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1	Primary Research Article
2	Title: Critical temperature and precipitation thresholds for the onset of xylogenesis of
3	Juniperus przewalskii in a semi-arid area of the northeastern Tibetan Plateau
4 5 6	Running head: Thresholds for the onset of xylogenesis
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29 Abstract

The onset of xylogenesis plays an important role in tree growth and carbon 30 31 sequestration at the level of ecosystems and is thus a key to modeling the responses of forest ecosystems to climate change. Temperature regulates the resumption of cambial 32 activity, but little is known about the effect of water availability on the onset of 33 xylogenesis. We monitored the onset of xylogenesis during 2009–2014 by weekly 34 microcoring Juniperus przewalskii trees at the upper and lower altitudinal limits of the 35 species on the northeastern Tibetan Plateau. A logistic regression was used to 36 37 calculate the probability of xylogenic activity at a given temperature. A two-dimensional reversed Gaussian model was used to fit the differences between the 38 observed date of onset of xylogenesis and days at given temperatures and 39 40 precipitation within a certain time window. The thermal thresholds at the beginning of the growing season were highly variable, providing additional evidence that 41 temperature was not the only factor initiating xylem growth under cold and dry 42 climatic conditions. The onset of xylogenesis was predicted well for climatic 43 thresholds characterized by a cumulative precipitation of  $17.0 \pm 5.6$  mm and an 44 average minimum temperature of  $1.5 \pm 1.4$  °C for a period of 12 days. Xylogenesis in 45 semi-arid regions with dry winters and springs can start when both critical 46 temperature and precipitation thresholds are reached. Such findings contribute to our 47 knowledge of the environmental drivers of growth resumption that were previously 48 49 investigated only in regions with abundant snow accumulation in winter and frequent precipitation in spring. Models of the onset of xylogenesis should include water 50

51	availability for more reliable predictions of xylem phenology in dry areas. A
52	mismatch of the thresholds of temperature and moisture for the onset of xylogenesis
53	may increase forest vulnerability in semi-arid areas under droughts due to global
54	climate change.
55	

- 56 Keywords: Phenology, xylem formation, Qilian juniper, two-dimensional Gaussian
- *model, drought, rain, altitudinal gradient*

Interest in xylem phenology (xylogenesis) and its sensitivity to climate change is 60 growing because wood is a major sink of carbon in terrestrial ecosystems (Babst et al., 61 2014; Cuny et al., 2015; Pérez-de-Lis et al., 2017). Temperature is increasingly 62 recognized as the primary driver of growth reactivation in cold climates (Rossi et al., 63 2007, 2008). Both observations and controlled experiments have demonstrated that 64 cambial activity is limited by low air temperatures in cold climates (Oribe et al., 2001; 65 Gricar et al., 2006; Rossi et al., 2008; Seo et al., 2008; Gruber et al., 2010; Begum et 66 67 al., 2013; Li et al., 2013). In addition, the onset of xylem production is delayed at higher latitudes and altitudes, confirming the role of temperature for xylogenesis 68 (Moser et al., 2010; Oladi et al., 2010; Huang et al., 2011). In particular, Rossi et al. 69 70 (2008) reported a critical daily minimum temperature for xylogenesis in conifers of 4-5 °C in cold climates. Shen et al. (2015), though, highlighted the impact of 71 precipitation on the starting date of vegetation phenology (canopy greening) in cold 72 73 and arid or semi-arid regions, indicating that cold and drought stress both affected the onset of growth. Ren et al. (2015) found a delay in the initiation of xylogenesis in 74 Qilian junipers (Juniperus przewalskii Kom.) under extremely dry spring conditions 75 in a cold and dry climate, which suggested a potential influence of water availability 76 on the start of xylogenesis, i.e. on the onset of cambial reactivation after the cold 77 dormant season (winter in the Northern Hemisphere). 78

The effect of precipitation on the growth dynamics of forest ecosystems needs tobe quantified to better understand the adaptation of plants to a changing climate,

which may be characterized by warmer and drier conditions (Allen et al., 2015). In 81 addition, the climatic thresholds for the resumption of xylem phenology may provide 82 keys to better understand mechanisms of forest resilience (e.g. post-drought recovery) 83 and the potential for tipping points under global change. Water acts on several 84 important growth processes in plants. The expansion of xylem cells is turgor-driven, 85 depending on the uptake of cellular water and on solute accumulation. Drought stress 86 affects the loss of turgor of differentiating cells (Kozlowski & Pallardy, 2002), so 87 shifts in the onset of xylogenesis might be potentially affected by variation in 88 89 moisture conditions, especially in the arid and semi-arid regions of the world. The available literature, however, is limited to studies conducted in regions characterized 90 by rains prior to the onset of xylogenesis (from winter to spring), such as the 91 92 Mediterranean basin (Camarero et al., 2010, 2015; Vieira et al., 2013), or by abundant water released during snowmelt, such as alpine valleys (Gruber et al., 2010; Eilmann 93 et al., 2011; Swidrak et al., 2011). Soil moisture could be a less important limiting 94 95 factor for the resumption of xylem formation at these sites than in arid or semi-arid areas. We investigated how cold and dry conditions could drive the onset of 96 xylogenesis by determining the relative influence of these two climatic stressors. 97

We selected a forested area on the northeastern Tibetan Plateau to test the effect of soil moisture on the onset of xylogenesis. The dry climate of this area is characterized by scarce winter precipitation, a very thin snowpack and the dependence of moisture availability for vegetation activity on the first rains of spring (Dai, 1990). The climate is described as cold and dry, with a mean annual temperature of 3.1 °C and a mean

annual precipitation of ca. 200 mm. Winter is extremely dry, and rain mainly falls 103 from May to September (Dai, 1990). The Qilian juniper forests in this area are 104 stressed by both drought and cold (Zheng et al., 2008). A recent study reported that 105 spring drought could delay the onset of xylogenesis in Oilian juniper despite optimal 106 thermal conditions (Ren et al., 2015). In addition, warmer spring conditions on the 107 plateau are increasing the vulnerability of forests to dry spells, indicated by a marked 108 decrease in growth and an increase in the frequency of missing tree rings (Liang et al., 109 2014, 2016). These findings suggested a potential interaction between precipitation 110 111 and temperature in the onset of xylogenesis under cold and dry conditions.

The objective of this study was to use Qilian juniper as a model species to investigate the onset of xylogenesis at the upper and lower altitudinal boundaries of its distribution during six growing seasons (2009-2014) and to identify the thresholds of temperature and precipitation controlling the onset of xylogenesis. We hypothesized that the onset of xylogenesis in Qilian juniper was constrained more by water deficit than by low temperatures.

118

#### 119 Materials and Methods

# 120 Study site, field sampling and sample preparation

The study was carried out in an undisturbed Qilian juniper forest near Dulan County on the northeastern Tibetan Plateau (36°00'N, 98°11'E). Two sites, at 3850 and 4210 m a.s.l. with slopes of 15°, were selected at the lower and upper timberline limits of the altitudinal distribution of the species. Five trees were randomly selected at each

site. The average diameter at breast height was 50-60 cm, and the average height was 125 8 m. Microcores were extracted weekly from 2009 to 2014 from the stems at a height 126 of 1.0-1.3 m using a Trephor microborer (Rossi et al., 2006) and stored in a 127 formalin-ethanol-acetic acid solution. The microcores were prepared to obtain 128 transverse sections (9-12 µm in thickness) using a Leica RM 2245 rotary microtome 129 (Leica Microsystems, Wetzlar, Germany), and the sections were stained using a 130 mixture of safranine, Astra Blue and ethanol and then permanently fixed. See Ren et 131 al. (2015) for more details on sampling strategy and slide preparation. 132

133

## 134 *Identification of the onset of xylogenesis*

The xylem sections were observed under a microscope at a magnification of  $100 \times$ 135 136 with visible and polarized light to distinguish the differentiating xylem cells. We concentrated on the radial-enlargement phase, which indicates the beginning of xylem 137 growth (Antonova & Stasova, 1993). Tracheids in the radial-enlarging phase 138 139 contained a protoplast enclosed in thin primary cell walls, and their radial diameters were at least twice that of a cambial cell (Rossi et al., 2006). The tracheids had 140 light-blue walls under normal light during this phase but were not visible under 141 polarized light due to the lack of a secondary wall. Xylogenesis was considered to 142 have begun for each tree when at least one radial file of enlarging cells was observed 143 in spring. 144

145

## 146 Meteorological data

147	Meteorological data were recorded at each site from October 2012 by automatic
148	stations (HOBO; ONSET, Pocasset, USA). Air temperature and precipitation were
149	measured every 30 min and stored in data loggers. Minimum, mean, and maximum
150	daily temperatures and daily precipitation were calculated for subsequent analyses.
151	Data for January 2009 to September 2012 were estimated using the measurements
152	collected from a meteorological station in Dulan (36°18'N, 98°06'E; 3190 m a.s.l.), 32
153	km from the study sites. The consistency of the estimates was based on the high
154	correlations ( $r > 0.92$ ) between the climatic data (temperature and precipitation) at the
155	two sites with those at the Dulan station (Supporting Information, Fig. S1).

# 157 Statistical analyses to predict climatic thresholds of xylogenesis

Based on previous research (Ren *et al.*, 2015), air temperature and precipitation were
selected as potential climatic drivers of the beginning of xylogenesis.

Logistic regression was used to calculate the probability of xylogenic activity at a given temperature using the LOGISTIC procedure in the SAS 9.4 statistical package [SAS Institute Inc., Cary, USA]). See Rossi *et al.* (2007, 2008) for more details on the calculation of temperature thresholds and model verification. The model was fitted with the minimum, mean, and maximum temperatures for each tree, site and year. None of 180 models applied was excluded because of a lack of fit (in all cases  $R^2 > 0.90$ ). Thermal thresholds were then compared between years using an ANOVA.

167 Two-dimensional reversed Gaussian models were used to calculate the difference 168 between the onset of xylogenesis and the day with a given temperature and precipitation within a certain time window. The Gaussian model generates afunnel-surface plot, with a circular-to-elliptical cross-section with the general form:

171 
$$Z_{xy} = Z_{\theta} - Aexp(-\frac{1}{2}(\frac{xcos\theta + ysin\theta - x_ccos\theta - y_csin\theta}{W_1})^2 - \frac{1}{2}(\frac{-xsin\theta + ycos\theta + x_csin\theta - y_ccos\theta}{W_2})^2)$$

where  $Z_{xy}$  is the mean absolute difference between the day of onset of xylogenesis and 172 the estimated day with a given average temperature x and cumulative precipitation y173 within the time window t across trees, sites and years,  $Z_0$  is the distance from the edge 174 of the surface to the plane z = 0, A is the height of the trough,  $x_0$  and  $y_0$  are the 175 coordinates defining the position of the center of the surface,  $W_1$  and  $W_2$  are the 176 spreads of the surface on the x- and y-axes, respectively, and  $\theta$  is the clockwise 177 rotation angle of the surface (see Supporting Information, Fig. S2). The model was 178 fitted with the corresponding temperature (minimum, mean and maximum air 179 temperatures) and precipitation series for each time window. The culmination of the 180 coefficient of determination  $(R^2)$  of the model was considered to correspond to the 181 optimal time window t. The critical average temperature (x) and cumulative 182 precipitation (y) were calculated when  $Z_{xy}$  was near 0 at the optimal time window t. 183 Standardized residuals were calculated for model verification. Model validation was 184 performed by comparing the observations (onset of xylogenesis) with the predicted 185 values calculated using data for precipitation and temperature as predictors. 186

187

188 **Results** 

#### 189 Spring meteorological conditions

190 The daily mean temperatures in March at the upper and lower sites were -4.7 and

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-3.2 °C, respectively, reaching 1.8 and 3.4 °C in May. Monthly precipitation, on
average, increased tenfold, from 5–6 mm in March to 50–60 mm in May (Supporting
Information, Fig. S3).

Spring (March to May) conditions varied between years (Fig. S3). The warmest spring during the study was in 2009, with daily mean temperatures reaching 1.4 and -0.2 °C at the lower and upper sites, respectively. The coldest and driest springs were in 2014, with mean temperatures of -0.2 and -1.8 °C and total precipitation of 25.8 and 31.0 mm at the lower and upper sites, respectively. Monthly precipitation in March 2014 ranged between 17.4 and 21.8 mm, which represented the highest amount of spring rain during the study period.

201

#### 202 Threshold temperatures

The threshold temperature with a probability of 0.5 for active xylogenesis was 203 calculated as an average for each year and site (Table 1). Thermal thresholds at the 204 lower site varied within large ranges, 0-5, 4-9 and 10-14 °C for the daily minimum, 205 mean and maximum temperatures, respectively. Thresholds were significantly higher 206 in 2010 than in other years and were lowest in 2014 (P<0.001). The thermal 207 thresholds were lower at the upper site, but also with large ranges, 0-5, 3-8 and 208 8-12 °C for the daily minimum, mean and maximum temperatures, respectively. The 209 thresholds at the upper site also differed significantly between years (P < 0.001). 210 Thresholds were significantly higher in 2010 than the other years and were lower in 211 2014 for the daily minimum and mean temperatures and in 2012 and 2014 for the 212

213 maximum temperature.

**Table 1** Threshold minimum, mean and maximum temperatures corresponding to

215 95% probability of active xylogenesis in *Juniperus przewalskii* estimated during

216 2009-2014 at the lower and upper study sites. Results from an ANOVA are reported as

217 *F* and *P*. Different letters within a row indicate significant differences at P < 0.05.

Site	Temperat ure (°C)	2009	2010	2011	2012	2013	2014	F	Р
	Minimum	$2.5\pm0.4^{\rm a}$	$5.0\pm0.7^{b}$	$2.4\pm0.7^{\rm a}$	$1.8\pm0.4^{\rm a}$	$2.0\pm0.9^{a}$	$0.0\pm0.7^{\rm c}$	29.93	< 0.001
Lower	Mean	$7.2\pm0.4^{\rm a}$	$9.3\pm0.8^{b}$	$7.5\pm0.7^{\rm a}$	$6.0\pm0.4^{a}$	$6.3\pm0.9^{a}$	$4.5\pm0.7^{\rm c}$	26.33	< 0.001
	Maximum	$12.0\pm0.5^{\rm a}$	$14.2\pm0.9^{\text{b}}$	$12.0 \pm 1.0^{\mathrm{a}}$	$10.0\pm0.4^{\rm c}$	$11.3\pm0.9^{\rm a,c}$	$9.8\pm0.8^{\rm c}$	21.80	< 0.001
	Minimum	$1.9\pm0.2^{a,b}$	$4.5\pm0.3^{\rm c}$	$2.4\pm0.4^{\rm a}$	$1.0\pm0.6^{b}$	$1.2\pm0.9^{\text{b}}$	$\textbf{-0.5} \pm 0.4^{d}$	53.83	< 0.001
Upper	Mean	$5.9\pm0.2^{a,b}$	$8.0\pm0.3^{\rm c}$	$6.4\pm0.3^{a}$	$4.5\pm0.6^{\text{d}}$	$5.0\pm0.9^{\text{b,d}}$	$3.4\pm0.4^{e}$	52.06	< 0.001
	Maximum	$10.1\pm0.2^{a,b}$	$12.2\pm0.3^{\rm c}$	$10.8\pm0.5^{a}$	$7.9\pm0.6^{\text{d}}$	$9.4\pm0.9^{b}$	$8.1\pm0.4^{d}$	51.31	< 0.001

218

#### 219 *Two-dimensional Gaussian models*

220  $R^2$  of the Gaussian models varied with the length of the time window (Fig. 1).  $R^2$ 221 increased for longer time windows, culminating with a time window of 12 days when 222  $R^2$  reached 0.97, 0.99 and 0.94 for the minimum, mean and maximum temperatures, 223 respectively.  $R^2$  decreased slightly (minimum and mean temperature) or substantially 224 (maximum temperature) for time windows longer than 12 days.



Fig. 1 Coefficient of determination ( $R^2$ ) for the two-dimensional Gaussian models within the time window from 1 to 18 days. Dotted lines indicate the time windows (in days) corresponding to maximum  $R^2$ .

230 The minimal  $Z_{xy}$  was 2.21 days for a time window of 12 days. The critical cumulative precipitation was  $17.0 \pm 5.6$  mm and the average minimum temperature 231 was  $1.5 \pm 1.4$  °C when  $Z_{xy}$  was <2.5 days (Fig. 2). The spreads of this trough on the x-232 233 and y-axes were 48.0 mm and 2.04 °C, respectively, with a counter-clockwise rotation of 4.73°. In the model with the average mean temperature, the minimal  $Z_{xy}$  was 1.85 234 days. The critical precipitation and temperature were  $26.9 \pm 3.9$  mm and  $4.6 \pm 1.8$  °C, 235 respectively when  $Z_{xy}$  was <2 days. The spreads of this trough on the x- and y-axes 236 were 44.1 mm and 1.94 °C respectively, with a counter-clockwise rotation of 8.22°. 237 The minimal  $Z_{xy}$  was 1.90 days in the model with the average maximum temperature. 238 The critical precipitation and temperature were 29.9  $\pm$  3.0 mm and 8.5  $\pm$  1.8 °C, 239 respectively, when  $Z_{xy}$  was <2 days. The spreads of this trough on the x- and y-axes 240 were 51.1 mm and 2.39 °C, respectively, with a counter-clockwise rotation of 9.52°. 241 Most of the standardized residuals of these three models converged from -2 to 2 242 (Supporting Information, Fig. S4). 243





Fig. 2 Surface plots and the corresponding level sets showing the two-dimensional
Gaussian distribution of the absolute difference between the day of onset of
xylogenesis and the estimated day with a given average temperature and cumulative
precipitation in the time window of 12 days. Note that the axes have different scales.

The absolute differences between the observed and predicted dates of onset of xylogenesis using average minimum temperature and cumulative precipitation in a time window of 12 days were smaller than the sampling interval by averages of 5.9 and 4.6 days at the lower and upper sites, respectively (Fig. 3). The predictions for 2009, 2011 and 2013 were the most reliable. The divergences between observations and predictions (16 days) were largest in 2014 at the lower site.



Fig. 3 Observed and predicted days of onset of xylogenesis (DOY, day of the year) in *Juniperus przewalskii* during 2009-2014 at the lower and upper study sites.
Predictions were obtained using the threshold average minimum temperature and
cumulative precipitation calculated by the models.

# 262 **Discussion**

263 We challenged the general opinion that temperature was the only driver of growth reactivation at high elevations by analyzing the onset of xylogenesis of Qilian juniper 264 subjected to cold and dry climatic conditions on the northeastern Tibetan Plateau. 265 Published threshold temperatures for the onset of xylogenesis in trees range from 2 to 266 3 °C (Rossi et al., 2007, 2008; Swidrak et al., 2011; Boulouf Lugo et al., 2012). The 267 range in the thermal thresholds of 5 °C for the onset of xylogenesis in Qilian juniper 268 provides additional evidence that temperature was not the only factor initiating xylem 269 growth under cold and dry climatic conditions. More reliable predictions were 270 attained when both thermal and precipitation thresholds for the onset of xylogenesis 271 were included in the fitted models. The interaction between temperature and 272 precipitation satisfactorily explained the day of onset of xylogenesis in 2010, which 273

was delayed by ca. three weeks compared with 2009 and 2011, despite the warm
conditions during that spring (Ren *et al.*, 2015). This finding suggests that spring
precipitation is also an important factor in the resumption of xylem formation in
Qilian juniper.

Water availability is an important determinant of xylem formation. Before the start 278 of xylem phenology, trees must compensate for the water lost during winter and 279 spring to recover an adequate water balance, because turgor is an important requisite 280 for xylem cell growth (Sevanto et al., 2006). Rehydration in spring can exceed six 281 282 weeks, and stems are fully rehydrated one month before the onset of radial growth (Turcotte et al., 2009). Both cell division and expansion in the xylem are sensitive to 283 changes in water potential (Abe & Nakai, 1999; Savidge, 2001). The water potential 284 285 in the cambium regulates mitosis and influences cell extension and the deposition of wall polymers (Abe & Nakai, 1999; Cosgrove, 2005; Arend & Fromm, 2007). Springs 286 were rainy or water was abundantly supplied by snowmelt in the cold regions of 287 288 previous studies, so the initiation of xylem growth was not limited by rehydration, and trees responded essentially to temperature rather than precipitation (Turcotte et al., 289 2009). Warmer springs in such areas can substantially advance xylem phenology 290 (Rossi et al., 2011). Winter and spring are similarly often wet in cold and 291 drought-prone regions such as continental Mediterranean forests, and moisture is not 292 considered the only factor in the resumption of xylem formation (Camarero et al., 293 2010). Winter is extremely dry in our study area, with scarce snow, and water 294 availability is consequently low before growth reactivation. Moreover, drought stress 295

would be higher under drier and warmer conditions, which would thus slow the onset 296 of xylogenesis, as observed in spring 2010. Soil moisture occasionally can be 297 increased by snowfall, such as the snowfall in 2014, as also occurs in boreal forests 298 (Vaganov et al., 1999). The amount of water available during the snowmelt in our 299 study increased soil moisture and possibly advanced the onset of xylogenesis, likely 300 explaining the difference of 16 days between observations and predictions in 2014 at 301 the lower site. This research found that the onset of xylogenesis in Qilian juniper 302 should meet the prerequisite for both critical temperature and precipitation. 303

304 This study is the first to demonstrate that the onset of xylogenesis is driven by an interaction between thermal and precipitation thresholds. The selected time window 305 of 12 days agrees with the period required for tracheid expansion and differentiation 306 307 (Vaganov et al., 2006; Cuny et al., 2015). Temperature is a well-recognized factor controlling the onset of xylem formation, but our findings provide new insights on the 308 climatic forcing of growth. The critical temperatures and precipitation provide keys 309 310 for modelling the response of forest ecosystems subjected to cold and dry constraints in response to climate change and would help our understanding of the regime shifts 311 in these ecosystems (Scheffer et al., 2001; Zhu et al., 2014). Our findings also support 312 the constraint of growth by drought stress in high-elevation forests or near the alpine 313 treeline, as indicated by previous studies (Liang et al., 2014; Piper et al., 2016). 314

Trees in semi-arid areas are generally limited by drought and high temperature at the beginning of the growing season, which increase rates of evapotranspiration (Allen *et al.*, 2015), and a similar constraint has been reported for the Tibetan Plateau

and other Asian mountains (Shao et al., 2005; Liang et al., 2006, 2016; Liu et al., 318 2006; Gou et al., 2014; Pederson et al., 2014; Yang et al., 2014; Zhang et al., 2015). 319 Warming-induced drought stress has been decreasing generalized tree growth and 320 increasing mortality in semi-arid areas across Asia (Dulamsuren et al., 2010; Liu et 321 al., 2013; Liang et al., 2016; Allen et al., 2015). In particular, the failure to produce 322 stem wood in a particular year (missing rings) is a response to dry and warm spring 323 conditions, and an increasing frequency of missing tree rings is also evident in 324 response to the warming in recent decades (Liang et al., 2014, 2016). Moreover, the 325 326 frequency of missing rings has been strongly linked to tree mortality (Liang et al., 2016). We hypothesize that a failure to reach critical water availability for growth 327 reactivation or a delay in cambial resumption in response to increasing drought stress 328 329 could be primary factors in the failure to form a complete ring and portend lower growth and forest dieback. A mismatch between critical temperatures and amounts of 330 moisture for the onset of xylogenesis under the drought conditions of global climate 331 332 change and the acceleration of dryland expansion (Peñuelas et al., 2007; Allen et al., 2015; Huang et al., 2015) will reduce forest resilience and risk regime shifts in 333 vulnerable semi-arid forests. Reyer et al. (2015) proposed the assessment of forest 334 resilience and potential tipping points at various levels, from leaf to biosphere, and 335 our study has stressed that climatic thresholds for the onset of xylogenesis can be key 336 indicators of forest resilience and tipping points under changing climates. 337

338

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517	Supporting Information captions
518	Fig. S1 Correlations between the daily minimum, mean and maximum temperatures
519	recorded at the Dulan meteorological station and the corresponding temperatures
520	recorded during 2012-2014 at the lower and upper study sites. ***, correlation
521	coefficients ( $r$ ) at $P < 0.001$ .
522	
523	Fig. S2 Sample surface plot and the corresponding level sets of a two-dimensional
524	Gaussian model. In the upper plot, $Z_0$ is the distance from the edge of the surface to
525	the plane ( $z = 0$ ), and A is the height of the trough. In the lower plot, $x_0$ and $y_0$ are the
526	coordinates defining the position of the center of the surface, and $\theta$ is the clockwise
527	rotation angle of the surface.
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529	Fig. S3 Daily air temperature (lines) and precipitation (bars) during 2009-2014 at the
530	lower and upper study sites.
531	
532	Fig. S4 The distribution of standardized residuals in the time window of 12 days as a
533	function of minimum, mean and maximum temperatures.
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Fig. S1 Correlations between the daily minimum, mean and maximum temperatures 541 recorded at the Dulan meteorological station and the corresponding temperatures 542 recorded during 2012-2014 at the lower and upper study sites. \*\*\*, correlation 543 coefficients (r) at P < 0.001. 544



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Fig. S3 Daily air temperature (lines) and precipitation (bars) during 2009-2014 at the

555 lower and upper study sites.





Fig. S4 The distribution of standardized residuals in the time window of 12 days as a 

function of minimum, mean and maximum temperatures.