Investigating the response of LAI to droughts in southern African vegetation using observations and model-simulations.

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Abstract

In many regions of the world, frequent and continual dry spells are exacerbating drought conditions, which have severe impacts on vegetation biomes. Vegetation in southern Africa is among the most affected by drought. Here, we assessed the spatio-temporal characteristics of meteorological drought in southern Africa using the Standardized Precipitation Evapotranspiration Index (SPEI) over a 30-year period (1982 – 2011). The severity and the effects of droughts on vegetation productiveness were examined at different drought time-scales (1- to 24-month time-scales). In this study, we characterized vegetation using the Leaf Area Index, after evaluating its relationship with the Normalized Difference Vegetation Index. Correlating the LAI with the SPEI, we found that the LAI responds strongly ($r = 0.6$) to drought over the central and southeastern parts of the region, with weaker impacts ($r < 0.4$) over parts of Madagascar, Angola and western parts of South Africa. Furthermore, the latitudinal distribution of LAI responses to drought indicates a similar temporal pattern but different magnitudes across timescales. The results of the study also showed that the seasonal response across different southern African biomes varies in magnitude and occurs mostly at shorter to intermediate timescales. The semi-desert biome strongly correlates ($r = 0.95$) to drought as characterized by the SPEI at 6-month timescale in the MAM (summer) season, while the tropical forest biome shows the weakest response ($r = 0.35$) at 6-month timescale in the DJF (hot and rainy) season. In addition, we found a stronger response (in the year 1983, $r = 0.84$ over Namibia and eastern parts of South African) of the LAI to drought during dry years as compared to wet years; and we found different temporal variability in global and regional responses across different biomes.

We also examined how well an ensemble of state-of-the-art dynamic global vegetation models (DGVMs) simulate the LAI and its response to drought. The spatial and seasonal response of the LAI to drought is mostly overestimated in the DGVM multi-model ensemble compared to the response calculated for the observation-based data. The correlation coefficient values for the multi-model ensemble are as high as 0.76 (annual) over South Africa, and 0.98 in MAM season over the temperate grassland biome. Furthermore, the DGVM model ensemble shows positive biases (3-month or longer) in the simulation of spatial distribution of drought timescales and overestimates the seasonal distribution timescales. The results of this study highlight the areas to target for further development of DGVMs and can be used to improve the models’ capability in simulating the drought-vegetation relationship.

Keywords: Drought intensity; Drought indices; Standardized precipitation evapotranspiration index; DGVMs; southern Africa; Drylands
1 Introduction

Drought can be described as a natural occurrence whereby the natural accessibility of water for a region is beneath the normal state over a long period of time (Xu et al., 2015). Globally, it is considered one of the world’s most important climate risks, with significant environmental, social and ecological impacts on different sectors (e.g. agriculture, forestry, hydrology) and human lives (Naumann et al., 2018). Increasing trends in the occurrence and severity of drought in West Africa and Mediterranean have huge impacts on water resources and agriculture (Sultan and Gaetani, 2016). In southern Africa, a region regarded as a climate hotspot because of the projected impacts of climate change on its numerous endemic vegetation, an understanding of these impacts is important for mitigation options in managing future drought events. Therefore, it is important to examine drought impacts on vegetation and evaluate how this is simulated in models.

Drought is a frequent occurrence in southern Africa, and has enormous impacts on vegetation in the region. For instance, drought has resulted in a significant loss of biomes and death of plants (Masih et al., 2014, Hoffman et al., 2009). It is reported that there has been a significant loss of vegetation cover over the region over the last 30 years (Driver et al., 2012; DEA, 2015). Drought has also impacted speciation of vegetation thereby causing significant changes to the region’s rich biomes, through the lack of formation of new species or even growth of species with underdeveloped morphological and physiological characteristics (Hoffman et al., 2009). Drought-induced vegetation loss has both ecological and socio-economic consequences on human lives. For instance, studies have shown that food security in the region is threatened due to the continual mortality of vegetation (FAO, 2000b; Müller et al., 2011). Other studies (e.g. Wang, 2010; Khosravi, 2017) have also reported that southern Africa could lose more than $200 billion of its GDP from the effects of drought on vegetation. The enormous impacts on vegetation have thus made it imperative to investigate how vegetation might respond to different drought intensities at varying timescales.

In order to monitor and quantify drought characteristics, drought indices are used (Wilhite and Glantz, 1985). Drought indices including the SPI (i.e. the standardized precipitation index), standardized water-level index and standardized anomaly index are derived from a single hydrological variable, which is rainfall (Kwon et al., 2019). Other indices such as the Palmer drought severity index, multivariate standardized drought index and standardized precipitation evapotranspiration index (SPEI) combine two or more variables related to other atmospheric or soil and environmental conditions that may predispose a plant to water stress (Palmer, 1965; Vicente-Serrano et al., 2010; Hao and AghaKouchak, 2013). Among the drought indices, the SPI is the most widely used because of the adjustable timescale and its relatively simple calculation (McKee et al., 1993). It is also recognized as appropriate for use in southern Africa (Hoffman et al., 2009). However, the SPI has a significant shortcoming, which is that its computation uses only rainfall without considering the effect of other meteorological variables in the development of drought occurrence (Teuling et al., 2013). In order to address this shortcoming, SPEI was developed for drought monitoring and it is regarded as a more suitable drought index in the region to investigate the spatiotemporal scale of drought (Ujeneza and Abiodun, 2014). SPEI is computed from the difference between potential evapotranspiration (PET) and rainfall (Vicente-Serrano et al., 2010). PET can be computed using different methods such as Hargreaves (HG) and Penman-Monteith (PM) method. Although studies (e.g. Vicente-Serrano et al., 2010) have found that PM
method captures drought better than HG, other studies (e.g. Lawal et al., 2019a) showed this difference is negligible over southern Africa.

Many studies have used different indices to quantify observed drought, characterize vegetation, and study drought effects on the productiveness of vegetation across different timescales. Several studies (e.g. Vicente-Serrano et al., 2015; Zhang et al., 2012; Lawal et al., 2019a,b) have shown that the satellite-derived normalized difference vegetation index (NDVI) is one the most important indicators of vegetation health and greenness. These studies applied NDVI in examining drought impacts on global vegetation biomes. However, other studies (Gitelson, 2004; Santin-Janin et al., 2009) have argued that while the NDVI is a true proxy for vegetation trends, its potential saturation makes it difficult to fully estimate biomass. In addition, because the NDVI parameters are not well calibrated and often missing in models, simulated NDVI can be biased. Due to its high correlation with NDVI, the leaf area index (LAI) is instead used to characterize vegetation conditions (Fan et al., 2008; Zhao et al., 2013). Although the LAI is an important vegetation proxy, it is rarely considered in the estimation of drought impacts on vegetation. Thus, quantifying the response of the LAI to drought over southern Africa is important for understanding the processes that modulate ecosystem services produced by vegetation which are crucial for human survival. (Melillo 2015).

Previous studies have also evaluated the performance of coupled climate models in simulating the response of vegetation to drought. For instance, Lawal et al. (2019a) reported that an ensemble of the Community Earth System Model (CESM) showed biases in response simulation of vegetation to drought. This was attributed to the parameterizations of the land component (i.e. Community Land Model, CLM) which poorly simulated observed NDVI. Given the poor replication of vegetation response to drought by a coupled climate model, there is a need to examine land-only models and whether they might better capture drought-vegetation relationship when the atmospheric forcings are derived from observations. The present study used Dynamic Global Vegetation models (DGVMs) to study vegetation response to drought, as little is known on how the LAI response to drought is simulated by DGVMs. The choice of DGVMs is because of their capability in simulating mostly accurate carbon exchange between the atmosphere and vegetation ecosystems (Lu et al., 2011).

The aim of this study is to investigate the response of LAI to droughts in southern African vegetation using observations. We also examined how well the responses are represented in model simulations. We used satellite-derived and simulated LAI to quantify vegetation responses to drought. We characterized the spatio-temporal extent of drought and its severity using the SPEI and then assessed the influence of drought using the LAI from satellite data and model simulations.

### 2 Data and Methodology

#### 2.1 Data

In this study, we used satellite-calculated (hereafter, observed LAI/observation-based LAI) and simulated LAI, and satellite-derived NDVI; gridded observation and reanalysis climate datasets. The gridded observation climate datasets include precipitation, maximum, mean and minimum temperature. These data were gotten from CRU (i.e. the Climate Research Unit; Mitchell & Jones, 2005; Harris et al., 2014). These are global monthly data which have 0.5° x 0.5° as spatial
resolution and spans 1901–2019 period. Here, we used the CRU data for the period 1982–2011 to compute observed drought indices (i.e. SPEI) to characterize the spatiotemporal severity of drought. CRU is a gridded observed dataset, which was used because of its suitable spatial and temporal resolutions. Previous studies (e.g. New 1999; New 2000; Wolski et al., 2018; Otto et al., 2018; Harris et al., 2020) have shown that there is a good and robust agreement between observation network and CRU over most parts of southern Africa. We should note that sparseness and missing data generally affect the correlation between CRU and station data in the region. Furthermore, with respect to inter-annual variability, CRU robustly captures the climate factors in southern Africa. The major exception is with the long-term trend of precipitation particularly over Western Cape province of South Africa as well as wetter than normal condition over the same province. These limitations do not affect the validity of our results because we are looking at below-normal precipitation, and temperature.

The reanalysis climate data we used are the CRUJRA, which is a combination of CRU and the Japanese Reanalysis data (JRA) (University of East Anglia Climatic Research Unit; Harris, I.C., 2019). It is a 6-hourly, land surface, gridded data with a spatial resolution 0.5° x 0.5°. CRUJRA was used to compute reanalysis drought indices and used for model simulations. Here we aggregated CRUJRA to monthly samples and used the data at the same spatial and temporal resolution as CRU.

For the satellite vegetation indices, first, we used the third generation of NDVI (hereafter, NDVI3g) from the Global Inventory Modelling and Mapping Studies (GIMMS), and spans period from 1981–2015, with a temporal resolution of biweekly and a spatial resolution of about 8km (Pinzon and Tucker, 2014; National Center for Atmospheric Research Climate Data Guide, accessed 2019). Here, we used the data for the period 1982–2011. Furthermore, we used the third generation of the GIMMS LAI (LAI3g) which also spans the period 1981–2015 and has a temporal resolution of biweekly and spatial resolution as GIMMS3g. The LAI data had been processed (at source) using a set of neural networks which were first trained on highest-quality and post-processed MODIS LAI and FPAR products and AVHRR GIMMS NDVI3g data for the overlapping period (2000 to 2009). The trained neural networks were then used to produce the LAI3g and FPAR3g data sets (Mao and Yan, 2019). For the study, LAI3g was also used for the period 1982–2011. We note that GIMMS LAI and the NDVI, used in this study, are two different indices. The LAI was post-processed using different data (MODIS LAI, FPAR, AVHRR NDVI) for the period of 2000-2009. GIMMS LAI product is superior here over the GIMMS NDVI, which is due to the information derived from the MODIS LAI. The additional properties on GIMMS LAI by MODIS differentiate the index from the NDVI. Thus, it was necessary to investigate how the two indices differ. In addition, other studies (Forkel et al., 2013, Schaefer et al., 2012, Rezaei et al., 2016, Lawal et al., 2019a) have investigated how well satellite derived LAI estimate actual and ground-measured LAI.

The simulated monthly LAI data were obtained from eleven Dynamic Global Vegetation Models (DGVMs) which are part of the Trendy-version 7 (Sitch et al., 2008; Le Quéré et al., 2014). These DGVMs are CABLE-POP (Haverd et al., 2018), CLM (Oleson et al., 2013), CLASS-CTEM (Melton and Arora, 2016), DLEM (Tian et al; 2015), JSBACH (Mauritsen et al., 2018), LPX (Lienert and Joos; 2018), OCN (Zaehe et al; 2011), ORCHIDEE (Goll et al., 2017), SURFEX (Joetzjer et al., 2015), JULES (Clark et al; 2011) and VISIT (Kato et al., 2013). LAI from the
models have a monthly temporal resolution spanning period from 1901 – 2017. We selected these DGVMs because they have been run with similar protocol (S3 simulations) and forcing datasets (i.e. CRUJRA).

2.2 Methods overview

2.2.1 Evaluation of DGVMs and the relationship between NDVI and LAI

The relationship between the NDVI and LAI was evaluated by computing the grid cell spatiotemporal correlation between GIMMS NDVI and GIMMS LAI. The spatiotemporal correlation between GIMMS LAI and simulated LAI from individual DGVMs was also calculated. This was necessary to show whether LAI is an appropriate estimator of NDVI, and how well the models simulate the LAI in the region. Furthermore, we made comparisons of seasonal of observed and modelled LAI. We note that the lack of available of data makes it difficult to compare GIMMS LAI and actual LAI. Nevertheless, the GIMMS LAI has been evaluated and agrees well with observations in other regions (Fan et al., 2019).

The climatology of observed and simulated climatic variables as well as LAI over six major biomes in southern Africa for the period 1982- 2011 were computed. These biomes are semi desert, Mediterranean, dry savanna, moist savanna, temperate grassland and tropical forest (Fig. 1; Sinclair & Beyers, 2015; Lawal et al., 2019a, b).

2.2.2 Description of Drought

For the present study, we adopted the definition of meteorological drought, “which is described as a period (e.g. a season) during which there is a deficit in the magnitude of precipitation in a particular area compared to the long-term normal (Palmer, 1965; Wilhite & Glantz, 1985)”. The deficit in magnitude of precipitation compared to long-term normal is mostly accounted for by temperature, and less by humidity, wind or other variables. Here, we used meteorological drought because it does not make any presumptions about soil characteristics or run-off. In addition, it is acknowledged to be a primary component in the depletion of vegetation productiveness and reduction of biomass (Vicente-Serrano, et al., 2010). Previous studies (Vicente-Serrano et al., 2006; Vicente-Serrano et al., 2013) have also used meteorological drought in the investigation of drought impacts on biomass and vegetation productiveness.

2.2.3 Drought computation and correlation with LAI

The analyses include calculating drought (i.e. SPEI) using CRU data over a 30-year (1982 – 2011) period for different drought timescales. The drought time-scale can be described as the aggregation of temporal duration (Vicente-Serrano et al., 2010). SPEI is an index that is used to quantify drought (see Table 1). Therefore, the quantified values of the index give the state of drought in a space. Our definition and approaches follow numerous previous studies (Vicente-Serrano et al., 2013; Khosravi et al., 2017; Zhao et al., 2013; Hao et al., 2013).
A time series of the evolution of drought for the 30-year period was plotted. The present study extends the timeframe for understanding drought impacts from 1982 to 2011 mainly because there were frequent droughts in the 2005 – 2011 window (Masih et al., 2014). The timeframe was then extended back to cover a 30-year period to be long enough to cover impacts of climate change, which is particularly important considering that southern Africa experiences more frequent droughts with impacts exacerbated by climate change. This information is important for considering adaptation measures and understanding the role of climate change.

The drought index, SPEI, is calculated from the deduction between precipitation (P) and potential evapotranspiration (PET) as shown below:

\[ D = P - PET \]  

(1)

where D-values represent a measurement of water deficit or surplus aggregated at different time scales. D values are obtained through aggregation over individual time-scales which span 1- to 24-months (i.e. 1-, 3-, 6-, 9-, 12-, 15-, 18-, 21- and 24-month). The time-scales were calculated by including the past values of the variable. For example, a time-scale of 15-month suggests that input from the preceding 15 months, which includes the present month, was used for calculating SPEI (Begueria et al., 2014). “For the 1-month timescale, only the current month data is used for the calculation. The D values were standardized by assuming a suitable statistical distribution (e.g. gamma, log-logistic). The log-logistic distribution was used to standardize the D values in this study” (see also Lawal et al., 2019a). For more details on the timescale computation, please see Vicente-Serrano et al. (2010); https://rdrr.io/cran/SPEI/man/spei.html. PET is computed from maximum temperature, minimum temperature and mean temperature, using Hargreaves technique (e.g. Vicente-Serrano et al., 2012; Begueria et al., 2014; Stagge et al., 2014).

We note that the PET in the SPEI was computed using Hargreaves (HG) method rather than Penman-Monteith (PM) because the data (e.g. vapour pressure, maximum and minimum humidity) required for computing PM over southern Africa are sometimes missing or not available at the needed gridded spatial resolutions and timespan. Although PM is considered better in most regions, Lawal et al. (2019a) showed that the variation between the PM and HG is negligible for southern African region. The study only considered observed SPEI_PM which was obtained from https://spei.csic.es/database.html and not modelled SPEI_PM due to the unavailability of simulated data required for its computation. Other studies (e.g. Begueria et al., 2014) have also found that there is an insignificant contrast in the strength of PM and HG for reproducing their divergence on measured variables such as vegetation indices. We should also note that the SPEI (unlike SPI or PDSI) is the most appropriate index for measuring drought in southern Africa, as it accounts for the effect of evaporative demand from the atmosphere in drought monitoring (Vicente-Serrano et al., 2010; Ujeneza et al., 2014). In addition, the SPEI is reported to be able to identify the geographical and temporal coverage of droughts (Vicente-Serrano et al., 2010; Ujeneza et al., 2014).
Table 1. Definition of drought thresholds based on the SPEI scale

<table>
<thead>
<tr>
<th>SPEI</th>
<th>Drought thresholds</th>
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<tbody>
<tr>
<td>2 or more</td>
<td>Extreme wet</td>
</tr>
<tr>
<td>1.5 to 1.99</td>
<td>Severe wet</td>
</tr>
<tr>
<td>1 to 1.49</td>
<td>Moderate wet</td>
</tr>
<tr>
<td>0 to 0.99</td>
<td>Mild wet</td>
</tr>
<tr>
<td>0 to -0.99</td>
<td>Mild drought</td>
</tr>
<tr>
<td>-1 to -1.49</td>
<td>Moderate drought</td>
</tr>
<tr>
<td>-1.5 to -1.99</td>
<td>Severe drought</td>
</tr>
<tr>
<td>-2 or less</td>
<td>Extreme drought</td>
</tr>
</tbody>
</table>


We deseasonalized GIMMS LAI by transforming monthly LAI series per pixel to symbolize the standardized deviations from extended mean. This was to make the sequence of LAI commensurate to SPEI (Vicente-Serrano et al., 2013) and eliminate the impact of periodicity on vegetation response. We note that the SPEI is intrinsically deseasonalized.

In order to reconcile the difference in spatial resolutions of CRU and GIMMS, we regirkd the data to the same spatial resolution using the bilinear interpolation method. We then computed the correlation per grid-cell between SPEI (based on CRU) and deseasonalized LAI over the 30-year period at the different drought timescales using Pearson correlations. We then compared the spatial distribution of maximum (peak) correlation and the comparable time-scales of drought for observed SPEI. Our analyses take into account the periods which LAI responds to the presence/absence as well as the severity of drought. This is referred to as “drought time scale” in the study.

Next, we investigated correlations at each grid cell between the drought (from reanalysis - CRUJRA) and an ensemble median of deseasonalized modelled LAI from individual DGVMs. The peak (maximum) correlations and equivalent time-scales from the complete 1- to 24-month time-scales were mapped for the ensemble median over the 30-year period. We used ensemble median because of its less sensitivity to independent outliers (Reuter et al., 2013). In summary, we calculated model ensemble drought from the median from individual members’ drought indices. The inter-annual variation of drought impacts on LAI by individual DGVM was also calculated for different timescales. We also examined the observed and an ensemble mean of simulated LAI response to drought across latitudes.

In summary, CRU was used to compute observed drought while CRUJRA was used to calculate modeled drought because it is what was used to force the models. This will allow for easy observation and model comparisons.

Similar to Lawal et al., 2019a, we calculated the seasonal mean for four seasons i.e. a) December-January-February (DJF); b) March-April-May (MAM); c) June-July-August (JJA); and d) September-October-November (SON) from the correlations of monthly series of drought and LAI.
These were computed from correlating monthly series (twelve series per year) per pixel of GIMMS-LAI and each monthly series of 1- to 24-months drought (SPEI) series over the 30-year period with Pearson correlation. The same technique was used for the model ensemble. In simpler terms, we calculated the correlations (twelve sequences in a year) of monthly LAI to monthly sequences of 1- to 24-months SPEI using 30-years of data. Subsequently, the seasonal mean of these correlations was calculated. The peak correlations and drought timescales of the models were calculated over six major biomes in southern Africa, namely (Fig. 1) – Temperate grassland, Tropical forest, Moist Savanna, Dry Savanna, Semi-desert and Mediterranean vegetation. These regions were selected because of their relative importance and are most affected by drought.

Finally, the impacts of extreme events (wet and dry years) at different time periods were compared; and the comparison of global and regional responses to drought across biomes for the period 1982 – 2011 was investigated.

3 Results

3.1 Grid cell correlations between NDVI and LAI

Figure 2 illustrates the relationships between the NDVI and LAI for observations, as well as the comparison between observed and model LAI. There is a strong linear relationship between
observed NDVI and LAI (Fig. 2a). The correlation (0.94) is high between both variables and the standard deviation is low (0.005). The standard deviation being referred to is for GIMMS LAI and individual DGVMs, as well as GIMMS LAI and GIMMS NDVI. From the figure, there is a log-like shape, where NDVI grows faster when LAI is low (<0.2) and becomes saturated when LAI goes higher (>0.3). A linear regression of the data shows a slope of 2.15. The low standard deviation indicates that the values from the two indices are close. Although there is a good agreement between observed NDVI and LAI, the 1:1 line shows that the datasets are not exactly equal.

Furthermore, there is good agreement between observed and the simulated LAI (Fig. 2). JULES has the highest correlation (0.97) with observation (Fig. 2i). CLM has the weakest (0.73) correlation with the observations (Fig. 2c). DLEM and LPX have the same correlation coefficient value of 0.87 with observation (Fig. 2e, 2g). The positive relationships between simulated and observed LAI indicate a general applicability in investigating the model’s performance of vegetation response to drought. It also shows that the correlation is strong enough to compare how the LAI reacts to drought in the ensemble. An aggregation of observation along the gradient of simulated LAI shows that most of the models have similar slopes with observation.
Figure 2. Scatterplots of correlations between vegetation indices (observation and model) for the period 1982 – 2011 over southern Africa. Inset values indicate the correlation coefficient (r) and standard deviation (Stdev) between GIMMS LAI and GIMMS NDVI, as well as GIMMS LAI and modelled LAI. The colour represents each grid cell. The pink solid line is the linear regression, while the dashed black line shows 1:1 line. The unequal x-axes are to visualize the detailed data for the models.
3.2 Seasonal and interannual variations of observed and modelled LAI

The comparison of seasonal and interannual variation of observed and modelled LAI is given in Fig. 3. The model shows a stronger positive bias in JJA and SON in comparison to summer and winter months; and a negative bias over the tropical forest region of Madagascar (Figs. 3A – 3D). In addition, model mostly overestimate the seasonal patterns of LAI in some regions during DJF and JJA, and underestimate LAI in MAM and SON (Figs. 3E – 3L). Over most parts of the region, there is a strong correlation and good agreement between observed and modelled LAI in DJF, MAM and JJA although it is weaker in SON (Figs. 3M – 3T). The strong correlation is more prevalent in JJA than other seasons (Fig. 3S); while it is weakest in southern parts of the region. However, the correlation is largely negative over Angola in DJF and SON (Figs. 3Q, 3T). Furthermore, the correlations between model and observed LAI is weaker in deseasonalized data (hereafter, Deseas. Correlation; Figs. 3M– 3P) than in original data (hereafter, ORIG. Correlation; Figs. 3Q– 3T), thereby showing the effects of seasonal patterns on time-series data. With respect to the period, 1982 – 2011 (hereafter, ANNUAL), the correlation between modelled and observed LAI are different for deseasonalized and original data (Figs. 3U – 3V). For the former (Fig. 3U), there is gradient in the correlation across the region, with higher values in central and southern parts than in Angola and Madagascar. However, with the original LAI data (Fig. 3V), the correlation is very high (about 0.85) and more prevalent, except in eastern Madagascar and Western Cape Province of South Africa.
Figure 3. Spatial seasonal distribution and inter-annual variability (IAV) of satellite-calculated and modelled LAI (multi-model mean) over southern Africa. (A) – (D) show the difference (bias); (E) – (H) and (I) – (L) show their standard deviation (Stdev); (M) – (P) show the correlations between deseasonalized GIMMS LAI and modelled LAI; (Q) – (T) show their correlations for original GIMMS LAI and modelled LAI and (U) – (V) show correlations...
between GIMMS LAI and modelled LAI but for the period 1982 – 2011. The inter-annual variability for observed and modelled LAI for the period 1982 – 2011 is shown in Fig. S8.

3.3 Climatology of observed and, simulated climate variables and LAI

This section compares the seasonal cycle of observational (CRU and CRUJRA) climate variables; as well as observed and simulated LAI from GIMMS LAI and TRENDY models, respectively. Precipitation and temperature are seasonally variable and their climatologies are mostly similar. For example, precipitation is higher in MAM and DJF over many of the biomes except in Mediterranean vegetation where precipitation is higher in JJA (Figs. 4a, 4d, 4g, 4m, 4p). The wettest month occurs over TF (i.e. tropical forest biome) where the precipitation is about 350mm. Conversely, during the dry season (JJA), there is little rainfall in the biome, although, it experiences some precipitation June and July. Over the Mediterranean vegetation (Fig. 4j), a winter (JJA) rainfall region, rainfall variability is lower and is mostly dry in the DJF and SON. Similarly, the highest minimum and maximum temperature in the region is observed in the DJF season, where the highest temperature value exceeds 30°C. Over the tropical forest biome, although the distribution pattern of precipitation and temperature are similar for most months, they differ during June and July months. The pattern of precipitation and temperature distribution generally differ over the Mediterranean vegetation. The pattern of temperature and precipitation from CRUJRA follows CRU, although the ensemble spread is much narrower (Fig. 4c, 4f, 4i, 4l, 4o, 4r). The spatial patterns of the climate variables from CRU and CRUJRA are shown in Fig. 5. CRUJRA well simulates the pattern of precipitation over southern Africa (Figs. 5a and 5b) as CRU although it shows some biases in magnitudes, as well as for minimum temperature (Figs. 5g and 5h).

There is not strong seasonality for LAI, with maximum observed LAI values less than 4 in all biomes. The models well reproduce the climatology of LAI over the southern African biomes with a few exceptions (Figs. 4b, 4e, 4h, 4k, 4n, 4q). For instance, the models simulate the drop in LAI over the semi desert, temperate grassland, tropical forest, dry savanna and moist savanna biomes in JJA. The highest increase in observed LAI occurs over the tropical forest in April, although the models simulate a decrease in LAI over tropical forest during this time. On the other hand, lowest amount (less than 0.1) of LAI is observed in September and this occurs over the semi-desert biome. Observations typically fall within the range of the model ensemble. In addition, the distribution pattern of simulated LAI is similar to observation in most biomes except in the Mediterranean and tropical forest biomes. The LAI pattern also follows that of the climatic variables although the former lag. The lag effect is accounted for in this study, and is known as drought time scale.
Figure 4. Annual cycle of observed climate variables (precipitation, mm/month; maximum, minimum and mean temperature, °C) and LAI for observation and multi-model mean (TRENDY) across six southern African biomes over the period 1982 – 2011. The annual cycle of the LAI for individual models are shown in Figure S5.
Figure 5. Spatial distribution of precipitation, mean temperature, maximum temperature and minimum temperature over southern Africa in CRU and CRUJR; for the periods 1982 – 2011.
### 3.4 The evolution of drought in southern Africa

Figure 6 shows the evolution of observed SPEI in southern Africa between 1982 and 2011. Here, drought indices from CRUJRA are not included because preliminary investigation showed close magnitudes for drought indices computed from CRU and CRUJRA. We note that CRU was used to calculate SPEI for observation while simulated SPEI was computed with CRUJRA.

There is inter-annual, seasonal and decadal variability of the drought indices during dry and wet conditions over southern Africa. Although 1-, 3- and 6-month SPEI indicate no trend in wet or dry spells, they show the intensity of drought event for the 30-year period (Figs. 6a, 6b, 6c). The highest magnitude of drought is captured by 1-month SPEI while the lowest is shown in 21-month SPEI. The severity of drought intensity is similar for all SPEI (i.e. 1-month to 24-month SPEI). The magnitude of the severity is on the y-axis of 1-month to 24-month SPEI.

We note an increasing trend in SPEI (9- to 24-month). Tables 2 and 3 illustrate when droughts occurred, as well as the severity of drought within the 30-year period. They also show, however, that droughts were most frequent and intense in the second decade, thus, indicating how climate change is expected to increase the frequency and severity of droughts.
Figure 6. Evolution of SPEI in southern Africa for the period 1982 – 2011. The trend is significant at 90% confidence interval for all timescales.


<table>
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<tr>
<th>Drought Timescale</th>
<th>Number of drought events</th>
<th>Year of moderate drought events</th>
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3.45 Spatial distribution of LAI response to drought and the timescales

Figure 7 presents the spatial distribution of the peak correlation between the SPEI and the LAI, and the timescales at which the correlation occurs. This is to show the magnitude of response of LAI to drought in southern Africa, and the length of the period for the response.

Observations show that southern African LAI can respond fairly strongly to droughts (peak correlation magnitudes of between 0.4 and 0.6), though the response is much weaker ($r < 0.4$) in eastern Madagascar, Angola and parts of South Africa (Fig. 7a). The TRENDY multi model median generally overestimate the observed magnitude of the LAI to drought response (Fig. 7c). Peak correlations for the models seem to be much stronger – in the 0.6-0.8 range for most of the regions. In addition, over the arid areas of Namibia, models simulate a LAI while observations depict no measurable LAI, indicating that models simulate the LAI in areas where observations show no measurable LAI.

The multi model median have a drought timescale that is mostly longer than the observations (Fig. 7b, d). For instance, drought response of simulated LAI occurs mostly over a longer time period (6-, 9-month timescale) than in the observation over eastern Madagascar. Over southern areas of Madagascar and central Zambia, the multi model median overestimates the drought timescale. Over central areas of South Africa and Mozambique, simulated LAI responds at intermediate (9-month) timescales. In comparison, similar drought timescales for the observation and the model ensemble median are shown in parts of Angola.
Figure 7. Spatial distribution of peak correlation between drought (SPEI) and LAI over the region of southern Africa in the observation and in the model ensemble median; for the period 1982–2011. Panels (a) and (c) show the peak correlation per pixel, which is independent of the timescale and the month of the year. Panels (b) and (d) indicate the timescales at which the peak correlation between SPEI and LAI is found. Areas with no significant correlation are white.

3.6 Latitudinal distributions of LAI response to drought and the timescales

The present study investigated and discussed the implications of drought on different vegetation/biome types across latitudes in the region. Here, we stratified the LAI response to drought based on latitude, because we intend to investigate and identify the shift in response based on the vegetation types across the latitudinal belt.

While the pattern of the latitudinal distribution of the LAI response to drought is identical across different timescales, the magnitudes generally differ (Fig. 8). The response is much weaker (less than 0.15) for the 1-month timescale than for other timescales. The strongest response is observed at 6-month timescale between latitudes 25 and 30°. The model ensemble mean generally agrees with the pattern of the observed LAI-SPEI correlations across all the timescales. However, the
magnitudes differ from the observation. Modeled correlation is stronger than observations for the longer (6-month or more) timescales. This means that the models are oversimplifying how LAI responds to drought by neglecting other climate factors, such that in models, LAI only correlates to water deficit (SPEI). Furthermore, there is an offset between observation and model mean which is consistent across most of the timescales, perhaps due to strong memory of the some of the models.

Figure 8. Mean correlation (observed and multi-model ensemble) of annual LAI and SPEI for 1982 – 2011 across latitudes over southern Africa for 1- to 24-month timescales.

3.7 Response of LAI to droughts across seasons

Observations show similar correlations between LAI and drought across all seasons in the biomes (Fig. 9). For the dry savanna, which is one of the most climate-impacted biomes in the region, LAI
response to drought is strong and the correlation is as high as 0.8 in MAM season and it occurs at 12-month timescale. The correlations between drought and LAI are also very strong in other seasons over the same biome and occur at 6- and 12-month drought timescale, except over the Mediterranean vegetation where the response occurs at 18-month in DJF season. Similarly, the peak correlations between drought and LAI are strong across the other biomes. With an exception of the tropical forest biome, the drought timescale is at longer time periods (> 6-months).

The model ensemble generally overestimates correlations across the biomes in different seasons. Whilst the correlation magnitude remains mostly larger than observation, nonetheless, models simulate closer correlation with observation in some biomes and seasons. For instance, over Mediterranean vegetation, models simulate fairly good response of LAI to drought in all seasons. Furthermore, in nearly all other biomes, the ensemble spread overlaps with the observations. In addition, simulations mostly overestimate drought timescale, except over dry savanna. A possible reason for the difference why the time scale for Dry savanna was underestimated may be because phenological triggers for dry savannah vegetation types respond differently to environmental variables, which the models do not capture. The African Dry savanna region is characterized by rapid vegetation changes due to fire, land-use among others, as well as senescence for prolonged dry periods (Rahimzadeh-Bajgiran et al., 2012; Zhu and Liu, 2015), which may have contributed to the underestimation in the response of the models. Similarly, models have different representations of fire, which could also indirectly contribute to the underestimated model responses to drought.
Figure 9. Seasonal correlations of drought (SPEI) and LAI across six southern African biomes. The values on the left axis show the peak correlation in observation and TRENDY models. The values on the right axis indicate the corresponding drought timescale.
Inter-annual variation of model simulation of drought impacts on LAI

Table 4 shows the correlations between observed mean SPEI and LAI for the period 1982 and 2011 as well simulations by individual DGVM across different timescales. Unlike Fig. 7 which shows the peak correlations, the table shows the mean correlation for the 30-year period.

There is variation in the inter-annual simulation of LAI response to drought across different timescales by individual models. For instance, on the 1-month timescale, JULES simulates the least correlation value while JSBACH shows the highest correlation value. Furthermore, JSBACH simulates the highest correlation value for most of the timescales while CLM simulates the least correlation values for most (3-, 9-, 12-, 15- and 18-month) of the timescales. A possible reason for the weak performance of CLM may be its representation of the canopy construction of the PFTs and of its foliage clumping representation. In addition, CLM is limited in its simulations of vegetation with regards to transpiration, due to rooting depth among others (Dahlin et al., 2020). Furthermore, CLM does not well simulate savanna ecosystems, but instead uses a combination of grasses, shrubs, and trees. There are also some problems (such as an unusual green-up in dry season) identified with stress deciduous responses (Dahlin et al., 2015).

Table 4. Model simulation of mean SPEI and LAI correlations between 1982 and 2011. * indicates the model with the lowest mean correlation.

<table>
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<tr>
<th>Correl.</th>
<th>GIMMS LAI</th>
<th>CABLE-Pop</th>
<th>CLM</th>
<th>CLASS-CTEM</th>
<th>DLEM</th>
<th>JSBACH</th>
<th>LPX</th>
<th>OCN</th>
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3.89 Impacts of extreme events on LAI

The impact of extreme events on LAI is shown in Fig. 10. The objective was to discuss the impacts of extreme drought events, and compound influences of drought on LAI during extreme hot and dry years. Here, extreme events are the wet years - i.e. the periods with precipitation higher than normal; and the dry years which include the periods of very high dry spells. To achieve this, we computed the correlation between LAI and corresponding dry/wet years at the different time periods. The maps of this correlation are then plotted.

The magnitudes of response of LAI are generally stronger during the dry years than in wet years (Fig. 10). The distribution pattern of the magnitude of response however, varies across the region. With the exception of the year 2010, the strongest drought response is observed in parts of Namibia, Botswana, Tanzania and South Africa. During the 2010 wet year period, the response is weaker in most parts except for the northern part of Madagascar and southwestern part of South Africa (Figure 10F).

Figure 10. Spatial pattern changes of observed LAI response during extreme wet and dry years. For (A) – (C) changes were calculated as a difference between the wet year and the 30-year mean and for (D) – (F) between the dry year and the 30-year mean. White areas indicate no correlation.

3.9.10 Comparison of global and regional distribution of LAI response to droughts (1982 – 2011)

There is variability in the global and regional temporal distribution of LAI response to drought (at 12-month timescale) when global vegetation biomes are split into regional biomes (Fig. 11). The map of the global biomes is shown in Fig. S1 in the supplementary material. The observed global response, indicates a decreasing trend of LAI response to drought while the model mean shows an increasing estimate.
The semi-desert biome dominates the LAI response as higher drought-vegetation correlations are observed (Figures 11G – I). Over the biome, there is more marked interannual variability which makes the biome an important player in global carbon cycling (Poulter et al., 2014). The response over the semi-desert in southern Africa is however weaker in comparison to the other semi-desert biomes.

The response over the Mediterranean vegetation in Australia is stronger than the Mediterranean vegetation over the rest of inland southern Africa. Over the biome, model simulates closer magnitudes in the latter than over Australia (Figures 11B and 9C).

Over the tropical forest biomes, there is a weaker response in Central Africa compared to Southern Africa and South America; the model simulates a closest response magnitude in South America (Figures 11D – F).

Figure 11. Correlations between SPEI (12-month timescale) and ensemble mean of LAI from TRENDY (blue line), GIMMS LAI (black line) for (A) Global (B) Mediterranean vegetation over Australia (C) Mediterranean vegetation over southern Africa (D) Tropical Forest over southern Africa (E) Tropical Forest over Central Africa (F) Tropical Forest over South America (G) Semi-desert biome over southern Africa (H) Semi-desert over Asia and (I) Semi-desert over South America.
4 Discussion

4.1 Relationship of LAI to phenological changes

LAI is a variable that is needed for global modelling of biogeochemistry, climate, ecology and hydrology and different primary production models (e.g., Running & Coughlan, 1988; Sellers et al., 1996; Bonan, 2002). In view of the need to run biogeochemical models at regional and global scales, accurate LAI data at moderate – high resolutions are crucial (Wang et al., 2004). The relationship between NDVI and LAI is applied as a support algorithm in MODIS LAI. Thus, from the viewpoint of availability of data, retrieving LAI from analyzing NDVI-LAI relationship remain the main perspective for high temporal resolution in regional and global-wide studies (Wang et al., 2004).

LAI showed a linear relationship with NDVI. This suggests that the NDVI is associated with the phenological changes of plants, the parts of surface cover class which contribute to the general reflectance as well as the variations in the angle of solar zenith (Wang et al., 2004). Studies (e.g. Myoung et al., 2013) have however, found that the relationship between NDVI and LAI varies intra- and inter-annually; and both vegetation indices differ temporally and seasonally over deciduous forests, which are sometimes not accounted for in models that test their relationship (Wang et al., 2004). For instance, while the relationship is strong during periods of leaf production and senescence, no relationship is observed during the period of leaf constant due to NDVI saturation above certain LAI values (Xue and Su, 2017).

Although the present study found a strong linear relationship between the NDVI and LAI in southern Africa, other studies (Potithep et al., 2010; Towers et al., 2019) have shown the two indices are not always directly proportional. For example, both indices do not exhibit the same relationships over different eco-regions such as the Evergreen Broadleaf Forest, Deciduous Needleleaf Forest. Furthermore, other studies (Fan et al., 2008; Tian et al., 2017) found that the LAI may be better indicator of plant biomass and health because of the saturation associated with the NDVI, particularly in drylands. This makes the LAI more applicable in monitoring vegetation response to drought. Evaluating how the LAI differs from the NDVI over different biomes (such as dry savanna, tropical forest, etc), with regards to temporal difference is shown in Fig. S7. Both the LAI and NDVI show similar annual cycles over southern Africa, except for the Tropical forest and Mediterranean vegetation.

4.2 The importance of sub-monthly data in drought computation and monitoring

The data used to evaluate drought indices is CRUJRA. JRA is a reanalysis and has 6-hourly temporal resolution. Additionally, CRUJRA is the data used to force the DGVMs, so the drought indices are being calculated based on the same data the models use for their simulations. JRA is a reanalysis but the combined CRUJRA product uses the sub-monthly information from JRA, and is constrained to the monthly CRU observations. The comparisons of the data are shown in Figs. 5 and S6. It is useful to use data with shorter times because the study focuses on an evaluation of drought impact, which is sensitive to timescale. In drylands, for instance, the uncertainties associated with monthly data in drought monitoring are reduced when sub-monthly data are used.
We note that, over different parts of the world, CRU has been widely validated against station data (Harris et al., 2020) and there is a high accuracy of the validation. Therefore, observed SPEI gives a high accuracy of measured drought. Another major advantage of using the observational-based CRU data is its spatial and temporal coverage. Station data are available for very few points and for limited times in the region of interest. The few data that are available are fraught with missing data, rendering them an unreliable data source (Harris et al., 2020).

4.3 Annual cycle of climate and vegetation in southern Africa

Climatologies of meteorological variables show that precipitation drops in JJA and SON seasons over the biomes except over Mediterranean vegetation. The dry condition that is experienced during these seasons could be attributed to the subtropical high pressure system which suppresses rainfall by shifting the ITCZ (Inter-Tropical Convergence Zone) away from these regions (Naik and Abiodun, 2016).

Observations also show low LAI over some parts of southern Africa (please see Figure S2). The weak gradient in some parts of the region may be due to low winter rains produced by the frontal system which is not sufficient for growth of expanse vegetation (Lange et al., 1999). The aridification of the western part of southern Africa may be attributed to the influence of cold sea surface temperature (SSTs) of the Namibian Upwelling system along the Namibian coasts (Ward et al., 1983). The aridification does not only result in cessation of river discharge but also sediments that would have favored the growth of drier vegetation (Dupont, 2006).

For Fig. S5, some of the models do not capture this peak around September over the Mediterranean and Tropical biomes. One possible explanation is that the models do not well reproduce the changes in the biomass and leaf area cover around that period (due to phenological responses to environmental variables). For both biomes, spring rainfall contribute to vegetation growth in the region, which may not be well reproduced by the models. For Fig. 4, the magnitudes (< 0.5) of the observation-based LAI over (southern Africa) semi-desert biome is quite “high” because the region is a pseudo-desert, which experiences very high summer temperatures but does receives some rainfall, and the Okavango river is flowing through it permanently. This region is rich in biodiversity such as Acacia spp (trees) and Aristida and Schmidita spp (savanna) (see WWF 2001; Street and Prinsloo, 2013; Lawal et al., 2018.) Thus, a 0.5 LAI in the biome, which may be higher than other desert biomes, is reasonable in this region.

4.4 LAI response to drought in observation

Drought is becoming frequent and more intense in southern Africa (Masih et al., 2014). The frequent and stronger dry spells is observed in Fig. 6. Climate change is expected to increase the frequency and severity of droughts (Tables 2 and 3). The severity and longer durations of drought have enormous impacts on the already endangered vegetation biomes in the region (Hoffman et al., 2009). The results show that drought impacts on vegetation occur across the different seasons in the region. The seasonal difference in the response of vegetation to drought across biomes is
influenced by numerous factors such as vegetation adaptive capacity and resilience, reproduction process and growth stage among others (Zeppel et al., 2014; Corlett, 2016). For instance, over the tropical forest biome, drought has the least impact on vegetation in the region, which could be because of the deeper rooting system of the vegetation which allows them access to soil at the deeper water table (El-Vilaly et al., 2017). It is reported that major drivers of vegetation resilience and productivity are precipitation and temperature which control the evapo-transpirative rate (Allen et al., 2010). It is worth noting that vegetation in southern Africa will be severely impacted if the trends continue in the same trajectory. For instance, the regions where there is a strong vegetation response to drought are experiencing wood encroachment and thus, will likely worsen based on the current trajectory of drought occurrence.

We note that the performance of drought indices is not only limited by the variables used in their computation but also by biomes and location where they are used (Xu et al., 2015). For example, SPI which is calculated using simple methods and has adaptable timescales performs better than SPEI in arid regions (Begueria et al., 2014). However, SPEI which requires more variables for its computation captures drought better in relatively humid zones (Begueria et al., 2014). SPEI is however limited by the potential evapotranspiration (PET) because of its sensitivity to the variable (Xu et al., 2015).

The vegetation situated at the borders of Botswana-Namibia and Mozambique-Zambia respond to droughts at an intermediate timescale (i.e. 9-month). The types of vegetation inhabiting these regions, which are well adapted to water shortage because of their physiological and morphological characteristics, takes prolonged period to respond to drought and thus, do not easily shows symptoms of water strain (Vicente-Serrano et al., 2013). The activities through which vegetation minimize water loss include reduction in photosynthesis, reduced canopy cover (Schwinning & Sala, 2004). The capacity for vegetation to store water is one adaptation for low water ecosystems, as is reduced daytime stomatal conductance and CAM photosynthesis. The disparity in the timescales spatial distribution in models from observation might be because the parameters are represented and estimated in the models (Murray et al., 2011). In addition, the models are not similar in their drought timescales simulations.

The varying response of the tropical forests in different regions may be because of the interplay of precipitation and temperature at different longitudes. Ahlstrom et al (2015) showed that temperature is particularly a strong factor in the response of this vegetation. Wang et al (2008) also reported that soil moisture variations play a key role in the magnitude of vegetation response. The weak response of LAI to drought over Madagascar and Angola may be attributed to the fact that the vegetation in these regions is able to store water for a long time which it uses during deficit (Chapotin et al., 2006). It may also be because rainfall is not the main regulatory component in the growth of vegetation in these regions (Fuller and Prince, 1996). In regions such as Botswana and Namibia, vegetation is highly dependent on water availability for their ecosystem functions (Anyamba et al., 2003). Please see Fig. S3 in the supplementary material for observed correlations at different timescales.

The seasonal response of LAI to drought varies and this could be attributed to many factors. For instance, the sensitivity of the semi-desert to water shortage makes them show quick response to drought (New, 2015). The response of vegetation to drought is particularly stronger in the MAM
season, because it is during this period that fruit, leaves, and biomass are produced by vegetation (Zeppel et al., 2014). Critical water requirements by vegetation for these developmental activities in the MAM season is the reason for the vegetation respond to droughts at a short timescale (Zeppel et al., 2014). The strongest vegetation response (a correlation of about 0.92) is observed over the semi desert biome (which is in the semi-arid environment) in the JJA season. This is a region with vegetation which heavily depend on water for all their ecosystems functioning without which they would not survive (New, 2015). The drought response of tropical forest is weaker compared to all other biomes. Over the tropical forest biome, the fairly subtle drought response may be because the biome can be tolerant to drought, have stronger robust capacity and is therefore, not extremely impacted by droughts as are the other biomes (Gilgen et al., 2005; Corlett, 2016).

The different response by the global biomes in different geographical locations may be because of climate variations and the sensitivity of LAI to climate variations (Ahlstrom et al., 2015). The dominance of LAI response to drought over the semi-desert biome could be attributed to the global bush encroachment and is in consonance with the increasing greenness (Donohue et al., 2009; Fensholt et al., 2012; Andela et al., 2013). Studies (Cai et al., 2014; Trenberth et al., 2014; Dai et al., 2013; Wang et al., 2014; Ahlstrom et al., 2015) have also found that increased and frequent ENSO events due to climate change have not only led to the expansion of LAI but could increase the water demand by semi-desert vegetation. This comparison is particularly important because until now, there has been little or no study on this.

4.5 How well the seasonal and interannual variations of LAI are captured in DGVM simulations

The models exhibit biases in the simulation of seasonal and interannual variations of LAI over southern Africa. Two major factors may be given for performance of the models. First, the influence of precipitation forcing data largely affect the seasonal cycle of LAI as well as the interannual variability (see Figs. 3 & S8). Although the difference between the forcing data for observed and modelled LAI (i.e. CRU and CRUJRA respectively) is small, it still influences the simulations by the DGVMs. The precipitation uncertainty has larger influence in some regions and biomes than others. For instance, the overestimation of LAI by models in arid biome of Namibia and Mediterranean vegetation of South Africa may be due to the irregularity in precipitation, whereby the precipitation changes are too little to be identified (in CRUJRA), as well as due the large spatial variability (Fekete et al., 2004; Greve et al., 2014). On the other hand, over the savanna biome of Angola, the underestimation is likely because of the sparse distribution of precipitation data, which permeates to CRUJRA (Fekete et al., 2004). Secondly, the differences in observed and simulated LAI may be due to the impacts of land use and land cover change on the latter (LULCC) on simulation, through soil moisture as well as evapotranspiration (Piao et al., 2015). LULCC exerts strong influence on water consumption, nutrient cycling, and root depth (Piao et al., 2007; Mango et al., 2011), and the extent of the influence also varies across biomes. For example, in MAM, over temperate grassland biome of South Africa, where the model underestimates LAI, the bias is likely because the models fail to simulate the frequency of evapotranspiration, which is about five times much less than the forest biome (Yang et al., 2015).
However, over most parts of the region, the correlation between observed and modelled LAI is strong, except in JJA season. This means that the models generally simulate the pattern of LAI distribution in southern Africa. We also note that, the lower correlations between observed and modelled LAI when the data were deseasonalized, is due to the sensitivity of LAI to variability and seasonality. These differences between observed and modelled system have impacts on how vegetation response to drought is captured, which we discuss in Section 4.6.

4.6 How well drought and LAI response is represented in DGVM simulations

The observed LAI is simulated within the models and calculated by GIMMS based on Mao and Yan, 2019. Lu et al., 2011 found that DGVMs perform better against observations than Earth system models (ESM) because they use observational-derived climate and can include more complex representations of vegetation processes. The ESM is a coupled model simulating its own climate, while the individual DGVMS models used in the present study are standalone, i.e. are applied with observational based meteorological forcing, and thus we remove one uncertainty. Since offline studies target the DGVM itself, removing one possible issue (incorrect climate drivers), it became imperative to use DGVM to study drought impacts.

DGVMs simulate the vegetation characteristics and impacts of climate on them. The validation of DGVM simulations of variables such as LAI is quite difficult. This is because of the unavailability of data on large spatiotemporal scale for the different vegetation classes (Potter and Klooster, 1998). Studies (e.g. Potter and Klooster, 1998) have also shown that errors present in the prediction of plant functional types (PFTs) tend to spread to biomass prediction in the model, thus possibly biasing estimates of carbon stored in terrestrial ecosystems. Nevertheless, the DGVMs used in this study simulate the spatial patterns of vegetation distribution though with a magnitude bias as shown in Fig. S2 in the supplementary material.

TRENDY models mostly simulate the temporal patterns of global and regional distributions of LAI response to drought. The biases shown by the models have been attributed to the fact that the models do not factor land use changes (Ahlstrom et al., 2015). This is evident in the simulation of LAI (please see Fig. S2).

The models’ weaker simulations might also be because some of the DGVMs do not well reproduce the LAI magnitude. The negligible difference in the spatial distributions of SPEI of the models could be due to fact that the model PET does not play a strong role in drought occurrence in the southern Africa and that precipitation is the main driver of drought in the region. The variations in the characterization of hydrological processes in the models are also a source of uncertainty because they reinforce the bifurcation in runoff outputs which has cascading effects on biospheric changes and evapotranspiration (Murray et al., 2011; Stewart et al., 2004). Also see Fig. S4 for correlations of the model ensemble median at different timescales. Another reason for the biases in the simulations may be to the design of the DGVM experimental set-up, which include the flux deviation between simulations without and with (Murray et al., 2011).

4.7 Variations in observed and simulated vegetation response to drought, and implications on model development
The biases shown by models could be attributed to the different limitations of individual DGVMs, and addressing these shortcomings would improve models’ performances. For example, the sub-optimal performance of CLM may be partly due to the inability of the model to capture foliage production and root system of vegetation for transpiration. The model is also unable to produce savanna ecosystems, which it simulates by approximating vegetation of forest and grassland ecoregions (Dahlin et al., 2020). In addition, the ineffective simulation of deciduousness would have contributed to the model biases in response simulations. Therefore, targeting these limitations is important for improving model’s performance in simulating morphology and physiological functioning of vegetation biomes. Furthermore, the DGVMs (e.g. JULES, DLEM) used in the study poorly replicate important ecological and physiological processes that are critical to capture the dynamics of savanna systems. Other DGVMs (e.g. JSBACH) poorly simulate significant environmental variables such as fire, which is very crucial for the vegetation growth cycle, particularly in the savanna biome (Thonicke et al., 2001; Romps et al., 2014; Kim et al., 2018; D’Onofrio et al., 2020). Also, over southern Africa, land use change (LUC) is a common and frequent occurrence, and is an important factor for vegetation turnover. However, most models do not well capture land management, which is an important driver of land cover change in the region. Thus, there is a need for future model development to account for rapid LUC over different regions. However, the disparity in observed and simulated response of vegetation to drought cannot be fully accounted by the DGVMs alone. The reanalysis (CRUJRA) has also shown some limitations in the simulation of climate variables. Compared to observation-based CRU, CRUJRA has closer magnitudes of maximum and minimum temperature, and addressing this would improve simulated response.

5 Summary and conclusions

Southern African vegetation is continually affected by drought. In this study, we estimated the spatiotemporal characteristics of meteorological drought in southern Africa using the Standardized Precipitation Evapotranspiration Index (SPEI) over a 30-year period (1982 – 2011). The severity of drought and its impacts on vegetation production were examined at various drought time-scales (1- to 24-month timescales) by correlating the drought index (SPEI) with GIMMS LAI at different timescales. We found that the LAI responds strongly ($r = 0.6$) to drought over the central and south eastern parts of the region, with weaker impacts ($r < 0.4$) over parts of Madagascar and Angola, mostly at a shorter time period (3-, 6-month timescale). We note that, comparing CRU to flux tower precipitation in Skukuza (Kwazulu Natal region) illustrates that CRU captures the timing of precipitation relatively well, though it underestimates the magnitude of dry season precipitation (Fig. S9). For seasonal responses, the semi-desert biome showed the strongest response ($r = 0.95$) to drought at 6-month timescale in the autumn season while the tropical forest biome shows the weakest response ($r = 0.35$) at 6-month timescale in the DJF season.

We assessed the relationship between the NDVI and LAI by computing a grid cell correlation of NDVI and LAI, and examined how well state-of-the-art dynamic global vegetation models (DGVMs) simulate LAI and its response to drought. The DGVM multi-model ensemble mostly overestimated the spatial and seasonal distribution of LAI response to drought in most parts of the region. The results also show that:
• The relationship between the NDVI and LAI is linear, implying that the vegetation index is connected to the changes in phenology of plant, reflectance and the angle of zenith variations from the surface cover class.

• The model ensemble simulates the correlation and timescale of LAI response to drought with biases.

• The model ensemble overestimates observed responses on the seasonal distribution of vegetation – drought correlations across different biomes.

• There is a stronger LAI response to rainfall deficit in dry years than in wet years (for instance, in the year 1983 which was a dry year, $r > 0.84$ over Namibia and northeastern parts of South Africa and $r < 0.70$ in 2000, a wet year in the same region); and a variability in the distribution patterns of global and regional response across biomes.

The present study has shown how the LAI responds to drought across the different southern African biomes. Given the present spatial coverage of space monitoring of vegetation in the region, the methods used in the study may be extended towards monitoring and characterizing the impacts of droughts on land cover change, as this may permit real-time monitoring of extreme events on terrestrial vegetation (Yin et al., 2020; Moore et al., 2018). The findings of this study (e.g. timescales of LAI response to drought) could also be used for the development of drought early warning systems in agriculture and forestry sectors. This could assist in the mitigation of direct and indirect costs associated with vegetation production.

Furthermore, this study has applied eleven DGVMs to study how well DGVMs can reproduce the response of LAI in southern African vegetation to drought. While this study may have provided an insight into the capability of DGVMs to simulate vegetation response to drought, the results of the study can however, be improved in some ways. For example, we applied eleven models from the TRENDY DGVMs. For future studies, the number of models should be increased, perhaps from other model intercomparison experiments, because using more models, as well as better quantification, might capture uncertainty better in their simulation of drought response by vegetation. The limitation of the DGVMs can be addressed by optimizing the models so that their capability in reproducing vegetation indices is enhanced and better quantified. In addition, there is a need to improve mechanistic relationships in the models, which could be achieved by enhancing the model approximations which had been done to achieve computational efficiencies (Transtrum et al., 2016). Furthermore, simple phenomenological model could be developed from the complex model. These simple models would use correlations among observations, unlike mechanistic relationships which exploit causative individual constituent and suffer over-fitting problems (Transtrum et al., 2016). Lastly, there may be the need for hybridizing machine learning and mechanistic models (Fayyad et al., 1996; Mitchell, 1997) to simulate vegetation parameters. This is because machine learning models have shown certain advantage in the prediction of outcomes of complex mechanisms by using databases of inputs and outputs for a given task (Fayyad et al., 1996; Mitchell, 1997).
Data availability. Sources of data used in this work are provided in Section 2.1 and comprise CRU, CRUJRA, NDVI3g and Trendy DGVMs.

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