

Adaptation of temperature optima of vegetation productivity across global biomes

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47 **The global distribution of the optimum air temperature of ecosystem-level gross primary**
48 **productivity (T_{opt}^{eco}) is poorly understood, despite its importance for ecosystem carbon**
49 **uptake under future warming. We provide empirical evidence for the existence of such an**
50 **optimum, using measurements of in situ eddy covariance and satellite-derived proxies, and**
51 **report its global distribution. T_{opt}^{eco} is consistently lower than the physiological optimum**
52 **temperature of leaf-level photosynthesis, which typically exceeds 30 °C. The global average**
53 **T_{opt}^{eco} is estimated to be 23 ± 6 °C, with warmer regions having higher T_{opt}^{eco} values than colder**
54 **regions. In tropical forests, particularly, T_{opt}^{eco} is close to growing-season air temperature and**
55 **is projected to fall below it under all scenarios of future climate, suggesting a limited safe**
56 **operating space for these ecosystems under future warming.**

57
58 Understanding how photosynthesis responds to warming has been a focus in plant research in
59 recent decades, and most of the existing knowledge comes from leaf-scale measurements¹⁻⁴. Most
60 leaf-scale temperature response curves show that photosynthesis increases with temperature up to
61 an optimum temperature (T_{opt}^{leaf}), which typically occurs in the 30-40°C temperature range^{5,6}.
62 Above this optimum temperature, foliar photosynthesis sharply declines as electron-transport and
63 Rubisco enzymatic capacities become impaired⁷. Field et al.⁸ first suggested that ecosystem-scale
64 optimum temperature (T_{opt}^{eco}) may differ from T_{opt}^{leaf} . At ecosystem scale, elevated air
65 temperatures do limit canopy photosynthesis by other processes than leaf carboxylation rates. For
66 instance, elevated air temperatures reduce hydraulic conductivities in the soil-root-stem-canopy
67 continuum (hydraulics; e.g. ref 6), accelerate leaf ageing and increase leaf thickness (phenology;
68 e.g. ref 9), and control stomatal closure, because a higher temperature usually comes with a
69 higher vapor pressure deficit (VPD)¹⁰.

70

71 Empirical leaf-scale photosynthesis-temperature relationships¹¹ have been directly incorporated
72 into global ecosystem models, with variants to account for acclimation, i.e. a temporal adjustment
73 of optimum photosynthetic temperature to air temperature during growth^{5,12,13}. This direct scaling
74 of temperature responses from leaves to ecosystems partly determines model projections of Gross
75 Primary Productivity (GPP) and CO₂ uptake by terrestrial ecosystems in climatic scenarios.
76 Verifying the existence of T_{opt}^{eco} in real world ecosystems, defining its spatial distribution across
77 and within biomes and understanding the relationships between T_{opt}^{eco} , prevailing air temperature
78 and T_{opt}^{leaf} are important for evaluating models and understanding the impacts of various targets of
79 climatic warming targets on ecosystem productivity.

80

81 In this study, we formulate and test the following hypotheses: (i) T_{opt}^{eco} is higher for biomes where
82 air temperature during growth is warmer (adaptation), (ii) T_{opt}^{eco} is lower than T_{opt}^{leaf} for any given
83 ecosystem, because the above mentioned limitations of plant hydraulics and phenology emerge
84 before temperature begins to impair foliar photosynthesis, and (iii) tropical forests already operate
85 near a high T_{opt}^{eco} above which canopy photosynthesis may decrease with even moderate air
86 temperature warming^{14,15}. Here we defined T_{opt}^{eco} as the daytime air temperature at which GPP is
87 highest over a period of several years, thus T_{opt}^{eco} can be empirically determined from productivity
88 observations and proxies (see Methods).

89

90 **Results and discussion**

91 We first applied this approach on time series of daily GPP derived from CO₂ flux measurements
92 at 153 globally distributed eddy covariance sites and found that a robust estimate of T_{opt}^{eco} could

93 be derived at 125 out of 153 sites (see Methods). T_{opt}^{eco} values derived from the FLUXNET data
94 range from 8.2°C to 35.8°C (Fig. 1a, Supplementary Table 1). Tropical sites have higher T_{opt}^{eco}
95 values than temperate and boreal sites (Supplementary Fig. 1), implying a dependence of T_{opt}^{eco} on
96 background climate. The FLUXNET multi-site analysis further indicates that across sites
97 T_{opt}^{eco} values are positively correlated with growing-season mean daily maximum air temperature
98 ($T_{max\ gs}^{air}$, see calculation in Methods) ($R=0.46$, $P<0.01$), with a spatial linear regression slope of
99 0.61 °C per °C across sites (Fig. 1a). Overall, these results confirm our first hypothesis stating
100 that higher T_{opt}^{eco} values occur where higher growth temperatures prevail, in support of findings of
101 Baldocchi et al.¹⁶.

102
103 Since eddy covariance measurements do not have a continuous spatial coverage, we also used
104 satellite observations known to be highly correlated with photosynthetic activity¹⁷, that is, GPP
105 proxies. The first proxy used is the near-infrared reflectance of terrestrial vegetation (NIR_V , the
106 product of total-scene NIR reflectance (NIR_T) by the Normalized Difference Vegetation Index
107 (NDVI). NIR_V was proven to have a high temporal correlation with GPP at flux tower sites¹⁷.
108 Satellite observations of NIR_T and NDVI from the Terra Moderate Resolution Imaging
109 Spectroradiometer (MODIS) were used to calculate NIR_V for the period of 2001-2013 (see
110 Methods). NIR_V -derived T_{opt}^{eco} is comparable with that estimated from eddy covariance flux tower
111 measurements (Fig. 1b), which gives support to using the NIR_V proxy for a global mapping of
112 T_{opt}^{eco} . The average T_{opt}^{eco} over the global vegetated areas is estimated to be 23 ± 6 °C (mean \pm 1 SD)
113 with large spatial gradients in latitude. As shown in Fig. 1c, maximum values close to 30°C
114 mainly appear over tropical forests, savannas and drylands and minimum values near 10°C
115 prevail at high-latitudes and in mountainous regions (Fig. 1c). This spatial pattern of T_{opt}^{eco} is

116 robust to the choice of a particular climate forcing dataset, or to the method used to estimate T_{opt}^{eco}
117 (Supplementary Fig. 2, see also Methods). Similar results are also found for other GPP proxies
118 (vegetation greenness (NDVI)¹⁸, Enhanced Vegetation Index (EVI)¹⁹, sun-induced vegetation
119 fluorescence (Sun-induced Chlorophyll Fluorescence, SIF)²⁰), or when daily mean air
120 temperature (T_{mean}^{air}) is used instead of daily maximum air temperature (T_{max}^{air}) to calculate T_{opt}^{eco}
121 (Supplementary Figs. 3-6, see also Methods). Note that although the covariance between air
122 temperature, atmospheric VPD and solar radiation may confuse the direct effect of air
123 temperature on vegetation productivity, we verified that neither VPD nor radiation is the
124 dominant factor determining the pattern of T_{opt}^{eco} at the global scale (see Methods).

125
126 In order to test the second hypothesis, we compared satellite-derived T_{opt}^{eco} with T_{opt}^{leaf} from the
127 responses of maximum Rubisco-limited carboxylation rates (V_{cmax}) to temperature from leaf-
128 scale measurements for 36 species⁵. We found that T_{opt}^{eco} is indeed lower than T_{opt}^{leaf}
129 (Supplementary Fig. 7), with a larger difference between T_{opt}^{leaf} and T_{opt}^{eco} for needle-leaved than
130 for broad-leaved forests. This difference may originate from the fact that T_{opt}^{eco} is additionally
131 limited by high VPD during hot and dry periods⁶ and by soil-moisture deficits during extensive
132 dry episodes²¹, under real world conditions. In contrast, leaf-level photosynthesis measurements
133 to determine V_{cmax} are never performed on drought-affected leaves. Moreover, tree hydraulic
134 conductance is limited, and may not be able to keep up with the increasing evaporative demands
135 at higher temperatures, especially for needle-leaved species that have a lower stem-water
136 conductivity²². Based on NIR_v-derived results, we did diagnose a lower T_{opt}^{eco} for needle-leaved
137 than for broad-leaf forests under the same growth temperature conditions (Supplementary Table
138 2). In addition, plant phenology controls leaf age, vitality (photosynthetic rates) and foliar density

139 (e.g. Leaf Area Index, LAI)²³, and may therefore co-determine ecosystem-level temperature
140 limitations and the optimum temperature for canopy photosynthesis²⁴. It is also important to
141 remark when comparing T_{opt}^{leaf} with T_{opt}^{eco} that leaf-scale measurements are often limited to sunlit
142 leaves, which could lead to a positive bias of existing in-situ T_{opt}^{leaf} measurements. What's more,
143 the tree species database used by Kattge & Knorr⁵ from which T_{opt}^{leaf} data were collected does not
144 include any tropical species. This may explain why global models prescribed with T_{opt}^{leaf} give very
145 uncertain results for tropical biomes.

146
147 The relationship between T_{opt}^{eco} and background climate is shown in Fig. 1d. The sampling of leaf-
148 scale studies does not provide consistent evidence about the dependence of T_{opt}^{leaf} on climate, with
149 positive correlations between T_{opt}^{leaf} and growing season air temperature in a set of studies^{1, 5, 25-27}
150 attributed to evolutionary adaptation⁷, but no clear relationship between T_{opt}^{leaf} and growing season
151 temperature in the review from Sage & Kubien⁷. In contrast, T_{opt}^{eco} inferred from satellite GPP
152 proxies in our study increases with $T_{max\ gs}^{air}$ across the globe. In temperature-precipitation space,
153 the spatial sensitivity of T_{opt}^{eco} to $T_{max\ gs}^{air}$ (the slope of the linear regression between these two
154 variables) is lower than 1 for any precipitation bin (Fig. 1d), suggesting that spatial gradients of
155 T_{opt}^{eco} are smaller than those of $T_{max\ gs}^{air}$, possibly because hydraulic and phenological limitations
156 further limit T_{opt}^{eco} across spatial gradients. In fact, the spatial sensitivity of T_{opt}^{eco} to
157 $T_{max\ gs}^{air}$ generally increases with increasing mean annual precipitation (Fig. 1d), even though T_{opt}^{eco}
158 is not significantly correlated with precipitation after controlling for the effect of $T_{max\ gs}^{air}$ (Fig.
159 1d). This thermal adaptation of T_{opt}^{eco} , suggested by the positive spatial slope of the T_{opt}^{eco} -air
160 temperature relation, is also observed across biomes. As shown in Fig. 2, there is a significant

161 positive correlation between T_{opt}^{eco} and $T_{max\ gs}^{air}$ with a slope of 0.76 across different biomes.
162 Among biomes, the largest mean T_{opt}^{eco} is found in tropical evergreen broad-leaved forest (29 ± 3
163 $^{\circ}\text{C}$), and the smallest mean T_{opt}^{eco} (13 ± 3 $^{\circ}\text{C}$) in cold grasslands covering the Tibetan Plateau (Fig. 2
164 and Supplementary Fig. 8).

165
166 Both model results and very limited observational studies suggest a decrease in canopy
167 photosynthesis of tropical forests at high temperature^{15, 28-31} which led us to formulate the third
168 hypothesis of tropical forests already operating at T_{opt}^{eco} being close to $T_{max\ gs}^{air}$ implying that
169 canopy photosynthesis may decrease in case of future warming^{14, 15}. This hypothesis is verified
170 from the data shown in Fig. 3 (see also Supplementary Fig. 9). T_{opt}^{eco} is indeed only slightly (1.4
171 $^{\circ}\text{C}$) lower than $T_{max\ gs}^{air}$ over tropical evergreen forests, suggesting a small ‘safety margin’ for
172 photosynthesis under future warming. In contrast, arctic (north of 65°N) and boreal (50°N - 65°N)
173 ecosystems exhibit substantially larger safety margins, i.e. a larger positive difference between
174 T_{opt}^{eco} and $T_{max\ gs}^{air}$ (Fig. 3a and Supplementary Fig. 9). Analysis of the 16-day averaged T_{max}^{air}
175 distribution during the period when T_{opt}^{eco} is observed further shows that the rank of T_{opt}^{eco} in the
176 T_{max}^{air} distribution is already near the highest quantile of T_{max}^{air} ($>70\%$) for tropical evergreen
177 forests (Supplementary Fig. 10). Based on this result, one may expect that rising air temperature
178 in the future may limit or decrease vegetation productivity in tropical forests, but not in temperate
179 or boreal ecosystems.

180
181 Global terrestrial daily maximum air temperature is projected to rise by 1.9 $^{\circ}\text{C}$ under the RCP2.6
182 low warming climate scenario and by 5.6 $^{\circ}\text{C}$ under the RCP8.5 scenario by 2100³². We compared
183 these $T_{max\ gs}^{air}$ projections with the present-day distribution of T_{opt}^{eco} with a focus on tropical

184 evergreen forests, where optimum temperature is currently just below the limit of $T_{max\ gs}^{air}$ (see
185 Methods; Fig. 3b and 3c). The key uncertainty in this discussion is, however, whether or not T_{opt}^{eco}
186 will acclimate and follow the increase in $T_{max\ gs}^{air}$. We therefore looked at possible acclimation
187 from time series of T_{opt}^{eco} retrieved from the Advanced Very High Resolution Radiometer
188 (AVHRR) NDVI, which spans the last 30 years and comprises almost a 1°C temperature range.
189 NDVI-derived T_{opt}^{eco} did not have a significant trend over the last three decades (Supplementary
190 Fig. 11), suggesting that either the recent 1°C warming is not large enough to elicit an
191 acclimation response from ecosystems, given decadal variability³³, or that acclimation does not
192 occur at multi-decadal timescales. Also the annual T_{opt}^{eco} derived from FLUXNET estimates of
193 GPP did not exhibit a positive trend and was not significantly correlated with annual variations of
194 $T_{max\ gs}^{air}$, although the FLUXNET time series are probably too short to properly evaluate trends of
195 T_{opt}^{eco} related to possible acclimation processes (Supplementary Fig. 12). Because we detected no
196 indication for its existence, we first assumed no acclimation in the comparison of future $T_{max\ gs}^{air}$
197 projections from climate models with the current distribution of T_{opt}^{eco} . Under this assumption, the
198 average $T_{max\ gs}^{air}$ of tropical evergreen forests will exceed the current value of T_{opt}^{eco} for RCP2.6 by
199 2.6°C, and by 5.7 °C for RCP8.5 (Fig. 3c). On the other hand, boreal and arctic biomes will still
200 remain within the safety margin, with T_{opt}^{eco} staying above $T_{max\ gs}^{air}$ except under the RCP8.5 high
201 warming scenario (Fig. 3b and Supplementary Fig. 13).

202
203 Despite the lack of in situ observational evidence for GPP acclimation to the ongoing warming
204 trend, we tested a simple future acclimation law based on the space-for-time substitution
205 assumption that T_{opt}^{eco} will evolve proportionally to $T_{max\ gs}^{air}$, following on the spatial temperature

206 sensitivity of T_{opt}^{eco} to $T_{max\ gs}^{air}$ in Fig. 1d. We took the differences in precipitation levels into
207 account, so that areas that become wetter also exhibit faster acclimation. Even with this assumed
208 acclimation law, $T_{max\ gs}^{air}$ will still surpass T_{opt}^{eco} by 1.7 °C under RCP2.6 and by 2.5 °C under
209 RCP8.5 for tropical evergreen forests (Fig. 3c). Not accounting for precipitation levels in the
210 acclimation rates produced similar results (Supplementary Figs. 14 and 15).

211
212 Our global-scale analysis of T_{opt}^{eco} derived from globally distributed point measurements of eddy
213 covariance and space-borne observations of proxies of vegetation productivity is a first attempt to
214 diagnose the global distribution of ecosystem-scale temperature optima of photosynthesis. It
215 should be noted, however, that hypotheses about that thermal acclimation of T_{opt}^{eco} are still highly
216 uncertain, because ecosystem adjustments can lag substantially behind the rate of future warming,
217 particularly for forests. The acclimation of plants to increasing atmospheric CO₂ concentration
218 and to changes in other environmental factors (e.g. VPD) was also not considered in the current
219 analyses. Constraining the spatially observed temperature sensitivity of T_{opt}^{eco} over time is a
220 priority for future studies. Continuous monitoring and dedicated manipulative experiments could
221 improve our understanding of the features of T_{opt}^{eco} and thermal acclimation in earth system
222 models³⁴.

223

224 **Methods**

225 **FLUXNET data** The half-hourly eddy-covariance Gross Primary Productivity (GPP) data were
226 obtained from FLUXNET datasets, and were quality-controlled, filtered against low turbulence,
227 and gap-filled using consistent methods, as described by Papale et al.³⁵. Only freely available
228 FLUXNET data were used in this study. All the half-hourly GPP data were aggregated into daily

229 accumulated GPP for further estimates of the optimal temperature for vegetation productivity. We
230 included only site-years with more than 80% of half-hourly data available. A total of 153
231 individual FLUXNET sites with 663 site-years of GPP data were used in this study.

232
233 **Near-infrared reflectance of terrestrial vegetation** Badgley et al.¹⁷ have recently proposed a
234 new approach for estimating vegetation photosynthetic capacity by remote sensing, i.e. the near-
235 infrared reflectance (NIR) of terrestrial vegetation (NIR_V), which can differentiate between the
236 confounding effects of background brightness, leaf area and the distribution of photosynthetic
237 capacity with depth in canopies¹⁷. NIR_V is calculated as the product of total scene NIR
238 reflectance (NIR_T) and Normalized Difference Vegetation Index (NDVI)¹⁷. As a proxy of
239 photosynthesis, NIR_V is suggested to be strongly correlated with Solar-induced Chlorophyll
240 Fluorescence (SIF), a direct index of photons intercepted by chlorophyll, and shows higher
241 correlation with observed GPP than NDVI¹⁷. We used satellite-derived NIR_V to calculate and
242 map the optimal air temperature for vegetation productivity at an ecosystem scale (T_{opt}^{eco}).
243 Following Badgley et al.¹⁷, we calculated 16-day NIR_V for 2001-2013 as the product of Moderate
244 Resolution Imaging Spectroradiometer (MODIS) 16-day NIR reflectance and MODIS 16-day
245 NDVI, both of which were derived from the MOD13A2 Vegetation Index Product with a spatial
246 resolution of 1 km. Only positive NIR_V values were used in the analysis. Pixels with a mean
247 annual NDVI>0.1 were defined as the vegetated area.

248
249 **Normalized Difference Vegetation Index** The Normalized Difference Vegetation Index (NDVI)
250 is a vegetation index defined as the ratio of the difference between NIR and red visible
251 reflectance to their sum, and is widely used to represent vegetation greenness³⁶. To account for
252 uncertainties from different satellite datasets, three independent NDVI datasets were utilized,

253 including Global Inventory Modeling and Mapping Studies (GIMMS) Advanced Very High
254 Resolution Radiometer (AVHRR) biweekly NDVI, Terra Moderate Resolution Imaging
255 Spectroradiometer (MODIS) 16-day NDVI and Satellite Pour l'Observation de la Terre
256 Vegetation (SPOT Vegetation) 10-day NDVI. The three NDVI datasets spanned the last three
257 decades: 1982-2009 for AVHRR NDVI datasets, 2000-2009 for MODIS NDVI datasets, and
258 1999-2009 for SPOT NDVI datasets, with the spatial resolutions of 8 km, 1 km, and 1 km,
259 respectively. All NDVI datasets have been corrected to reduce the effects of volcanic aerosols,
260 solar angle, and sensor errors^{18,37,38}. Pixels with a mean annual NDVI>0.1 were defined as the
261 vegetated area for each dataset.

262
263 **Enhanced Vegetation Index** The Enhanced Vegetation Index (EVI) is another vegetation index
264 designed to enhance the vegetation signal by minimizing canopy-soil variations and to improve
265 sensitivity over dense vegetation conditions¹⁹, and is found to correlate well with estimated GPP
266 on a site-by-site basis³⁹. We used a 16-day EVI dataset for the period of 2000-2009 with a spatial
267 resolution of 1 km from the MOD12A1 Vegetation Index Product. Effects from aerosols, solar
268 angle and sensor error have all been corrected¹⁹.

269
270 **Sun-induced Chlorophyll Fluorescence** Chlorophylls in plant would absorb shortwave radiation
271 and dissipate the excess energy as light or heat. The longwave radiation re-emitted by
272 chlorophylls is referred as chlorophyll fluorescence. Recent studies have reported that remotely
273 sensed Sun-induced Chlorophyll Fluorescence (SIF) could serve as an indicator of photosynthesis
274 rate and it is well correlated with model-simulated GPP⁴⁰. Following the previous studies^{40,41}, we
275 retrieved SIF from two different retrieval windows, 757 nm and 771 nm, as well as the two
276 polarization states, S and P using observation of Fourier Transform Spectrometer (FTS) on the

277 Japanese Greenhouse gases Observing SATellite (GOSAT)²⁰. These diverse SIF samples of were
278 then aggregated into monthly gridded data at a spatial resolution of 2° from June 2009 to June
279 2012.

280
281 **Vegetation distribution** We used MODIS land cover with the classification scheme of the
282 International Geosphere-Biosphere Programme (IGBP). The MODIS IGBP land cover data were
283 derived from the MOD12Q1 Land Cover Science Data Product at a spatial resolution of 1 km and
284 an updated digital Köppen-Geiger world map of climatic classification⁴². Within the vegetated
285 area defined by NDVI thresholds, the 17 land cover types were reclassified into 9 vegetation
286 types, namely evergreen needle-leaved forest (ENF), evergreen broad-leaved forest (EBF),
287 deciduous needle-leaved forest (DNF), deciduous broad-leaved forest (DBF), mixed forest (MF),
288 savannas, cropland, grassland and shrubland. Based on the main climates in the world Map of the
289 Köppen-Geiger climatic classification⁴², grassland was further subdivided into temperate
290 grasslands, boreal and arctic tundra, and shrubland was further subdivided into temperate and
291 boreal shrubland. The grassland over the Tibetan Plateau was considered separately.

292
293 **Climate dataset** The gridded air temperature and precipitation data for the period of 1982-2013
294 were obtained from the CRU/NCEP 6-hourly dataset with a spatial resolution of 0.5°. It should be
295 noted that near-surface air temperature (the thermodynamic temperature of the air at a height of 2
296 m) rather than land surface temperature (the temperature of underlying canopy surface) were used
297 to calculate T_{opt}^{eco} considering the fact that for tropical regions, especially tropical forests, daytime
298 surface temperature can be much lower than daytime air temperature as a result of the
299 evapotranspiration effects⁴³. Furthermore, the climate forcing data of global ecosystem models
300 are generally air temperature instead of surface temperature (e.g. ref 44, 45). Corresponding to

301 the temporal resolutions of MODIS, AVHRR and SPOT datasets, respectively, the 6-hourly
302 climate data were aggregated into 16-day, biweekly, and 10-day values before further analyses.
303 Given the different spatial resolutions of satellite observations and climate data, we simply
304 extracted time series of daily maximum air temperature and precipitation from the aggregated
305 CRU/NCEP data for each pixel of the sets of remotely sensed data. The daily maximum air
306 temperature (T_{max}^{air}) of the growing season averaged over 2001-2013 was calculated as the current
307 mean growing-season daily maximum air temperature ($T_{max\ gs}^{air}$). Information on the growing
308 season was derived from the study by Zhu et al.⁴⁶, which was first determined from the GIMMS
309 Leaf Area Index data set (GIMMS LAI_{3g}) using a Savitzky-Golay filter and then refined by
310 excluding the ground-freeze period identified by the Freeze/Thaw Earth System Data Record (see
311 details in ref 46). We also documented the temperature thresholds at which the growing season
312 begins and ends for each year. Temperature thresholds were averaged over 2001-2013 for the
313 onset and end of the growing season, respectively. We also used WATCH Forcing Data
314 Methodology to ERA-Interim data with a temporal resolution of three hours (WFDEI)⁴⁷.

315
316 We also used climate projections for the end of the 21st century (2091-2100) under the RCP2.6,
317 RCP4.5 and RCP.85 scenarios³² to determine the impact of future warming on vegetation
318 productivity. Considering the mismatch between CRU/NCEP datasets and outputs from General
319 Circulation Models (GCM) for current climate conditions, we generated future temperature and
320 precipitation data by adding the relative changes in GCM-derived climate projections to the
321 current climate. $T_{max\ gs}^{air}$ for the late 21st century was estimated using the same temperature
322 thresholds as for the current $T_{max\ gs}^{air}$. All GCM projections were resampled to a resolution of 1°
323 using first-order conservative interpolation method⁴⁸.

324

325 **Analysis** We estimated local T_{opt}^{eco} by examining the temperature response curve of MODIS
326 NIR_V . Following Yuan et al.²⁷ and Niu et al.⁴⁹, NIR_V time series throughout the entire monitoring
327 period and the corresponding temperature data were grouped into 1°C-temperature bins for each
328 pixel within vegetated areas. We used the 90% quantile of the NIR_V data as the response of NIR_V
329 within each temperature bin due to the potential influences of other environmental constraints like
330 clouds and droughts. We next calculated the running means of every three temperature bins to
331 develop the temperature response curve of NIR_V . The T_{opt}^{eco} was then determined from the
332 response curve at which NIR_V was maximized (Supplementary Fig. 16). Note that T_{opt}^{eco} may not
333 be detected for some pixels where the maximum NIR_V was only attained at either end of the
334 response curve. The derivation of T_{opt}^{eco} is robust to the choice of a particular climate-forcing
335 dataset (Supplementary Fig. 2). Instead of using the temperature corresponding to the maximum
336 90th quantile NIR_V to calculate T_{opt}^{eco} , we also applied nonlinear regression of the photosynthetic
337 temperature response data (Eq.1) to estimate T_{opt}^{eco} , which produced similar results
338 (Supplementary Fig. 2):

$$339 \quad NIR_{V(T)} = NIR_{V(OPT)} - b(T - T_{opt}^{eco})^2 \quad (\text{Eq.1})$$

340 where $NIR_{V(T)}$ is the NIR_V value at a daily maximum temperature T, b is a parameter describing
341 the spread of the parabola^{33,50}. T_{opt}^{eco} is the vertex of each fit and $NIR_{V(OPT)}$ is the NIR_V value at
342 T_{opt}^{eco} . Finally, we used daily mean air temperature (T_{mean}^{air}) instead of T_{max}^{air} to calculate T_{opt}^{eco} . In
343 this test, T_{opt}^{eco} derived from T_{mean}^{air} is smaller than T_{opt}^{eco} estimated from T_{max}^{air} , but the two variables
344 were strongly spatially correlated (Supplementary Fig. 6).

345

346 We investigated the relationship between T_{opt}^{eco} and climate variables by averaging T_{opt}^{eco} in the
347 climate space with 1-°C intervals of mean annual T_{max}^{air} averaged over the growing season
348 ($T_{max\ gs}^{air}$) and 100-mm intervals of mean annual precipitation (MAP) (Fig. 1d). For each MAP
349 interval, we calculated the ‘apparent’ spatial sensitivity of T_{opt}^{eco} in response to changes in $T_{max\ gs}^{air}$
350 using bootstrapping method. We performed the linear regression analysis 1000 times by
351 randomly selecting a subset of 80% of the samples from pairs of T_{opt}^{eco} and $T_{max\ gs}^{air}$ within each
352 MAP interval. The mean and SD of the 1000 temperature sensitivity of T_{opt}^{eco} were subsequently
353 estimated along the MAP gradient.

354

355 Air temperature, atmospheric vapor pressure deficit (VPD) and solar radiation usually co-vary in
356 time and space, so that the empirical observation of spatial patterns of T_{opt}^{eco} in this study cannot
357 be unambiguously attributed to air temperature as a single explaining factor of T_{opt}^{eco} . Under
358 conditions of high temperature, atmospheric VPD increases, soil moisture decreases with a lag,
359 stomatal conductance, xylem conductance and hence carbon assimilation rates (GPP at the
360 ecosystem-scale) decrease to prevent exceedingly low leaf water potentials and resulting plant
361 tissue damage from cavitation⁵¹. We show that across climatic gradients T_{opt}^{eco} is systematically
362 higher at high maximum air temperatures but not systematically lower at high VPD conditions
363 (Supplementary Fig. 17). To examine whether VPD can substantially affect the relationship
364 between T_{opt}^{eco} and $T_{max\ gs}^{air}$, we further calculated the partial (‘intrinsic’) sensitivity of T_{opt}^{eco} to
365 $T_{max\ gs}^{air}$ in each grid point based on the following bilinear regression:

366
$$T_{opt}^{eco} = k_0 + k_1 \times T_{max\ gs}^{air} + k_2 \times VPD + k_3 \times VPD \times T_{max\ gs}^{air} \text{ (Eq.2)}$$

367 where the partial sensitivity of T_{opt}^{eco} to $T_{max\ gs}^{air}$ is defined as k_1 in Eq. 2 under each VPD bin.
368 Then we compared the partial sensitivity with the apparent sensitivity of T_{opt}^{eco} to $T_{max\ gs}^{air}$
369 estimated using abovementioned linear regression between T_{opt}^{eco} and $T_{max\ gs}^{air}$ for each VPD bin.
370 As shown in Supplementary Fig. 18, although the apparent sensitivity of T_{opt}^{eco} to $T_{max\ gs}^{air}$ is
371 generally lower than the partial ('intrinsic') sensitivity of T_{opt}^{eco} to $T_{max\ gs}^{air}$ the apparent sensitivity
372 to $T_{max\ gs}^{air}$ remains positive even when VPD is taken into account excepted under very high VPD
373 bins (higher than ~4.5 kPa) representing less than 1% of the study area. These results suggest that
374 the patterns of T_{opt}^{eco} are not dominated by high VPD reducing photosynthesis, as an indirect effect
375 of higher air temperature increasing VPD. Moreover, we also calculated the percentiles of
376 downward shortwave solar radiation (Rad) at the time of year when T_{opt}^{eco} is observed for the 16-
377 day averaged Rad distribution. As shown in Supplementary Fig. 19, the Rad value when T_{opt}^{eco}
378 was retrieved from global observations were below the 95th percentile in the 16-day Rad
379 distribution for ~80% of the study area, which is mainly in mid and low latitudes such as Africa,
380 India, Australia, eastern Brazil and southern and southwestern of North America. By comparison,
381 for most part of boreal regions, part of south China, southeastern US, as well as part of South
382 America, the timing of T_{opt}^{eco} is consistent the time of maximum solar radiation. This is because
383 T_{opt}^{eco} in these regions generally appears in summer, which is also the period when solar radiation
384 displays its maximum during the year.

385
386 The NIR_v-derived T_{opt}^{eco} was compared with T_{opt}^{eco} estimated using GPP data from 153 eddy
387 covariance sites. For each site, we extracted and averaged T_{opt}^{eco} values within a 3×3 pixel window
388 around the site from the NIR_v-derived T_{opt}^{eco} map, and calculated the SD of the nine T_{opt}^{eco} values

389 within the window. T_{opt}^{eco} was determined for each site-year with daily-accumulated GPP and
390 corresponding temperature data from flux tower observations, applying the same method to
391 estimate local T_{opt}^{eco} using NIR_V datasets. A robust estimate of T_{opt}^{eco} can be derived for 125 sites
392 (Supplementary Table 1). For each site, we calculated the mean and SD of T_{opt}^{eco} across different
393 years. The results show that NIR_V-derived T_{opt}^{eco} is comparable with that estimated independently
394 from measurements of flux-tower eddy covariance (Fig. 1b).

395
396 We compared the spatial distribution of T_{opt}^{eco} derived from NIR_V with the one obtained from
397 NDVI datasets. Consistent spatial patterns of T_{opt}^{eco} are derived from each of the three NDVI
398 datasets (Supplementary Fig. 20). A global composite map of T_{opt}^{eco} (Supplementary Fig. 3) was
399 then generated by averaging over estimates derived from the three NDVI datasets. Given the
400 inconsistent spatial resolutions of the different products, we resampled T_{opt}^{eco} to a common grid of
401 8 km before averaging. T_{opt}^{eco} from NDVI datasets generally show a spatial pattern similar to that
402 from NIR_V, but with smaller NDVI-derived T_{opt}^{eco} values for central Australia and southern South
403 America (Supplementary Fig. 3). We compared the spatial distribution of T_{opt}^{eco} derived from
404 NIR_V with that from MODIS EVI data during 2001-2013, and found that the EVI derived T_{opt}^{eco}
405 shows very similar spatial pattern to that of NIR_V derived T_{opt}^{eco} (Supplementary Fig. 4). The
406 distribution of T_{opt}^{eco} derived from NIR_V and from GOSAT SIF datasets also have similar spatial
407 patterns, even though the NIR_V-derived T_{opt}^{eco} is higher in tropical regions, particularly in
408 cultivated areas of southeastern Brazil (Supplementary Fig. 5).

409

410 The NIR_V-derived T_{opt}^{eco} was compared with leaf-scale optimal temperature of maximum Rubisco-
411 limited carboxylation rates (V_{cmax}) (T_{opt}^{leaf}) from leaf-scale measurements⁵. The compilation of
412 data from Kattge and Knorr⁵ contains 48 available measurements of T_{opt}^{leaf} values for 36 species.
413 For T_{opt}^{leaf} measurements with site location (longitude and latitude) reported, we extracted and
414 averaged T_{opt}^{eco} values within a 3×3 pixel window around the site from the NIR_V-derived T_{opt}^{eco}
415 map. For T_{opt}^{leaf} measurements without the information of site location, we calculated the average
416 NIR_V-derived T_{opt}^{eco} values across pixels with both the same growing season mean temperature
417 and the same plant functional type as the corresponding site. As shown in Supplementary Fig. 7,
418 T_{opt}^{eco} is generally lower than T_{opt}^{leaf} .

419
420 To account for potential changes in T_{opt}^{eco} under future warming, we estimated the acclimated T_{opt}^{eco}
421 for vegetation productivity by the end of the 21st century (2091-2100), using recent IPCC climate
422 projections³². Given the relatively large uncertainties of precipitation projections, we considered
423 two future precipitation scenarios. For the first scenario, we estimated acclimated T_{opt}^{eco} pixel by
424 pixel using the temperature sensitivity of T_{opt}^{eco} under the present MAP level, assuming that MAP
425 does not change between the end of the 21st century. For the second scenario, we accounted for in
426 MAP, and the acclimated T_{opt}^{eco} was calculated pixel by pixel using the temperature sensitivity of
427 T_{opt}^{eco} under the projected MAP level for 2091-2100. Acclimated T_{opt}^{eco} was averaged across the
428 GCMs under each scenario.

429

430 **Data Availability**

431 All data is available in the main text or the supplementary information. All computer codes used
432 in this study can be provided by the corresponding author upon reasonable requests.

433

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563

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582

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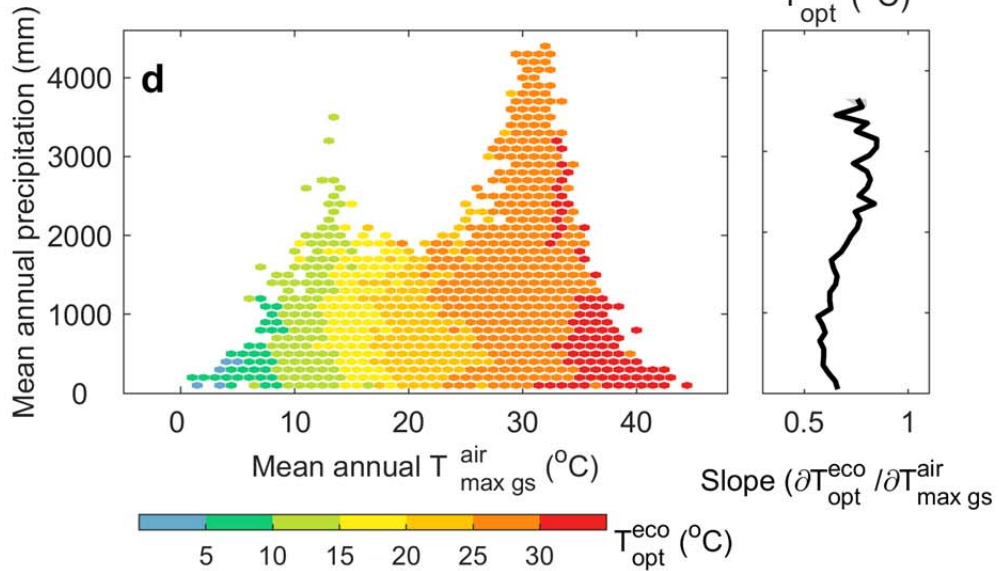
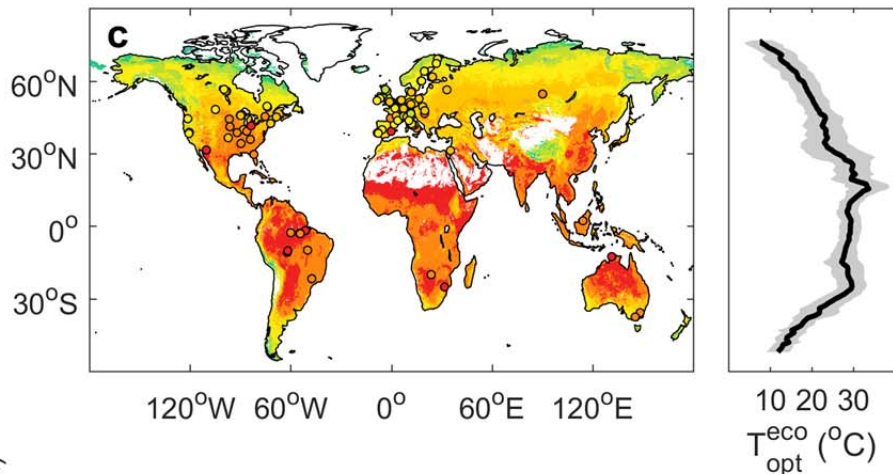
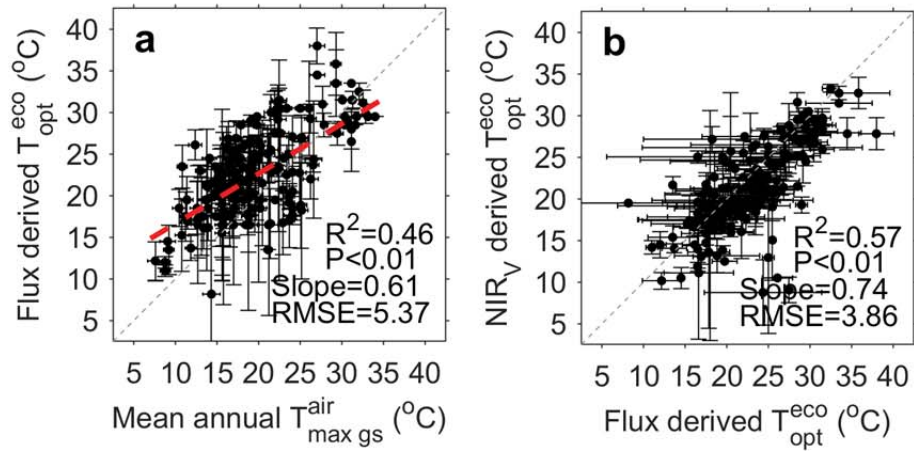
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586 **Competing interests:** Authors declare no competing interests.

587

588 **Fig. 1 | Distribution of ecosystem-scale optimal temperature (T_{opt}^{eco}) for vegetation**
589 **productivity derived from FLUXNET and satellite-based data for near-infrared reflectance**
590 **of vegetation (NIR_V).** **a,** Relationship between mean annual daily maximum air temperature
591 during the growing season ($T_{max\ gs}^{air}$) and T_{opt}^{eco} derived from daily measurements of
592 photosynthesis across FLUXNET eddy-covariance sites. The dotted gray line represents $y=x$. **b,**
593 Relationship between T_{opt}^{eco} derived from FLUXNET data and T_{opt}^{eco} derived from NIR_V data. For
594 each site, we extracted and averaged T_{opt}^{eco} values within a 3×3 pixel window around the site from
595 NIR_V-derived T_{opt}^{eco} map, and calculated the SD of the nine T_{opt}^{eco} values within the window. The
596 dotted gray line represents $y=x$. **c,** Spatial distribution of T_{opt}^{eco} for vegetation productivity (left
597 panel), and T_{opt}^{eco} averaged by latitude (right panel). T_{opt}^{eco} is determined using NIR_V data calculated
598 based on satellite observations from Moderate Resolution Imaging Spectroradiometer (MODIS).
599 Areas of tropical forests based on current vegetation distribution are indicated by hatching. The
600 circles on the map are colored according to the local value of T_{opt}^{eco} retrieved from GPP at the
601 location of each FLUXNET site. The solid line and shaded area in the right panel indicate the
602 mean and SD, respectively, of T_{opt}^{eco} summarized by latitude. **d,** T_{opt}^{eco} in the climate space (left
603 panel) and the temperature sensitivity of T_{opt}^{eco} along the precipitation gradient (right panel). Each
604 climate bin is defined by 1-°C intervals of $T_{max\ gs}^{air}$ and 100-mm intervals of mean annual
605 precipitation, based on current climate conditions averaged over 2001-2013. The solid line in the
606 right panel represents the temperature sensitivity of T_{opt}^{eco} along the precipitation gradient,
607 calculated as the slope of the linear regression between T_{opt}^{eco} and $T_{max\ gs}^{air}$ for a given precipitation
608 level. The shaded area indicates the SD of temperature sensitivity of T_{opt}^{eco} estimated by
609 bootstrapping.

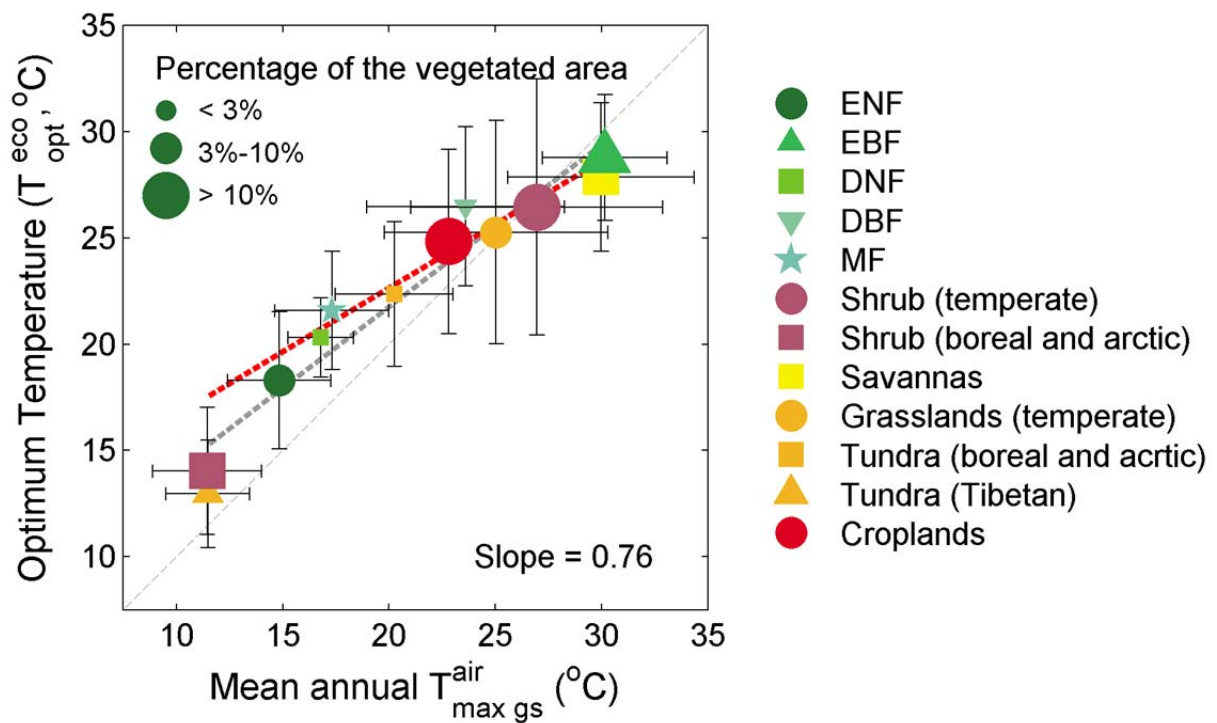
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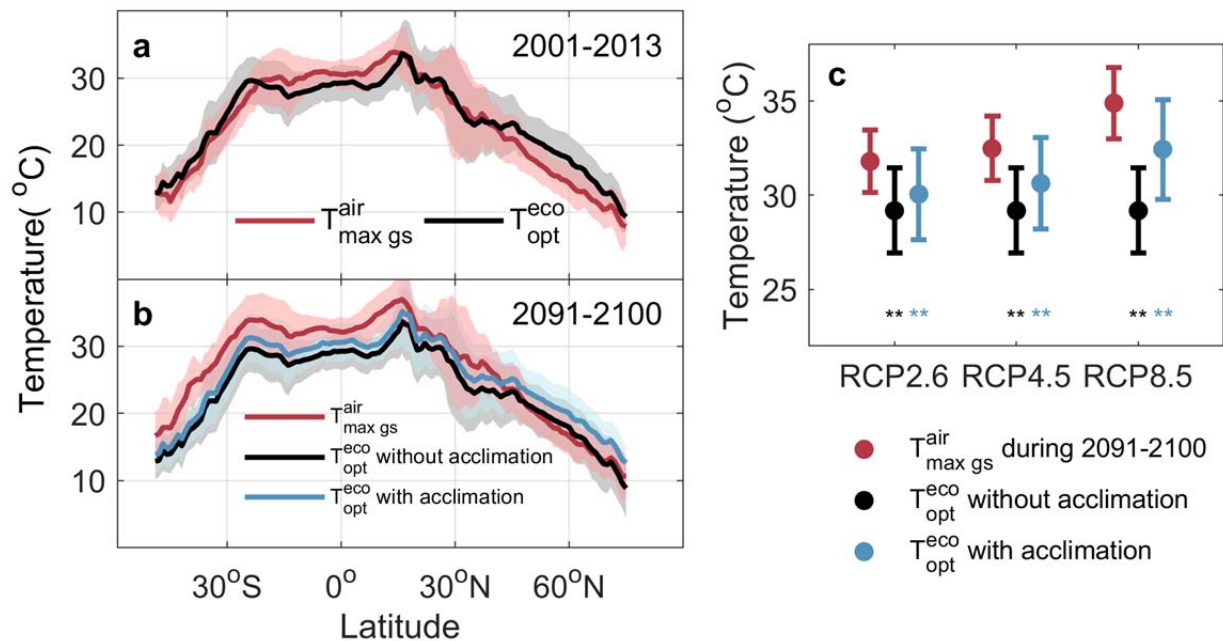
613 **Fig. 2 | Relationship between mean annual daily maximum air temperature during the**
 614 **growing season ($T_{max\ gs}^{air}$) and ecosystem-scale optimum temperature for vegetation**
 615 **productivity (T_{opt}^{eco}) across vegetation types.** The error bars indicate the SDs of $T_{opt}^{eco}/T_{max\ gs}^{air}$ for
 616 each vegetation type: ENF, evergreen needle-leaved forest; EBF, evergreen broad-leaved forest;
 617 DNF, deciduous needle-leaved forest; DBF, deciduous broad-leaved forest; MF, mixed forest;
 618 Shrub, closed and open shrublands. The light-gray dotted line represents $y=x$. The dark-gray
 619 dotted line is $y=0.76x+6.48$ with the slope value shown in the bottom right. The red dotted line is
 620 the flux tower derived slope (0.61) from Fig. 1a.



621

622

623 **Fig. 3 | Change with latitude in ecosystem-scale optimal temperature for vegetation**
624 **productivity (T_{opt}^{eco}) and daily maximum air temperature averaged over the growing season**
625 **($T_{max\ gs}^{air}$).** **a**, Current T_{opt}^{eco} versus current $T_{max\ gs}^{air}$; **b**, Future T_{opt}^{eco} versus future $T_{max\ gs}^{air}$. Current
626 T_{opt}^{eco} and $T_{max\ gs}^{air}$ are calculated using current temperature for 2001-2013, whereas acclimated
627 T_{opt}^{eco} and future $T_{max\ gs}^{air}$ are calculated using temperature for 2091-2100 projected by General
628 Circulation Models (GCMs) under the RCP4.5 scenario. Acclimated T_{opt}^{eco} is determined based on
629 the projected temperature and temperature sensitivity of T_{opt}^{eco} using the annual precipitation level
630 predicted for 2091-2100. The solid line and shaded area in each panel indicate the mean and SD,
631 respectively, of T_{opt}^{eco} or $T_{max\ gs}^{air}$ summarized by latitude. **c**, Future T_{opt}^{eco} versus future $T_{max\ gs}^{air}$ for
632 tropical evergreen forests. ** indicates that T_{opt}^{eco} is significantly lower than $T_{max\ gs}^{air}$ at $P < 0.01$ in a
633 paired t-test.



634