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Contrasting responses of autumn leaf senescence to daytime and nighttime warming

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Plant phenology is a sensitive indicator of climate change\textsuperscript{1-4}, and plays a significant role in regulating carbon uptake by plants\textsuperscript{5-7}. Previous studies have focused on spring leaf-out by daytime temperature and the onset of snowmelt time\textsuperscript{8-9}, but the drivers controlling leaf senescence date (LSD) in autumn remain largely unknown\textsuperscript{10-12}. Using long-term ground phenological records (14536 time series since the 1900s) and satellite greenness observations dating back to the 1980s, we show that rising preseason maximum daytime (T\textsubscript{day}) and minimum nighttime (T\textsubscript{night}) temperatures had contrasting effects on the timing of autumn LSD in the Northern Hemisphere (>20°N). If higher T\textsubscript{day} leads to an earlier or later LSD, an increase in T\textsubscript{night} systematically drives LSD to occur oppositely. Contrasting impacts of daytime and nighttime warming on drought stress may be the underlying mechanism. A new LSD model considering these opposite effects improved autumn phenology modeling, and predicted an overall earlier autumn LSD by the end of this century compared with traditional projections. These results challenge the notion of prolonged growth under higher autumn temperatures, suggesting instead that leaf senescence in the Northern Hemisphere will begin earlier than currently expected, causing a positive climate feedback.

Climate change over the last several decades has modified the dates of plant flowering, leaf emergence, growth stages, and senescence, collectively termed phenology\textsuperscript{13} with substantial ecological and environmental consequences\textsuperscript{4}. Both observations and model simulations have found that air temperature has a positive influence on the onset of plant growth in the Northern Hemisphere (NH), e.g., higher spring temperature triggers earlier
leaf-out and flowering dates and hence extends the growing season\textsuperscript{8,14-15}. In contrast to those extensive research efforts on spring phenology, autumn phenology, particularly leaf senescence date (LSD), is more challenging to understand, and has not received sufficient attention\textsuperscript{16,17}, while also serving as an important indicator of changing foliar physiological properties. Yet, autumn phenology may be as important as spring in regulating the interannual variability of carbon balance\textsuperscript{7}.

LSD has been occurring later in most regions over the last few decades\textsuperscript{18}, but providing an explanation for this change is difficult\textsuperscript{9}. An increase in global temperature is assumed to be a driver of LSD trends\textsuperscript{19}, but studies indicated that the contribution of temperature to LSD variability is low, especially compared to spring phenology\textsuperscript{20,21}. We argue that ignoring the asymmetric effects\textsuperscript{22} of daytime maximum temperature ($T_{\text{day}}$) versus nighttime minimum temperature ($T_{\text{night}}$) and their differing impacts on LSD, contributes to the reported overall low contribution of temperature to LSD variability. To test this, we used measured and gridded preseason (defined as months from June to LSD) $T_{\text{day}}$ and $T_{\text{night}}$ in the NH, together with LSD data from three different datasets: (a) long-term phenological observations at ground sites from 14536 time series since the 1900s (Fig. S1), (b) the latest third generation of the Normalized Difference Vegetation Index (NDVI, GIMMS3g.v1) for 1982-2015, and (c) NDVI and enhanced vegetation index (EVI) from the Moderate Resolution Imaging Spectroradiometer (MODIS) products for 2001-2015.

Preseason forcing had a better predictive strength on LSD than either summer or autumn
climate forcing alone (Fig. S2). Because preseason $T_{\text{day}}$ and $T_{\text{night}}$ were highly correlated, we used a partial correlation to remove the effects of $T_{\text{night}}$ and of precipitation and radiation (similarly for $T_{\text{night}}$) to investigate the response of LSD to $T_{\text{day}}$. Correlations were classified into four types, (A) $T_{\text{day}}^{+} T_{\text{night}}^{+}$, (B) $T_{\text{day}}^{-} T_{\text{night}}^{-}$, (C) $T_{\text{day}}^{+} T_{\text{night}}^{-}$ and (D) $T_{\text{day}}^{-} T_{\text{night}}^{+}$, where $T^{+}$, $T^{-}$ represent positive and negative partial correlation coefficient of T with LSD.

Overall, all three datasets suggested that the onset of autumn LSD responded oppositely to $T_{\text{day}}$ and $T_{\text{night}}$. The proportions of ground sites of Types A and B were significantly lower than those of Type C and D (Fig. 1a). More significant R values for both $T_{\text{day}}$ and $T_{\text{night}}$ were found within Types C and D, with only two and one records out of 2231 time series having significant R within Type A and B, respectively. These results from ground sites are consistent with those for the two satellite greenness products (Fig. 1b, c). Types C and D together accounted for 83.7 and 80.0% for GIMMS3g and MODIS pixels, respectively. Only 0.8 and 1.5% of the pixels had the same sign of response of LSD to $T_{\text{day}}$ or $T_{\text{night}}$ (i.e. significant pixels for Types A+B) for GIMMS3g and MODIS, respectively. The GIMMS3g dataset contained different fractions of Types C and D (45.6% vs. 38.1%), but the compositions of Type C and D in GIMMS3g (i.e., contrasting effects of night and day temperatures) became more consistent with the MODIS results when the overlapping periods between the two sensors is considered (Figs. S3-4). More details on the fractions of the four correlation types for different vegetation types are provided in Figs. S5-6.

Figure 1 Frequency of the partial correlation coefficient (R) between leaf senescence date
(LSD) and daytime ($T_{\text{day}}$), nighttime ($T_{\text{night}}$) temperatures. (a) Data for 14536 time series of ground sites, (b) the GIMMS3g dataset for 1982-2015, and (c) the MODIS product for 2001-2015. $T^+$, $T^-$ represent positive and negative partial correlation coefficient of $T$ with LSD. Significance was set at $P < 0.05$.

The satellite greenness products also allowed us to evaluate spatial patterns of LSD changes in response to variations in $T_{\text{day}}$ and $T_{\text{night}}$ (Fig. 2). For the GIMMS3g data, higher $T_{\text{day}}$ was associated with a delayed LSD for 10.7% of the pixels (mostly boreal regions) and with an earlier LSD for 7.5% of the pixels (central North America, borders of Eurasia and central China). $T_{\text{night}}$ had evident opposite influences on LSD than $T_{\text{day}}$. The patterns of opposite effects from $T_{\text{day}}$ and $T_{\text{night}}$ on LSD were highly spatially consistent in all regions where $T_{\text{day}}$ and $T_{\text{night}}$ were significantly correlated with LSD. Similar results were obtained with MODIS observations (Fig. 2b, d). LSD for approximately 20% all pixels was significantly correlated with $T_{\text{day}}$, of which 60.1 and 39.9% were negatively and positively correlated, respectively. The area where LSD was positively correlated with $T_{\text{night}}$ was larger (9.4%) than the area with negative correlations (6.5%).

Vegetation grouped into Köppen-Geiger zones showed contrasting patterns between the effects of $T_{\text{day}}$ and $T_{\text{night}}$ on LSD (Fig. 2e, f). Type D was more widely distributed, while Type C was more common for continental climates. Monsoon-influenced but not extremely cold regions and mild climates also had higher proportions of Type C. Grouping these correlation types by vegetation types lead to similar results (Fig. 2g, h). In theory,
we would expect to find Type C more in wet vegetation types and Type D in dry types. The real world seems to show the same thing but still there could be many locations that do not neatly fall into that continuum and suggests additional mechanisms may work, probably adaption.

Figure 2 Spatial distributions of the partial correlation coefficient (R) between leaf senescence date (LSD) and daytime (T_{day}), nighttime (T_{night}) temperatures. $R^+$, $R^-$ represent positive and negative partial correlation coefficient of T with LSD. (a) LSD vs. T_{day} for GIMMS3g, (b) LSD vs. T_{day} for MODIS, (c) LSD vs. T_{night} for GIMMS3g, and (d) LSD vs. T_{night} for MODIS. (e), (f), represent distributions of correlation types in Köppen-Geiger climatic classification using GIMMS3g and MODIS, respectively. (g) and (h) represent distributions of correlation types for vegetation types (see Methods) using GIMMS3g and MODIS, respectively. Significance was set at $P < 0.05$.

Our results suggest that ecological trade-offs, particularly those driven by regional differences in water stress, may underlie the contrasting relationships between LSD and T_{day} and T_{night}. Type C was mostly found in humid regions where water is a less limiting factor for plant growth. In these cases, a higher T_{day}, in the likely absence of severe water stress, benefits photosynthesis while elevated T_{night} increases nighttime leaf respiration.

Explanations for the prevalence of Type D relations in dry regions are more complicated. The Standardized Precipitation Evapotranspiration Index (SPEI)\textsuperscript{23}, an indicator of drought
stress, accounted for the contrasting effects of increases in $T_{\text{day}}$ and $T_{\text{night}}$ on LSD for Type D (Fig. 3). Partial correlation data indicate that increased $T_{\text{day}}$ is negatively correlated with SPEI (Fig. 3a), a stronger sensitivity to drought in dry regions that negatively affects plant growth and consequently leads to an earlier LSD. In contrast, we found that an increase in $T_{\text{night}}$ is associated with a higher SPEI, that is, wetter conditions, and arguably reduced water stress, which could extend the duration of photosynthesis and lead to delayed LSD (Fig. 3b). The latter mechanism is consistent with the generally positive partial correlation values between evapotranspiration (ET) and $T_{\text{night}}$, that is, more soil moisture being available for ET in the late season, and sustaining delayed LSD (Fig. 3f), and with studies showing that water stress accelerates leaf drop in dry ecosystems more so than in humid ecosystems\textsuperscript{24}. The responses of radiation to $T_{\text{day}}$ and $T_{\text{night}}$ may also be viewed as a further evidence for the linkage between leaf senescence and plant water status to support the contrasting patterns (Fig. S7), given that a higher $T_{\text{day}}$ was associated with stronger radiation and potentially a higher chance of water stress. These findings suggest that dry regions, in which Type D dominates, may be especially vulnerable to earlier onset of LSD if climate change reduces local precipitation and increases daytime evaporation with rising $T_{\text{day}}$.

Apart from physiological mechanisms relating to water stress, ecological processes may also contribute to these patterns. Warmer daytime versus nighttime temperature may have contrasting effects on different species since species adaptations lead to intrinsic differences in their timing of leaf emergence and senescence that are optimized to
maximize carbon gain and minimize water losses\textsuperscript{25-27}. The ecosystem-scale responses of
phenology reflects the scaled responses of ecological dynamics of multiple individual
species gaining or losing a competitive advantage in a changing climate, or presenting an
induced advantage as a result of land-use change and planting\textsuperscript{17,26}. Recent results
suggest that the magnitude of phenological change to effects by shifts in plant species
composition may be similar as that by climate change\textsuperscript{27}, and the autumn phenology may
thus change accordingly.

\textbf{Figure 3 Partial correlation coefficient (R) between the Standardized Precipitation
Evapotranspiration Index (SPEI), evapotranspiration (ET), and daytime (T_{\text{day}}), nighttime
(T_{\text{night}}) temperatures.} (a) SPEI vs. T_{\text{day}} for GIMMS3g, (b) SPEI vs. T_{\text{night}} for GIMMS3g, (c)
SPEI vs. T_{\text{day}} for MODIS, (d) SPEI vs. T_{\text{night}} for MODIS, (e) ET vs. T_{\text{day}} for MODIS, and (f)
ET vs. T_{\text{night}} for MODIS. Significance was set at $P < 0.05$.

We tested the implications of the observation analysis on future trends in autumn LSD by
developing a weighted day-night-temperature growing-degree-day (DN\textsubscript{GDD}) algorithm
incorporating these opposite changes in LSD to $T_{\text{day}}$ and $T_{\text{night}}$ (see Methods). The new
model substantially improved LSD modeling (in terms of R (Figs. S8-10), RMSE (Figs.
S11-13) and percentage of significant pixels (Figs. S14-15)) compared with the currently
used threshold or GDD methods both for the overall dataset and for vegetation types.

Spatial patterns of improvements using MODIS and GIMMS3g were also investigated
(Figs. S16-17). The results from MODIS and the ground sites (Fig. S18) were more consistent with the new model, and the accuracy of the threshold method was much lower, so we used the coefficients from the MODIS data to predict LSD variability by the end of this century using traditional GDD and DN$_{GDD}$ algorithms under two Representative Concentration Pathways (RCP) scenarios (RCP 4.5 and RCP 8.5) (Fig. S19, and Fig. 4).

LSD from the DN$_{GDD}$ method was overall earlier than conventional predictions across Köppen-Geiger climatic classification types. Globally, LSD was earlier for about 68% of the terrestrial biosphere under RCP 4.5 and for about 70% under RCP 8.5. LSD was mostly later for central North America, western Russia, and southwestern China. Most vegetation types showed earlier LSD estimates under two RCP scenarios while the temperate grasslands were expected to have later senescence dates.

Figure 4 Leaf senescence date (LSD) differences from the weighted day-night-temperature growing-degree-day (DN$_{GDD}$) and traditional GDD (LSD$_{DNGDD}$-LSD$_{GDD}$) models under two RCP scenarios. (a), (b), (c) represent LSD$_{DNGDD}$-LSD$_{GDD}$ under RCP 4.5, and averages of differences for the Köppen-Geiger climatic classification, and vegetation types, respectively. (d), (e), (f) represent LSD$_{DNGDD}$-LSD$_{GDD}$ under RCP 8.5, and averages of differences for the Köppen-Geiger climatic classification, and vegetation types, respectively.
Climatic variability, particularly temperature, has driven phenological changes over the last several decades but has been challenging to predict. Our ability to predict autumn LSD is particularly limited. We are the first to report, using 14536 ground time series and more than 30 years of remotely sensed observations, the opposite responses of LSD to daytime and nighttime warming, providing a new perspective to account for the previous low estimation accuracy of autumn LSD when relying solely on mean temperature. A model based on mean temperature cannot correctly predict LSD changes, because LSD responds to $T_{\text{day}}$ and $T_{\text{night}}$ in opposite directions. Our results also provide a perspective to account for the carry-over effects between spring and autumn phenology, i.e. the start and end of a growing season always move in the same direction\textsuperscript{28}. An earlier start of a season is mainly triggered by higher spring temperatures, with increased growth depleting soil water\textsuperscript{29}, which is then associated with autumn drought, inducing a reduction in growth and consequently leading to an earlier end to the growing season\textsuperscript{30}. Our improvement in modeling autumn phenology is a strong and convincing evidence for the value of incorporating daytime and nighttime temperatures in terrestrial models, rather than mean temperature alone. The application of this model projects an overall earlier than currently expected start of autumn senescence in the NH by the end of this century, particularly in dry regions. The earlier data of autumn senescence may be a potentially unrecognized positive feedback to climate change and consequently a weakening in the capability of terrestrial carbon uptake.
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Contributions: C.W., H.W. and Q.G. designed the research. C.W. wrote the first draft of the paper. J.P. and C.P. extensively revised the writing. H.W. performed the site model simulations. X.W. performed remote sensing model simulations. All the authors contributed to writing the paper.

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Competing interests: The authors declare no competing financial interests.

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Methods

1. Phenological observation data

We used observations of leaf senescence date (LSD) from three independent phenological datasets.

1) The Pan European Phenological Database (PEP725; http://www.pep725.eu/), an open-access database with long term plant phenological observations (since 1868) from 19608 sites and 78 species across 25 European countries.

2) The China Phenological Observation Network (CPON), with data since 1963 for >100 species at 42 sites across China.

3) LSD data for two tree species (Acer palmatum and Ginkgo biloba) at 54 meteorological stations in South Korea for 1989-2007.

The definitions of LSD notably differ among the datasets. LSD for the PEP725, CPON, and Korean datasets is defined as the date when 50, 90, and 20% of the tree leaves, respectively, change color from green to red or yellow. We removed outliers using the methods to exclude potential biases and inadequate degrees of freedom and focused on time series with at least 15 years of records for 1900-2015. We thus analyzed 14536 LSD
2. LSD derived from satellite data

LSD in the Northern Hemisphere was determined using two satellite-derived vegetation indices, the Normalized Difference Vegetation Index (NDVI) and the Enhanced Vegetation Index (EVI)\textsuperscript{33}. Both NDVI and EVI are direct indicators of vegetation growth and have been widely applied for investigating vegetation phenology\textsuperscript{34}. We used two datasets to reduce the uncertainties caused by a single data source: GIMMS NDVI third-generation (NDVI3g) data derived from the Advanced Very High Resolution Radiometer (AVHRR) and NDVI and EVI derived from the Moderate Resolution Imaging Spectroradiometer (MODIS). The GIMMS NDVI3g data have a spatial resolution of 1/12°, a half-month interval, and a 34-year time span (1982-2015). The MODIS 16-day composite product MOD13C1 (Collection 6) includes both NDVI and EVI with a 0.05° resolution for 2001-2015.

We eliminated the impacts of areas with sparse vegetation on the results by first excluding pixels with annual NDVI <0.1 or annual EVI <0.08\textsuperscript{35}. A Savitzky-Golay filter was then used to smooth the NDVI (EVI) time series\textsuperscript{36}. We then estimated LSD using two methods.

The first method was a dynamic-threshold approach, which uses an annually defined threshold for each pixel based on the NDVI ratio:

\[
NDVI_{\text{ratio}} = \left( \frac{NDVI - NDVI_{\text{min}}}{NDVI_{\text{max}} - NDVI_{\text{min}}} \right)
\]  

(1)

where NDVI is the daily NDVI and NDVI\textsubscript{max} and NDVI\textsubscript{min} are the annual daily maximum
and minimum of NDVIs. The NDVIRatio ranges from 0 to 1. LSD is determined when
NDVIRatio decreases to 0.5 in autumn.37,38

The second method was based on a series of piecewise logistic functions. The NDVI
time series were first divided into two sections by the maximum daily NDVI in each year,
and a double logistic function was applied to fit each section:

\[
y(t) = a_1 + (a_2 - a_1) \left[ \frac{1}{1 + e^{(a_3 - t)/a_4}} - \frac{1}{1 + e^{(a_5 - t)/a_6}} \right]
\]  

(2)

LSD was then defined as the time when the curvature changing rate reached its last
local maximum value.

For GIMMS3g data, we calculated LSD using NDVI from both the dynamic-threshold
approach and the piecewise logistic function method. Since MODIS sensor provides EVI,
we further used EVI-based logistic function method to derive LSD. To sum up, for
GIMMS3g data, average LSD from threshold and logistic function method were used, and
for MODIS, an additional LSD from EVI-based logistic function method was used (not for
MODIS NDVI data).

At high latitudes (or elevations), snow cover is important for regional climate and
arrives early in autumn and potentially masking evergreen vegetation. However, we
suggested that using a Savitzky-Golay filter could solve the noise from a "sudden" change
in the time series of NDVI due to snow.36 In particular, a study showed that snowfall had
little influence on determining EOS in western Arctic Russia.41 For high elevations, our
previous analysis on Tibetan Plateau showed that for more than 98% of regions snow
happened later than LSD.42
3. Climatic data

We used the CRU-TS 4.00 dataset with a spatial resolution of 0.5°×0.5° for 1901-2015. We extracted monthly data for $T_{\text{max}}$, $T_{\text{min}}$, $T_{\text{mean}}$, precipitation, and cloud cover from this dataset for analyzing LSD from in-situ observations and the two remote sensing data. We modeled past and future LSD by temperature by acquiring daily gridded data for maximum and minimum temperature with a spatial resolution of 0.5° from NOAA Earth System Research Laboratory Physical Sciences Division for 1982-2015. We used daily $T_{\text{max}}$ and $T_{\text{min}}$ simulated by the CCSM 4 model under two climatic scenarios (RCP4.5 and RCP8.5) for future climatic data (2081-2100). These data were from an open-access database of the Coupled Model Intercomparison Project Phase 5.

4. Analyses

We used partial correlation analyses to determine the responses of LSD to $T_{\text{day}}$ and $T_{\text{night}}$. The reason is that directly correlating LSD to $T_{\text{day}}$ would give misleading results since $T_{\text{night}}$ is a confounding variable that is numerically related to both LSD and $T_{\text{day}}$, violating independence of variables in multiple correlation tests. Thus, using the partial correlation between LSD and $T_{\text{day}}$ will measure the degree of association with the effect of a set of controlling random variables removed (e.g., $T_{\text{night}}$, precipitation, radiation), given that these factors have shown strong influences on LSD. Since Echer and Silva (2014) demonstrated that clouds are the main atmospheric factor modulating the surface incidence of solar radiation, cloud cover data was used to model the effect of radiation on LSD, as similarly conducted in previous analyses. Correlations were classified into
four types, (A) $T_{\text{day}+T_{\text{night}+}}$, (B) $T_{\text{day}+T_{\text{night}-}}$, (C) $T_{\text{day}+T_{\text{night}-}}$ and (D) $T_{\text{day}-T_{\text{night}+}}$, where $T^+$, $T^-$ represent positive and negative partial correlation coefficient of $T$ with LSD. An $R$ of at least 0.514 for MODIS is required for the significance test ($p = 0.05$), but this value decreases to 0.339 for the longer GIMMS3g data. These analyses were investigated for both Köppen-Geiger climatic classifications and vegetation types (Table S2). Crops were excluded because their signal may result from changes in cropping or harvest cycles, rather than from climate change. Furthermore, since at low latitudes plant phenology of tropical and subtropical areas responds to other factors than temperature, regions with latitudes < 20° N were also excluded.

Current phenology algorithms in most terrestrial-biosphere models are solely based on temperatures in the preceding months. We determined the length of the preseason whose average $T_{\text{day}}$ had the most influence on LSD by calculating the partial correlation coefficients between LSD and mean $T_{\text{day}}$ during 0, 1, 2, … $n$ months prior to LSD, controlling for corresponding mean $T_{\text{night}}$, total precipitation, and radiation. The maximum range ($n$) of the preseason is generally from June to the multiyear mean date of LSD (see Fig. S20 for example). The partial correlation coefficients with the highest absolute values were then used in the following analysis. We obtained the relationship between LSD and $T_{\text{night}}$ the same way but replacing $T_{\text{day}}$ with $T_{\text{night}}$. This analytical procedure was applied for observed LSD from ground sites and derived LSD from the MODIS and GIMMS NDVI3g data.

5. Models for predicting LSD
Our results indicated that LSD responded oppositely to $T_{\text{day}}$ and $T_{\text{night}}$, so we developed a weighted day-night-temperature growing-degree-day (DNGDD) algorithm from observations to model LSD, and compared the algorithm with currently used threshold and GDD models based on $T_{\text{mean}}$.

The threshold model was the simplest method. We calculated average $T_{\text{mean}}$ for five days before LSD in each year and used the multiyear mean value as the threshold to model LSD. If $T_{\text{mean}}$ was lower than the threshold for five consecutive days from 1st July, the last date was considered the LSD.

GDD was calculated as:

\[
GDD(d) = \max\left(T_b - T_{\text{mean}}(d), 0\right)
\]  

(3)

\[
GDD_{\text{threshold}} = \sum_{d=d_0}^{LCD} GDD(d)
\]  

(4)

where $T_b$ is the base temperature set to 15, 20, 25, and 30 °C, $T_{\text{mean}}(d)$ is the mean daily temperature, and $d_0$ is the date on which the calculation begins (1st July in this study). LSD is the observed or derived date of leaf coloring in each year. The date when $GDD(d)$ exceeded the multiyear average GDD threshold was defined as LSD.

Our DNGDD model improved upon the original GDD model and was calculated by:

\[
GDD(d) = k \cdot \max\left(T_b - T_{\text{day}}(d), 0\right) + (1 - k) \cdot \max\left(T_b - T_{\text{night}}(d), 0\right)
\]  

(5)

where $T_{\text{day}}(d)$ is the daily maximum temperature, $T_{\text{night}}(d)$ is the daily minimum temperature, and $k$ is the weighting factor. When $0 < k < 1$, the effects of $T_{\text{day}}$ and $T_{\text{night}}$ on LSD are consistently positive; When $k > 1$ or $k < 0$, the effects of $T_{\text{day}}$ and $T_{\text{night}}$ on LSD are opposite. In order to determine the value of $k$, we first calculated the ratio of $R_{\text{day}}$ and $R_{\text{night}}$ for each station or pixel, and found 99.9% of the ratio values were between -10 and 10 for each station or pixel.
both ground and satellite data (Figure S21). In other words, the level of T\textsubscript{day}(T\textsubscript{night}) effect could be 1 to 10 times than the level of T\textsubscript{night}(T\textsubscript{day}) effect (note that T\textsubscript{day} represents T\textsubscript{day} with the effects of T\textsubscript{night} removed). Therefore, the values of $k$ ranged from –1 to 2 (see Table S3). In addition, when $k$ tends to infinity, the effects of T\textsubscript{day} and T\textsubscript{night} on LSD are opposite with same level.

We evaluated the accuracy and obtained the most appropriate parameters of the models by calculating the correlation coefficient (R) and the root mean square error (RMSE) between modeled and observed LSD. $T_b$ and $k$ with the lowest RMSE were considered the most appropriate values for each site or pixel.

**Data Availability**

The data that support the findings of this study are available from the corresponding author upon request.

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