable moisture from the equatorial Atlantic Ocean over to Amazon by passing through the Guiana Shield  
448 and ultimately carrying it across the La Plata Basin via the South American low-level jet (Bovolo et al., 2018; van  
449 der Ent et al., 2010; Zemp et al., 2014). Similarly, for Africa, south-eastern trade winds bring moisture from the  
450 Indian Ocean over the centre of the African continent (Mamalakis et al., 2021).

451 Under a warmer climate, sea surface temperature over the equatorial Atlantic and the northern Indian  
452 Ocean is projected to increase (Pascale et al., 2019; Zilli et al., 2019), leading to a southward shift in ITCZ over  
453 the eastern Pacific and Atlantic Oceans, and northward over east Africa and the Indian Ocean (Mamalakis et al.,  
454 2021; Xie et al., 2010). Previous studies also acknowledge that the intense surface warming over the Sahara  
455 under future climate can also attract ITCZ northwards in Africa (Cook and Vizy, 2012; Dunning et al., 2018;  
456 Mamalakis et al., 2021). Since these shifts in ITCZ can potentially both mitigate and aggravate (especially critical  
457 for highly water-limited forests) the impact of (accumulated) water-deficit on the forest ecosystem, including  
458 those caused by localised deforestation (Leite-Filho et al., 2021; Schumacher et al., 2022; Staal et al., 2018;  
459 Wunderling et al., 2022); it warrants the need to include changes in atmospheric circulation for studies analysing  
460 the impact of future climate on the resilience of forest ecosystems (Staal et al., 2020; Zemp et al., 2017).

461





462  
 463 **Figure 4: Minimal and maximal extent of potential ecosystem transitions under future climate change in the**  
 464 **entire study region over South America and Africa.** The three transition types are: forest-savanna transition,  
 465 from any class to highly water-limited forests, and to a less water-limited state (see definitions in Fig. 2 and 3).  
 466 For better visualisation of these transitions, in this figure, we first converted all grid cells to shape, merged  
 467 them, and then smoothed them using the 'polynomial approximation with exponential kernel' function (with a  
 468 tolerance value of 1) in ArcGIS pro. The unsmoothed version of the transitions is shown in Fig. 3. The minimal  
 469 and maximal represent the minimum and maximum possible extent of transitions (as quantified in Fig. 3) based  
 470 on changes between current (empirical; 2001-2012) and future (SSPs; 2086-2100) climate conditions regardless  
 471 of the SSP scenarios.

472  
 473 **4.3 Discrepancy between prescribed future land-use and projected transitions**

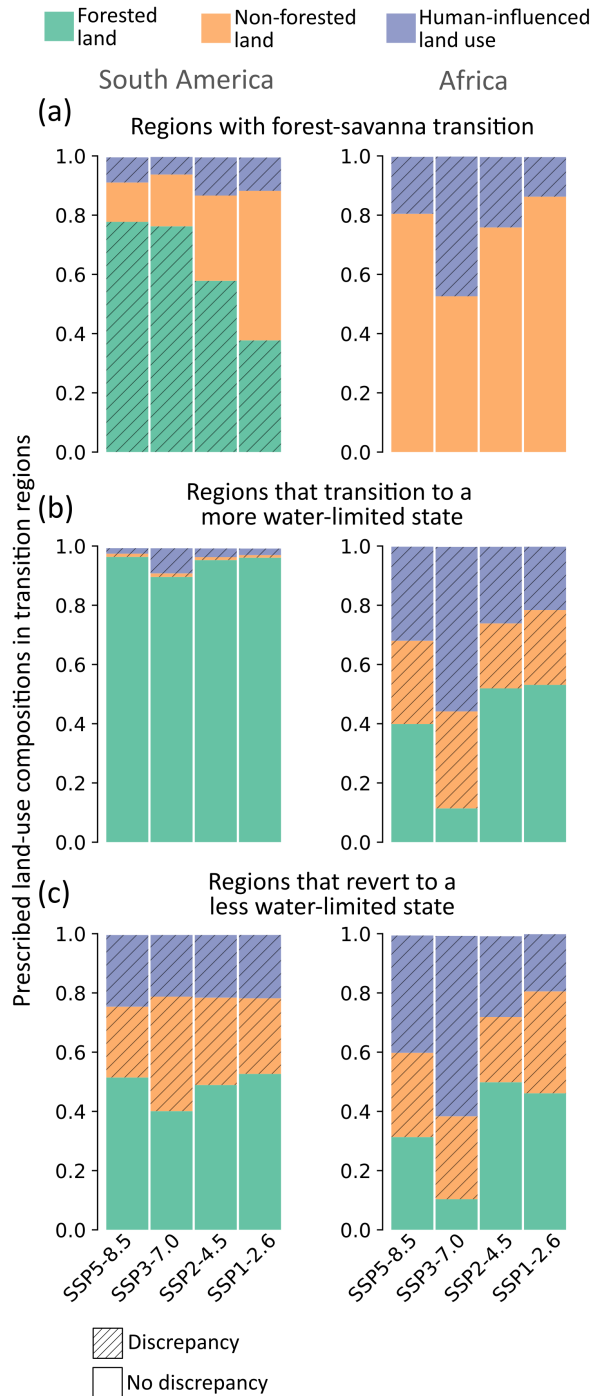
474 The land-use information in CMIP6-ESMs is not biophysically simulated, but prescribed based on simulations  
 475 from Integrated Assessment Models (IAMs) for each SSP scenario (Hurtt et al., 2020). Therefore, it is valuable  
 476 to examine whether these prescribed land-use scenarios agree or conflict with the changes projected (assuming  
 477 equilibrium between hydroclimate and the ecosystem) by our  $S_r$ -based ecosystem transitions (Fig. 5 and Fig.  
 478 S15-S17).

479 The most noticeable discrepancies are observed in South America, where the extent of forest-savanna  
 480 transitions is underestimated in prescribed land-use scenarios compared to those projected in this study (i.e.,  
 481 prescribed land-use predicts forests in the region whose hydroclimate can't support forest; Fig. 4 and 5a).  
 482 Additionally, in South America, our analysis highlights the potential of some forests reverting to a 'less water-  
 483 limited state' in places where the prescribed land-use in the ESMs suggest non-forest landscape (Fig. 4 and 5c).

484 These discrepancies arise because the prescribed land-use in CMIP6-ESMs do not shift in response to  
485 hydroclimatic changes. Despite our approach assuming equilibrium and overlooking the temporal dynamics of  
486 transitions, based on broad climate change pattern (Sect 4.2), we believe it more accurately represents the  
487 ecohydrological state of the ecosystems.

488           However, these prescribed land-uses can introduce errors in subsequent biophysical processes  
489 simulated in ESMs (Ma et al., 2020), affecting the accuracy of projected transitions. For example, prescribing a  
490 region as a forest that would be grassland in the future will lead to the extraction of deeper subsoil moisture in  
491 ESMs, which (actual) grasslands do not have the capacity to access (Ahlström et al., 2017; Yu et al., 2022). This  
492 will result in an overestimation of the ecosystem's evaporation, potentially altering precipitation patterns  
493 downwind and leading to inaccurate water budget assessments for these ecosystems. Consequently, causing  
494 erroneous projections of the ecosystem state. These discrepancies underscore the urgent need for  
495 enhancements in the land surface components of ESMs, enabling dynamic simulations of vegetation-climate  
496 feedbacks. Such improvements would provide a more accurate representation of the ecohydrology of  
497 terrestrial ecosystems and their response to changing climate conditions.

498



499  
 500 **Figure 5: Prescribed land-use composition for each transition region under different SSP scenarios (median**  
 501 **2086-2100), calculated as the ratio between the prescribed land use area and the projected transition area.**  
 502 Regions where IAM prescribed land use are same as the projected transitions (from Fig. 3) are shown in plain  
 503 colours (i.e., no discrepancy). Whereas regions where IAM-prescribed land use differs from projected  
 504 transitions are hatched (i.e., discrepancy).

505

#### 506 4.4 Limitations

507 This study assumes that the  $S_r$ -derived thresholds used to classify terrestrial ecosystems under current climate  
 508 conditions remain valid under future climate change. However, forests themselves are dynamically adapting

509 their structure and functions in response to climate change, altering their critical thresholds (Doughty et al.,  
510 2023). Thus, assuming a static critical threshold may lead to inaccuracies in estimating forests' resilience to  
511 future climate change. For instance, under the CO<sub>2</sub> fertilisation effect, forests may become more water-use  
512 efficient (i.e., transpire less and therefore need for a lower  $S_r$ ) (Xue et al., 2015), potentially delaying their tipping  
513 under warming scenarios compared to those projected in this study. Conversely, factors such as nutrient  
514 limitation (Condit et al., 2013) or extensive human influence (van Nes et al., 2016) in the ecosystem might lead  
515 to an earlier tipping than anticipated.

516 However, the uncertainty surrounding the effect of CO<sub>2</sub> fertilisation, nutrient limitation, and human  
517 influence on vegetation remain significant research frontiers for enhancing our understanding of rainforest  
518 tipping under future climate change (Fleischer et al., 2019; Hofhansl et al., 2016). Additionally, factors such as  
519 precipitation variability, species composition, soil properties, and topography can contribute to varied local-  
520 scale forest responses to future climate change (Staal et al., 2020). It should also be noted that though these  
521 uncertainties may hinder our understanding of local-scale forest resilience, the influence of future hydroclimatic  
522 changes on forests still constitutes major prediction uncertainties. Therefore, in this study, regardless of how  
523 these influences are parametrised or simulated in each ESM, we assume that hydroclimatic estimates projected  
524 by the ESMs represent the actual climate.

525 Of course, this assumption opens us and other studies projecting forest conditions to future climate  
526 change to certain limitations. Our ability to project forest-savanna transitions (or any transition) relies on the  
527 model's capacity to simulate complex feedbacks. Some models capture complex vegetation-atmosphere  
528 interaction, simulating local and regional scale feedbacks across time (Ferreira et al., 2011; Jach et al., 2020);  
529 others rely on simpler parametrisation (Nof, 2008) (e.g., parametrisation of CO<sub>2</sub> fertilisation; Koch et al., 2021).  
530 However, caution should be taken to not overgeneralise the functioning of tropical forests just from the analysis  
531 presented in this study, and also realise the current potential of ESMs to simulate them (Staal et al., 2020). We  
532 believe that by considering simulations from multiple ESMs under different SSP scenarios, not only do we  
533 highlight the agreements and conflicts between potential transitions; but also allow future studies to  
534 disentangle vegetation-climate feedbacks and improve the modelling of local-scale interactions (e.g.,  
535 vegetation's water-uptake profile, species response to CO<sub>2</sub> fertilisation) in the ESMs.

536

## 537 **5 Conclusions**

538 Classifying terrestrial ecosystems based on empirical and CMIP6 ESMs-derived  $S_r$  allowed us to assess the future  
539 transitions in the rainforest ecosystems. Our findings indicate that climate under the lowest emission scenarios  
540 significantly reduces the risk of rainforest tipping and maximises reversion to a less water-limited state, while  
541 climate under the high emission scenarios have the opposite effect on the forest ecosystem. Specifically, in the

542 Amazon rainforest, the risk of forest-to-savanna transition increases considerably with incremental increase in  
543 warming. Conversely, in the Congo, the variation in transition risk across different emission scenarios is  
544 relatively minor.

545 Notably, our analysis suggests very limited tipping risk that is ‘unavoidable’ (i.e., regions prone to a  
546 forest-savanna transition in all scenarios), and the vast majority of potential transition risks can still be avoided  
547 by steering towards a less severe climate scenario, thereby underscoring the critical window of opportunity.  
548 Moreover, regions projected to revert to a less water-limited state could potentially become more amenable  
549 to restoration and responsive to deforestation prevention efforts. This study highlights the importance of  
550 restricting global temperature change below 1.5-2°C warming relative to the pre-industrial levels to prevent  
551 forest tipping risks and provide the best conditions for effective ecosystem stewardship.

552

## 553 **Appendix A: Methodology**

### 554 **A1. Root zone storage capacity calculation**

555 Our method to calculate  $S_r$  is adopted from Singh et al. (2020). For estimating  $S_r$ , we first obtained the water  
556 deficit ( $D_t$ ) at daily time step from the daily estimates of precipitation ( $P_t$ ) and evaporation ( $E_t$ ) (Fig. A1) using:

$$557 \quad D_t = E_t - P_t \quad (\text{A1})$$

558 Here,  $t$  denotes the day count since the start of the simulation, with simulation for each grid starting in  
559 the month with maximum precipitation. Second, we calculated the accumulated water deficit integrated at  
560 each one-day timestep for one year using:

$$561 \quad D_{a(t+1)} = \max\{0, D_{a(t)} + D_{t+1}\} \quad (\text{A2})$$

562 Where  $D_{a(t+1)}$  is the accumulated water deficit at each time step (Fig. A1). Here, an increase in the  
563 accumulated water deficit will occur when  $E_t > P_t$ , and a decrease when  $E_t < P_t$ . However, since this algorithm  
564 estimates a running estimate of root zone storage reservoir size, we use a maximum function to calculate the  
565 accumulated water deficit, which by definition can never be below zero. Not allowing  $D_{a(t+1)}$  to be negative also  
566 means that excess moisture from precipitation will either contribute to deep drainage or runoff. Lastly, the  
567 maximum accumulated annual water deficit ( $D_{a,y}$ ) will represent the maximum storage required by the  
568 vegetation to respond to the critical dry periods (Fig. A1).

$$569 \quad D_{a,y} = \max\{D_{a(t+1)}\} \quad t = 1 : n - 1 \quad (\text{A3})$$

570 This simulation runs for a whole year, with  $n$  denoting the number of days in year  $y$ .

571 Although different terrestrial ecosystems (e.g., forest, savanna and grasslands) adapt to different  
572 drought return periods (de Boer-Euser et al., 2016; Gao et al., 2014; Wang-Erlandsson et al., 2016). For instance,  
573 grasslands and savanna adapt to shorter drought return periods (i.e., <10 years and 10-20 years, respectively).  
574 In contrast, forests adapt to long drought return periods (>40 years) (Wang-Erlandsson et al., 2016). For this  
575 study, we use a uniform 20-year drought return period (following Bouaziz et al., 2020; Nijzink et al., 2016) to  
576 avoid any artificially introduced transitions between different ecosystems. Thus, this 20-year drought return  
577 period  $S_r$  refers to the maximum amount of root zone moisture accessible to vegetation for transpiration during  
578 the largest accumulated annual water deficit expected every twenty years under static climate conditions. This  
579 we analyse using on the Gumbel extreme value distribution (Gumbel, 1958) and apply it to normalise all  $D_{a,y}$ .  
580 The Gumbel distribution ( $F(x)$ ) is given by:

$$581 \quad F(x) = \exp \left[ -\exp \left[ -\frac{(x - \mu)}{\alpha} \right] \right] \quad (A4)$$

582 Where  $\mu$  and  $\alpha$  are the location and scale parameters, respectively. We calculate this using the python  
583 package 'skextremes'(skextremes Documentation):

$$584 \quad S_r = \overline{D_{a,y}} + K \times \sigma_{n-1} \quad (A5)$$

585 Where  $K$  is the frequency factor given by:

$$586 \quad K = \frac{y_t - y_n}{S_n} \quad (A6)$$

587 And  $y_t$  is the reduced variate given by:

$$588 \quad y_t = - \left[ \ln \left[ \ln \left( \frac{T}{T-1} \right) \right] \right] \quad (A7)$$

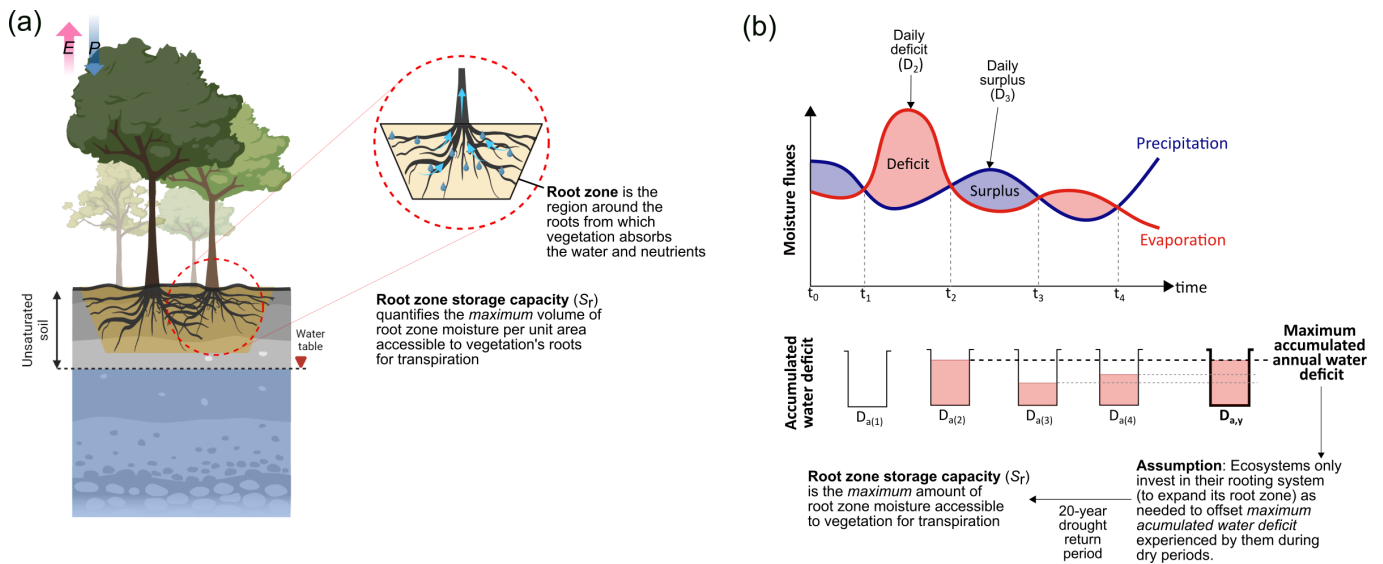
589 Where  $T$  is the drought return period (i.e., 20 years used in this study),  $\overline{D_{a,y}}$  is the mean annual  
590 accumulated deficit for the years 2001-2012,  $\sigma_{n-1}$  is the standard deviation of the sample. Also,  $y_n$  is the  
591 reduced mean and  $S_n$  is the reduced standard deviation, which for  $n = 11$  years (since we are calculating  $S_r$  in a  
592 hydrological year – simulation starts mid-year – we therefore lose one year) is equal to 0.4996 and 0.9676,  
593 respectively (Gumbel, 1958).

594 Since the CMIP6 (-historical and -SSP estimates, the timeframe considered are 2000-2014 and 2086-  
595 2100, respectively) doesn't have daily estimates of evaporation and precipitation for all Earth System Models  
596 (ESMs), we directly use the monthly estimates of precipitation and evaporation to modify Eq. (A1) as:

$$597 \quad D_t = E_{t(monthly)} - P_{t(monthly)} \quad (A8)$$

598 Here,  $t(\text{monthly})$  denotes the month count since the start of the simulation. The rest of the steps (Eq.  
 599 A2-A7) remain the same for CMIP6 datasets. For CMIP6 runs,  $\mathcal{Y}_n$  and  $S_n$  in Eq. (6) are calculated for  $n = 14$  years  
 600 (Eq. A7) equal to 0.5100 and 1.0095, respectively. The  $S_r$  estimates derived from daily and monthly empirical  
 601 estimates (from Eq. A1 and A8) are compared in Fig. S8 to evaluate uncertainty.

602



603

604 **Figure A1:** The figure illustrates the root zone storage capacity ( $S_r$ ) of the ecosystem. (a) We show the difference  
 605 between the ecosystem's root zone and how that constitutes its  $S_r$ . (b) Conceptual illustration of how the  
 606 ecosystem's precipitation and evaporation fluxes constitute the maximum accumulated annual water deficit  
 607 ( $D_{a,y}$ ) and  $S_r$ . The figure is adopted from Singh (2023) and Wang-Erlandsson et al. (2016).

608

609 **A2. Abiotic and biotic factors influence soil moisture availability**

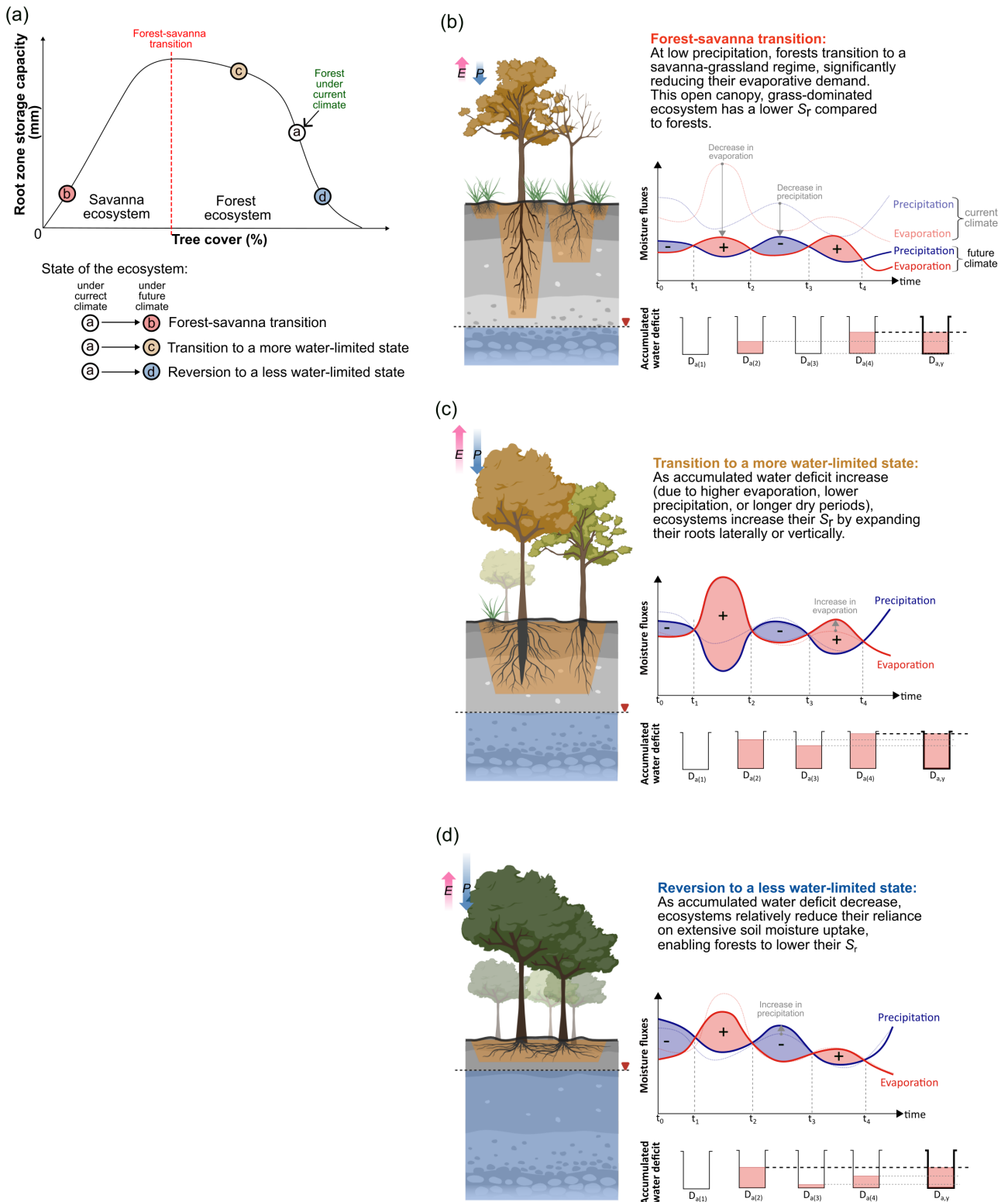
610 In this study,  $S_r$  quantifies the hydrological buffer necessary for an ecosystem to maintain its structure and  
 611 functions, reflecting the amount of root zone soil moisture available to vegetation for transpiration. Our mass-  
 612 balance-based  $S_r$  methodology, while not directly distinguishing between the biotic and abiotic influences on  
 613 soil moisture and root characteristics, does incorporate their critical role in shaping the ecohydrology of the  
 614 ecosystem under climate change. By utilising empirical precipitation and evaporation data, our approach  
 615 theoretically captures the combined impact of these biotic and abiotic factors on the actual hydrological regime  
 616 (including soil moisture) of the ecosystem (Sect. 2.3.2).

617 We acknowledge that abiotic factors such as soil texture, structure, and depth profoundly affect soil  
 618 water-holding capacity (Fayos, 1997). For instance, field studies suggest that clay and organic-rich soils exhibit  
 619 superior water retention capabilities due to their fine textures and high surface areas, which is crucial to  
 620 vegetation for moisture uptake during extended dry periods (Bronick and Lal, 2005; Fayos, 1997). Additionally,

621 the depth and porosity of soil also dictate its ability to absorb and store water in the soil, with deeper, less  
622 compacted soils providing a higher buffer against drought by allowing greater water infiltration (Indoria et al.,  
623 2020; Smith et al., 2001). Climate change, by altering temperature and precipitation patterns, can modify these  
624 abiotic soil properties, potentially leading to loss in soil water retention capacity through erosion and  
625 compaction (Dexter, 2004).

626 Moreover, biotic factors, including plant-root dynamics and microbial activity, also play essential roles  
627 in shaping the ecosystem (Brunner et al., 2015; Sveen et al., 2024). Deep and extensive root systems not only  
628 directly improve access to deeper soil moisture, but also physically modify the soil to enhance its permeability  
629 and storage (Canadell et al., 1996; Jackson et al., 1996). Additionally, microbial processes contribute by breaking  
630 down organic matter, thereby improving the soil's structural integrity and ability to retain water (Dittert et al.,  
631 2006). These biotic interactions, coupled with changing abiotic factors under climate change, underscore the  
632 complex dynamics that govern soil moisture availability and ecosystem resilience. However, this study does not  
633 consider the direct impact of future climate change on biotic and abiotic factors, nor their influence on  
634 ecosystems, beyond changes to  $S_r$ .





635

636 **Figure A2:** (a) The figure compares the root zone storage capacity ( $S_r$ ) with the ecosystem state (i.e., tree cover).  
 637 This figure expands on the conceptual illustration from Fig. A1, showing how the ecosystem's precipitation and  
 638 evaporation fluxes contribute to  $S_r$  under different forest transition scenarios: (b) forest-savanna transition, (c)  
 639 transition to a more water-limited state, and (d) reversion to a less water-limited state.

640 **A3. Using precipitation to discern savanna from forests under future climate change**

641 Under future climate change, some ecosystems will remain forest, while others may transition to  
642 savanna. In our  $S_r$ -based framework, without information about above-ground forest structure, it is difficult to  
643 discern whether an ecosystem is a forest or savanna just with  $S_r$  (for instance, an ecosystem with  $S_r$  of 200 mm  
644 can either be a moderately water-limited forest or savanna; Sect. 2.3.2). Differentiating these ecosystems is  
645 easier under the current climate, where we have several remote sensing products capturing vegetation  
646 structure (e.g., tree cover density, tree height, floristic patterns) (Aleman et al., 2020; Hirota et al., 2011; Xu et  
647 al., 2016). However, under future climate, we must find a proxy, since land-use information in ESMs are  
648 prescribed (i.e., not biophysically simulated) (Ma et al., 2020).

649 To address this, previous studies have either relied on vegetation structure proxies provided by ESMs  
650 (e.g., net primary productivity) (Boulton et al., 2013; Jones et al., 2009), or assumed that terrestrial ecosystems  
651 are in equilibrium with their climate (Staal et al., 2020) (see Supplement). In this study, we adopted the latter  
652 approach and utilised climate variables, specifically (bias-corrected) mean annual precipitation and the  
653 precipitation seasonality index, as proxies to make this distinction (Fig. S4). The climate conditions (or range)  
654 necessary for forest ecosystems to sustain themselves are determined by comparing empirical estimates of  
655 mean annual precipitation and precipitation seasonality index with  $S_r$ . These estimates are then bias-corrected  
656 (following the same methods described in Sect. 2.3.3) before applying them to future climate scenarios. This  
657 (revised) classification of terrestrial ecosystems is then used to assess forest transitions under future climate  
658 change scenarios.

659

## 660 **Appendix B: Results**

### 661 **B1. Sensitivity analysis reveals robust performance of the framework**

662 Sensitivity analysis reveals that by setting an extreme  $S_r$  threshold – signifying a forest-savanna transition for  
663 ecosystems that cannot maintain their above-ground structure at high  $S_r$  – we observe some shifts near the  
664 already projected risk regions and coastal areas (Fig. 3 and Fig. S18). However, the transition risk identified in  
665 the coastal regions may be an artefact of interpolating hydroclimate estimates to higher resolution. Additionally,  
666 since evaporation is more prevalent over oceans than land, this could result in high  $S_r$  values, thereby projecting  
667 an elevated tipping risk in these coastal areas.

668 We also discover that variations in the evaporation datasets and return periods used for calculating  $S_r$   
669 have minimal effect on forest transitions (Fig. S19 and S20). Although the forest classification thresholds may  
670 shift with different evaporation products under current climate conditions (Singh et al., 2020), our histogram  
671 equivalence method ensures that forest classifications under future climates adjust accordingly, resulting in  
672 only minor alterations to the final outcome (Fig. 1b and Fig. S19). Furthermore, while  $S_r$  values tend to increase

673 with increase with shorter return periods, the impact of these changes becomes less significant with longer  
674 return periods (Wang-Erlandsson et al., 2016); leading to minor variations in the end results (Fig. S20).

675 Moreover, lowering the forest-savanna transition thresholds can reduce the risk of forest-savanna  
676 transition since it expands the associated range of climate conditions (i.e., mean annual precipitation and  
677 seasonality) necessary for forests to sustain their structure and functions (Fig. S21). Conversely, increasing the  
678 forest-savanna transition threshold leads to an opposite trend, where the risk of transition increases (Fig. S22).  
679 Despite these sensitivity analyses, the variation in transition magnitudes is minor, and the trends across  
680 different SSP scenarios for both continents remain consistent (Fig. 2 and Fig. S18-S22). Therefore, the  
681 conclusions drawn from this study remain robust, even with variations in factors that could potentially affect  
682 forest transitions.

## 683 **Data availability**

684 All the data generated during this study is made publicly available at Zenodo:  
685 <https://zenodo.org/record/7706640>. Other datasets that support the findings of this study are publicly  
686 available at: (CMIP6; citations referred to in Table S2) <https://aims2.llnl.gov/>, (Root zone storage capacity;  
687 empirical) <https://github.com/chandrakant6492/Drought-coping-strategy>, (P-CHIRPS)  
688 <https://data.chc.ucsb.edu/products/CHIRPS-2.0/>, (E-BESS) <ftp://147.46.64.183/>, (E-FLUXCOM) [ftp.bgc-](ftp.bgc-jena.mpg.de)  
689 [jena.mpg.de](ftp.bgc-jena.mpg.de), (E-PML) <https://data.csiro.au/collections/#collection/CIcsiro:17375v2>, (E-ERA5)  
690 <https://cds.climate.copernicus.eu/cdsapp#!/dataset/reanalysis-era5-single-levels>, (Globcover)  
691 [http://due.esrin.esa.int/page\\_globcover.php](http://due.esrin.esa.int/page_globcover.php). Potential transitions for each ESM based on the comparison  
692 between empirical (2001-2012) and SSP (2086-2100) scenarios are presented in the Supplement.

## 693 **Code availability**

694 The python-language scripts used for the analyses presented in this study are available from GitHub:  
695 <https://github.com/chandrakant6492/Future-forest-transitions-CMIP6>. The python-language code for  
696 calculating (empirical) root zone storage capacity is available from GitHub:  
697 <https://github.com/chandrakant6492/Drought-coping-strategy>.

## 698 **Acknowledgements**

699 C.S., I.F. and L.W.-E. acknowledge funding support from the European Research Council (ERC) project 'Earth  
700 Resilience in the Anthropocene', project number ERC-2016-ADG-743080. L.W.-E. also acknowledges funding  
701 support from the Swedish Research Council for Sustainable Development (FORMAS), project number 2019-  
702 01220 and the IKEA Foundation. R.v.d.E. acknowledges funding support from the Netherlands Organisation for  
703 Scientific Research (NWO), project number 016.Veni.181.015. The authors also acknowledge the computational  
704 support provided by Microsoft Planetary Computer (<https://planetarycomputer.microsoft.com>) for performing  
705 the analyses.

## 706 **Author contributions**

707 All authors contributed to the conceptualisation of this research. CS performed the analyses and wrote the  
708 initial draft. All authors contributed to the discussion and revisions, leading to the final version of the  
709 manuscript.

## 710 **Competing interests**

711 The authors declare no competing interests.

712

713

714 **References**

- 715 Ahlström, A., Canadell, J. G., Schurgers, G., Wu, M., Berry, J. A., Guan, K., and Jackson, R. B.: Hydrologic  
716 resilience and Amazon productivity, *Nature Communications*, 8, 387, [https://doi.org/10.1038/s41467-017-](https://doi.org/10.1038/s41467-017-00306-z)  
717 00306-z, 2017.
- 718 Albasha, R., Mailhol, J.-C., and Cheviron, B.: Compensatory uptake functions in empirical macroscopic root  
719 water uptake models – Experimental and numerical analysis, *Agricultural Water Management*, 155, 22–39,  
720 <https://doi.org/10.1016/j.agwat.2015.03.010>, 2015.
- 721 Aleman, J. C., Fayolle, A., Favier, C., Staver, A. C., Dexter, K. G., Ryan, C. M., Azihou, A. F., Bauman, D., Beest,  
722 M. te, Chidumayo, E. N., Comiskey, J. A., Cromsigt, J. P. G. M., Dessard, H., Doucet, J.-L., Finckh, M., Gillet, J.-F.,  
723 Gourlet-Fleury, S., Hempson, G. P., Holdo, R. M., Kirunda, B., Kouame, F. N., Mahy, G., Gonçalves, F. M. P.,  
724 McNicol, I., Quintano, P. N., Plumptre, A. J., Pritchard, R. C., Revermann, R., Schmitt, C. B., Swemmer, A. M.,  
725 Talila, H., Woollen, E., and Swaine, M. D.: Floristic evidence for alternative biome states in tropical Africa,  
726 *PNAS*, 117, 28183–28190, <https://doi.org/10.1073/pnas.2011515117>, 2020.
- 727 Armstrong McKay, D. I., Staal, A., Abrams, J. F., Winkelmann, R., Sakschewski, B., Loriani, S., Fetzer, I., Cornell,  
728 S. E., Rockström, J., and Lenton, T. M.: Exceeding 1.5°C global warming could trigger multiple climate tipping  
729 points, *Science*, 377, eabn7950, <https://doi.org/10.1126/science.abn7950>, 2022.
- 730 Arora, V. K., Seiler, C., Wang, L., and Kou-Giesbrecht, S.: Towards an ensemble-based evaluation of land  
731 surface models in light of uncertain forcings and observations, *Biogeosciences*, 20, 1313–1355,  
732 <https://doi.org/10.5194/bg-20-1313-2023>, 2023.
- 733 Baker, J. C. A., Garcia-Carreras, L., Buermann, W., Souza, D. C. de, Marsham, J. H., Kubota, P. Y., Gloor, M.,  
734 Coelho, C. A. S., and Spracklen, D. V.: Robust Amazon precipitation projections in climate models that capture  
735 realistic land–atmosphere interactions, *Environ. Res. Lett.*, 16, 074002, [https://doi.org/10.1088/1748-](https://doi.org/10.1088/1748-9326/abfb2e)  
736 9326/abfb2e, 2021.
- 737 Barros, F. de V., Bittencourt, P. R. L., Brum, M., Restrepo-Coupe, N., Pereira, L., Teodoro, G. S., Saleska, S. R.,  
738 Borma, L. S., Christoffersen, B. O., Penha, D., Alves, L. F., Lima, A. J. N., Carneiro, V. M. C., Gentine, P., Lee, J.-  
739 E., Aragão, L. E. O. C., Ivanov, V., Leal, L. S. M., Araujo, A. C., and Oliveira, R. S.: Hydraulic traits explain  
740 differential responses of Amazonian forests to the 2015 El Niño-induced drought, *New Phytologist*, 223,  
741 1253–1266, <https://doi.org/10.1111/nph.15909>, 2019.
- 742 Bauman, D., Fortunel, C., Delhay, G., Malhi, Y., Cernusak, L. A., Bentley, L. P., Rifai, S. W., Aguirre-Gutiérrez,  
743 J., Menor, I. O., Phillips, O. L., McNellis, B. E., Bradford, M., Laurance, S. G. W., Hutchinson, M. F., Dempsey, R.,  
744 Santos-Andrade, P. E., Ninantay-Rivera, H. R., Chambi Paucar, J. R., and McMahon, S. M.: Tropical tree  
745 mortality has increased with rising atmospheric water stress, *Nature*, 1–6, [https://doi.org/10.1038/s41586-](https://doi.org/10.1038/s41586-022-04737-7)  
746 022-04737-7, 2022.
- 747 de Boer-Euser, T., McMillan, H. K., Hrachowitz, M., Winsemius, H. C., and Savenije, H. H. G.: Influence of soil  
748 and climate on root zone storage capacity, *Water Resources Research*, 52, 2009–2024,  
749 <https://doi.org/10.1002/2015WR018115>, 2016.
- 750 Bouaziz, L. J. E., Steele-Dunne, S. C., Schellekens, J., Weerts, A. H., Stam, J., Sprokkereef, E., Winsemius, H. H.  
751 C., Savenije, H. H. G., and Hrachowitz, M.: Improved Understanding of the Link Between Catchment-Scale  
752 Vegetation Accessible Storage and Satellite-Derived Soil Water Index, *Water Resources Research*, 56,  
753 e2019WR026365, <https://doi.org/10.1029/2019WR026365>, 2020.
- 754 Boulton, C. A., Good, P., and Lenton, T. M.: Early warning signals of simulated Amazon rainforest dieback,  
755 *Theor Ecol*, 6, 373–384, <https://doi.org/10.1007/s12080-013-0191-7>, 2013.

- 756 Boulton, C. A., Booth, B. B. B., and Good, P.: Exploring uncertainty of Amazon dieback in a perturbed  
757 parameter Earth system ensemble, *Global Change Biology*, 23, 5032–5044,  
758 <https://doi.org/10.1111/gcb.13733>, 2017.
- 759 Boulton, C. A., Lenton, T. M., and Boers, N.: Pronounced loss of Amazon rainforest resilience since the early  
760 2000s, *Nat. Clim. Chang.*, 12, 271–278, <https://doi.org/10.1038/s41558-022-01287-8>, 2022.
- 761 Bovolo, C. I., Wagner, T., Parkin, G., Hein-Griggs, D., Pereira, R., and Jones, R.: The Guiana Shield rainforests—  
762 overlooked guardians of South American climate, *Environ. Res. Lett.*, 13, 074029,  
763 <https://doi.org/10.1088/1748-9326/aacf60>, 2018.
- 764 Brienen, R. J. W., Phillips, O. L., Feldpausch, T. R., Gloor, E., Baker, T. R., Lloyd, J., Lopez-Gonzalez, G.,  
765 Monteagudo-Mendoza, A., Malhi, Y., Lewis, S. L., Vásquez Martínez, R., Alexiades, M., Álvarez Dávila, E.,  
766 Alvarez-Loayza, P., Andrade, A., Aragão, L. E. O. C., Araujo-Murakami, A., Arets, E. J. M. M., Arroyo, L., Aymard  
767 C., G. A., Bánki, O. S., Baraloto, C., Barroso, J., Bonal, D., Boot, R. G. A., Camargo, J. L. C., Castilho, C. V., Chama,  
768 V., Chao, K. J., Chave, J., Comiskey, J. A., Cornejo Valverde, F., da Costa, L., de Oliveira, E. A., Di Fiore, A., Erwin,  
769 T. L., Fauset, S., Forsthofer, M., Galbraith, D. R., Grahame, E. S., Groot, N., Hérault, B., Higuchi, N., Honorio  
770 Coronado, E. N., Keeling, H., Killeen, T. J., Laurance, W. F., Laurance, S., Licona, J., Magnussen, W. E., Marimon,  
771 B. S., Marimon-Junior, B. H., Mendoza, C., Neill, D. A., Nogueira, E. M., Núñez, P., Pallqui Camacho, N. C.,  
772 Parada, A., Pardo-Molina, G., Peacock, J., Peña-Claros, M., Pickavance, G. C., Pitman, N. C. A., Poorter, L.,  
773 Prieto, A., Quesada, C. A., Ramírez, F., Ramírez-Angulo, H., Restrepo, Z., Roopsind, A., Rudas, A., Salomão, R.  
774 P., Schwarz, M., Silva, N., Silva-Espejo, J. E., Silveira, M., Stropp, J., Talbot, J., ter Steege, H., Teran-Aguilar, J.,  
775 Terborgh, J., Thomas-Caesar, R., Toledo, M., Torello-Raventos, M., Umetsu, R. K., van der Heijden, G. M. F.,  
776 van der Hout, P., Guimarães Vieira, I. C., Vieira, S. A., Vilanova, E., Vos, V. A., and Zagt, R. J.: Long-term decline  
777 of the Amazon carbon sink, *Nature*, 519, 344–348, <https://doi.org/10.1038/nature14283>, 2015.
- 778 Bronick, C. J. and Lal, R.: Soil structure and management: a review, *Geoderma*, 124, 3–22,  
779 <https://doi.org/10.1016/j.geoderma.2004.03.005>, 2005.
- 780 Brooks, P. D., Chorover, J., Fan, Y., Godsey, S. E., Maxwell, R. M., McNamara, J. P., and Tague, C.: Hydrological  
781 partitioning in the critical zone: Recent advances and opportunities for developing transferable understanding  
782 of water cycle dynamics, *Water Resources Research*, 51, 6973–6987,  
783 <https://doi.org/10.1002/2015WR017039>, 2015.
- 784 Brum, M., Vadeboncoeur, M. A., Ivanov, V., Asbjornsen, H., Saleska, S., Alves, L. F., Penha, D., Dias, J. D.,  
785 Aragão, L. E. O. C., Barros, F., Bittencourt, P., Pereira, L., and Oliveira, R. S.: Hydrological niche segregation  
786 defines forest structure and drought tolerance strategies in a seasonal Amazon forest, *Journal of Ecology*, 107,  
787 318–333, <https://doi.org/10.1111/1365-2745.13022>, 2019.
- 788 Brunner, I., Herzog, C., Dawes, M. A., Arend, M., and Sperisen, C.: How tree roots respond to drought,  
789 *Frontiers in Plant Science*, 6, 2015.
- 790 Bruno, R. D., Rocha, H. R. da, Freitas, H. C. de, Goulden, M. L., and Miller, S. D.: Soil moisture dynamics in an  
791 eastern Amazonian tropical forest, *Hydrological Processes*, 20, 2477–2489, <https://doi.org/10.1002/hyp.6211>,  
792 2006.
- 793 Canadell, J., Jackson, R. B., Ehleringer, J. B., Mooney, H. A., Sala, O. E., and Schulze, E.-D.: Maximum rooting  
794 depth of vegetation types at the global scale, *Oecologia*, 108, 583–595, <https://doi.org/10.1007/BF00329030>,  
795 1996.
- 796 Canadell, J. G., Monteiro, P. M. S., Costa, M. H., Cunha, L. C. D., Cox, P. M., Eliseev, A. V., Henson, S., Ishii, M.,  
797 Jaccard, S., Koven, C., Lohila, A., Patra, P. K., Piao, S., Syampungani, S., Zaehle, S., Zickfeld, K., Alexandrov, G.  
798 A., Bala, G., Bopp, L., Boysen, L., Cao, L., Chandra, N., Ciais, P., Denisov, S. N., Dentener, F. J., Douville, H., Fay,

799 A., Forster, P., Fox-Kemper, B., Friedlingstein, P., Fu, W., Fuss, S., Garçon, V., Gier, B., Gillett, N. P., Gregor, L.,  
800 Haustein, K., Haverd, V., He, J., Hewitt, H. T., Hoffman, F. M., Ilyina, T., Jackson, R., Jones, C., Keller, D. P.,  
801 Kwiatkowski, L., Lamboll, R. D., Lan, X., Lauffkötter, C., Quéré, C. L., Lenton, A., Lewis, J., Liddicoat, S.,  
802 Lorenzoni, L., Lovenduski, N., Macdougall, A. H., Mathesius, S., Matthews, D. H., Meinshausen, M., Mokhov, I.  
803 I., Naik, V., Nicholls, Z. R. J., Nurhati, I. S., O'sullivan, M., Peters, G., Pongratz, J., Poulter, B., Sallée, J.-B.,  
804 Saunois, M., Schuur, E. A. G., Seneviratne, S., Stavert, A., Suntharalingam, P., Tachiiri, K., Terhaar, J.,  
805 Thompson, R., Tian, H., Turnbull, J., Vicente-Serrano, S. M., Wang, X., Wanninkhof, R. H., Williamson, P.,  
806 Brovkin, V., Feely, R. A., and Lebehot, A. D.: Global Carbon and other Biogeochemical Cycles and Feedbacks,  
807 in: IPCC AR6 WGI, Final Government Distribution, chapter 5, 2021.

808 Chai, Y., Martins, G., Nobre, C., von Randow, C., Chen, T., and Dolman, H.: Constraining Amazonian land  
809 surface temperature sensitivity to precipitation and the probability of forest dieback, *npj Clim Atmos Sci*, 4, 1–  
810 7, <https://doi.org/10.1038/s41612-021-00162-1>, 2021.

811 Cheng, S., Huang, J., Ji, F., and Lin, L.: Uncertainties of soil moisture in historical simulations and future  
812 projections, *Journal of Geophysical Research: Atmospheres*, 122, 2239–2253,  
813 <https://doi.org/10.1002/2016JD025871>, 2017.

814 Cole, L. E. S., Bhagwat, S. A., and Willis, K. J.: Recovery and resilience of tropical forests after disturbance,  
815 *Nature Communications*, 5, 3906, <https://doi.org/10.1038/ncomms4906>, 2014.

816 Condit, R., Engelbrecht, B. M. J., Pino, D., Pérez, R., and Turner, B. L.: Species distributions in response to  
817 individual soil nutrients and seasonal drought across a community of tropical trees, *PNAS*, 110, 5064–5068,  
818 <https://doi.org/10.1073/pnas.1218042110>, 2013.

819 Cook, K. H. and Vizzy, E. K.: Impact of climate change on mid-twenty-first century growing seasons in Africa,  
820 *Clim Dyn*, 39, 2937–2955, <https://doi.org/10.1007/s00382-012-1324-1>, 2012.

821 Cooper, G. S., Willcock, S., and Dearing, J. A.: Regime shifts occur disproportionately faster in larger  
822 ecosystems, *Nature Communications*, 11, 1175, <https://doi.org/10.1038/s41467-020-15029-x>, 2020.

823 skextremes Documentation: <https://github.com/kikocorreoso/scikit-extremes>.

824 Cox, P. M., Betts, R. A., Collins, M., Harris, P. P., Huntingford, C., and Jones, C. D.: Amazonian forest dieback  
825 under climate-carbon cycle projections for the 21st century, *Theor Appl Climatol*, 78, 137–156,  
826 <https://doi.org/10.1007/s00704-004-0049-4>, 2004.

827 Dai, A.: Drought under global warming: a review, *WIREs Climate Change*, 2, 45–65,  
828 <https://doi.org/10.1002/wcc.81>, 2011.

829 Davidson, E. A., de Araújo, A. C., Artaxo, P., Balch, J. K., Brown, I. F., C. Bustamante, M. M., Coe, M. T., DeFries,  
830 R. S., Keller, M., Longo, M., Munger, J. W., Schroeder, W., Soares-Filho, B. S., Souza, C. M., and Wofsy, S. C.:  
831 The Amazon basin in transition, *Nature*, 481, 321–328, <https://doi.org/10.1038/nature10717>, 2012.

832 Dexter, A. R.: Soil physical quality: Part II. Friability, tillage, tilth and hard-setting, *Geoderma*, 120, 215–225,  
833 <https://doi.org/10.1016/j.geoderma.2003.09.005>, 2004.

834 Dittert, K., Wätzel, J., and Sattelmacher, B.: Responses of *Alnus glutinosa* to Anaerobic Conditions -  
835 Mechanisms and Rate of Oxygen Flux into the Roots, *Plant Biology*, 8, 212–223, [https://doi.org/10.1055/s-](https://doi.org/10.1055/s-2005-873041)  
836 2005-873041, 2006.

837 Doughty, C. E., Keany, J. M., Wiebe, B. C., Rey-Sanchez, C., Carter, K. R., Middleby, K. B., Cheesman, A. W.,  
838 Goulden, M. L., da Rocha, H. R., Miller, S. D., Malhi, Y., Fauset, S., Gloor, E., Slot, M., Oliveras Menor, I., Crous,

- 839 K. Y., Goldsmith, G. R., and Fisher, J. B.: Tropical forests are approaching critical temperature thresholds,  
840 *Nature*, 621, 105–111, <https://doi.org/10.1038/s41586-023-06391-z>, 2023.
- 841 Drijfhout, S., Bathiany, S., Beaulieu, C., Brovkin, V., Claussen, M., Huntingford, C., Scheffer, M., Sgubin, G., and  
842 Swingedouw, D.: Catalogue of abrupt shifts in Intergovernmental Panel on Climate Change climate models,  
843 *Proceedings of the National Academy of Sciences*, 112, E5777–E5786,  
844 <https://doi.org/10.1073/pnas.1511451112>, 2015.
- 845 Dunning, C. M., Black, E., and Allan, R. P.: Later Wet Seasons with More Intense Rainfall over Africa under  
846 Future Climate Change, *Journal of Climate*, 31, 9719–9738, 2018.
- 847 van der Ent, R. J., Savenije, H. H. G., Schaefli, B., and Steele-Dunne, S. C.: Origin and fate of atmospheric  
848 moisture over continents, *Water Resources Research*, 46, <https://doi.org/10.1029/2010WR009127>, 2010.
- 849 GlobCover land-use map: [http://due.esrin.esa.int/page\\_globcover.php](http://due.esrin.esa.int/page_globcover.php), last access: 27 February 2022.
- 850 Esquivel-Muelbert, A., Baker, T. R., Dexter, K. G., Lewis, S. L., Brienen, R. J. W., Feldpausch, T. R., Lloyd, J.,  
851 Monteagudo-Mendoza, A., Arroyo, L., Álvarez-Dávila, E., Higuchi, N., Marimon, B. S., Marimon-Junior, B. H.,  
852 Silveira, M., Vilanova, E., Gloor, E., Malhi, Y., Chave, J., Barlow, J., Bonal, D., Cardozo, N. D., Erwin, T., Fauset,  
853 S., Hérault, B., Laurance, S., Poorter, L., Qie, L., Stahl, C., Sullivan, M. J. P., Steege, H. ter, Vos, V. A., Zuidema,  
854 P. A., Almeida, E., Oliveira, E. A. de, Andrade, A., Vieira, S. A., Aragão, L., Araujo-Murakami, A., Arets, E., C, G.  
855 A. A., Baraloto, C., Camargo, P. B., Barroso, J. G., Bongers, F., Boot, R., Camargo, J. L., Castro, W., Moscoso, V.  
856 C., Comiskey, J., Valverde, F. C., Costa, A. C. L. da, Pasquel, J. del A., Fiore, A. D., Duque, L. F., Elias, F., Engel, J.,  
857 Llampazo, G. F., Galbraith, D., Fernández, R. H., Coronado, E. H., Hubau, W., Jimenez-Rojas, E., Lima, A. J. N.,  
858 Umetsu, R. K., Laurance, W., Lopez-Gonzalez, G., Lovejoy, T., Cruz, O. A. M., Morandi, P. S., Neill, D., Vargas, P.  
859 N., Camacho, N. C. P., Gutierrez, A. P., Pardo, G., Peacock, J., Peña-Claros, M., Peñuela-Mora, M. C., Petronelli,  
860 P., Pickavance, G. C., Pitman, N., Prieto, A., Quesada, C., Ramírez-Angulo, H., Réjou-Méchain, M., Correa, Z. R.,  
861 Roopsind, A., Rudas, A., Salomão, R., Silva, N., Espejo, J. S., Singh, J., Stropp, J., Terborgh, J., Thomas, R.,  
862 Toledo, M., Torres-Lezama, A., Gamarra, L. V., Meer, P. J. van de, Heijden, G. van der, et al.: Compositional  
863 response of Amazon forests to climate change, *Global Change Biology*, 25, 39–56,  
864 <https://doi.org/10.1111/gcb.14413>, 2019.
- 865 Fan, Y., Miguez-Macho, G., Jobbágy, E. G., Jackson, R. B., and Otero-Casal, C.: Hydrologic regulation of plant  
866 rooting depth, *Proceedings of the National Academy of Sciences*, 114, 10572–10577,  
867 <https://doi.org/10.1073/pnas.1712381114>, 2017.
- 868 Fayos, C. B.: The roles of texture and structure in the water retention capacity of burnt Mediterranean soils  
869 with varying rainfall, *CATENA*, 31, 219–236, [https://doi.org/10.1016/S0341-8162\(97\)00041-6](https://doi.org/10.1016/S0341-8162(97)00041-6), 1997.
- 870 Ferreira, D., Marshall, J., and Rose, B.: Climate Determinism Revisited: Multiple Equilibria in a Complex  
871 Climate Model, *Journal of Climate*, 24, 992–1012, <https://doi.org/10.1175/2010JCLI3580.1>, 2011.
- 872 Fleischer, K., Rammig, A., De Kauwe, M. G., Walker, A. P., Domingues, T. F., Fuchslueger, L., Garcia, S., Goll, D.  
873 S., Grandis, A., Jiang, M., Haverd, V., Hofhansl, F., Holm, J. A., Kruijt, B., Leung, F., Medlyn, B. E., Mercado, L.  
874 M., Norby, R. J., Pak, B., von Randow, C., Quesada, C. A., Schaap, K. J., Valverde-Barrantes, O. J., Wang, Y.-P.,  
875 Yang, X., Zaehle, S., Zhu, Q., and Lapola, D. M.: Amazon forest response to CO<sub>2</sub> fertilization dependent on  
876 plant phosphorus acquisition, *Nat. Geosci.*, 12, 736–741, <https://doi.org/10.1038/s41561-019-0404-9>, 2019.
- 877 Flores, B. M., Montoya, E., Sakschewski, B., Nascimento, N., Staal, A., Betts, R. A., Levis, C., Lapola, D. M.,  
878 Esquivel-Muelbert, A., Jakovac, C., Nobre, C. A., Oliveira, R. S., Borma, L. S., Nian, D., Boers, N., Hecht, S. B., ter  
879 Steege, H., Arieira, J., Lucas, I. L., Berenguer, E., Marengo, J. A., Gatti, L. V., Mattos, C. R. C., and Hirota, M.:  
880 Critical transitions in the Amazon forest system, *Nature*, 626, 555–564, <https://doi.org/10.1038/s41586-023-06970-0>,  
881 2024.



882 Funk, C., Peterson, P., Landsfeld, M., Pedreros, D., Verdin, J., Shukla, S., Husak, G., Rowland, J., Harrison, L.,  
883 Hoell, A., and Michaelsen, J.: The climate hazards infrared precipitation with stations—a new environmental  
884 record for monitoring extremes, *Scientific Data*, 2, 150066, <https://doi.org/10.1038/sdata.2015.66>, 2015.

885 Gao, H., Hrachowitz, M., Schymanski, S. J., Fenicia, F., Sriwongsitanon, N., and Savenije, H. H. G.: Climate  
886 controls how ecosystems size the root zone storage capacity at catchment scale: Root zone storage capacity in  
887 catchments, *Geophysical Research Letters*, 41, 7916–7923, <https://doi.org/10.1002/2014GL061668>, 2014.

888 Grimm, N. B., Chapin III, F. S., Bierwagen, B., Gonzalez, P., Groffman, P. M., Luo, Y., Melton, F., Nadelhoffer, K.,  
889 Pairis, A., Raymond, P. A., Schimel, J., and Williamson, C. E.: The impacts of climate change on ecosystem  
890 structure and function, *Frontiers in Ecology and the Environment*, 11, 474–482,  
891 <https://doi.org/10.1890/120282>, 2013.

892 Gumbel, E. J.: *Statistics of extremes.*, Columbia University Press, New York, 1958.

893 Guswa, A. J.: The influence of climate on root depth: A carbon cost-benefit analysis, *Water Resources*  
894 *Research*, 44, W02427, <https://doi.org/10.1029/2007WR006384>, 2008.

895 Hahm, W. J., Rempe, D. M., Dralle, D. N., Dawson, T. E., Lovill, S. M., Bryk, A. B., Bish, D. L., Schieber, J., and  
896 Dietrich, W. E.: Lithologically Controlled Subsurface Critical Zone Thickness and Water Storage Capacity  
897 Determine Regional Plant Community Composition, *Water Resources Research*, 55, 3028–3055,  
898 <https://doi.org/10.1029/2018WR023760>, 2019.

899 Hall, A., Cox, P., Huntingford, C., and Klein, S.: Progressing emergent constraints on future climate change,  
900 *Nat. Clim. Chang.*, 9, 269–278, <https://doi.org/10.1038/s41558-019-0436-6>, 2019.

901 Hersbach, H., Bell, B., Berrisford, P., Hirahara, S., Horányi, A., Muñoz-Sabater, J., Nicolas, J., Peubey, C., Radu,  
902 R., Schepers, D., Simmons, A., Soci, C., Abdalla, S., Abellan, X., Balsamo, G., Bechtold, P., Biavati, G., Bidlot, J.,  
903 Bonavita, M., Chiara, G. D., Dahlgren, P., Dee, D., Diamantakis, M., Dragani, R., Flemming, J., Forbes, R.,  
904 Fuentes, M., Geer, A., Haimberger, L., Healy, S., Hogan, R. J., Hólm, E., Janisková, M., Keeley, S., Laloyaux, P.,  
905 Lopez, P., Lupu, C., Radnoti, G., Rosnay, P. de, Rozum, I., Vamborg, F., Villaume, S., and Thépaut, J.-N.: The  
906 ERA5 Global Reanalysis, *Quarterly Journal of the Royal Meteorological Society*, 245, 111840,  
907 <https://doi.org/10.1002/qj.3803>, 2020.

908 Higgins, S. I. and Scheiter, S.: Atmospheric CO<sub>2</sub> forces abrupt vegetation shifts locally, but not globally, *Nature*,  
909 488, 209–212, <https://doi.org/10.1038/nature11238>, 2012.

910 Hildebrandt, A., Kleidon, A., and Bechmann, M.: A thermodynamic formulation of root water uptake,  
911 *Hydrology and Earth System Sciences*, 20, 3441–3454, <https://doi.org/10.5194/hess-20-3441-2016>, 2016.

912 Hirota, M., Holmgren, M., Van Nes, E. H., and Scheffer, M.: Global Resilience of Tropical Forest and Savanna to  
913 Critical Transitions, *Science*, 334, 232–235, <https://doi.org/10.1126/science.1210657>, 2011.

914 Hirota, M., Flores, B. M., Betts, R., Borma, L. S., Esquivel-Muelbert, A., Jakovac, C., Lapola, D. M., Montoya, E.,  
915 Oliveira, R. S., and Sakschewski, B.: Chapter 24: Resilience of the Amazon forest to global changes: Assessing  
916 the risk of tipping points, in: *Amazon Assessment Report 2021*, edited by: Nobre, C., Encalada, A., Anderson,  
917 E., Roca Alcazar, F. H., Bustamante, M., Mena, C., Peña-Claros, M., Poveda, G., Rodriguez, J. P., Saleska, S.,  
918 Trumbore, S. E., Val, A., Villa Nova, L., Abramovay, R., Alencar, A., Rodriguez Alza, A. C., Armenteras, D.,  
919 Artaxo, P., Athayde, S., Barretto Filho, H. T., Barlow, J., Berenguer, E., Bortolotto, F., Costa, F. de A., Costa, M.  
920 H., Cuvi, N., Fearnside, P., Ferreira, J., Flores, B. M., Frieri, S., Gatti, L. V., Guayasamin, J. M., Hecht, S., Hirota,  
921 M., Hoorn, C., Josse, C., Lapola, D. M., Larrea, C., Larrea-Alcazar, D. M., Lehm Ardaya, Z., Malhi, Y., Marengo, J.  
922 A., Melack, J., Moraes R., M., Moutinho, P., Murmis, M. R., Neves, E. G., Paez, B., Painter, L., Ramos, A.,

- 923 Rosero-Peña, M. C., Schminck, M., Sist, P., ter Steege, H., Val, P., van der Voort, H., Varese, M., and Zapata-  
924 Ríos, G., UN Sustainable Development Solutions Network (SDSN), <https://doi.org/10.55161/QPYS9758>, 2021.
- 925 Hofhansl, F., Andersen, K. M., Fleischer, K., Fuchslueger, L., Rammig, A., Schaap, K. J., Valverde-Barrantes, O.  
926 J., and Lapola, D. M.: Amazon Forest Ecosystem Responses to Elevated Atmospheric CO<sub>2</sub> and Alterations in  
927 Nutrient Availability: Filling the Gaps with Model-Experiment Integration, *Frontiers in Earth Science*, 4, 2016.
- 928 Hubau, W., Lewis, S. L., Phillips, O. L., Affum-Baffoe, K., Beeckman, H., Cuní-Sánchez, A., Daniels, A. K.,  
929 Ewango, C. E. N., Fauset, S., Mukinzi, J. M., Sheil, D., Sonké, B., Sullivan, M. J. P., Sunderland, T. C. H.,  
930 Taedoumg, H., Thomas, S. C., White, L. J. T., Abernethy, K. A., Adu-Bredu, S., Amani, C. A., Baker, T. R., Banin,  
931 L. F., Baya, F., Begne, S. K., Bennett, A. C., Benedet, F., Bitariho, R., Bocko, Y. E., Boeckx, P., Boundja, P.,  
932 Brienen, R. J. W., Brncic, T., Chezeaux, E., Chuyong, G. B., Clark, C. J., Collins, M., Comiskey, J. A., Coomes, D.  
933 A., Dargie, G. C., de Haulleville, T., Kamdem, M. N. D., Doucet, J.-L., Esquivel-Muelbert, A., Feldpausch, T. R.,  
934 Fofanah, A., Foli, E. G., Gilpin, M., Gloor, E., Gonmadje, C., Gourlet-Fleury, S., Hall, J. S., Hamilton, A. C., Harris,  
935 D. J., Hart, T. B., Hockemba, M. B. N., Hladik, A., Ifo, S. A., Jeffery, K. J., Jucker, T., Yakusu, E. K., Kearsley, E.,  
936 Kenfack, D., Koch, A., Leal, M. E., Levesley, A., Lindsell, J. A., Lisingo, J., Lopez-Gonzalez, G., Lovett, J. C.,  
937 Makana, J.-R., Malhi, Y., Marshall, A. R., Martin, J., Martin, E. H., Mbayu, F. M., Medjibe, V. P., Mihindou, V.,  
938 Mitchard, E. T. A., Moore, S., Munishi, P. K. T., Bengone, N. N., Ojo, L., Ondo, F. E., Peh, K. S.-H., Pickavance, G.  
939 C., Poulsen, A. D., Poulsen, J. R., Qie, L., Reitsma, J., Rovero, F., Swaine, M. D., Talbot, J., Taplin, J., Taylor, D.  
940 M., Thomas, D. W., Toirambe, B., Mukendi, J. T., Tuagben, D., Umunay, P. M., et al.: Asynchronous carbon sink  
941 saturation in African and Amazonian tropical forests, *Nature*, 579, 80–87, <https://doi.org/10.1038/s41586-020-2035-0>, 2020.
- 943 Huntingford, C., Zelazowski, P., Galbraith, D., Mercado, L. M., Sitch, S., Fisher, R., Lomas, M., Walker, A. P.,  
944 Jones, C. D., Booth, B. B. B., Malhi, Y., Hemming, D., Kay, G., Good, P., Lewis, S. L., Phillips, O. L., Atkin, O. K.,  
945 Lloyd, J., Gloor, E., Zaragoza-Castells, J., Meir, P., Betts, R., Harris, P. P., Nobre, C., Marengo, J., and Cox, P. M.:  
946 Simulated resilience of tropical rainforests to CO<sub>2</sub>-induced climate change, *Nature Geosci*, 6, 268–273,  
947 <https://doi.org/10.1038/ngeo1741>, 2013.
- 948 Hurtt, G. C., Chini, L., Sahajpal, R., Frohling, S., Bodirsky, B. L., Calvin, K., Doelman, J. C., Fisk, J., Fujimori, S.,  
949 Klein Goldewijk, K., Hasegawa, T., Havlik, P., Heinemann, A., Humpenöder, F., Jungclaus, J., Kaplan, J. O.,  
950 Kennedy, J., Krisztin, T., Lawrence, D., Lawrence, P., Ma, L., Mertz, O., Pongratz, J., Popp, A., Poulter, B., Riahi,  
951 K., Shevliakova, E., Stehfest, E., Thornton, P., Tubiello, F. N., van Vuuren, D. P., and Zhang, X.: Harmonization  
952 of global land use change and management for the period 850–2100 (LUH2) for CMIP6, *Geoscientific Model  
953 Development*, 13, 5425–5464, <https://doi.org/10.5194/gmd-13-5425-2020>, 2020.
- 954 Indoria, A. K., Sharma, K. L., and Reddy, K. S.: Chapter 18 - Hydraulic properties of soil under warming climate,  
955 in: *Climate Change and Soil Interactions*, edited by: Prasad, M. N. V. and Pietrzykowski, M., Elsevier, 473–508,  
956 <https://doi.org/10.1016/B978-0-12-818032-7.00018-7>, 2020.
- 957 Jach, L., Warrach-Sagi, K., Ingwersen, J., Kaas, E., and Wulfmeyer, V.: Land Cover Impacts on Land-Atmosphere  
958 Coupling Strength in Climate Simulations With WRF Over Europe, *Journal of Geophysical Research:  
959 Atmospheres*, 125, e2019JD031989, <https://doi.org/10.1029/2019JD031989>, 2020.
- 960 Jackson, R. B., Canadell, J., Ehleringer, J. R., Mooney, H. A., Sala, O. E., and Schulze, E. D.: A global analysis of  
961 root distributions for terrestrial biomes, *Oecologia*, 108, 389–411, <https://doi.org/10.1007/BF00333714>,  
962 1996.
- 963 Jehn, F. U., Kemp, L., Ilin, E., Funk, C., Wang, J. R., and Breuer, L.: Focus of the IPCC Assessment Reports Has  
964 Shifted to Lower Temperatures, *Earth's Future*, 10, e2022EF002876, <https://doi.org/10.1029/2022EF002876>,  
965 2022.

- 966 Jiang, C. and Ryu, Y.: Multi-scale evaluation of global gross primary productivity and evapotranspiration  
967 products derived from Breathing Earth System Simulator (BESS), *Remote Sensing of Environment*, 186, 528–  
968 547, <https://doi.org/10.1016/j.rse.2016.08.030>, 2016.
- 969 Jones, C., Lowe, J., Liddicoat, S., and Betts, R.: Committed terrestrial ecosystem changes due to climate  
970 change, *Nature Geosci*, 2, 484–487, <https://doi.org/10.1038/ngeo555>, 2009.
- 971 Jung, M., Koirala, S., Weber, U., Ichii, K., Gans, F., Camps-Valls, G., Papale, D., Schwalm, C., Tramontana, G.,  
972 and Reichstein, M.: The FLUXCOM ensemble of global land-atmosphere energy fluxes, *Sci Data*, 6, 74,  
973 <https://doi.org/10.1038/s41597-019-0076-8>, 2019.
- 974 Koch, A., Hubau, W., and Lewis, S. L.: Earth System Models Are Not Capturing Present-Day Tropical Forest  
975 Carbon Dynamics, *Earth's Future*, 9, e2020EF001874, <https://doi.org/10.1029/2020EF001874>, 2021.
- 976 Kooperman, G. J., Chen, Y., Hoffman, F. M., Koven, C. D., Lindsay, K., Pritchard, M. S., Swann, A. L. S., and  
977 Randerson, J. T.: Forest response to rising CO<sub>2</sub> drives zonally asymmetric rainfall change over tropical land,  
978 *Nature Clim Change*, 8, 434–440, <https://doi.org/10.1038/s41558-018-0144-7>, 2018.
- 979 Körner, C.: A matter of tree longevity, *Science*, 355, 130–131, <https://doi.org/10.1126/science.aal2449>, 2017.
- 980 Küçük, Ç., Koirala, S., Carvalhais, N., Miralles, D. G., Reichstein, M., and Jung, M.: Characterizing the Response  
981 of Vegetation Cover to Water Limitation in Africa Using Geostationary Satellites, *Journal of Advances in*  
982 *Modeling Earth Systems*, 14, e2021MS002730, <https://doi.org/10.1029/2021MS002730>, 2022.
- 983 Kukal, M. S. and Irmak, S.: Can limits of plant available water be inferred from soil moisture distributions?,  
984 *Agricultural & Environmental Letters*, 8, e20113, <https://doi.org/10.1002/ael2.20113>, 2023.
- 985 Lammertsma, E. I., Boer, H. J. de, Dekker, S. C., Dilcher, D. L., Lotter, A. F., and Wagner-Cremer, F.: Global CO<sub>2</sub>  
986 rise leads to reduced maximum stomatal conductance in Florida vegetation, *PNAS*, 108, 4035–4040,  
987 <https://doi.org/10.1073/pnas.1100371108>, 2011.
- 988 Lawrence, D., Coe, M., Walker, W., Verchot, L., and Vandecar, K.: The Unseen Effects of Deforestation:  
989 Biophysical Effects on Climate, *Frontiers in Forests and Global Change*, 5, 2022.
- 990 Leite-Filho, A. T., Soares-Filho, B. S., Davis, J. L., Abrahão, G. M., and Börner, J.: Deforestation reduces rainfall  
991 and agricultural revenues in the Brazilian Amazon, *Nat Commun*, 12, 2591, <https://doi.org/10.1038/s41467-021-22840-7>, 2021.
- 993 Lenton, T. M.: Early warning of climate tipping points, *Nature Clim Change*, 1, 201–209,  
994 <https://doi.org/10.1038/nclimate1143>, 2011.
- 995 Lenton, T. M., Rockström, J., Gaffney, O., Rahmstorf, S., Richardson, K., Steffen, W., and Schellnhuber, H. J.:  
996 Climate tipping points — too risky to bet against, *Nature*, 575, 592–595, <https://doi.org/10.1038/d41586-019-03595-0>, 2019.
- 998 Lewis, S. L., Edwards, D. P., and Galbraith, D.: Increasing human dominance of tropical forests, *Science*, 349,  
999 827–832, <https://doi.org/10.1126/science.aaa9932>, 2015.
- 1000 Li, Y., Brando, P. M., Morton, D. C., Lawrence, D. M., Yang, H., and Randerson, J. T.: Deforestation-induced  
1001 climate change reduces carbon storage in remaining tropical forests, *Nat Commun*, 13, 1964,  
1002 <https://doi.org/10.1038/s41467-022-29601-0>, 2022.

- 1003 Liu, W., Sun, F., Lim, W. H., Zhang, J., Wang, H., Shiogama, H., and Zhang, Y.: Global drought and severe  
1004 drought-affected populations in 1.5 and 2 °C warmer worlds, *Earth System Dynamics*, 9, 267–283,  
1005 <https://doi.org/10.5194/esd-9-267-2018>, 2018.
- 1006 Liu, Y., Kumar, M., Katul, G. G., Feng, X., and Konings, A. G.: Plant hydraulics accentuates the effect of  
1007 atmospheric moisture stress on transpiration, *Nat. Clim. Chang.*, 10, 691–695,  
1008 <https://doi.org/10.1038/s41558-020-0781-5>, 2020.
- 1009 Ma, L., Hurtt, G. C., Chini, L. P., Sahajpal, R., Pongratz, J., Frohling, S., Stehfest, E., Klein Goldewijk, K., O’Leary,  
1010 D., and Doelman, J. C.: Global rules for translating land-use change (LUH2) to land-cover change for CMIP6  
1011 using GLM2, *Geoscientific Model Development*, 13, 3203–3220, <https://doi.org/10.5194/gmd-13-3203-2020>,  
1012 2020.
- 1013 Malhi, Y., Roberts, J. T., Betts, R. A., Killeen, T. J., Li, W., and Nobre, C. A.: Climate Change, Deforestation, and  
1014 the Fate of the Amazon, *Science*, 319, 169–172, <https://doi.org/10.1126/science.1146961>, 2008.
- 1015 Malhi, Y., Gardner, T. A., Goldsmith, G. R., Silman, M. R., and Zelazowski, P.: Tropical Forests in the  
1016 Anthropocene, *Annu. Rev. Environ. Resour.*, 39, 125–159, <https://doi.org/10.1146/annurev-environ-030713-155141>, 2014.
- 1018 Mamalakis, A., Randerson, J. T., Yu, J.-Y., Pritchard, M. S., Magnusdottir, G., Smyth, P., Levine, P. A., Yu, S., and  
1019 Foufoula-Georgiou, E.: Zonally contrasting shifts of the tropical rain belt in response to climate change, *Nature  
1020 Climate Change*, 11, 143–151, <https://doi.org/10.1038/s41558-020-00963-x>, 2021.
- 1021 Maslin, M. and Austin, P.: Climate models at their limit?, *Nature*, 486, 183–184,  
1022 <https://doi.org/10.1038/486183a>, 2012.
- 1023 McCormick, E. L., Dralle, D. N., Hahn, W. J., Tune, A. K., Schmidt, L. M., Chadwick, K. D., and Rempe, D. M.:  
1024 Widespread woody plant use of water stored in bedrock, *Nature*, 597, 225–229,  
1025 <https://doi.org/10.1038/s41586-021-03761-3>, 2021.
- 1026 McFarlane, N.: Parameterizations: representing key processes in climate models without resolving them,  
1027 *WIREs Climate Change*, 2, 482–497, <https://doi.org/10.1002/wcc.122>, 2011.
- 1028 Nepstad, D. C., Verssimo, A., Alencar, A., Nobre, C., Lima, E., Lefebvre, P., Schlesinger, P., Potter, C., Moutinho,  
1029 P., Mendoza, E., Cochrane, M., and Brooks, V.: Large-scale impoverishment of Amazonian forests by logging  
1030 and fire, *Nature*, 398, 505–508, <https://doi.org/10.1038/19066>, 1999.
- 1031 van Nes, E. H., Arani, B. M. S., Staal, A., van der Bolt, B., Flores, B. M., Bathiany, S., and Scheffer, M.: What Do  
1032 You Mean, ‘Tipping Point’?, *Trends in Ecology & Evolution*, 31, 902–904,  
1033 <https://doi.org/10.1016/j.tree.2016.09.011>, 2016.
- 1034 Nijzink, R., Hutton, C., Pechlivanidis, I., Capell, R., Arheimer, B., Freer, J., Han, D., Wagener, T., McGuire, K.,  
1035 Savenije, H., and Hrachowitz, M.: The evolution of root-zone moisture capacities after deforestation: a step  
1036 towards hydrological predictions under change?, *Hydrology and Earth System Sciences*, 20, 4775–4799,  
1037 <https://doi.org/10.5194/hess-20-4775-2016>, 2016.
- 1038 Nippert, J. B. and Holdo, R. M.: Challenging the maximum rooting depth paradigm in grasslands and savannas,  
1039 *Functional Ecology*, 29, 739–745, <https://doi.org/10.1111/1365-2435.12390>, 2015.
- 1040 Nof, D.: Simple Versus Complex Climate Modeling, *Eos, Transactions American Geophysical Union*, 89, 544–  
1041 545, <https://doi.org/10.1029/2008EO520006>, 2008.

- 1042 Oliveira, R. S., Dawson, T. E., Burgess, S. S. O., and Nepstad, D. C.: Hydraulic redistribution in three Amazonian  
1043 trees, *Oecologia*, 145, 354–363, <https://doi.org/10.1007/s00442-005-0108-2>, 2005.
- 1044 Parry, I. M., Ritchie, P. D. L., and Cox, P. M.: Evidence of localised Amazon rainforest dieback in CMIP6 models,  
1045 *Earth System Dynamics*, 13, 1667–1675, <https://doi.org/10.5194/esd-13-1667-2022>, 2022.
- 1046 Pascale, S., Carvalho, L. M. V., Adams, D. K., Castro, C. L., and Cavalcanti, I. F. A.: Current and Future Variations  
1047 of the Monsoons of the Americas in a Warming Climate, *Curr Clim Change Rep*, 5, 125–144,  
1048 <https://doi.org/10.1007/s40641-019-00135-w>, 2019.
- 1049 Piani, C., Weedon, G. P., Best, M., Gomes, S. M., Viterbo, P., Hagemann, S., and Haerter, J. O.: Statistical bias  
1050 correction of global simulated daily precipitation and temperature for the application of hydrological models,  
1051 *Journal of Hydrology*, 395, 199–215, <https://doi.org/10.1016/j.jhydrol.2010.10.024>, 2010.
- 1052 Poorter, L., Bongers, F., Aide, T. M., Almeyda Zambrano, A. M., Balvanera, P., Becknell, J. M., Boukili, V.,  
1053 Brancalion, P. H. S., Broadbent, E. N., Chazdon, R. L., Craven, D., de Almeida-Cortez, J. S., Cabral, G. A. L., de  
1054 Jong, B. H. J., Denslow, J. S., Dent, D. H., DeWalt, S. J., Dupuy, J. M., Durán, S. M., Espírito-Santo, M. M.,  
1055 Fandino, M. C., César, R. G., Hall, J. S., Hernandez-Stefanoni, J. L., Jakovac, C. C., Junqueira, A. B., Kennard, D.,  
1056 Letcher, S. G., Licona, J.-C., Lohbeck, M., Marín-Spiotta, E., Martínez-Ramos, M., Massoca, P., Meave, J. A.,  
1057 Mesquita, R., Mora, F., Muñoz, R., Muscarella, R., Nunes, Y. R. F., Ochoa-Gaona, S., de Oliveira, A. A., Orihuela-  
1058 Belmonte, E., Peña-Claros, M., Pérez-García, E. A., Piotta, D., Powers, J. S., Rodríguez-Velázquez, J., Romero-  
1059 Pérez, I. E., Ruíz, J., Saldarriaga, J. G., Sanchez-Azofeifa, A., Schwartz, N. B., Steininger, M. K., Swenson, N. G.,  
1060 Toledo, M., Uriarte, M., van Breugel, M., van der Wal, H., Veloso, M. D. M., Vester, H. F. M., Vicentini, A.,  
1061 Vieira, I. C. G., Bentos, T. V., Williamson, G. B., and Rozendaal, D. M. A.: Biomass resilience of Neotropical  
1062 secondary forests, *Nature*, 530, 211–214, <https://doi.org/10.1038/nature16512>, 2016.
- 1063 Rammig, A.: Tropical carbon sinks are saturating at different times on different continents, *Nature*, 579, 38–  
1064 39, <https://doi.org/10.1038/d41586-020-00423-8>, 2020.
- 1065 Reyer, C. P. O., Brouwers, N., Rammig, A., Brook, B. W., Epila, J., Grant, R. F., Holmgren, M., Langerwisch, F.,  
1066 Leuzinger, S., Lucht, W., Medlyn, B., Pfeifer, M., Steinkamp, J., Vanderwel, M. C., Verbeeck, H., and Vilella, D.  
1067 M.: Forest resilience and tipping points at different spatio-temporal scales: approaches and challenges,  
1068 *Journal of Ecology*, 103, 5–15, <https://doi.org/10.1111/1365-2745.12337>, 2015.
- 1069 Rosas, T., Mencuccini, M., Barba, J., Cochard, H., Saura-Mas, S., and Martínez-Vilalta, J.: Adjustments and  
1070 coordination of hydraulic, leaf and stem traits along a water availability gradient, *New Phytologist*, 223, 632–  
1071 646, <https://doi.org/10.1111/nph.15684>, 2019.
- 1072 Schenk, H. J.: Soil depth, plant rooting strategies and species' niches, *New Phytologist*, 178, 223–225,  
1073 <https://doi.org/10.1111/j.1469-8137.2008.02427.x>, 2008.
- 1074 Schenk, H. J. and Jackson, R. B.: The Global Biogeography of Roots, *Ecological Monographs*, 72, 311–328,  
1075 [https://doi.org/10.1890/0012-9615\(2002\)072\[0311:TGBOR\]2.0.CO;2](https://doi.org/10.1890/0012-9615(2002)072[0311:TGBOR]2.0.CO;2), 2002.
- 1076 Schumacher, D. L., Keune, J., Dirmeyer, P., and Miralles, D. G.: Drought self-propagation in drylands due to  
1077 land–atmosphere feedbacks, *Nat. Geosci.*, 15, 262–268, <https://doi.org/10.1038/s41561-022-00912-7>, 2022.
- 1078 Singh, C.: Rooting for forest resilience : Implications of climate and land-use change on the tropical  
1079 rainforests, 2023.
- 1080 Singh, C., Wang-Erlandsson, L., Fetzer, I., Rockström, J., and van der Ent, R.: Rootzone storage capacity reveals  
1081 drought coping strategies along rainforest-savanna transitions, *Environ. Res. Lett.*, 15, 124021,  
1082 <https://doi.org/10.1088/1748-9326/abc377>, 2020.

- 1083 Singh, C., van der Ent, R., Wang-Erlandsson, L., and Fetzer, I.: Hydroclimatic adaptation critical to the resilience  
1084 of tropical forests, *Global Change Biology*, 28, 2930–2939, <https://doi.org/10.1111/gcb.16115>, 2022.
- 1085 Slik, J. W. F., Franklin, J., Arroyo-Rodríguez, V., Field, R., Aguilar, S., Aguirre, N., Ahumada, J., Aiba, S.-I., Alves,  
1086 L. F., K. A., Avella, A., Mora, F., Aymard C., G. A., Báez, S., Balvanera, P., Bastian, M. L., Bastin, J.-F., Bellingham,  
1087 P. J., van den Berg, E., da Conceição Bispo, P., Boeckx, P., Boehning-Gaese, K., Bongers, F., Boyle, B.,  
1088 Brambach, F., Brearley, F. Q., Brown, S., Chai, S.-L., Chazdon, R. L., Chen, S., Chhang, P., Chuyong, G., Ewango,  
1089 C., Coronado, I. M., Cristóbal-Azkarate, J., Culmsee, H., Damas, K., Dattaraja, H. S., Davidar, P., DeWalt, S. J.,  
1090 Din, H., Drake, D. R., Duque, A., Durigan, G., Eichhorn, K., Eler, E. S., Enoki, T., Ensslin, A., Fandohan, A. B.,  
1091 Farwig, N., Feeley, K. J., Fischer, M., Forshed, O., Garcia, Q. S., Garkoti, S. C., Gillespie, T. W., Gillet, J.-F.,  
1092 Gonmadje, C., Granzow-de la Cerda, I., Griffith, D. M., Grogan, J., Hakeem, K. R., Harris, D. J., Harrison, R. D.,  
1093 Hector, A., Hemp, A., Homeier, J., Hussain, M. S., Ibarra-Manríquez, G., Hanum, I. F., Imai, N., Jansen, P. A.,  
1094 Joly, C. A., Joseph, S., Kartawinata, K., Kearsley, E., Kelly, D. L., Kessler, M., Killeen, T. J., Kooyman, R. M.,  
1095 Laumonier, Y., Laurance, S. G., Laurance, W. F., Lawes, M. J., Letcher, S. G., Lindsell, J., Lovett, J., Lozada, J., Lu,  
1096 X., Lykke, A. M., Mahmud, K. B., Mahayani, N. P. D., Mansor, A., Marshall, A. R., Martin, E. H., Calderado Leal  
1097 Matos, D., Meave, J. A., Melo, F. P. L., Mendoza, Z. H. A., et al.: Phylogenetic classification of the world's  
1098 tropical forests, *Proceedings of the National Academy of Sciences*, 115, 1837–1842,  
1099 <https://doi.org/10.1073/pnas.1714977115>, 2018.
- 1100 Smith, C. W., Johnston, M. A., and Lorentz, S. A.: The effect of soil compaction on the water retention  
1101 characteristics of soils in forest plantations, *South African Journal of Plant and Soil*, 18, 87–97,  
1102 <https://doi.org/10.1080/02571862.2001.10634410>, 2001.
- 1103 Staal, A., Tuinenburg, O. A., Bosmans, J. H. C., Holmgren, M., van Nes, E. H., Scheffer, M., Zemp, D. C., and  
1104 Dekker, S. C.: Forest-rainfall cascades buffer against drought across the Amazon, *Nature Climate Change*, 8,  
1105 539–543, <https://doi.org/10.1038/s41558-018-0177-y>, 2018.
- 1106 Staal, A., Fetzer, I., Wang-Erlandsson, L., Bosmans, J. H. C., Dekker, S. C., van Nes, E. H., Rockström, J., and  
1107 Tuinenburg, O. A.: Hysteresis of tropical forests in the 21st century, *Nat Commun*, 11, 4978,  
1108 <https://doi.org/10.1038/s41467-020-18728-7>, 2020.
- 1109 Stevens, B. and Bony, S.: What Are Climate Models Missing?, *Science*, 340, 1053–1054,  
1110 <https://doi.org/10.1126/science.1237554>, 2013.
- 1111 Still, C. J., Berry, J. A., Collatz, G. J., and DeFries, R. S.: Global distribution of C3 and C4 vegetation: Carbon  
1112 cycle implications, *Global Biogeochemical Cycles*, 17, 6-1-6–14, <https://doi.org/10.1029/2001GB001807>,  
1113 2003.
- 1114 Stocker, B. D., Tumber-Dávila, S. J., Konings, A. G., Anderson, M. C., Hain, C., and Jackson, R. B.: Global  
1115 patterns of water storage in the rooting zones of vegetation, *Nat. Geosci.*, 1–7,  
1116 <https://doi.org/10.1038/s41561-023-01125-2>, 2023.
- 1117 Sveen, T. R., Hannula, S. E., and Bahram, M.: Microbial regulation of feedbacks to ecosystem change, *Trends in*  
1118 *Microbiology*, 32, 68–78, <https://doi.org/10.1016/j.tim.2023.06.006>, 2024.
- 1119 Trumbore, S., Brando, P., and Hartmann, H.: Forest health and global change, *Science*, 349, 814–818,  
1120 <https://doi.org/10.1126/science.aac6759>, 2015.
- 1121 Valdes, P.: Built for stability, *Nature Geosci*, 4, 414–416, <https://doi.org/10.1038/ngeo1200>, 2011.
- 1122 Wang, E., Smith, C. J., Wang, E., and Smith, C. J.: Modelling the growth and water uptake function of plant  
1123 root systems: a review, *Aust. J. Agric. Res.*, 55, 501–523, <https://doi.org/10.1071/AR03201>, 2004.

- 1124 Wang-Erlandsson, L., Bastiaanssen, W. G. M., Gao, H., Jägermeyr, J., Senay, G. B., van Dijk, A. I. J. M.,  
 1125 Guerschman, J. P., Keys, P. W., Gordon, L. J., and Savenije, H. H. G.: Global root zone storage capacity from  
 1126 satellite-based evaporation, *Hydrology and Earth System Sciences*, 20, 1459–1481,  
 1127 <https://doi.org/10.5194/hess-20-1459-2016>, 2016.
- 1128 Wang-Erlandsson, L., Tobian, A., van der Ent, R. J., Fetzer, I., te Wierik, S., Porkka, M., Staal, A., Jaramillo, F.,  
 1129 Dahlmann, H., Singh, C., Greve, P., Gerten, D., Keys, P. W., Gleeson, T., Cornell, S. E., Steffen, W., Bai, X., and  
 1130 Rockström, J.: A planetary boundary for green water, *Nat Rev Earth Environ*, 3, 380–392,  
 1131 <https://doi.org/10.1038/s43017-022-00287-8>, 2022.
- 1132 Wolfe, B. T., Sperry, J. S., and Kursar, T. A.: Does leaf shedding protect stems from cavitation during seasonal  
 1133 droughts? A test of the hydraulic fuse hypothesis, *New Phytologist*, 212, 1007–1018,  
 1134 <https://doi.org/10.1111/nph.14087>, 2016.
- 1135 Wunderling, N., Staal, A., Sakschewski, B., Hirota, M., Tuinenburg, O. A., Donges, J. F., Barbosa, H. M. J., and  
 1136 Winkelmann, R.: Recurrent droughts increase risk of cascading tipping events by outpacing adaptive capacities  
 1137 in the Amazon rainforest, *Proceedings of the National Academy of Sciences*, 119, e2120777119,  
 1138 <https://doi.org/10.1073/pnas.2120777119>, 2022.
- 1139 Xie, S.-P., Deser, C., Vecchi, G. A., Ma, J., Teng, H., and Wittenberg, A. T.: Global Warming Pattern Formation:  
 1140 Sea Surface Temperature and Rainfall, *Journal of Climate*, 23, 966–986,  
 1141 <https://doi.org/10.1175/2009JCLI3329.1>, 2010.
- 1142 Xu, C., Hantson, S., Holmgren, M., van Nes, E. H., Staal, A., and Scheffer, M.: Remotely sensed canopy height  
 1143 reveals three pantropical ecosystem states, *Ecology*, 97, 2518–2521, <https://doi.org/10.1002/ecy.1470>, 2016.
- 1144 Xue, B.-L., Guo, Q., Otto, A., Xiao, J., Tao, S., and Li, L.: Global patterns, trends, and drivers of water use  
 1145 efficiency from 2000 to 2013, *Ecosphere*, 6, art174, <https://doi.org/10.1890/ES14-00416.1>, 2015.
- 1146 Yang, Y., Saatchi, S. S., Xu, L., Yu, Y., Choi, S., Phillips, N., Kennedy, R., Keller, M., Knyazikhin, Y., and Myneni, R.  
 1147 B.: Post-drought decline of the Amazon carbon sink, *Nat Commun*, 9, 3172, <https://doi.org/10.1038/s41467-018-05668-6>, 2018.
- 1149 Yu, Z., Chen, X., Zhou, G., Agathokleous, E., Li, L., Liu, Z., Wu, J., Zhou, P., Xue, M., Chen, Y., Yan, W., Liu, L., Shi,  
 1150 T., and Zhao, X.: Natural forest growth and human induced ecosystem disturbance influence water yield in  
 1151 forests, *Commun Earth Environ*, 3, 148, <https://doi.org/10.1038/s43247-022-00483-w>, 2022.
- 1152 Yuan, K., Zhu, Q., Riley, W. J., Li, F., and Wu, H.: Understanding and reducing the uncertainties of land surface  
 1153 energy flux partitioning within CMIP6 land models, *Agricultural and Forest Meteorology*, 319, 108920,  
 1154 <https://doi.org/10.1016/j.agrformet.2022.108920>, 2022.
- 1155 Zemp, D. C., Schleussner, C.-F., Barbosa, H. M. J., van der Ent, R. J., Donges, J. F., Heinke, J., Sampaio, G., and  
 1156 Rammig, A.: On the importance of cascading moisture recycling in South America, *Atmospheric Chemistry and  
 1157 Physics*, 14, 13337–13359, <https://doi.org/10.5194/acp-14-13337-2014>, 2014.
- 1158 Zemp, D. C., Schleussner, C.-F., Barbosa, H. M. J., Hirota, M., Montade, V., Sampaio, G., Staal, A., Wang-  
 1159 Erlandsson, L., and Rammig, A.: Self-amplified Amazon forest loss due to vegetation-atmosphere feedbacks,  
 1160 *Nature Communications*, 8, 14681, <https://doi.org/10.1038/ncomms14681>, 2017.
- 1161 Zhang, Y., Peña-Arancibia, J. L., McVicar, T. R., Chiew, F. H. S., Vaze, J., Liu, C., Lu, X., Zheng, H., Wang, Y., Liu, Y.  
 1162 Y., Miralles, D. G., and Pan, M.: Multi-decadal trends in global terrestrial evapotranspiration and its  
 1163 components, *Scientific Reports*, 6, 19124, <https://doi.org/10.1038/srep19124>, 2016.

1164 Zilli, M. T., Carvalho, L. M. V., and Lintner, B. R.: The poleward shift of South Atlantic Convergence Zone in  
1165 recent decades, *Clim Dyn*, 52, 2545–2563, <https://doi.org/10.1007/s00382-018-4277-1>, 2019.

1166