### Global distribution of the rooting zone water storage capacity reflects plant adaptation to the environment

Benjamin D. Stocker<sup>1,2,\*</sup>, Shersingh Joseph Tumber-Dávila<sup>3,4</sup>, Alexandra G. Konings<sup>3</sup>, Martha B. Anderson<sup>5</sup>, Christopher Hain<sup>6</sup>, Robert B. Jackson<sup>3,7,8</sup>

<sup>a</sup> Department of Environmental Systems Science, ETH, Universitätsstrassee 2, 8092 Zürich, Switzerland

<sup>b</sup> Swiss Federal Institute for Forest, Snow and Landscape Research WSL, Zürcherstrasse 111, 8903 Birmensdorf, Switzerland

 $^{\rm c}$  Department of Earth System Science, Stanford University, Stanford, CA 94305-4216, USA

<sup>d</sup> Harvard Forest, Harvard University, Petersham, MA 01366, USA

 $^{\rm e}$  USDA-ARS Hydrology and Remote Sensing Laboratory, Beltsville, MD 20705 USA  $^{\rm f}$  NASA Marshall Space Flight Center, AL 35808, USA

 $^{\rm g}$  Woods Institute for the Environment, Stanford University, Stanford, CA 94305-4216, USA

<sup>h</sup> Precourt Institute for Energy, Stanford University, Stanford, CA 94305-4216, USA

\* bestocke@ethz.ch

### Abstract

The rooting zone water storage capacity  $(S_0)$  extends from the soil surface to the weathered bedrock (the Critical Zone) and determines land-atmosphere exchange during dry periods. Despite its importance to land-surface modeling, variations of  $S_0$  across space are largely unknown as they cannot be observed directly. We developed a method to diagnose global variations of  $S_0$  from the relationship between vegetation activity (measured by sun-induced fluorescence and by the evaporative fraction) and the cumulative water deficit (CWD). We then show that spatial variations in  $S_0$  can be predicted from the assumption that plants are adapted to sustain CWD extremes occurring with a return period that is related to the life form of dominant plants and the large-scale topographical setting. Predicted biome-level  $S_0$  distributions, translated to an apparent rooting depth  $(z_r)$  by accounting for soil texture, are consistent with observations from a comprehensive  $z_r$  dataset. Large spatial variations in  $S_0$  across the globe reflect adaptation of  $z_r$  to the hydroclimate and topography and implies large heterogeneity in the sensitivity of vegetation activity to drought. The magnitude of  $S_0$ inferred for most of the Earth's vegetated regions and particularly for those with a large seasonality in their hydroclimate indicates an important role for plant access to water stored at depth - beyond the soil layers commonly considered in land-surface models.

# Introduction

Water availability to vegetation exerts a strong control on the terrestrial carbon cycle [2,30], surface-atmosphere exchanges [60], and the development of heat waves [46]. Under dry conditions and increasing cumulative water deficits (CWD, defined as the

difference between evapotranspiration, ET, and precipitation, P, summed over continuous dry periods), water availability to vegetation becomes increasingly limiting, leading to reduced CO<sub>2</sub> assimilation and transpiration [10], to accelerated heating of near-surface air [68], and - in extreme cases - to premature defoliation of water-stressed vegetation [5] and enhanced tree mortality [9,25].

To sustain activity during dry periods and resist impacts of droughts, plants rely on water stored below the surface [26, 44]. The larger the rooting zone water storage capacity ( $S_0$ ), the longer plants can withstand soil moisture limitation during dry periods [66] and the lower their sensitivity to an increasing CWD.  $S_0$  is therefore a key factor determining drought impacts, land-atmosphere exchanges, and runoff regimes, particularly in climates with seasonal asynchrony in radiation and precipitation [23, 26, 44]. In models,  $S_0$  is commonly conceived as a function of the soil texture and the plants' rooting depth ( $z_r$ ), limited to the soil depth [44, 59]. Recent research has revealed a substantial component of  $S_0$  and contributions to ET by water stored beneath the soil [48], in weathered and fractured bedrock [12, 43, 53] or groundwater [18, 28, 67]. Plant access to such deep moisture plays an important role in controlling near-surface climate [42, 57, 65], runoff regimes [26], global patterns of vegetation cover [37], and mitigating impacts of droughts [17].

However,  $S_0$  is impossible to observe directly across large scales and its spatial variations are poorly understood [39]. Gravimetric remote sensing provides information of terrestrial water storage variations but only at  $\sim 100$  km spatial resolution [30]. Information from microwave remote sensing is limited to the top few centimeters of the soil, and derived root zone soil moisture estimates rely on extrapolations and assumptions for  $z_r$  [13]. Global compilations of local  $z_r$  measurements [7,55] have resolved this observational challenge only partly because of their limited size (owing to prohibitively laborious data collection efforts), the importance of the (often unknown) groundwater table depth (WTD) in relation to  $z_r$  [18], and large documented variations in  $z_r$  across multiple scales.  $z_r$  varies across large-scale climatic gradients and the Earth's biomes [7,55], along topographical gradients and catchment-scale variations of the WTD [18], between individuals growing under different conditions [69], and among different growth forms and species growing in close vicinity [8,55]. Modelling approaches for predicting  $z_r$  have conceived their spatial variations as the result of eco-evolutionary pressure and plants' adaptation to the prevailing hydro-climate [21, 34, 58, 62, 73]. Other approaches for estimating the global  $z_r$ distribution made use of empirical relationships between in-situ observations and climatic factors [54]. Catchment-scale water balance analyses [23] have shown that the seasonality of potential evapotranspiration and precipitation, its relative timing, and the ensuing cumulative water deficit (CWD) are important driving factors for plants genetic or phenotypic adaptation of  $z_r$  and hence  $S_0$ .

Despite its crucial role in controlling water and carbon fluxes and the scarcity of observations, practically all land surface models, climate models, terrestrial biosphere models, numerical weather prediction models, or hydrological models, and several widely used remote sensing-based models of vegetation productivity and evapotranspiration rely on a specification of  $S_0$  either directly as the depth of a "water bucket" (a simple model for water storage along the rooting zone), or indirectly through a prescribed rooting depth  $z_r$  and soil texture across the profile, or the shape of an empirical water stress function. Typically, water stored along the entire Critical Zone - the Earth's outermost layer of biota and rocks that interacts with the atmosphere, the hydrosphere, and biogeochemical cycles - is not fully represented in models [28,53]. As a consequence, spatial variations in  $z_r$  and  $S_0$  arise in models predominantly from prescribed variations in soil texture [38], from distinct parameter choices of  $z_r$  for different plant functional types (PFTs), and from the PFTs' spatial distribution [14].

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

The evident plasticity of  $z_r$  and  $S_0$  along climatic and topographic gradients is typically ignored. Implications of this simplification may be substantial for the simulation of land-atmosphere coupling and drought impacts [28, 42, 57].

Here, we estimate the global distribution of effective  $S_0$  by fusing multiple time series of Earth observation data streams at a resolution of  $0.05^{\circ}$  and validate global patterns against a globally distributed dataset of  $z_r$ , observed in the field. We demonstrate that the sensitivity of vegetation activity to an increasing CWD is correlated with the magnitude of local CWD extremes, and that this reflects adaptation of the rooting depth to the local hydroclimate.

# Methods

### Cumulative water deficit estimation

The cumulative water deficit (CWD) is determined here from the cumulative difference of actual evapotranspiration (ET) and the liquid water infiltration to the soil  $(P_{in})$ . ET is based on thermal infrared remote sensing, provided by the global ALEXI data product at daily and  $0.05^{\circ}$  resolution, covering years 2003-2018. Values in energy units of the latent heat flux are converted to mass units accounting for the temperature and air pressure dependence of the latent heat of vaporization following [11].  $P_{\rm in}$  is based on daily reanalysis data of precipitation in the form of rain and snow from WATCH-WFDEI [70]. A simple snow accumulation and melt model [50] is applied to account for the effect of snowpack as a temporary water storage that supplies  $P_{\rm in}$ during spring and early summer. Snow melt is assumed to occur above 1°C and with a rate of 1 mm  $d^{-1}$  (°C)<sup>-1</sup>. The CWD is derived by applying a running sum of (ET - $P_{\rm in}$ ) for days where (ET -  $P_{\rm in}$ ) is positive (net water loss from the soil), and terminating the summation after rain has reduced the running sum to zero. All precipitation and snowmelt  $(P_{in})$  is assumed to contribute to reducing the CWD. This implicitly assumes that no runoff occurs while the CWD is above zero. The period between the start and end of accumulation is referred to as a CWD event. Within each event, data are removed after rain has reduced the CWD to below 90% of its maximum value within the same event. This concerns the analysis of SIF and EF (see below) and avoids effects of relieved water stress by re-wetting topsoil layers before the CWD is fully compensated. The algorithm to determine daily CWD values and events is implemented by the R package cwd [63].

### Estimating $S_0$ from SIF and EF

As the ecosystem-level CWD increases, both gross primary production (GPP, ecosystem-level photosynthesis) and evapotranspiration (ET) are limited by the availability of plant access to water. Below, we refer to GPP and ET as a generic "vegetation activity" variable X(t). This principle can be formulated, in its simplest form, as a model of X(t) being a linear function of the remaining water stored along the rooting zone S(t), expressed as a fraction of the total rooting zone water storage capacity  $S_0$  [40]:

$$X(t) = X_0 \cdot S(t) / S_0 \tag{1}$$

Following Eq. 1,  $S_0$  can be interpreted as the total rooting zone water storage capacity, or the depth of a "water bucket" that supplies moisture for evapotranspiration. Following [66] and with X(t) representing ET, the temporal dynamics during rain-free periods (where runoff can be neglected), are described by the differential equation 100

57

58

59

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

86

87

89

91

92

93

94

95

96

97

98

$$dS/dt = -X(t) \Rightarrow dS/dt = -X_0 \cdot S(t)/S_0 \tag{2}$$

and solved by an exponential function with a characteristic decay time scale  $\lambda$ :

$$X(t) = X_0 \cdot \exp(-[t - t_0]/\lambda) \tag{3}$$

101

102

103

104

112

113

114

115

116

117

118

119

120

135

 $\lambda$  is related to  $S_0$  as  $S_0 = \lambda X_0$ , where  $X_0$  is the initial ET at  $S(t_0) = S_0$ . In other words, the apparent observed exponential ET decay time scale  $\lambda$ , together with  $X_0$ , reflects the total rooting zone water storage capacity  $S_0$ .

Fitting exponentials from observational data is subject to assumptions regarding stomatal responses to declines in S(t) and is relatively sensitive to data scatter. Hence, resulting estimates of  $S_0$  may not be robust. With  $\text{CWD} = S_0 - S(t)$  and Eq. 1, the relationship of X(t) and CWD(t) can be expressed as a linear function

$$X(t) = X_0 - X_0 / S_0 \cdot \text{CWD}(t) \tag{4}$$

and observational data for X(t) can be used to fit a linear regression model. Its intercept *a* and slope *b* can then be used as an alternative, and potentially more robust estimate for  $S_0$ :

$$S_0 = -a/b \tag{5}$$

This has the further advantage that estimates for  $S_0$  can be derived using any observable quantity of vegetation activity X(t) (not just ET as in [66]) under the assumption that activity attains zero at the point when the CWD reaches the total rooting zone water storage capacity; i.e.,  $X(t^*) = 0$  for  $\text{CWD}(t^*) = S_0$ .

2

Here, we use sun-induced fluorescence (SIF, [16]), normalised by incident shortwave radiation (WATCH-WFDEI data [70]), and the evaporative fraction (EF), defined as the ratio of evapotranspiration (ALEXI-ET data [27]) over net radiation (GLASS data [32]), as two alternative proxies for water-constrained vegetation activity X. ET and net radiation data are both provided at  $0.05^{\circ}$  and daily resolution.

 $S_{\rm dSIF}$  and  $S_{\rm dEF}$  were derived based on the linear relationship of SIF and EF versus 121 CWD by applying Eq. 5. The relationship was derived for each pixel with pooled data 122 belonging to the single largest CWD event of each year, and using the 90% quantile of 123 EF and normalised SIF within 50 evenly spaced bins along the CWD axis. Binning and 124 considering percentiles were chosen to reduce effects of vegetation activity reduction due 125 to factors other than water stress (CWD). We then tested for each pixel whether the 126 data support the model of a single linear decline of SIF (EF) with increasing CWD (Eq. 127 5), or, alternatively, a segmented regression model with one or two change points. The 128 R package segmented was used [47]. The model with the lowest BIC was chosen and 129  $S_{\rm dSIF}$  and  $S_{\rm dEF}$  were quantified only for pixels where no significant change point was 130 detected and where the regression of EF (SIF) vs. CWD had a significantly negative 131 slope. "Flattening" EF (SIF) vs. CWD relationships were identified where a significant 132 change point was detected and where the slope of the second regression segment was 133 significantly less negative (p = 0.05 of t-test) compared to the slope of the first segment. 134

#### Rooting zone water storage capacity and rooting depth model

Following ref. [23], the rooting zone water storage capacity  $S_0$  is modelled based on the assumption that plants size their rooting depth such that the corresponding  $S_0$  is just large enough to maintain function under the expected maximum cumulative water deficit (CWD) occurring with a return period of T years. Magnitudes of CWD extremes are estimated by fitting an extreme value distribution (Gumbel) to the annual maximum CWD values for each pixel separately, using the *extRemes* R package [24]. Magnitudes of extremes with a given return period T are derived and yield  $S_{\text{CWDXT}}$ .  $S_{\text{CWDXT}}$  are translated into an effective depth  $z_{\text{CWDXT}}$  using estimates of the plant-available soil water holding capacity, based on soil texture data from a gridded version of the Harmonized World Soil Database [20,71] and pedo-transfer functions derived by [4]. A global map of  $z_{\text{CWDX80}}$  is provided in Fig. S13.

### Estimating return periods

Diagnosed values of  $S_{\rm dSIF}$  and  $S_{\rm dEF}$  provide a constraint on the return period T. To yield stable estimates of T and avoid effects of the strong non-linearity of the function to derive T from the fitted extreme value distributions and magnitudes estimated by  $S_{\rm dSIF}$  and  $S_{\rm dEF}$ , we pooled estimates  $S_{\rm dSIF}$  ( $S_{\rm dEF}$ ) and  $S_{\rm CWDXT}$  values within 1° pixels ( $\leq 400$  values). A range of discrete values T was screened (10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250, 300, 350, 400, 450, 500 years) and the best estimate T was chosen based on comparison to  $S_{\rm dSIF}$  ( $T_{\rm SIF}$ ) and to  $S_{\rm dEF}$  ( $T_{\rm EF}$ ), i.e., where the absolute value of the median of the logarithm of the bias was minimal.

### Rooting depth observations

The rooting depth data set (N = 5524) was compiled by combining and complementing 157 published datasets from [54] and [18]. The data includes observations of the maximum 158 rooting depth of plants taken from 361 published studies plus additional environmental 159 and climate data.  $z_r$  was taken as the plant's maximum rooting depth. Data were 160 aggregated by sites (N = 359) based on longitude and latitude information. Sites were 161 classified into biomes using maps of terrestrial ecoregions [49]. Quantiles (10%, 90%)162 were determined for each biome. For a subset of the data, where parallel measurements 163 of the water table depth (WTD) was available, we conducted the same analysis, but 164 took the minimum of WTD and  $z_r$ . 165

# Results

### Estimating ET

Unbiased estimates of ET during rain-free periods are essential for determining CWD and estimating  $S_0$  and implied  $z_r$ . We tested different remote sensing-based ET products and found that the ALEXI-TIR product, which is based on thermal infrared remote sensing [3,27], exhibits no systematic bias during progressing droughts (Fig. S1). The stability in ET estimates from ALEXI-TIR during drought are enabled by its effective use of information about the surface energy partitioning, allowing inference of ET rates without reliance on a priori specified and inherently uncertain surface conductances [45] or shapes of empirical water stress functions [19], and without assumptions of rooting depth or effective  $S_0$ . ALEXI-TIR is thus well-suited for estimating actual ET behaviour during drought without introducing circularity in inferring  $S_0$ .

### Diagnosing $S_0$ from SIF and EF

By employing first principles for the constraint of rooting zone water availability on ET and photosynthesis [66], we developed a method to derive how the sensitivity of these fluxes to increasing CWD relates to  $S_0$  and how this sensitivity can be used to reveal effects of access to extensive deep water stores (see Methods). Following this method, we determined the apparent  $S_0$  as the CWD at which vegetation "activity" ceases. The parallel information of evapotranspiration, precipitation, and snow mass balance enables

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

166

167

168

169

170

171

172

173

174

175

176

177

178

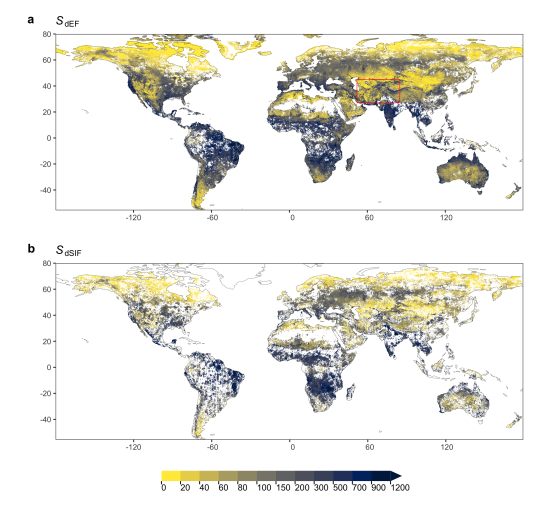


Figure 1. Rooting zone water storage capacity (mm) estimated from the sensitivity of the evaporative fraction ( $S_{dEF}$ , a) and sun-induced fluorescence ( $S_{dSIF}$ , b) to the cumulative water deficit (CWD). Data is aggregated to 0.1° resolution. The red box in (a) shows the outline of the magnified map provided in Fig. 2. Blank cells mark areas where all native cells at 0.05° resolution did not exhibit a significant and single, linearly declining relationship with increasing CWD.

a quantification of CWD over time. Vegetation activity was estimated from two alternative approaches: from the evaporative fraction (EF, defined as ET over net radiation), and from sun-induced fluorescence (SIF, normalised by incident shortwave radiation). SIF is a proxy for ecosystem photosynthesis [74] and is taken here from a spatially downscaled data product [16] based on GOME-2 data [33,36]. Since net radiation and shortwave radiation are first-order controls on ET and SIF, respectively, and to avoid effects by seasonally varying radiation inputs, we used EF instead of ET, and considered the ecosystem-level fluorescence yield, quantified as SIF divided by shortwave radiation (henceforth referred to as 'SIF') for all analyses.

Fig. 1 reveals large global variations in  $S_0$ . Estimates based on EF and SIF agree closely ( $R^2$ =0.78, Fig. S2). The lowest sensitivity of vegetation activity to an increasing CWD, and thus the largest apparent  $S_0$ , is found in regions with a strong seasonality in radiation and water availability and substantial vegetation cover - particularly in monsoonal climates. In contrast, the lowest  $S_0$  values appear not only in regions where seasonal water deficits are limited due to short inter-storm duration (e.g., inner tropics) and/or low levels of potential evapotranspiration (e.g., high latitudes), but also in deserts and arid grasslands. This reflects the limited water surplus accumulating during rain events from which vegetation can draw during dry periods. A rapid decline of ET and SIF with an increasing CWD is related to vegetation cover dynamics, governed by greening after rain pulses and browning during dry periods [35].

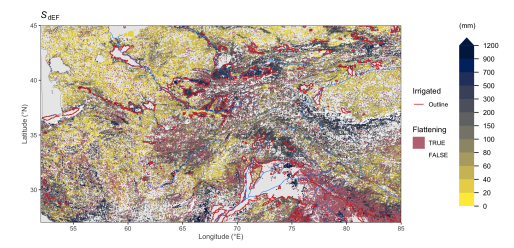


Figure 2. Rooting zone water storage capacity in Central Asia, estimated from the evaporative fraction  $(S_{dEF})$ . Mauve areas ("flattening") show grid cells where a significant reduction in the slope in EF vs. CWD was identified beyond a certain threshold.  $S_{dEF}$  values are not calculated for gridcells classified as "Flattening". Red lines show outlines of major irrigated areas, i.e. where the irrigated land area fraction is above 30% [61]. Information about irrigated areas was used only for mapping here, but is not used for other parts of the analysis. Blank grid cells indicate areas with a sustained imbalance of ET being greater than P. Additional regional maps are provided by Figs. S3-S5.

Clear patterns emerge also at smaller scales (Fig. 2).  $S_{dSIF}$  and  $S_{dEF}$  consistently (Fig. S6) reveal how the sensitivity of photosynthesis and transpiration to drought stress varies across different topographical settings, indicating generally larger  $S_0$  in mountain regions and along rivers (Amu Darya) and deltas (Volga). We note however, that ALEXI ET estimates over mountainous terrain may be biased high where low incident net radiation and surface temperatures are caused not by high evaporative fractions but rather by topography effects and local shading. The maps of  $S_{dSIF}$  and

212

186

187

188

189

190

191

192

193

194

195

196

197

198

199

200

201

202

203

204

 $S_{\text{dEF}}$  also bear strong imprints of human land use. Major irrigated cropland areas are 213 congruent with some of the highest apparent  $S_0$  values recorded here. In these areas, 214 our analysis yields particularly high CWD values and a low sensitivity of SIF and EF to 215 CWD based on remotely sensed apparent ET and precipitation time series, without 216 using information about the location and magnitude of irrigation. Other major irrigated 217 areas appear as blank cells in Fig. 2 because the algorithm used to calculate CWD (see 218 Methods) fails due to a long-term imbalance between P and ET and a "runaway CWD". 219 This indicates a sustained over-use of local water resources, caused by lateral water 220 redistribution at scales beyond  $\sim 5$  km, either via streamflow diversion or groundwater 221 flow and extraction. 222

Regressing vegetation activity against CWD also identifies locations where a decoupling of the two variables appears, i.e., where the sensitivity EF or SIF significantly decreases beyond a certain CWD threshold ("Flattening" in Fig. 2, see Methods). Such areas are particularly common in the vicinity of mountain regions, in areas with irrigated croplands, and in savannahs (Fig. S7). Related mechanisms may be at play. A flattening of the EF (SIF) vs. CWD relationship is likely due to the different portions of the vegetation having access to distinct water resources and respective storage capacities. In areas with large topographic gradients, this may be due to within-gridcell heterogeneity in plant access to the saturated zone. In savannahs, a shift in ET contributions from grasses and trees and a related shift in transpiration occurs as grasses, which are often more shallow-rooted than trees, senesce. In irrigated cropland areas, the flattening likely reflects land use heterogeneity within  $\sim$ 5 km grid cells and the persistent water access of irrigated fields while EF and SIF are reduced more rapidly in surrounding vegetation.

### Inferring $S_0$ and $z_r$ from CWD distributions

What controls spatial variations in  $S_0$  and  $z_r$  and the sensitivity of vegetation activity 238 to water stress? Following ref. [23], we hypothesized that plants are adapted to the local 239 hydroclimate and size their  $z_r$  to maintain transpiration and photosynthesis in the face 240 of water deficits during dry periods commonly experienced over the course of a plant's 241 lifetime. Specifically, we hypothesized that  $S_0$  can be inferred from the distribution of 242 annual CWD maxima at a given location and that  $S_0$  corresponds to a CWD extreme 243 recurring with a return period of T years, where T is on the same order of magnitude as 244 the typical lifetime of dominant plants (see Methods). We start by using T = 80 years 245 for estimating  $S_0$  and  $z_r$  here (termed  $S_{\text{CWDX80}}$  and  $z_{\text{CWDX80}}$ ) and fit CWD extreme 246 value distributions separately for each pixel. 247

223

224

225

226

227

228

229

230

231

232

233

234

235

236

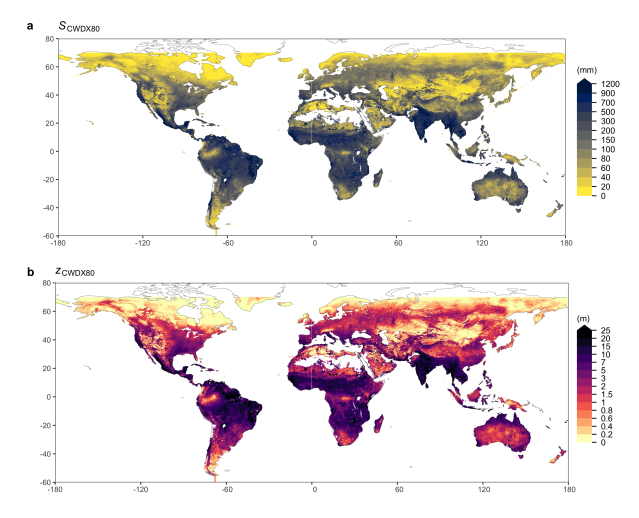


Figure 3. Spatial variations of (a) the rooting zone water storage capacity, estimated by  $S_{\text{CWDX80}}$  (mm), and (b) the apparent plant rooting depth  $z_{\text{CWDX80}}$  (m). Values are remapped to a 0.1° resolution. Blank grid cells are either permanent inland water bodies and ocean, or locations with long-term accumulation of water deficits. Values are removed in gridcells where more than 80% is non-vegetation surface based on MODIS Landcover [22].

Fig. 3a shows the global distribution of  $S_{\text{CWDX80}}$  and reveals patterns across 253 multiple scales - in close agreement with  $S_{\text{dSIF}}$  and  $S_{\text{dEF}}$  ( $R^2 = 0.76$  and  $R^2 = 0.84$ , 254 respectively, Fig. S6). This indicates that the sensitivity of vegetation activity to an 255 increasing CWD is strongly controlled by the magnitude of local CWD extremes. 256 Perennially moist tropical regions around the Equator (northwestern Amazon, Congo, 257 southeast Asia) where persistent cloud cover prohibits the derivation of  $S_{dSIF}$  and  $S_{dEF}$ , 258 emerge here as characterised by particularly low  $S_{\text{CWDX80}}$ . Fine granularity and large 259 spatial heterogeneity of  $S_{\rm CWDX80}$  at regional scales, particularly in arid and semi-arid 260 regions, reveal the importance of the local topographical setting (and irrigation, see Fig. 261 2) for determining plant-available water storage capacities (Figs. S8-S11). Variations 262 are likely to extend to even smaller scales along the hillslope topography [18] and within 263 individual forest stands [1]. These scales lie beyond the resolution of the satellite remote 264 sensing data used here to calculate CWD. 265

248

249

250

251

### Validation with rooting depth observations

Next, we investigate whether the strong control of CWD extremes on the sensitivity of 267 vegetation activity to water deficits reflects the adaptation of plant rooting depth to the 268 hydroclimate. For this, we compare  $S_{CWDX80}$  against (fully independent) field 269 observations of  $z_r$  from a novel compilation of tree-level data with global coverage, 270 extended from previous work [18,56], now encompassing 5524 observations (see also 271 Methods). Data was aggregated by taking the mean of individual observations for 1705 272 globally distributed sites (SI Fig. S12). To avoid the inevitable scale mis-match between 273 in situ ecosystem observations and global remote sensing [39], we focused our evaluation 274 against observed rooting depth at the scale of biomes. We converted estimates of 275  $S_{\rm CWDX80}$  into a corresponding depth ( $z_{\rm CWDX80}$ ) by accounting for soil texture along the 276 rooting profile (see Methods, Fig. 3b). 277

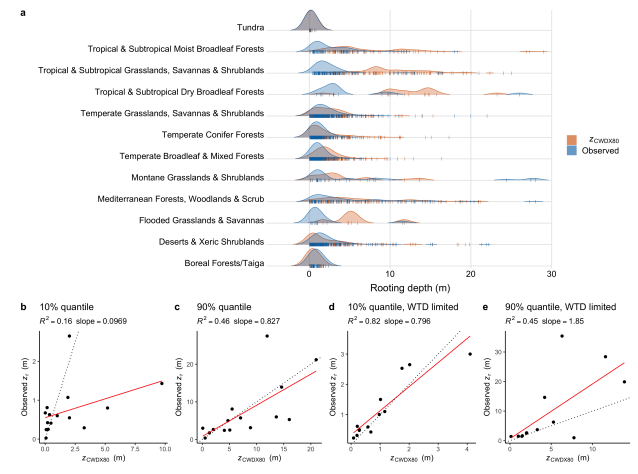


Figure 4. Modelled and observed rooting depth by biomes. (a) Kernel density estimates of observed and predicted  $(z_{CWDX80})$  rooting depth by biomes, based on data aggregated by sites, shown by vertical colored tick marks. 10% and 90% quantiles of observed vs. predicted  $(z_{CWDX80})$  rooting depth by biome of all data (b,c), and of a subset of the data where water table depth was measured along with rooting depth (d,e). Classification of sites into biomes was done based on [49]. Dotted lines in b-e represent the 1:1 line.



266

10/20

to be overestimated by  $z_{\rm CWDX80}$  (Fig. 4b). Using a subset of the data where 287 information about the water table depth (WTD) is provided (490 entries from 359 sites), 288 we limited predicted  $z_{\rm CWDX80}$  to the value of the observed local WTD (53% of all 289 observations). This yields an improved agreement (Fig. 4d) and suggests that our 290 predictive model for  $z_r$  overestimates values where the roots access groundwater. While 291 acting as a constraint on the rooting depth [18], plant access to groundwater implies 292 sustained transpiration during dry periods, correspondingly large cumulative water 293 deficits and, by implication of the model design, large  $S_{\rm CWDX80}$  and  $z_{\rm CWDX80}$ . In other 294 words, large inferred  $S_{\rm CWDX80}$  and  $z_{\rm CWDX80}$  reflect sustained transpiration, enabled not 295 by the exceptional depth of roots, but by roots tapping into the groundwater, and CWD 296 is limited not by the supply of water, but by the demand (effectively radiation-limited 297 ET). This highlights that estimates of  $S_0$  and  $z_r$  presented here are to be interpreted as 298 apparent quantities, subject to groundwater influence (and irrigation, as shown above). 299 It also demonstrates that knowledge about the groundwater depth and whether 300 vegetation taps into it is key for explaining ET during dry periods at a large proportion 301 of sites investigated here. 302

### Estimating return periods

Underlying the estimates of  $S_{\rm CWDX80}$  is the assumption that plant rooting strategies are 304 adapted to sustain CWD extremes with a return period T = 80 years.  $S_{\rm dSIF}$  and  $S_{\rm dEF}$ 305 provide an independent constraint to test this assumption. Fig. 3 suggests that T is not 306 a global constant and that a general pattern emerges, largely in consistency between the 307 independent estimates  $T_{\rm EF}$  and  $T_{\rm SIF}$ . A clear tendency towards higher T emerges with 308 an increasing gridcell average forest cover fraction (Fig. 3d).  $S_0$  appears to be adapted 309 to 500-year and even rarer events in forested regions but to lower T outside. A return 310 period on the order of multiple centuries is consistent with the typical lifetime of 311 trees [6] and suggests that optimal plant adaptation of life history strategies to a 312 stochastic environment may be understood as being governed by frequencies of climate 313 extremes in relation to the lifetime of affected organisms. Interestingly, substantial 314 variations in T remain even within land cover types (e.g. the boreal forests of Russia). 315 This variation may be related to the large-scale topographical setting and the tendency 316 towards shallow groundwater table depths, as measured by the Compound Topographic 317 Index ([41], Fig. 3c). In such areas, vegetation appears to sustain particularly rare 318 CWD extremes (diagnosed here from its low sensitivity to CWD during the observation 319 period), possibly enabled by roots' access to a relatively shallow saturated zone. 320

### Discussion and conclusion

 $S_0$  is a central quantity in determining land-atmosphere exchange and vegetation resistance to drought, but its spatial variations cannot be observed directly and have largely been ignored in models. Using first-principles modelling and integrating multiple data streams at daily and ~5 km resolution, we diagnosed a hydrologically effective ecosystem-level  $S_0$  from the sensitivity of vegetation activity to CWD. Our analysis revealed extensive spatial heterogeneity of plant-available moisture storage in the Critical Zone, with apparent variations extending from the continental to the regional scale. While large-scale variations in  $S_0$  are mainly driven by the hydroclimate, more fine-grained variations within regions are linked to topography and how vegetation and land use is distributed across the landscape and determines actual ET.

By detecting where the sensitivity of vegetation activity to an increasing CWD <sup>332</sup> declines beyond a certain CWD threshold, we identified locations where a portion of the <sup>333</sup> vegetation may have access to deep water stores and where groundwater may contribute <sup>334</sup>

303

321

322

323

324

325

326

327

328

329

330

bioRxiv preprint doi: https://doi.org/10.1101/2021.09.17.460332; this version posted September 19, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

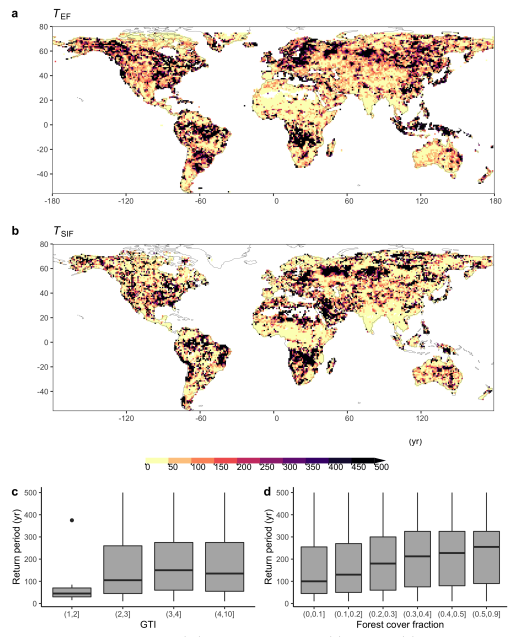


Figure 3. Return periods T (yr), diagnosed from EF (a) and SIF (b). To diagnose T, a range of alternative values of T are screened and the corresponding range of values  $S_{\text{CWDXT}}$  are compared to  $S_{\text{dEF}}$  ( $S_{\text{dSIF}}$ ) within 1° grid cells (resolution of maps shown here). The best matching T was retained for each gridcell, yielding a global distribution of  $T_{\text{EF}}$  ( $T_{\text{SIF}}$ ). The lower panel shows the distribution of diagnosed return periods T (mean of  $T_{\text{EF}}$  and  $T_{\text{SIF}}$ ) within bins of the Compound Topographical Index [41] (c) and forest cover fractions (MOD44B [29]) (d).

to land-atmosphere coupling [42]. Traditional land surface models, which typically employ the assumption of free drainage out of the bottom soil layer and simulate continuously increasing water stress effects as CWD increases, may not be suitable for reliably simulating land-atmosphere exchange during prolonged dry periods in these regions.

Diagnosed values of  $S_0$  are representative of water stored across the entire rooting zone - from the soil surface to the weathered bedrock and groundwater. Particularly in regions with pronounced dry seasons, magnitudes of  $S_{\rm CWDX80}$  greatly exceed typical values of the total soil water holding capacity when considering the top 1 or 2 meters of the soil column and texture information from global databases ([20], Fig. S13) - as commonly done in models [11,28,44]. The discrepancy in magnitude and spatial patterns underlines the critical role of plant access to deep water and the need to extend the focus beyond soil moisture for simulating land-atmosphere exchange [12]. We note that using  $S_{\rm CWDX80}$  ( $z_{\rm CWDX80}$ ) for directly parameterizing  $S_0$  ( $z_r$ ) in global models may be misleading in forested areas with particularly small  $S_{\rm CWDX80}$ . Additional effects of how  $z_r$  determines access to belowground resources and function (e.g., nutrients, mechanical stability) should be considered.

Our estimate of  $S_0$  implicitly includes water intercepted by leaf and branch surfaces, internal plant water storage, and moisture stored in the topsoil and supplied to evaporation. These components are generally smaller in magnitude compared to moisture storage supplied to transpiration [64], and their contribution to ET declines rapidly as CWD increases. (Moreover, readily available global evaporation data is lacking [39]). Hence, spatial variations in  $S_0$  primarily reflect variations mediated by moisture stored across the root zone.

Our analysis identified mountain regions as "water towers" with particularly high  $S_0$ , in spite of shallow soil and regolith depths along upland hillslopes [51]. This could be due to hillslope-scale variations in groundwater depth where lateral flow pushes the saturated zone closer to the surface in valley bottoms and enables sustained transpiration during prolonged rain-free periods. Lateral subsurface flow is not explicitly accounted for, but can be considered relatively small at the scale investigated here ( $\sim 5$ km). Nevertheless, local convergence (divergence) acts to supply (remove) moisture and sustain (reduce) ET, leading to larger (smaller) CWD values. This effect should be captured by the relatively accurate remote sensing of ET by TIR (Fig. S1), and likely contributes to the strong contrast in  $S_{CWDX80}$  along topographic gradients (Figs. S8-S11). However, further research should assess the accuracy of spatial variations in annual mean ET and potential effects of terrain, where lower temperature rise signals on shaded slopes may be mis-interpreted by the ALEXI algorithm as signatures of higher ET. Since ET data are used both in  $S_{\text{CWDX80}}$  and in  $S_{\text{dEF}}$  and  $S_{\text{dSIF}}$ , these quantities are not entirely independent, potentially leading to an overestimation of their correlation strengths (Fig. S7). Nevertheless, our predictions of continental-scale variations in  $S_0$  and implied  $z_r$  agree well with patterns derived from an extended dataset of tree-level  $z_r$  observations - an entirely independent observational constraint.

In contrast to earlier studies [18,34,39,54,73], here we mainly focused on  $S_0$ , instead of  $z_r$ . This makes our quantification independent of large variations in porosity along depths extending beneath the soil and focuses on the quantity relevant for the surface water balance. Nevertheless, where vegetation accesses the saturated zone, effective  $S_0$ may be even larger than numbers derived here based on past CWD time series. Similarly, the concept of a constrained water storage capacity is inappropriate where irrigation supplies water to ET. For comparison to observations and to earlier studies, we converted  $S_0$  into  $z_r$  using soil texture information provided for the top 1 m of the soil profile [20,71]. This was chosen due to a lack of information about porosity of the subsoil.

335

336

337

338

339

340

341

342

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

384

385

Our  $z_r$  map (Fig. S13) yields patterns that are mostly consistent with previous estimates of the global distribution of  $z_r$  [18, 34, 39, 54, 73], but differ in important aspects. Deepest roots are predicted here in monsoonal climates with substantial vegetation cover, particularly in south and southeast Asia, but are predicted along the arid edges of vegetated zones elsewhere [18, 73]. This may be due to differences between TIR-derived ET versus ET derived from other approaches used in [18,73]. Our evaluation against observations suggests that the distribution of  $z_r$  is reliably predicted by  $z_{\rm CWDX80}$  for deserts and xeric shrublands and in tropical and subtropical broadleaf forests (Fig. 4). Suggested deep rooting in dry steppe regions of Kazakhstan [18,73] and further east [39,54] are not corroborated here. Mismatches may be due to the rapid dynamics of green foliage cover and ensuing water losses in these regions. In contrast to other studies [73], the ET and P time series used here capture these dynamics at a daily resolution. Common to most [18, 34, 54, 73], but not all [39], global  $z_r$  estimates, including the present one, is the characteristic contrast between  $z_r$  in the northwestern and northeastern Amazon. Taken together, constraints available from local  $z_r$ observations and from global remote sensing of vegetation activity converge on consistent patterns across multiple spatial scales and suggest that belowground vegetation structure can be sensed from space.

Using methods developed here, combined with emerging data from recently launched satellite missions for thermal infrared remote sensing (e.g., ECOSTRESS) [72] bears promise for resolving  $S_0$  variations at the hillslope scale (100 m - 1 km). This will provide critical information for reliably estimating water deficits and spatial heterogeneity of ET across the landscape at scales relevant to multiple stakeholders. Contrasting future CWD extremes with their past distribution will yield insight into the severity of droughts with respect to levels to which  $z_r$  is adapted today. Our study revealed a tight control of the climatology of water deficits on vegetation sensitivity to drought stress, and demonstrated how land-atmosphere interactions and the Critical Zone water storage capacity are governed by vegetation and its adaptation to the hydroclimate and topography across the globe. It remains to be shown whether plasticity in  $z_r$  is sufficiently rapid to keep pace with a changing climate with strong and wide-spread increases in rainfall variability [52], and to what degree rising CO<sub>2</sub> alters plant water use and their carbon economy and thereby the costs and benefits of deep roots [15,31].

# Acknowledgments

B.D.S was funded by the Swiss National Science Foundation grant no. PCEFP2\_181115 and by the generosity of Eric and Wendy Schmidt by recommendation of the Schmidt Futures program. We acknowledge ICDC, CEN, University of Hamburg for data support. SJTD was funded by the NSF GRFP and LTER, and the NASEM Ford Foundation Fellowship Program. AGK was supported by NSF DEB-1942133.

# References

- E. Agee, L. He, G. Bisht, V. Couvreur, P. Shahbaz, F. Meunier, C. M. Gough, A. M. Matheny, G. Bohrer, and V. Ivanov. Root lateral interactions drive water uptake patterns under water limitation. *Advances in Water Resources*, 151:103896, 2021.
- A. Ahlström, M. R. Raupach, G. Schurgers, B. Smith, A. Arneth, M. Jung, M. Reichstein, J. G. Canadell, P. Friedlingstein, A. K. Jain, E. Kato, B. Poulter, S. Sitch, B. D. Stocker, N. Viovy, Y. P. Wang, A. Wiltshire, S. Zaehle, and

387

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

407

408

409

410

411

412

413

414

415

416

417

418

419

N. Zeng. The dominant role of semi-arid ecosystems in the trend and variability of the land CO2 sink. *Science*, 348(6237):895–899, 2015.

- M. Anderson. A Two-Source Time-Integrated model for estimating surface fluxes using thermal infrared remote sensing. *Remote Sensing of Environment*, 60(2):195–216, 1997.
- 4. V. Balland, J. A. P. Pollacco, and P. A. Arp. Modeling soil hydraulic properties for a wide range of soil conditions. *Ecological Modelling*, 219(3-4):300-316, 2008.
- A. Bastos, R. Orth, M. Reichstein, P. Ciais, N. Viovy, S. Zaehle, P. Anthoni, A. Arneth, P. Gentine, E. Joetzjer, S. Lienert, T. Loughran, P. C. McGuire, S. O, J. Pongratz, and S. Sitch. Increased vulnerability of european ecosystems to two compound dry and hot summers in 2018 and 2019. *Earth System Dynamics Discussions*, 2021:1–32, 2021.
- 6. R. J. W. Brienen, L. Caldwell, L. Duchesne, S. Voelker, J. Barichivich, M. Baliva, G. Ceccantini, A. Di Filippo, S. Helama, G. M. Locosselli, L. Lopez, G. Piovesan, J. Schöngart, R. Villalba, and E. Gloor. Forest carbon sink neutralized by pervasive growth-lifespan trade-offs. *Nat. Commun.*, 11(1):4241, Sept. 2020.
- J. Canadell, R. B. Jackson, J. B. Ehleringer, H. A. Mooney, O. E. Sala, and E.-D. Schulze. Maximum rooting depth of vegetation types at the global scale. *Oecologia*, 108(4):583–595, 1996.
- R. Chitra-Tarak, C. Xu, S. Aguilar, K. J. Anderson-Teixeira, J. Chambers, M. Detto, B. Faybishenko, R. A. Fisher, R. G. Knox, C. D. Koven, L. M. Kueppers, N. Kunert, S. J. Kupers, N. G. McDowell, B. D. Newman, S. R. Paton, R. Pérez, L. Ruiz, L. Sack, J. M. Warren, B. T. Wolfe, C. Wright, S. J. Wright, J. Zailaa, and S. M. McMahon. Hydraulically-vulnerable trees survive on deep-water access during droughts in a tropical forest. *New Phytologist*, 231(5):1798–1813, 2021.
- B. Choat, T. J. Brodribb, C. R. Brodersen, R. A. Duursma, R. López, and B. E. Medlyn. Triggers of tree mortality under drought. *Nature*, 558(7711):531–539, June 2018.
- I. R. Cowan and G. D. Farquhar. Stomatal function in relation to leaf metabolism and environment. Symp. Soc. Exp. Biol., 31:471–505, 1977.
- T. W. Davis, I. C. Prentice, B. D. Stocker, R. T. Thomas, R. J. Whitley, H. Wang, B. J. Evans, A. V. Gallego-Sala, M. T. Sykes, and W. Cramer. Simple process-led algorithms for simulating habitats (SPLASH v.1.0): robust indices of radiation, evapotranspiration and plant-available moisture. *Geoscientific Model Development*, 10(2):689–708, 2017.
- T. E. Dawson, W. Jesse Hahm, and K. Crutchfield-Peters. Digging deeper: what the critical zone perspective adds to the study of plant ecophysiology. *New Phytologist*, 226(3):666–671, 2020.
- W. Dorigo, W. Wagner, C. Albergel, F. Albrecht, G. Balsamo, L. Brocca, D. Chung, M. Ertl, M. Forkel, A. Gruber, E. Haas, P. D. Hamer, M. Hirschi, J. Ikonen, R. de Jeu, R. Kidd, W. Lahoz, Y. Y. Liu, D. Miralles, T. Mistelbauer, N. Nicolai-Shaw, R. Parinussa, C. Pratola, C. Reimer, R. van der Schalie, S. I. Seneviratne, T. Smolander, and P. Lecomte. ESA CCI soil moisture for improved earth system understanding: State-of-the art and future directions. *Remote Sens. Environ.*, 203:185–215, Dec. 2017.

- 14. B. A. Drewniak. Simulating dynamic roots in the energy exascale earth system land model. *Journal of Advances in Modeling Earth Systems*, 11(1):338–359, 2019.
- 15. R. A. Duursma, C. V. M. Barton, D. Eamus, B. E. Medlyn, D. S. Ellsworth, M. A. Forster, D. T. Tissue, S. Linder, and R. E. McMurtrie. Rooting depth explains [CO2] x drought interaction in eucalyptus saligna. *Tree Physiol.*, 31(9):922–931, Sept. 2011.
- G. Duveiller, F. Filipponi, S. Walther, P. Köhler, C. Frankenberg, L. Guanter, and A. Cescatti. A spatially downscaled sun-induced fluorescence global product for enhanced monitoring of vegetation productivity. *Earth System Science Data*, 12(2):1101–1116, May 2020.
- E. J. L. Esteban, C. V. Castilho, K. L. Melgaço, and F. R. C. Costa. The other side of droughts: wet extremes and topography as buffers of negative drought effects in an amazonian forest. *New Phytologist*, 229(4):1995–2006, 2021.
- Y. Fan, G. Miguez-Macho, E. G. Jobbágy, R. B. Jackson, and C. Otero-Casal. Hydrologic regulation of plant rooting depth. *Proc. Natl. Acad. Sci. U. S. A.*, 114(40):10572–10577, Oct. 2017.
- J. B. Fisher, B. Lee, A. J. Purdy, G. H. Halverson, M. B. Dohlen, K. Cawse-Nicholson, A. Wang, R. G. Anderson, B. Aragon, M. A. Arain, D. D. Baldocchi, J. M. Baker, H. Barral, C. J. Bernacchi, C. Bernhofer, S. C. Biraud, G. Bohrer, N. Brunsell, B. Cappelaere, S. Castro-Contreras, J. Chun, B. J. Conrad, E. Cremonese, J. Demarty, A. R. Desai, A. De Ligne, L. Foltýnová, M. L. Goulden, T. J. Griffis, T. Grünwald, M. S. Johnson, M. Kang, D. Kelbe, N. Kowalska, J. Lim, I. Maïnassara, M. F. McCabe, J. E. C. Missik, B. P. Mohanty, C. E. Moore, L. Morillas, R. Morrison, J. W. Munger, G. Posse, A. D. Richardson, E. S. Russell, Y. Ryu, A. Sanchez-Azofeifa, M. Schmidt, E. Schwartz, I. Sharp, L. Šigut, Y. Tang, G. Hulley, M. Anderson, C. Hain, A. French, E. Wood, and S. Hook. ECOSTRESS: NASA's next generation mission to measure evapotranspiration from the international space station. *Water Resour. Res.*, 56(4):e2019WR026058, Apr. 2020.
- Food and Agriculture Organization and Food and Agriculture Organization of the United Nations. *Harmonized World Soil Database: Fao/Iiasa/isric/isscas/jrc* (Version 1.0). Food & Agriculture Org, Dec. 2008.
- O. Franklin, S. P. Harrison, R. Dewar, C. E. Farrior, Å. Brännström, U. Dieckmann, S. Pietsch, D. Falster, W. Cramer, M. Loreau, H. Wang, A. Mäkelä, K. T. Rebel, E. Meron, S. J. Schymanski, E. Rovenskaya, B. D. Stocker, S. Zaehle, S. Manzoni, M. van Oijen, I. J. Wright, P. Ciais, P. M. van Bodegom, J. Peñuelas, F. Hofhansl, C. Terrer, N. A. Soudzilovskaia, G. Midgley, and I. C. Prentice. Organizing principles for vegetation dynamics. *Nat Plants*, 6(5):444–453, May 2020.
- M. A. Friedl, D. Sulla-Menashe, B. Tan, A. Schneider, N. Ramankutty, A. Sibley, and X. Huang. MODIS collection 5 global land cover: Algorithm refinements and characterization of new datasets. *Remote Sensing of Environment*, 114(1):168–182, 2010.
- H. Gao, M. Hrachowitz, S. J. Schymanski, F. Fenicia, N. Sriwongsitanon, and H. H. G. Savenije. Climate controls how ecosystems size the root zone storage capacity at catchment scale. *Geophysical Research Letters*, 41(22):7916–7923, 2014.

- E. Gilleland and R. W. Katz. extremes2.0: An extreme value analysis package inr. Journal of Statistical Software, 72(8), 2016.
- M. L. Goulden and R. C. Bales. California forest die-off linked to multi-year deep soil drying in 2012–2015 drought. *Nature Geoscience*, 12(8):632–637, 2019.
- 26. W. J. Hahm, D. N. Dralle, D. M. Rempe, A. B. Bryk, S. E. Thompson, T. E. Dawson, and W. E. Dietrich. Low subsurface water storage capacity relative to annual rainfall decouples mediterranean plant productivity and water use from rainfall variability. *Geophysical Research Letters*, 46(12):6544–6553, 2019.
- C. R. Hain and M. C. Anderson. Estimating morning change in land surface temperature from MODIS Day/Night observations: Applications for surface energy balance modeling. *Geophys. Res. Lett.*, 44(19):9723–9733, Oct. 2017.
- C. R. Hain, W. T. Crow, M. C. Anderson, and M. Tugrul Yilmaz. Diagnosing neglected soil moisture Source–Sink processes via a thermal Infrared–Based Two-Source energy balance model. *Journal of Hydrometeorology*, 16(3):1070–1086, 2015.
- M. C. Hansen, J. R. G. Townshend, R. S. DeFries, and M. Carroll. Estimation of tree cover using MODIS data at global, continental and regional/local scales. *International Journal of Remote Sensing*, 26(19):4359–4380, 2005.
- V. Humphrey, J. Zscheischler, P. Ciais, L. Gudmundsson, S. Sitch, and S. I. Seneviratne. Sensitivity of atmospheric CO 2 growth rate to observed changes in terrestrial water storage. *Nature*, 560(7720):628–631, Aug. 2018.
- C. M. Iversen. Digging deeper: fine-root responses to rising atmospheric CO concentration in forested ecosystems. New Phytol., 186(2):346–357, Apr. 2010.
- 32. B. Jiang, S. Liang, A. Jia, J. Xu, X. Zhang, Z. Xiao, X. Zhao, K. Jia, and Y. Yao. Validation of the surface daytime net radiation product from version 4.0 GLASS product suite. *IEEE Geoscience and Remote Sensing Letters*, 16(4):509–513, 2019.
- 33. J. Joiner, L. Guanter, R. Lindstrot, M. Voigt, A. P. Vasilkov, E. M. Middleton, K. F. Huemmrich, Y. Yoshida, and C. Frankenberg. Global monitoring of terrestrial chlorophyll fluorescence from moderate-spectral-resolution near-infrared satellite measurements: methodology, simulations, and application to GOME-2. Atmospheric Measurement Techniques, 6(10):2803-2823, 2013.
- A. Kleidon and M. Heimann. A method of determining rooting depth from a terrestrial biosphere model and its impacts on the global water and carbon cycle. *Global Change Biology*, 4(3):275–286, 1998.
- A. K. Knapp and M. D. Smith. Variation among biomes in temporal dynamics of aboveground primary production. *Science*, 291(5503):481–484, Jan. 2001.
- P. Köhler, L. Guanter, and J. Joiner. A linear method for the retrieval of sun-induced chlorophyll fluorescence from GOME-2 and SCIAMACHY data. *Atmospheric Measurement Techniques*, 8(6):2589–2608, 2015.
- 37. S. Koirala, M. Jung, M. Reichstein, I. E. M. de Graaf, G. Camps-Valls, K. Ichii, D. Papale, B. Ráduly, C. R. Schwalm, G. Tramontana, and N. Carvalhais. Global distribution of groundwater-vegetation spatial covariation. *Geophysical Research Letters*, 44(9):4134–4142, 2017.

- 38. G. J. M. D. Lannoy, G. J. M. De Lannoy, R. D. Koster, R. H. Reichle, S. P. P. Mahanama, and Q. Liu. An updated treatment of soil texture and associated hydraulic properties in a global land modeling system. *Journal of Advances in Modeling Earth Systems*, 6(4):957–979, 2014.
- Y. Liu, A. G. Konings, D. Kennedy, and P. Gentine. Global coordination in plant physiological and rooting strategies in response to water stress. *Global Biogeochemical Cycles*, 35(7):NA, 2021.
- S. Manabe. Climate and the ocean circulation. Monthly Weather Review, 97(11):739–774, 1969.
- T. R. Marthews, S. J. Dadson, B. Lehner, S. Abele, and N. Gedney. High-resolution global topographic index values for use in large-scale hydrological modelling. *Hydrology and Earth System Sciences*, 19(1):91–104, 2015.
- 42. R. M. Maxwell and L. E. Condon. Connections between groundwater flow and transpiration partitioning. *Science*, 353(6297):377–380, July 2016.
- E. L. McCormick, D. N. Dralle, W. J. Hahm, A. K. Tune, L. M. Schmidt, K. D. Chadwick, and D. M. Rempe. Widespread woody plant use of water stored in bedrock. *Nature*, 597(7875):225–229, Sep 2021.
- P. C. D. Milly. Climate, soil water storage, and the average annual water balance. Water Resources Research, 30(7):2143–2156, 1994.
- 45. Q. Mu, F. A. Heinsch, M. Zhao, and S. W. Running. Development of a global evapotranspiration algorithm based on MODIS and global meteorology data. *Remote Sensing of Environment*, 111(4):519–536, 2007.
- B. Mueller and S. I. Seneviratne. Hot days induced by precipitation deficits at the global scale. *Proceedings of the National Academy of Sciences*, 109(31):12398–12403, 2012.
- V. M. R. Muggeo. Estimating regression models with unknown break-points. Stat. Med., 22(19):3055–3071, Oct. 2003.
- 48. D. Nadal-Sala, B. E. Medlyn, N. K. Ruehr, C. V. M. Barton, D. S. Ellsworth, C. Gracia, D. T. Tissue, M. G. Tjoelker, and S. Sabaté. Increasing aridity will not offset CO2 fertilization in fast-growing eucalypts with access to deep soil water. *Global Change Biology*, 27(12):2970–2990, 2021.
- 49. D. M. Olson, E. Dinerstein, E. D. Wikramanayake, N. D. Burgess, G. V. N. Powell, E. C. Underwood, J. A. D'amico, I. Itoua, H. E. Strand, J. C. Morrison, C. J. Loucks, T. F. Allnutt, T. H. Ricketts, Y. Kura, J. F. Lamoreux, W. W. Wettengel, P. Hedao, and K. R. Kassem. Terrestrial ecoregions of the world: A new map of life on earth. *BioScience*, 51(11):933, 2001.
- R. Orth, R. D. Koster, and S. I. Seneviratne. Inferring soil moisture memory from streamflow observations using a simple water balance model. *Journal of Hydrometeorology*, 14(6):1773–1790, 2013.
- 51. J. D. Pelletier, P. D. Broxton, P. Hazenberg, X. Zeng, P. A. Troch, G. Niu, Z. Williams, M. A. Brunke, and D. Gochis. A gridded global data set of soil, intact regolith, and sedimentary deposit thicknesses for regional and global land surface modeling. *Journal of Advances in Modeling Earth Systems*, 8(1):41–65, 2016.

- 52. A. G. Pendergrass, R. Knutti, F. Lehner, C. Deser, and B. M. Sanderson. Precipitation variability increases in a warmer climate. *Scientific Reports*, 7(1), 2017.
- D. M. Rempe and W. E. Dietrich. Direct observations of rock moisture, a hidden component of the hydrologic cycle. *Proc. Natl. Acad. Sci. U. S. A.*, 115(11):2664–2669, Mar. 2018.
- H. J. Schenk and R. B. Jackson. Mapping the global distribution of deep roots in relation to climate and soil characteristics. *Geoderma*, 126(1-2):129–140, 2005.
- H. J. Schenk, H. Jochen Schenk, and R. B. Jackson. The global biogeography of roots. *Ecological Monographs*, 72(3):311–328, 2002.
- H. J. Schenk, H. Jochen Schenk, and R. B. Jackson. Rooting depths, lateral root spreads and below-ground/above-ground allometries of plants in water-limited ecosystems. *Journal of Ecology*, 90(3):480–494, 2002.
- 57. L. Schlemmer, C. Schär, D. Lüthi, and L. Strebel. A groundwater and runoff formulation for weather and climate models. *Journal of Advances in Modeling Earth Systems*, 10(8):1809–1832, 2018.
- 58. S. J. Schymanski, M. Sivapalan, M. L. Roderick, L. B. Hutley, and J. Beringer. An optimality-based model of the dynamic feedbacks between natural vegetation and the water balance. *Water Resources Research*, 45(1), 2009.
- S. I. Seneviratne, T. Corti, E. L. Davin, M. Hirschi, E. B. Jaeger, I. Lehner, B. Orlowsky, and A. J. Teuling. Investigating soil moisture-climate interactions in a changing climate: A review. *Earth-Science Reviews*, 99(3-4):125-161, 2010.
- S. I. Seneviratne, D. Luthi, M. Litschi, and C. Schar. Land-atmosphere coupling and climate change in europe. *Nature*, 443(7108):205–209, Sept. 2006.
- S. Siebert, P. Döll, J. Hoogeveen, J.-M. Faures, K. Frenken, and S. Feick. Development and validation of the global map of irrigation areas. *Hydrology and Earth System Sciences*, 9(5):535–547, 2005.
- G. Sivandran and R. L. Bras. Dynamic root distributions in ecohydrological modeling: A case study at walnut gulch experimental watershed. *Water Resources Research*, 49(6):3292–3305, 2013.
- 63. B. Stocker. stineb/cwd: v1.0 initial release, Sept. 2021.
- 64. P. C. Stoy, T. S. El-Madany, J. B. Fisher, P. Gentine, T. Gerken, S. P. Good, A. Klosterhalfen, S. Liu, D. G. Miralles, O. Perez-Priego, A. J. Rigden, T. H. Skaggs, G. Wohlfahrt, R. G. Anderson, A. M. J. Coenders-Gerrits, M. Jung, W. H. Maes, I. Mammarella, M. Mauder, M. Migliavacca, J. A. Nelson, R. Poyatos, M. Reichstein, R. L. Scott, and S. Wolf. Reviews and syntheses: Turning the challenges of partitioning ecosystem evaporation and transpiration into opportunities. *Biogeosciences*, 16(19):3747–3775, 2019.
- 65. A. J. Teuling, S. I. Seneviratne, R. Stöckli, M. Reichstein, E. Moors, P. Ciais, S. Luyssaert, B. van den Hurk, C. Ammann, C. Bernhofer, E. Dellwik, D. Gianelle, B. Gielen, T. Grünwald, K. Klumpp, L. Montagnani, C. Moureaux, M. Sottocornola, and G. Wohlfahrt. Contrasting response of european forest and grassland energy exchange to heatwaves. *Nature Geoscience*, 3(10):722–727, 2010.

- A. J. Teuling, S. I. Seneviratne, C. Williams, and P. A. Troch. Observed timescales of evapotranspiration response to soil moisture. *Geophysical Research Letters*, 33(23), 2006.
- 67. S. E. Thompson, C. J. Harman, A. G. Konings, M. Sivapalan, A. Neal, and P. A. Troch. Comparative hydrology across AmeriFlux sites: The variable roles of climate, vegetation, and groundwater. *Water Resources Research*, 47(10), 2011.
- 68. M. M. Vogel, R. Orth, F. Cheruy, S. Hagemann, R. Lorenz, B. J. J. Hurk, and S. I. Seneviratne. Regional amplification of projected changes in extreme temperatures strongly controlled by soil moisture-temperature feedbacks. *Geophysical Research Letters*, 44(3):1511–1519, 2017.
- J. E. Weaver and R. W. Darland. Soil-Root relationships of certain native grasses in various soil types. *Ecological Monographs*, 19(4):303–338, 1949.
- 70. G. P. Weedon, G. Balsamo, N. Bellouin, S. Gomes, M. J. Best, and P. Viterbo. The WFDEI meteorological forcing data set: WATCH forcing data methodology applied to ERA-Interim reanalysis data. *Water Resour. Res.*, 50(9):7505–7514, Sept. 2014.
- 71. W. WIEDER. Regridded harmonized world soil database v1.2, 2014.
- J. Xiao, J. B. Fisher, H. Hashimoto, K. Ichii, and N. C. Parazoo. Emerging satellite observations for diurnal cycling of ecosystem processes. *Nat Plants*, July 2021.
- Y. Yang, R. J. Donohue, and T. R. McVicar. Global estimation of effective plant rooting depth: Implications for hydrological modeling. *Water Resources Research*, 52(10):8260–8276, 2016.
- 74. Y. Zhang, L. Guanter, J. A. Berry, C. van der Tol, X. Yang, J. Tang, and F. Zhang. Model-based analysis of the relationship between sun-induced chlorophyll fluorescence and gross primary production for remote sensing applications. *Remote Sensing of Environment*, 187:145–155, 2016.