

RESEARCH ARTICLE

EARTH SCIENCES

Global variations in critical drought thresholds that impact vegetation

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Abstract

Identifying the thresholds of drought that, if crossed, suppress vegetation functioning is vital for accurate quantification of how land ecosystems respond to climate variability and change.

We present a globally applicable framework to identify drought thresholds for vegetation responses to different levels of known soil-moisture deficits using four remotely-sensed

vegetation proxies spanning 2001-2018. The thresholds identified represent critical inflection points for changing vegetation response from highly resistant to highly vulnerable in response to drought stress, and as a warning signal for substantial vegetation impacts. Drought thresholds varied geographically, with much lower percentiles of soil moisture anomalies in vegetated areas covered by more forests, corresponding to a comparably stronger capacity to mitigate soil water deficit stress in forested ecosystems. Generally, those lower thresholds are detected in more humid climates. State-of-the-art land models, however, overestimated thresholds of soil moisture (i.e. overestimating drought impacts), especially in more humid areas with higher forest covers and arid areas with few forest covers. Based on climate model projections, we predict that the risk of vegetation damage will increase by the end of 21st century in some hotspots like East Asia, Europe, Amazon, southern Australia, eastern and southern Africa. Our data-based results will inform projections on future drought impacts on terrestrial ecosystems and provide an effective tool for drought management.

Keywords: Drought Threshold, Inflection Points, Vegetation Response, Soil Moisture, Drought Impacts

Main text

Drought will become more intense, frequent, and longer lasting in many regions around the world with increasing global warming¹, posing risks to ecosystem health and societal welfare²⁻⁴. Yet, not all droughts reduce biome productivity, as vegetation resistance can mitigate negative drought impacts for some ecosystem types more so than others⁵⁻⁷. On the other hand, resistance is limited up to the highest drought stress that vegetation can tolerate. Once drought increases further and crosses a critical threshold, an amplification of vegetation response is expected. The divergent vegetation responses to drought are known to vary across biomes⁷ and induce nonlinear responses e.g., due to carry-over effects⁸⁻⁹, both making it challenging to assess drought impacts on global terrestrial ecosystems.

Many previous studies evaluate the ability of an ecosystem to resist drought by linking vegetation growth to potential driving factors or detecting impacts on vegetation growth relative to normal conditions. A previous study¹⁰ proposed a vegetation sensitivity index associated with vegetation memory effects and climate forcings based on an autoregressive modelling approach, empirically estimating the resilience of an ecosystem to global climate variability. However, while advancing the general understanding of ecological response to environmental perturbations, that study did not derive specific thresholds for drought onset. Moreover, many metrics on resistance or resilience are developed and applied to assess vegetation response to drought, which have made much-needed investigations on the divergent ecosystems' ability to withstand drought stress and promoted the understanding of the underlying mechanisms¹¹⁻¹³. Unfortunately, there is still a knowledge gap in distinguishing the climate effects and vegetation capacity by considering coupled climate-ecosystem dynamics. The aim of our study is to overcome the limitation via a percentile-based threshold that considers both the climate anomaly and vegetation response. Moreover, in contrast to some of the metrics conditioned for use in a specific drought event, we intend to provide the general applicable critical threshold for the shifts of vegetation response to drought. As such, our study is expected to reduce uncertainty in assessing drought impacts on vegetation growth in a changing climate.

To examine the global response of biome productivity to drought, we first investigate different vegetation responses to drought using anomalies of soil moisture from the Global Land Evaporation Amsterdam Model (GLEAM) dataset and multiple remotely sensed vegetation proxies that describe vegetation "greenness" and productivity (Fig. 1; Methods). They are the normalized difference vegetation index (NDVI) and its nonlinear generalization, kernel NDVI (kNDVI) (ref¹⁴); the near-infrared reflectance of vegetation (NIRv) and solar-induced chlorophyll fluorescence (SIF). Using 10th percentile to define both drought occurrence and a pronounced vegetation suppression in response to drought in the growing season, we derive the coincidence probability for vegetation suppression per drought year (Fig. 1a). If there is a simple one-to-one mapping between levels of drought intensity and

greenness suppression, then the coincidence probabilities would be near 100%. Instead, our results suggest that major drought does not necessarily translate into a noticeable vegetation response. Indeed, vegetation shows a sizeable reduction in greenness and productivity only for <30% on average of these 1-in-10 drought events during 2001-2018, and even lower in regions like the Amazonia, the Congo basin and boreal areas (Fig. 1).

Hence, a globally unified quantitative definition of detrimental drought impacts on vegetation based on percentiles alone does not work. In addition, the severity of drought that suppresses vegetation varies considerably across different regions and biomes¹⁵⁻¹⁷. For instance, vegetation activity might be highly adapted to a permanent state of water scarcity leading to a low probability of reduced greenness during drought in some arid zones. For a global assessment of drought ecosystem impacts, a critically important question is therefore: what are the drought thresholds that are associated with a high coincidence of vegetation growth suppression potentially leading to widespread vegetation mortality, and how do they vary spatially?

We create an observation-based function that links vegetation response to different percentile levels of negative soil moisture anomalies (Equations 1, 2 in Methods and Supplementary Fig. 1). We use < 10th as the threshold for defining vegetation response to drought because the 10th percentile is a threshold universally set in several studies about vegetation productivity extremes¹⁸⁻¹⁹, and corresponds to a wide occurrence of vegetation suppression. As illustrated in Supplementary Fig. 2, a workflow combining principal component analysis (PCA), the coincidence analysis, and the detection of inflection points using segment regression are carried out in an orderly manner in this study, all are based on experiences from earlier studies of extreme events and detecting ecological thresholds²⁰⁻²². Feature extraction for global lands is first undertaken through PCA (Supplementary Figs. 3-4), which is designed to overcome the difficulty in limited years of data for point-by-point detection and single global thresholds that lose geographical information²¹. By doing so, the dominant modes are determined at regional to global scales to find grid cells with comparable climate conditions and vegetation dynamics.

The applicable framework to identify drought thresholds

To describe drought thresholds, we investigate the non-linear relationship between vegetation response and drought intensity. The main functional feature of this relationship is an inflection point, which is the threshold that marks the start of intensifying drought impacts and defines the drought stress at which level vegetation suppression starts shifting (Supplementary Figs. 1, 5 and 6). We base this assumption of non-linearity on both physiological theory and emerging multiple reports of different phases for vegetation responses to increasing drought stress²⁴⁻³⁰. For those areas with nonlinear relationships (82%-86% of global vegetated areas), we derive the response function as a fractional coincidence r of vegetation suppression when soil moisture is at or below different soil moisture deficit levels (Methods, Eqns (1 ,2)). According to Eq (1), the values of r range from 0 to 1, describing a coincidence between drought and the anomalous vegetation response from none to a complete drought response²¹.

In our framework, a relatively stable and low coincidence rate of vegetation response to increasing drought severity characterizes a *phase A* response. In this phase, we suggest vegetation have a strong capacity to withstand slight-to-mild soil water deficits. We also have a *phase B* response, which is a phase of “rapid response” when the coincidence rate of drought-vegetation-anomaly increases substantially and steeply as droughts intensify beyond T_{Smsurf} (Supplementary Fig. 1). The inflection point T_{Smsurf} that delimits the two phases is a threshold capturing the nonlinear change of vegetation response from Phase A to Phase B, as vegetation adjusts from a slow to a rapid rate of increasing impacts for rising drought. We examine the performance of the non-linear response trajectory for different biomes (Supplementary Fig. 5; Methods), and verify the T_{Smsurf} can be detected.

Spatial patterns of drought thresholds

We find that T_{Smsurf} from satellite-derived vegetation indices occur at values $> 10^{\text{th}}$ percentile of soil water anomalies in $\sim 70\%$ of areas (Fig. 2a; Supplementary Fig. 7). This suggests the

shifts of vegetation response are prone to occur even the drought stress is smaller than the severity of those 1-in-10 drought events in many locations. Our satellite-based T_{SMsurf} , although local consistent between nearby points, does have strongly geographical variation (Fig. 2a). For places having thresholds of lower percentile values, a transition to a vegetation response to drought is induced by higher levels of drought stress. This higher resistance corresponds to a stronger capacity for vegetation to mitigate soil water deficit and reduce sensitivity to drought. Generally, we find the observe-based T_{SMsurf} have more valid and stronger gradient pattern along the forest cover than the aridity, decreasing from low to high fractions of forest covers (Fig. 3a-b), demonstrating that forests are more drought-resistant. The highest percentile of drought thresholds ($> 15^{\text{th}}$ on average) appears where there is $< 25\%$ of trees, irrespective of if in arid or humid areas. This interesting finding suggests that vegetated areas covered by little forest have limited capacity to cope with additional water stress due to the universality that the roots of other vegetation have insufficient depth to access deeper soil water³¹⁻³². Meanwhile, this within-year response may not capture the lag effects while it is well-known that many forest ecosystems may react with substantial lag times to stress events³³⁻³⁴. By contrast, the lowest drought thresholds are for more humid regions covered by medium ($> 30\%$) to high tree cover ($> 50\%$), such as in the Amazon, the Congo, eastern US, southern China, and part of Siberia (Figs. 2a and 3a-b).

Furthermore, drought thresholds are suggested to vary in different biomes (Supplementary Fig. 8). The slight difference between T_{SMsurf} of forests (13.0th percentile on average) and grasses (14.4th percentile on average, Supplementary Fig. 9a) is in line with the expectation that woody structures and deep forest roots³⁵⁻³⁶ allow for a better ability to cope with drought years³⁶⁻³⁸. Irrespective of whether located in tropical, temperate or boreal climatic regions, the T_{SMsurf} values for forests are slightly lower (i.e. less likely to cross a stress threshold) than those of grasses, additionally supporting our finding of decreasing thresholds with increasing forest covers. Notably, for the same vegetation type, both forest and grass in tropical climates have much higher T_{SMsurf} than in temperate and boreal climates, possibly associated with a higher sensitivity of vegetation in tropical areas to drought stress.

Moreover, crop T_{Smsurf} decreases with increasing irrigation, which reveals that agricultural management is an effective measure for alleviating the negative drought impacts. Although intuitive, we verify that higher levels of irrigation can postpone the arrival of the shift of vegetation response to drought (Supplementary Fig. 8b).

Nevertheless, the observationally-based T_{Smsurf} values have some uncertainties, and especially in some tropical regions (e.g. the Amazonia) and cold areas with larger difference among different vegetation proxies (Supplementary 7, 9a). We find much smaller coincidence rates (maximum value <0.3 , Supplementary Fig. 10) in those areas than other regions. Given that drought thresholds are estimated tightly depending on the observation records, the shift of vegetation response to drought may not actually occur if the coincidence rates keep low, and thereby adding to the uncertainty. According to our finding in Fig. 3, we infer strong resistance of vegetation to drought is the main reason for the low coincidence rates, consistent with having high forest covers in these areas.

In addition to vegetation resistance, there are several possible reasons for uncertainties in tropical regions. First, some tropical regions are generally featured by water surplus throughout the year so low coincidence rates are detected even though those humid regions are theoretically more sensitive to drought³⁹. Second, an increase in greenness during the dry season can result from the structural change of forest canopies (e.g. decreased shadow of high trees and leaf abscission). Thus, decrease in vegetation greenness may not timely be observed under drought⁴⁰⁻⁴², and the response of greenness and photosynthesis to drought stress might decouple. Consequently, mean T_{Smsurf} for greenness (NDVI and kNDVI) might underestimate drought thresholds when comparing T_{Smsurf} for vegetation photosynthesis (SIF, GPP and LAI, Supplementary Figs. 7 and 11a-b). In particular, we find much higher percentiles of T_{Smsurf} for GPP and LAI in the Congo (Supplementary Fig. 11a-b), possibly because long-term increase in dry season length reduced the photosynthesis of the Congo rainforest⁴³. By contrast, T_{Smsurf} for NDVI, kNDVI, NIRv and SIF are lower in the Congo rainforest owing to observed increase in greenness under drought (e.g. NDVI, ref⁴⁰) or a response lag (e.g. SIF, ref⁴⁴) to drought.

To verify our main finding on the drought threshold map, we also obtain a similar spatial pattern of drought thresholds over a longer time period (1982-2018) using GIMMS NDVI_{3g} data (Supplementary Fig.11c). While the systematic biases in the satellite sensor for the pre-2000 time series⁴⁵ probably cause the difference (i.e. lower magnitudes of thresholds in >60% of grid cells for GIMMS NDVI_{3g}). Compared to the entire growing-season T_{SMsurf} (Fig. 2a), the averaged T_{SMsurf} for each month slot within the growing season has higher magnitudes in ~54% of grid cells (Supplementary Fig.12). We thus infer that vegetation response to drought might vary at different time scales since drought durations can affect the drought intensities of the whole growing season. Besides, the mean response of the vegetation during the growing season could be partly offset by changing month-to-month vegetation responses to the same levels of drought. In the other direction, areas like Amazon and the Congo show higher growing-season T_{SMsurf} than averaged monthly T_{SMsurf} , which instead suggests a stronger capacity for humid rainforests to buffer against short-term drought stress.

Models underestimate drought thresholds

T_{SMsurf} estimated by an ensemble of global dynamic vegetation models (DGVMs) (Supplementary Table 1) show large uncertainty among different models (Supplementary Fig. 9b). By comparison, modelled T_{SMsurf} is suggested to overestimate satellite-derived T_{SMsurf} in ~58% of vegetated areas (Figs. 2a vs. 2b), mainly in semi-humid to humid areas with medium to high forest cover and in the arid areas with very few forest covers (Fig. 3). This suggests DGVMs may overestimate drought impacts on vegetation as modelled vegetation suppression will occur even if the drought stress is much slighter compared to observations.

This finding indicates the poor characterization of drought-related physiological mechanisms in models (e.g. ref⁴⁶). Modelled T_{SMsurf} have a weaker forest cover gradient than satellite-derived T_{SMsurf} (Fig. 3c). The spatial variations of modelled T_{SMsurf} are thus suggested to be less driven by self-regulation from different forest covers relative to climatic water availability. In areas with high forest covers, model simulations tend to underestimate the positive effects of physiological regulations on mitigating drought damage, e.g. by setting a fixed modelled rooting depth. The relevant empirical parameters of modelled root depth may

cause the calculated overestimates of vegetation sensitivity to soil water stress⁴⁷⁻⁴⁸. Whereas in arid and semi-arid areas with low forest covers, model simulations are likely to underestimate the role of the adaption or acclimation of vegetation to regular soil water shortage, causing an easier response shift to increasing drought stress. Notably, higher modelled T_{SMsurf} in humid regions with non-forest land covers is likely to result from very limited grid cells instead of implying a larger difference of modelled T_{SMsurf} deviating from observations (Fig. 3c-d). Meanwhile, DGVMs are known to underestimate the sensitivity of vegetation transpiration to high VPD levels in warm droughts, predominantly due to their lack of detailed plant hydraulic function⁴⁹. We suggest this is a contributing factor to why modelled T_{SMsurf} is frequently too low (i.e. having less drought damage) in some semi-arid regions at mid-to-low latitudes that have less forest cover (15%~35%).

Future risks of crossing T_{SMsurf}

As we verify above, in many instances DGVMs fail to reproduce the measurement-based values of T_{SMsurf} but it is still worthwhile to derive the changes in soil-moisture-measured droughts from Earth-system models (ESMs) based on the observe-based T_{SMsurf} . Twelve ESMs are used from CMIP6 to quantify any probabilistic change of droughts, defined by crossing the data-based threshold T_{SMsurf} (hereafter referred to as T_{SMsurf} -inferred droughts), between two decades of history (2001-2020) and future (2081-2100) under SSP2-4.5 and SSP5-8.5 (Fig. 4)⁵⁰.

Under SSP2-4.5, T_{SMsurf} -inferred droughts are projected to increase in half of the vegetated areas where T_{SMsurf} is robustly detectable in present-day measurements. We find particularly high risks of raised T_{SMsurf} -inferred droughts in key hotspots of Europe, East Asia, eastern and northern of North America and some areas of the Amazon, Australia, and Africa (Fig. 4). By the end of the 21st century, we estimate that the largest increase of the occurrence probability of T_{SMsurf} -inferred droughts will reach >3% in those hotspots (Fig. 4a). While, T_{SMsurf} -inferred droughts will generally increase in more humid regions, especially where the aridity index values >1.0, projected by >70% of ESMs in humid regions (Fig. 4b). The increases in drought frequency in more humid regions could be caused by the

rising-temperature-induced intensification of evapotranspiration (ET), which exacerbates the soil water deficit. Whereas ET is suppressed in more arid areas, in tandem with increased water use efficiency under higher atmospheric CO₂.

Of particular interest is that the occurrence probability for T_{SMsurf} -inferred droughts does not become higher in regions like Europe and western North America under the high-emission scenario of SSP5-8.5 (Fig. 4c-d), probably due to a balance between increased temperatures and precipitation levels. This finding indicates that increases in atmospheric CO₂ concentrations may partly ameliorate the impacts of future drought stress by increasing the water use efficiency of vegetation⁵¹. This CO₂ effect thus impacts the projected soil moisture and partially offsets any increases in crossing the drought thresholds of soil moisture⁵². The exceptions to this lack of change are for some hotspots like Amazon, Australia and eastern Africa, as already noted (Fig. 4c-d). Future drought conditions are thus suggested to have a relatively smaller change when characterized by soil moisture alone⁵³⁻⁵⁴.

We note that the changes in occurrence probability for future droughts are underestimated using a universal 10th percentile instead of using T_{SMsurf} , especially marked in Europe, eastern North America, and South America (Supplementary Fig.13). Our framework for threshold detection therefore has important implications for accurately quantifying the impacts of future climates on vegetation development. The understanding of climate-ecosystem dynamics under drought is particularly much-needed for those regions that are projected to have rapid increases in extreme heat event frequency and for dryland locations that, by definition, are already subjected to drought stress. However, caveats concern the impacts of data quality, response scaling, different durations of drought within the growing season, and possible acclimation and adaptation of vegetation to climate change (see more in the Supplementary Discussion). Besides, this data-led method for identifying thresholds is limited by observations so the complicated non-linearity between drought stress and vegetation response needs further explorations.

In summary, our study develops a diagnostic drought threshold that describes the critical inflection point for vegetation responses to drought. Drought thresholds tend to vary geographically with fraction of forest cover. The global variations of drought thresholds provide a unique perspective on investigating drought impacts on vegetation worldwide. Moreover, model evaluation on thresholds revealed that the state-of-the-art vegetation models tended to overestimate the negative drought impacts on ecosystem structure and functioning, such as their vegetation productivity and terrestrial carbon cycle, particularly in humid ecosystems with high forest covers. Although our results find humid ecosystems are less affected if they are covered by more forest, future increased risks of drought in those hotspots call for more attention towards humid ecosystems, especially to those contributing a lot to global carbon sinks, human livelihood and economic development under future climate change. We recommend advancing satellite observations and improving model projections by refining biological processes in order to develop effective ecosystem policies and drought management strategies to alleviate future drought stress.

Methods

Drought and aridity metrics

Drought indicators The moisture contents of surface soil are obtained from the satellite-derived Global Land Evaporation Amsterdam Model (GLEAM) data set (version v3.3a)⁵⁵. The monthly soil moisture data is aggregated from the original spatial resolution of $0.25^{\circ} \times 0.25^{\circ}$ to the unified spatial resolution of $0.5^{\circ} \times 0.5^{\circ}$ that we use throughout for all variables, and averaged for the growing seasons of each year. We define the growing season use the existing dataset developed by Zhu et al. (ref⁵⁶, see Supplementary Fig. 14), which is first determined from the GIMMS LAI_{3g} data set in 1981-2009 using a Savitzky-Golay filter and then refined by excluding the ground-freeze period identified by the Freeze/Thaw Earth System Data record⁵⁷. The anomalies of growing-season drought indicators are detrended by subtracting the linear trend for 2001-2018.

Aridity index We calculate a simple index of background aridity to assess the projections of historical and future vegetation responses to drought. Our index is a function of the amount of background dryness or wetness (Fig. 3-4). The aridity index is derived as the multiyear average (2001-2018) of the ratio of annual P to annual PET and using values from the CRU TS4.04 data set⁵⁸. Values of the aridity index increase in value between arid and humid regions. Values <0.65 we refer to as drylands, with 0.2-0.5 defined as semi-arid regions and 0.5-0.65 defined as semi-humid regions⁵⁹.

Vegetation indicators

Satellite observations NDVI is used as a reliable surrogate measure of vegetation greenness, broadly representing chlorophyll content and canopy structure. Monthly NDVI for 2001-2018 is derived from the Moderate Resolution Imaging Spectroradiometer (MODIS) Collection 6 MOD13C2 with the native spatial resolution $0.05 \times 0.05^\circ$ (<https://lpdaac.usgs.gov/products/mod13c2v006/>). MODIS NDVI is aggregated to $0.5 \times 0.5^\circ$ grid resolution, averaging within each such grid, the NDVI values in all subpixels, which is also applied for other satellite data. Very low monthly values (<0.1) are removed from our analysis to exclude areas of barren, rock, sand (i.e. deserts) or snow. We also use kNDVI, simply written as $kNDVI = \tanh(NDVI)^2$. This vegetation index is a nonlinear version of NDVI, developed by a kernel methods framework. kNDVI is believed to reduce saturation effects and enhance robustness to noise¹⁴.

Our third estimate of vegetation dynamics uses data from NIRv⁶². The NIRv version is MODIS 16-day NDVI and NIR reflectance ($NDVI \times NIR$), derived from the MCD43C4 Vegetation Index Product with a spatial resolution of 0.05° . All values where $NIRv \leq 0$ are excluded, following the threshold of Badgley et al. (ref⁶⁰).

We obtain a fourth estimate of the state of the vegetation from SIF, which is a measure of the amount of light emitted by chlorophyll, itself a proxy of photosynthetic activity⁶¹. We use a gridded global contiguous SIF (CSIF) data set, reconstructed from SIF observations from the Orbiting Carbon Observatory-2 (OCO-2) and using a machine-learning approach⁶².

This clear-sky set of instantaneous CSIF data has a 0.05° spatial and four-day temporal resolution.

To remove the possibility that the different thresholds for satellite observations and model simulations are caused only by using inconsistent variables, we also used MODIS LAI from MCD15A2Hv006 to compare with modelled LAI. This LAI product is an 8-day composite dataset with 500m spatial resolution⁶³. It is produced using a main algorithm based on the radiative transfer model and a backup algorithm using the relationship between NDVI and LAI.

We also apply gross primary production (GPP) from Vegetation Photosynthesis Model (VPM) (ref²³) as a proxy of vegetation photosynthesis. This GPP dataset is based on an improved light use efficiency theory and is driven by remotely sensed MODIS Enhanced Vegetation Index and climate data from NCEP Reanalysis II.

In addition, we use the NOAA Global Inventory Monitoring and Modeling System third-generation global dataset (GIMMS NDVI_{3g}, 1982-2018, original resolution $1/12 \times 1/12^\circ$ and 15-day) observed by the Very High Resolution Radiometer (AVHRR) sensors for a longer period estimation⁶⁴⁻⁶⁵.

Simulations by Dynamic global vegetation models (DGVMs) We study modelled responses to drought using simulated leaf area index (LAI), defined as the amount of leaf area per unit of ground area. The gridded estimates of LAI from nine process-oriented DGVMs in the TRENDY-v8 model intercomparison project are used (Supplementary Table 1), including JSBACH, JULES-ES, LPI-GUESS, LPX-Bern, ORCHIDEE-CNP, SDGVM, OCN, ISBA-CTRIP, and VISIT. We apply model simulations from the TRENDY S3 experiment, which has the full set of temporally-changing forcings, including CO₂ levels, climate and land use.

Soil drought estimated by CMIP6 models. We select 12 Earth-system model (ESM) simulations, operated with “all forcings”, from the CMIP6 archive (Supplementary Table 2).

These forcings are the combination of shared socioeconomic pathways (SSPs) and forcing levels of the representative concentration pathway (RCP) in the Scenario Model

Intercomparison Project⁵⁰. We select only the ESMs where the original spatial resolutions $\leq 2^\circ$ to avoid the potentially large uncertainty caused by lower resolutions. We use these multi-model climatic projections under two emission scenarios of SSP2-4.5 for intermediate emissions of air pollutants and greenhouse gases and SSP5-8.5 for high emissions.

We use the soil-moisture content for the 0-10 cm layer (CMIP6 variable name: “mrsos”) from the CMIP6 simulations to indicate future drought, mirroring our similar satellite-based measurements. All ESM variables are interpolated to a spatial resolution of $0.5^\circ \times 0.5^\circ$ to match the satellite-derived drought thresholds.

Statistics

Feature extraction using principal component analysis (PCA)

Earth Observed (EO) data enable major new insights into the functioning of the land surface, offering incredible high-resolution spatial information. However, the remote sensing technique only provides satellite observations for a relatively short period (i.e., 18 years in this study). Here, we focus on detecting vegetation response to rarely occurring drought. With only 18 years of satellite observations, obtaining sufficient samples of anomalous events for statistical analysis at an individual grid is difficult and has large uncertainty. At the high spatial resolution of EO data, adjacent points are not independent, and so a method is needed to identify the dominant geographical modes of variation, although not scales (such as global) that remove spatial signals. A method strongly suited to these requirements is PCA, as used elsewhere for feature extraction in anomalous event detection^{21, 66-67}.

In the first step, we apply PCA-based feature extraction, such that in the low-dimensional space of principal components, similar grid cells are close to each other, even if they may be geographically distant (step i in Supplementary Fig. 2). This method finds a compromise between fully local and global thresholding and refrains from (i) inaccurately estimating drought impacts on vegetation growth by setting a global unified threshold for each grid, (ii) setting an equal distribution of drought occurrence for each grid at the spatial extent against the reality that drought events do not occur equally everywhere. Using PCA,

we identify areas characterized by climate and vegetation similarities and break through the limitations for short-term satellite observation and rare occurring anomalous events.

We select a set of variables that are believed to meet the need that (i) they impact the characteristics of drought occurrence and related vegetation growth levels, and (ii) contribute to the geographical distinction of climate conditions and vegetation distribution, composition, and growth condition. Here, seven variables are used for our PCA derivation; mean annual precipitation (MAP), mean annual temperature (MAT), interannual variability of the vapor pressure deficit (VPDvar), the fraction of tree cover (Treefrac), interannual variability of NDVI and NIRv (NDVIvar and NIRvvar) and species richness (SpeciesN) (see details in Supplementary Table 3). Among them, the interannual variabilities of VPD, NDVI and NIRv are calculated as the standard deviation of variable anomalies.

MAP, MAT and VPDvar are used as climate forcing factors. MAP and MAT represent the mean climatological condition for vegetation growth. We also use VPDvar as a metric of the interannual variability of atmospheric water demand. Furthermore, we include the Treefrac variable because it provides information on land cover types and is known to characterise divergent ecosystem responses to water availability^{38, 67}. Biodiversity is also believed to affect the response of the ecosystem to drought⁶⁸⁻⁶⁹ so we also use SpeciesN to indicate the ecosystem functional diversity and stability of the ecosystem. Finally, NDVIvar and NIRvvar are used to reflect the natural variability of vegetation greenness and photosynthesis. Note that variations of kNDVI and CSIF are not considered in our PCA analysis. We exclude kNDVI to avoid information redundancy because its variability is similar to NDVI. We also exclude CSIF due to the short record of OCO-2 SIF 2014-2017 used to reconstruct CSIF, which may not show a full representation of the interannual variability of vegetation growth. Additionally, variables that describe soil conditions, such as soil fertility are not included because they did not provide improved explanatory power (Supplementary Fig. 4). This is perhaps because the impacts of soil characteristics may have been reflected in vegetation covers, natural variability of vegetation growth, and ecosystem biodiversity.

Accordingly, seven principal components (PCs) of PCA are obtained, contributed by all original variables and ranked by explained variances in decreasing order. The PCs replace the seven original variables to become new features so each grid cell possesses values of PC1-PC7. By comparing the cumulative explanation of the PCs (Supplementary Fig. 3), we chose to retain the first three principal components (PCs), as they explain, globally, 83% of the variance, yet reduce dimension and classify global lands into different clusters with comparable climate and vegetation conditions. As shown in Supplementary Fig. 3, taking the first 3 PCs substantially increases the cumulative variance explained (compared to taking the first 2 PCs (66%) or 1 PC (42%)), and allows the distinguishing of feature differentiation of global lands driven by vegetation cover and climates.

In the next step, we link a target grid cell $x(lat, lon)$ in the geographical domain and find its location point $p(u, v, m)$ in the three-dimensional (3D) space of PC1-PC3, based on their first three PC scores (Supplementary Fig. 2). Taking $p(u, v, m)$ as a centre in the PC space, we find all the neighbor points in a window of $3 \times 3 \times 3$ meshes whose width is within 4% of the total length of the corresponding PCs²¹. Although $p(u, v, m)$ may not be geographically close to the grid cells that are located as the neighbour points in the 3D PC space, they are divided into the same clustering based on comparable climate and vegetation conditions. In such a PCA clustering (i.e. all points in a set of $3 \times 3 \times 3$ meshes), we define droughts (<10th percentile for anomalies surface or root-zone soil water content) and vegetation response to drought (<10th percentile for anomalies of vegetation indices). For example, if one clustering owns 50 points (point $p + 49$ neighbour points), the anomalies of 10th percentile will be derived based on 900 values (50 grid cells \times 18 years) of drought or vegetation indicators. We then have available the coincidence rates, as well as the drought threshold based on those 50 grid cells in the clustering, and the final results are assigned to the grid cell $x(lat, lon)$.

Here, splitting up the space of PCs allows us to obtain more regionally relevant event thresholds (for details, see ref²¹), and allows us to achieve the compromise between a completely global scale and overly local scales. The global map of drought thresholds in this study can reflect the spatial heterogeneity because our approach to derive thresholds retain

more information than those for the regional scale. On average, there are only ~0.2% of all grid cells located in the same centered mesh of a $3 \times 3 \times 3$ window in the PCs' space so that these grid cells would be assigned by the same threshold values (see the spatial pattern b in the diagram, Supplementary Fig. 2). This ensures that different grid cells would lose less information on differentiating the divergent vegetation responses to drought.

Response trajectories for vegetation to different levels of drought stress

Coincidence analysis. The main overall feature of our calculations is coincidence analysis²⁰ to identify droughts and the resulting suppression of vegetation greenness and photosynthetic activity. In particular, this method allows us to build response curves for vegetation activity at different levels of drought stress. We assume vegetation responses to be independent between years and including when major drought happens in consecutive years. That is, we assume no interannual memory that may cause lagged effects as well as post-drought legacy effects. For all grid cells in each moving $3 \times 3 \times 3$ set based on the first three leading PCs (see details above), we identify drought occurrence indicated by SMSurf anomalies and vegetation response indicated by anomalies of different satellite observations or model simulations.

We quantify the coincidence rate (r) using the single vegetation threshold of any fraction less than the 10th percentile of vegetation anomaly ($\theta_{veg < 10th}$). We applied < 10th percentile to define vegetation response to drought because, at this level, the impacts of drought on vegetation have widely occurred, and can be detected and assessed in different locations under current climate change¹⁹. We suggest that any drought frequently crossing this threshold would accumulatively impact the vegetation development and may provide an important insight into the potential change of ecosystem structure and functioning under future climate change. The crossing of this threshold is recorded per drought year when the drought level of the corresponding growing seasons is equal to or less than the percentile-based anomalies ($\theta_{dro \leq q}$) of SMSurf. Hence r is calculated as:

$$r = \frac{Fre(G < \theta_{veg < 10th} \forall t \text{ when } \theta_{dro} \leq \theta_{dro \leq q})}{Fre(\forall t \text{ when } \theta_{dro} \leq \theta_{dro \leq q})} \quad (1)$$

where G is a metric of anomalous vegetation greenness or photosynthetic capacity, t is the drought year and $\theta_{dro \leq q}$ is a soil moisture threshold expressed by percentile q . The denominator of Eq. (1) is the frequency of drought occurrence below percentile q . The numerator is similarly the frequency of when soil moisture is below percentile q , except we only now count this time when additionally, there is vegetation suppression.

Here, the drought threshold q corresponds to the anomalies of SMSurf as a function of the percentile ranges on the x -axis in Supplementary Fig. 1. Smaller q values correspond to more extreme soil moisture deficits. Quantity r for each location accordingly corresponds to the y -axis in Supplementary Fig. 1 (and subsequent plots). In each window, r is estimated only when sufficient samples (> 162 , 9 grid cells \times 18 years) are available.

The relationship of r and q We explore the relationship between percentile-based drought thresholds and the coincidence rates, r , of drought-vegetation anomalies. It is expected that the coincidence rates of concurrence between drought and vegetation response are low and change little when experiencing slight to mild droughts owing to vegetation's resistance (phase A in Supplementary Fig. 1). The coincidence rates then sharply increase once the drought stress levels exceed their tolerance limits (phase B). To confirm this form of response, we sampled r and $\theta_{dro \leq q}$ with q ranging from 1st - 50th percentiles (with an interval of 1 percentile) of soil moisture anomalies. By testing different nonlinear curve fitting, we believe an exponential-type curve is the optimal functional form to capture the relationship between drought threshold and drought-vegetation-anomaly coincidence rates (See details in the Supplementary Discussion 2 and Supplementary Figs. 15-16). We thus applied an exponential function, across the full range of 1st - 50th percentiles, written as $r = me^{\beta\theta_{dro \leq q}} + k$, to test whether the response trajectories of vegetation follow a generic response curve that a flexible non-linear fit is better than a linear fit. If so, the relationship between drought stress levels and vegetation decrease shows threshold behavior and vegetation response can be distinguished from two segments. Hence, a breakpoint is able to be searched.

We exclude areas where response trajectories of vegetation do not follow a generic format of the response curve if (i) a simple linear model outperforms a non-linear model. To select the better model, we apply the Akaike information criterion (AIC) to evaluate the fitness of linear and non-linear models, If AIC of the linear model is larger than that of the non-linear model unless the linear fitness performance is superior ($R^2 > 0.95$), we regard the response curve follows non-linear relationship between drought severity and vegetation response. (ii) poor fitness of the exponential function for the response curve (the adjusted R squared ≤ 0.5) (Supplementary Fig. 6). In order to avoid mistakenly identifying inflection points from satellite data, we consider thresholds in the areas where the maximum value r is < 0.3 (Supplementary Fig. 10) or thresholds detected only in a single vegetation proxy possess (not marked by dots in Fig. 2) are unrobust.

Identification of drought thresholds for vegetation response

We identify the percentile-based drought threshold (T_{SMsurf}), which characterize the initiation of vegetation response to rising drought levels. T_{SMsurf} are percentile values corresponding to the anomalous soil moisture $\theta_{dro \leq T_{SMsurf}}$ and $\theta_{dro \leq T_{SMroot}}$, respectively. Although we fit a function for r with an exponential continuous in its first derivation, here we search for T_{SMsurf} as an inflection point of an abrupt change by vegetation as drought intensifies. We use a piecewise linear regression^{22, 70} to obtain two distinct linear segments based on the samples r with $\theta_{dro \leq q}$ when $q=50$ th, 49th, 48th, ..., 3th, 2th, 1th of SMsurf:

$$r = \begin{cases} \beta_0 + \beta_1 \theta_{dro \leq q} + \varepsilon & \text{for } \theta_{dro \leq q} \geq \theta_{dro \leq T_{SMsurf}} \\ \beta_0 + \beta_1 \theta_{dro \leq q} + \beta_2 (\theta_{dro \leq q} - \theta_{dro \leq T_{SMsurf}}) + \varepsilon & \text{for } \theta_{dro \leq q} < \theta_{dro \leq T_{SMsurf}} \end{cases} \quad (2)$$

where $\theta_{dro \leq T_{SMsurf}}$ is the inflection point of the two-segmented linear regression between $\theta_{dro \leq q}$ and r , β_0 , β_1 and β_2 are fitted regression coefficients of the two distinct linear segments, from which $\theta_{dro \leq T_{SMsurf}}$ is inferred and ε is the residual of the fit. Note that β_1 is non-zero when $\theta_{dro \leq q}$ is $< \theta_{dro \leq T_{SMsurf}}$. We determine the optimal segmentation with a best fit was determined by the minimal square error of this linear model. To minimize the possibility

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of incorrectly identifying T_{SMsurf} , we exclude cases where the slope (β_1) of the first segment of the line regression (the upper line in Eq. 2) is larger than the slope of the second segment (the lower line in Eq. 2) to avoid deriving the wrong inflection point. We present the sample locations for estimating percentile-based thresholds in the Supplementary Information (Supplementary Fig. 5).

Using surface soil moisture, the global patterns of T_{SMsurf} based on NDVI, kNDVI, NIRv, and SIF verify the consistency among the vegetation indicators (Supplementary Fig. 7). We thus average drought thresholds across all satellite observations at each grid and present histograms of all data combinations (Fig. 2a). To evaluate the impacts of drought durations on the thresholds, we also detect the T_{SMsurf} for each month (Supplementary Fig. 12) similar to Fig. 2a. Mean T_{SMsurf} are averaged by T_{SMsurf} of January to December only if those months are in the growing season for that location. We make a comparison between the average monthly T_{SMsurf} for months during the growing season only and the whole growing season T_{SMsurf} .

For the model evaluation of T_{SMsurf} , we exclude the model simulations from JULES-ES, ORCHIDEE-CNP and SDGVM. Among all nine vegetation models (Supplementary Table 2), JULES-ES, ORCHIDEE-CNP and SDGVM have very low coincidence rates (maximum values <0.3) for exceeding 90% of grid cells (Supplementary Fig. 17). We thus consider JULES-ES, ORCHIDEE-CNP and SDGVM as having failed to capture the spatial patterns of thresholds. To further investigate the difference between observation-based thresholds and model-derived thresholds, we also display the drought thresholds along the gradients of aridity index and tree cover for areas overlapped by both observation-derived and model-derived thresholds (Fig. 3). We exclude those meshes where averages are calculated by thresholds from fewer than five grid cells in Fig. 3.

Shuffling test to remove randomness

We perform a statistical “shuffle” test to assess the significance of the critical thresholds for testing the robustness of our findings. We create 500 surrogate time series of the vegetation time series by randomly shuffling the original dates of the vegetation time series, and estimate the T_{SMsurf} based on the time series of the drought indicators and the surrogate

vegetation time series. Our null hypothesis is that T_{SMsurf} for the original and surrogate vegetation time series do not differ systematically. This hypothesis applies to “nonsignificant T_{SMsurf} ”. An alternative hypothesis is that the thresholds T for the original and surrogate vegetation time series differed systematically, and so our discovered links between driver and response are valid. The estimates of T_{SMsurf} are assumed to reject the null hypothesis and be significant if they were 2.5-97.5% outside the surrogate distribution, corresponding to $P < 0.05$. When we perform such an analysis, we find we reject the nonsignificant T_{SMsurf} ($P > 0.05$) possibility and so adopt the alternative hypothesis that there is a nonlinear relationship, as based on both satellite observations and model simulations from DGVMs.

Assessing future changes in T_{SMsurf} -inferred drought frequencies

For each grid, we first derive anomalies of the moisture content of surface soil for these two periods by subtracting their linear trends of 2001-2020 and 2081-2100, respectively. We then derive the anomalies of soil-moisture corresponding to the percentiles of satellite-derived T_{SMsurf} and calculate the probabilities of the occurrence of droughts smaller than T_{SMsurf} -based anomalies at historical (2001-2020) and the future time (2081-2100). Since model simulations are suggested to have a poor characterization of drought-related physiological mechanisms (Fig. 3c), we only predict the changes in probabilities of drought occurrence along the future aridity index. Considering that those satellite-derived T_{SMsurf} still maintain large uncertainty in some tropical, boreal and arctic areas, we exclude these areas when predicting the future probability of drought occurrence.

Acknowledgements

This study was supported by the National Natural Science Foundation of China (41988101).

Author contributions

S. Piao and X. Li designed the research; M. Mahecha, S. Piao and X. Li contributed to the methods; X. Li performed analysis and created all figures. X. Li, C. Huntingford and S. Piao created the first draft of the paper; and all authors contributed to the interpretation of the results and to the text.

Data availability. MOD13C2 is available at <https://lpdaac.usgs.gov/products/mod13c2v006/>; MCD43C4 is available at <https://lpdaac.usgs.gov/products/mcd43c4v006/>; MCD15A2H Version 6 is available at <https://lpdaac.usgs.gov/products/mcd15a2hv006/>; CSIF is available at <https://osf.io/8xqy6/>; GLEAM soil moisture is available at <https://www.gleam.eu/>; Model simulations of CMIP6 is available at <https://pcmdi.llnl.gov/CMIP6/>.

Code availability. All computer codes for the analysis of the data are available from the corresponding author on reasonable request.

Competing interests

The authors declare no competing interests.

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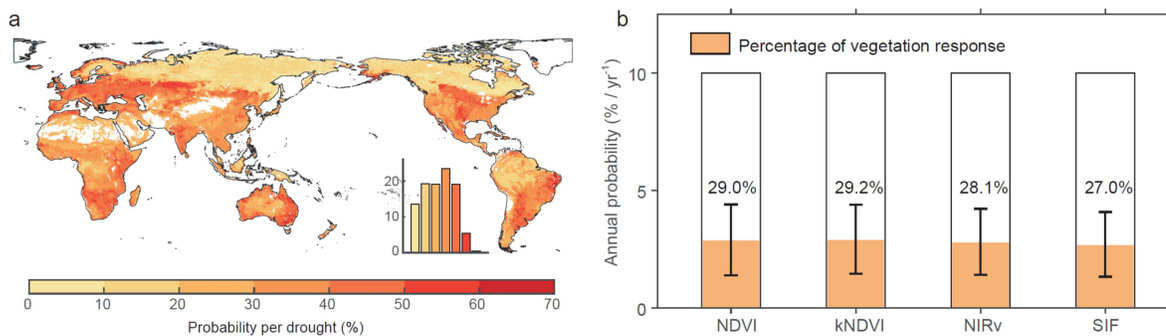


Figure 1. Drought-induced decreases in vegetation in years of drought occurrence during 2001-2018. A. Average probability of growing-season vegetation anomalies lower than the threshold of 10th percentile when drought years are identified by < 10th percentile of anomalies of growing-season drought indicators. Drought occurrences are identified by the moisture contents of surface soil (Smsurf). Vegetation declines are detected based on four satellite observations: normalized difference vegetation index (NDVI), kernel NDVI (kNDVI), near-infrared reflectance of vegetation (NIRv) and solar-induced chlorophyll fluorescence (SIF). The white areas in **a** represent the absence of values due to the lack of vegetation cover. **B.** Overall probabilities of drought occurrence per year (open bars) and of being in both a drought year and when the vegetation has declined (colors) for kNDVI, NDVI, NIRv, and SIF (27.0%-29.2% on average for occurrence of vegetation response per drought year). The probabilities are global and derived from all grids for which values are available. The errorbars are indicated by the standard deviation of all probability values. The texts on bars in **b** represent coincidence probabilities for vegetation response in each drought year.

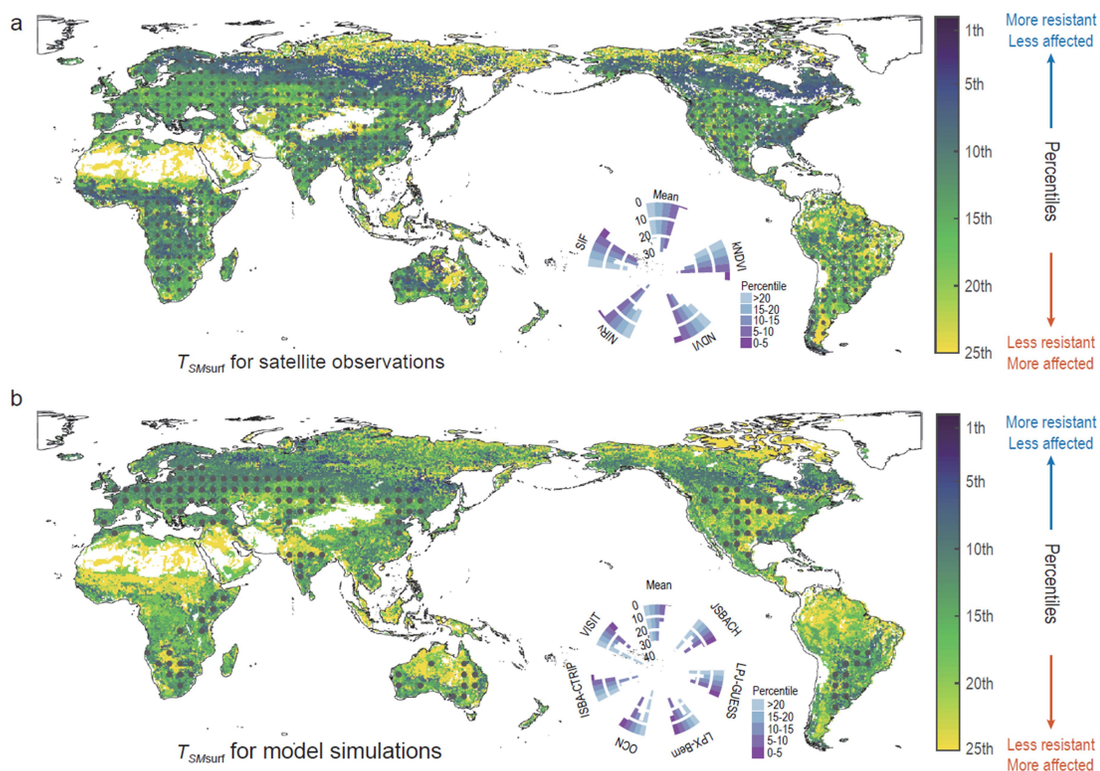


Figure 2. Global map of drought threshold (T_{SMSurf}) for vegetation response to surface soil-moisture anomalies during the growing season. a. The main map of thresholds aggregated over different combinations of four satellite-based on vegetation indicators and moisture content of surface soil (SMSurf) for 2001-2018. Droughts are identified by anomalies of SMSurf, and changes in vegetation activity and greenness are identified by anomalies of NDVI, kNDVI, NIRv and SIF. The areas averaged by more than one single satellite dataset and model simulations with maximum coincidence rates exceeding 0.3 marked by dots. The color palette denotes the percentiles of SMSurf anomalies. Lower percentiles correspond to the definition of a more severe soil water deficit as drought. Under the same degree of vegetation suppression, lower percentiles for thresholds imply locations where vegetation is more resistant to drought and so affected less by arid conditions while the higher percentiles are the opposite. The inset on the right shows the distributions of satellite thresholds as small histograms for the individual combinations of SMSurf and vegetation indicators (NDVI, kNDVI, NIRv, and SIF). The numbers 0, 10, 20 and 30 in the small histograms represent the percentages of vegetated areas in different subranges, indicated by the colour bars on the left **b.** The same format as for **a**, but the spatial patterns of aggregated threshold are from model simulations during 2001-2018. Droughts are identified by the same anomalies of SMSurf as in **a**, but the vegetation activity is estimated by anomalies of leaf area index (LAI) from six Dynamic Global Vegetation Models (DGVMs) in the TRENDY ensemble (JABACH, LPJ-GUESS, LPX-Bern, OCN, ISBA-CTIP and VISIT).

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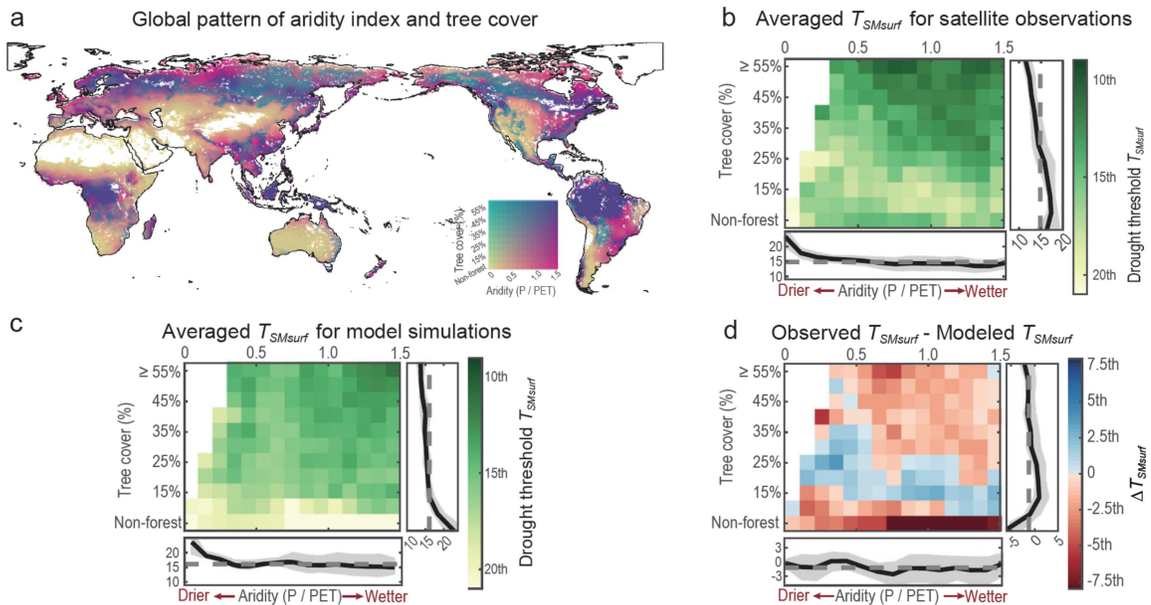


Figure 3. Patterns of drought threshold (T_{SMsurf}) along the gradients of aridity and tree cover.

a. Map for different levels of aridity (horizontal axis of the color palette) and fractions of tree cover (vertical axis of the color palette). **b.** main panel: variation of averaged drought thresholds derived from satellite observations as a function of different levels of aridity index measured by mean annual P / PET (x-axis) and the fraction of tree covers (y-axis). Non-forest is defined as <1% of tree cover. Right: change of drought thresholds along the fraction of tree cover for the means (line) and variations (shaded areas) across all levels of aridity (0-1.5). Lower: change of drought thresholds along the aridity index for the means (line) and variations (shaded areas) across all ranges of tree cover. Drier areas are denoted by smaller values of aridity index, and vice versa. Smaller percentiles of thresholds indicate that a higher drought stress level should be reached to cause a pronounced vegetation response. **c.** Same format as for **b**, but the variation of average drought thresholds derived from model simulations. **d.** Same format as for **a**, but the difference between satellite observations and model simulations. The positive and negative values suggest the underestimation and overestimation of drought thresholds by model simulations, respectively.

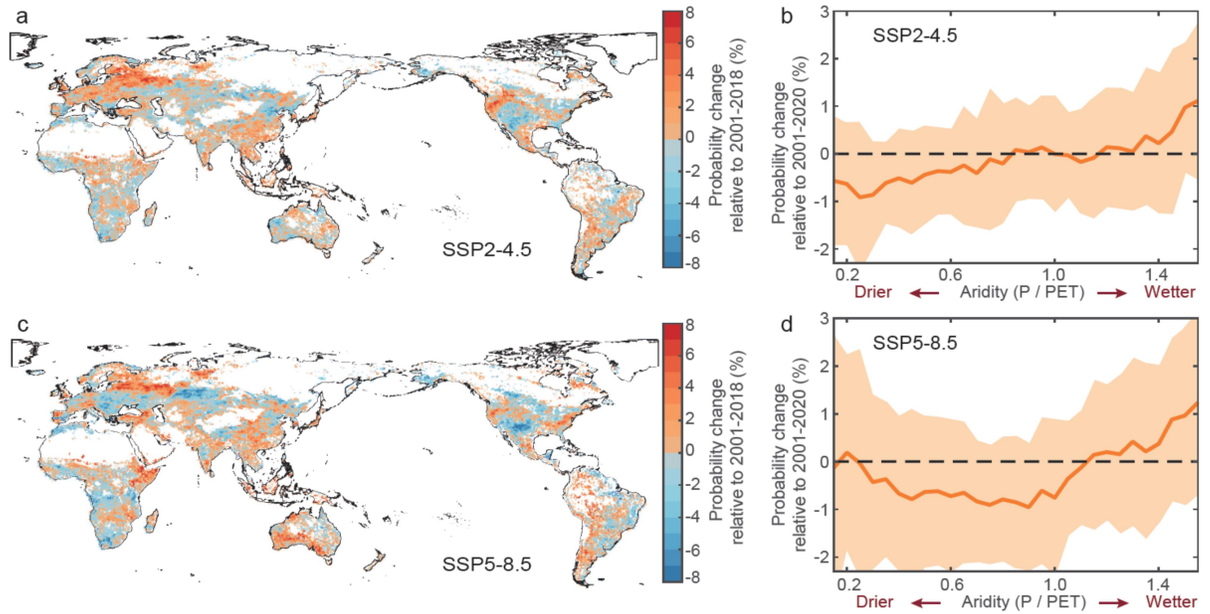


Figure 4. Patterns of annual probability for T-inferred drought occurrence in the future two decades 2081-2100 relative to the recent two decades 2001-2020 under SSP2-4.5 and SSP5-8.5. **a.** Spatial distribution of probability change for drought occurrence per year during 2081-2100 relative to the historical period of 2001-2020 under SSP2-4.5, derived from ensemble means of 12 Earth-system models (ESMs) (see Supplementary Table 2). **b.** The average change of probability for drought occurrence per year during 2081-2100 along the gradients of aridity (mean annual P / PET of 2081-2100, *x*-axis), relative to the historical period of 2001-2020 under SSP2-4.5. Non-forest is defined as <1% of tree cover. **c-d.** Same format as for **a-b**, but under SSP 5-8.5. Soil-moisture content for the 0-10 cm layer (CMIP6 variable name: “mrsos”) was used as an indicator of future drought. The variable “mrsos” was interpolated to a reference spatial resolution of $0.5 \times 0.5^\circ$ to match the satellite-derived drought thresholds and for mapping purposes.

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