Climate change drives widespread shifts in lake thermal habitat

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Lake surfaces are warming worldwide, raising concerns about lake organism responses to thermal habitat changes. Species may cope with temperature increases by shifting their seasonality or their depth to track suitable thermal habitats, but these responses may be constrained by ecological interactions, life histories or limiting resources. Here we use 32 million temperature measurements from 139 lakes to quantify thermal habitat change (percentage of non-overlap) and assess how this change is exacerbated by potential habitat constraints. Long-term temperature change resulted in an average 6.2% non-overlap between thermal habitats in baseline (1978–1995) and recent (1996–2013) time periods, with non-overlap increasing to 19.4% on average when habitats were restricted by season and depth. Tropical lakes exhibited substantially higher thermal non-overlap compared with lakes at other latitudes. Lakes with high thermal habitat change coincided with those having numerous endemic species, suggesting that conservation actions should consider thermal habitat change to preserve lake biodiversity.

Global warming increases lake surface temperatures worldwide1–2, which strongly influences lake functioning, thermal structures and ecosystem processes3–5. As lakes warm, the available thermal habitat over specific temperature ranges can shrink or expand, with consequences for organisms, depending on their thermal tolerances6–8. In some cases, suitable thermal habitats may shrink or expand to the extent that native species become poorly adapted and non-native species thrive9–11. Changes in thermal habitat may be especially impactful in lakes because many species are ectothermic and, as on islands12 and mountaintops13, they are partially restricted by the boundaries of lakes. Thus, the shrinkage and expansion of thermal habitats raise important concerns about how climate change affects lake ecosystems14 and the implications of these changes for the threatened biodiversity that lake ecosystems currently support15.

With climate change, lakes are generally assumed to gain warm and lose cold thermal habitats. However, thermal habitat change in lakes is complex, as temperatures and temperature trends can vary vertically16–18, horizontally19–21 and seasonally21,22 within lakes. Furthermore, a substantial proportion of lakes exhibit cooling in their deeper waters, at least during stratified periods16–18,21–23, resulting in volumetrically cooler lakes in some regions21. This bottom-water cooling can be itself a thermal response of lakes to surface warming via a strengthening of thermal stratification17,26.
which shields bottom waters from downward transmission of surface heating. Furthermore, reductions in water clarity, such as temperature-induced increases in phytoplankton biomass, can also cause deep-water cooling in lakes. These contrasting mechanisms of surface and deep-water temperature change make it difficult to predict how thermal habitat may shift in response to observed global warming. Such knowledge is essential to improve our understanding of the vulnerability of lake ecosystems to climate change.

Some aquatic species may cope with a changing climate by shifting their seasonality (that is, phenology) or their depth distributions to track their suitable thermal habitat. For example, some fishes with broad environmental tolerances may overcome local thermal habitat change by changing their depth and seasonality to take advantage of new resources and species interactions. However, specialists whose seasonality and depth are constrained by species interactions, life history or resources such as light, nutrients and oxygen, may be more susceptible to thermal habitat change. For example, Planktothrix rubescens, a filamentous cyanobacterium, is adapted to light and thermal stratification conditions which occur only at specific depths and seasons. Similarly, some species of Daphnia, a common herbivorous zooplankton genus, partially rely on photoperiod as a cue for diapausing eggs to develop in spring, thereby limiting their capacity to track earlier phytoplankton blooms as lake temperatures change. Anoxic zones may hinder...
the ability of aerobic species to move deeper in lakes to track their suitable thermal habitat. Assessing whether habitat constraints exacerbate the risk of thermal habitat change would improve our understanding of the capacity of lake ecosystems to cope with climate variation.

Here we quantify the observed long-term thermal change in 139 lakes distributed across six continents—representing 69.3% of the Earth’s surface freshwater habitat by water volume (70.6×10^3 km^3 out of the 101.8×10^3 km^3 global total)^41. Using decades of lake temperature depth profiles, including more than 32 million total temperature observations, we calculated thermal non-overlap as the core metric of change. We defined thermal habitat change as the difference between recent lake temperatures (second half of each lake’s time series) compared with an earlier baseline period (first half of each lake’s time series). Thermal habitat change was quantified as the non-overlapped area of the two temperature distributions (recent and baseline) as a percentage of the combined area of those distributions, following an established method^42. Temperature distributions were volume-weighted to best capture the volumetric habitat available for species. The resulting values of thermal non-overlap are a measure of relative thermal change standardized against temperature variation in the baseline period. To evaluate whether depth and seasonal habitat restrictions exacerbate thermal habitat changes, we recalculated thermal non-overlap over various potential restricted habitat ranges for each lake. These potential habitat restrictions provide a proxy for the broad range of possible species’ capacities to cope with temperature change (Fig. 1). We used boosted regression trees (BRT) to model lake-to-lake variability.

Fig. 2 | Long-term changes in cumulative annual thermal habitat in lakes by volume. a–h. Volumetric thermal habitats are summarized for Lakes Vänern, Sweden (a,c); Kinneret, Israel (b,d); Giles, United States (e,g); and Trout, United States (f,h) by the change in volume of lake water within 0.1°C temperature bins as a percentage of each lake’s total volume (c,d,g,h). The purple to yellow colour scale reinforces the temperature gradient on the x axis and is consistent across panels to facilitate the comparison across lakes. Lakes shown here were chosen for representativeness; the same plots for all 139 lakes are available at https://doi.org/10.5281/zenodo.4686874.
in thermal non-overlap to determine which lake characteristics are most strongly related to variability in thermal habitat change.

Over the past several decades, volume-weighted, whole-lake temperatures have increased in 77% of lakes with a mean trend of +0.12 °C per decade. However, whole-lake warming rates failed to reveal the complexity of thermal habitat change across each lake’s temperature spectrum—most lakes exhibited serial losses and gains along their temperature gradients (Fig. 2) with implications for which species would be most affected. After controlling for differences in the length and seasonal coverage of each lake’s temperature time series and standardizing against a null estimate (Methods), we found that the mean thermal non-overlap across lakes from 1978 to 2013 was 6.2% and the distribution was skewed (median = 5.2%, inner quartile range = 4.5–6.9%; Fig. 3). Thus, 6.2% of the cumulative temperature distributions across both time periods are composed of either thermal habitat losses or gains over specific temperature ranges.

To evaluate whether habitat restrictions exacerbate the risk of thermal habitat change, we recalculated thermal non-overlap over various potential restricted habitat ranges for each lake. We considered two dimensions of habitat restrictions—depth and seasonality. The intensity of habitat range restrictions was scaled on a continuous gradient from 0 to 0.95 to facilitate comparisons across lakes, where 0 corresponds to no habitat restriction and 0.95 corresponds to habitats that are restricted to 5% of the available depths or days of the year. Thermal non-overlap was highest for the most restricted habitats (habitat restrictions of 0.95 in Fig. 4). When hypothetical habitats were restricted to 5% of all available depths, average thermal non-overlap increased from 6.2% to 9.7% (difference of 3.5%; Fig. 4a). When habitats were restricted to 5% of all available days of the year, average thermal non-overlap increased from 6.2% to 11.0% (difference of 4.8%; Fig. 4a). When habitats were restricted by both depth and season, strong synergistic interactions resulted in an increase in thermal non-overlap from 6.2% to 19.4% (difference of 13.2%). This interactive effect exceeded the additive effect when temperatures were compared at either restricted depths or restricted days of the year alone (13.2% > 3.5% + 4.8%).

The relative importance of seasonal versus depth habitat restrictions for thermal non-overlap depended on lake characteristics—seasonal habitat restrictions affected thermal non-overlap most strongly in shallower lakes (for example, Pesiöjärvi, Müggelsee and Annie; Fig. 4b–d) whereas depth habitat restrictions affected thermal non-overlap most strongly in deeper lakes (for example, Tahoe, Zürich and Tanganyika; Fig. 4e–g). Although we expect the most
severe habitat restrictions (values of 0.95) to be rare in lakes, any organism with a more limited habitat range due to various ecological constraints would be likely to face greater thermal non-overlap. Conversely, generalist species whose seasonality or depth is unencumbered by strict ecological constraints would be markedly less likely to face thermal habitat changes.

The BRT analysis showed that patterns in the magnitude of thermal non-overlap among lakes were strongly associated with mean lake depth. Thermal non-overlap was higher in deeper lakes, especially when seasonal habitat restrictions were low (seasonal habitat restriction <0.65; Fig. 5). More intense seasonal habitat restrictions elevated thermal non-overlap values in shallow lakes, thereby minimizing the difference between thermal non-overlap in deep and shallow lakes (Fig. 5). This interaction may have been caused by the effect of baseline temperature variation on values of thermal non-overlap (Extended Data Fig. 1). Shallow lakes typically have larger seasonal temperature variation, which would tend to reduce thermal non-overlap. Thus, when seasonal habitat restrictions are applied to shallow lakes, the thermal non-overlap-enhancing effects of high seasonal temperature variation are minimized. In the light of this feature, we would expect the costs of various habitat restrictions in terms of thermal non-overlap to depend strongly on lake depth.

Patterns across lakes in the magnitude of thermal non-overlap were strongly associated with latitude (relative importance of all predictors shown in Extended Data Fig. 3). We expected temperate and arctic lakes to have greater thermal non-overlap because lake surface warming rates tend to be greater there. However, in contrast to our expectation, tropical lakes exhibited substantially higher thermal non-overlap compared with lakes at other latitudes (Figs. 4 and 5). High thermal non-overlap in the tropics may arise from the thermal habitat change metric we used, which is affected by the baseline temperature variability (Extended Data Fig. 1). Smaller inter- and intra-annual temperature variation in tropical lakes upweights their magnitude of temperature change and produces relatively high thermal non-overlap. Quantifying thermal habitat change in
this way may be more ecologically meaningful because the effects of temperature change on species depend on the breadth of species’ thermal tolerances, which tend to be narrower when environmental temperature variation is low43–45. Overall, these analyses suggest that lake ecosystem sensitivity to climate change, as characterized by thermal non-overlap, differs considerably from the global pattern in the magnitude of lake surface warming. Despite slower surface warming rates, habitat changes are likely to be felt most strongly in tropical lakes where biodiversity may be most affected.

The global pattern in thermal non-overlap may have differed from that of whole-lake warming rates (Extended Data Fig. 4) because transparency changes sometimes shield deep waters from downward transmission of heat25. Shielding would counterbalance calculations of whole-lake warming rates when integrating across the entire water column because some depths warm whereas others cool. When calculating whole-lake warming rates, lakes which exhibit deep-water cooling often have moderate or no significant differences in whole-lake mean annual temperature25. However, in our thermal non-overlap calculations, simultaneous warming and cooling over different parts of the water column would both elevate values of thermal non-overlap because the metric is agnostic toward the direction of change. Therefore, thermal non-overlap calculations reflect more than just heat content and, importantly, whole-lake temperature trends do not adequately describe the magnitude of thermal habitat change.

Some species may benefit from the changes in thermal habitat observed here. For instance, in Lake Zürich, the net growth rate of *P. rubescens* is maximized in warmer months (July–September) at specific depths (5–20 m) due to light, temperature and density stratification conditions. Restricting volumetric thermal habitat over this range increases the non-overlap value from 16% for the lake overall to 22% for the most suitable habitat for *P. rubescens* (Fig. 6). In response, *P. rubescens* growth is anticipated to increase in Lake Zürich35,36 because warmer temperatures and higher thermal stratification are typically more suitable for the species’ growth46. While some individual species with highly restricted habitats may benefit from thermal non-overlap45, thermal non-overlap is generally expected to increase the overall likelihood of species extinctions and community disruptions41. It has even been warned that climate change may drive certain species to local extirpations via changes to lake thermal habitat47. Our analysis provides a more global context for these concerns and goes further by showing that, in many cases,
species will require the absence of habitat restrictions to effectively track their suitable thermal conditions. Shifts across depth and season are already widely observed in lakes for a variety of taxa\textsuperscript{32,48,49}. However, the inability of most species to adopt these coping strategies\textsuperscript{34,39,48} may increase the likelihood of community disruptions due to the disappearance of habitat over their suitable thermal ranges.

Importantly, lakes with high thermal non-overlap closely coincide with those identified as containing a global heritage of freshwater biological diversity and endemism\textsuperscript{50}, including Lakes Baikal (thermal non-overlap without habitat restrictions = 5.5%), Biwa (9.9%), Tanganyika (15.4%) and Victoria (18.5%). In these lakes, elevated risks of extinction due to thermal habitat change are possible, as is the disruption and disaggregation of lake ecosystems due to mismatched shifts across species\textsuperscript{6,7,39}. Given the potential for the formation of new communities as thermal habitats change, the link between overall thermal habitat changes and ecological responses could fundamentally alter the nutrient and energy pathways that drive these ecosystems\textsuperscript{5,7,36}.

Changes in lake thermal habitat may be especially impactful because most lake taxa are ectothermic, and many are partially restricted by the boundaries of lakes. Given these vulnerabilities to thermal habitat changes, standard conservation measures such as in-lake protected areas\textsuperscript{51} may be ineffective at fully stemming the negative consequences of thermal habitat change\textsuperscript{51}. Instead, lake conservation efforts focused on enhancing the capacity for organisms to shift across depth and season (for example, by reducing the size of lake anoxic zones) may help some species avoid thermal non-overlap. Furthermore, species that are active year-round in lakes may be sensitive to the full annual cycle of temperature without a chance to avoid seasons with unsuitable thermal habitat. Thus, for species habitats that are restricted across season or depth, conservation actions focused on enhancing within-lake shifts will be insufficient. Instead, efforts focused on enhancing connectivity among lakes (for example, dam removal) may be more effective at ameliorating the negative consequences of thermal habitat change\textsuperscript{52}.

Lake temperature change can indirectly influence the habitat available for lake species aside from the direct temperature effect. For instance, lake temperature change influences underwater light availability and dissolved oxygen concentrations\textsuperscript{28,53–55}. Anoxic zones may increase in extent or duration as lakes warm, further restricting the depths and seasons that can be occupied by aerobic organisms. Light penetration may increase or decrease with changes in water transparency driven by climate change with added consequences for the habitat available for photosynthetic organisms. Future studies focused on specific taxa should include these other important determinants of the habitat space available to them. The modular nature of the non-overlap calculations used herein allows for such

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**Fig. 6 | Thermal habitat change for the restricted habitat of P. rubescens in Lake Zürich.** a. Epifluorescence microscopy photo of P. rubescens, a filamentous cyanobacterium present in Lake Zürich, with its optimal habitats occurring from July to September at depths of 5–20 m. b,c, Thermal habitat change (percentage of non-overlap) in this restricted range increases from 16% for Lake Zürich overall (b) to 22% for the habitat of P. rubescens (c). Credit: photo in a, Thomas Posch.
work incorporating a wider variety of environmental variables that determine habitat suitability14. The redistribution of life on Earth is a major ecological response to anthropogenic global warming. As Earth warms, lake thermal habitats may shrink, expand or shift to seasons or depths where ecological interactions, life histories or resources limit species’ growth and reproduction. These thermal shifts will inevitably have consequences for the species that lakes currently support. Forecasts of how species will respond to thermal non-overlap will always be less certain because they are often extrapolated from present conditions. However, with the results presented here, we have a framework for developing testable hypotheses relating lake thermal habitat change to biotic change over time for a large number of observed ecological time series. Given the risk of lake biodiversity loss, there is considerable societal value in resolving the connections between our results and real ecological responses in lakes.

Online content
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We used long-term time series of lake temperature profiles to determine the magnitude of thermal habitat change in 139 widely distributed lakes. Time series were interpolated across depth and season to generate data with consistent resolution and across lakes. To assess temperature change, we used a method, thermal non-overlap, based on the percentage of two kernel density estimations of lake temperature which are non-overlapping. We calculated the metric for a range of plausible seasonal and depth habitat restrictions for aquatic species in the face of climate change. We used BRT to explain variability across lakes in their thermal habitat non-overlap as a function of lake characteristics (mean depth and latitude) characteristics of the time series for each lake (starting day of the year, ending day of the year, starting year and ending year, average number of sampling dates per year, long-term trend in the number of sampling dates per year, long-term trend in the yearly seasonal range of sampling dates), the habitat restriction values (season and depth) and the location of the time series delineation for thermal non-overlap calculations (70th, 50th and/or 70th quantiles of the years included in each lake’s time series).

Study sites. We compiled long-term lake temperature data from 139 lakes across the globe. Temperature variations in many of these lakes have already been linked to climate change\(^1\)\(^\text{–}\)\(^3\), but temperature change in at least one lake may be partially due to background climate variation in addition to anthropogenic climate change (Atlantic Multidecadal Oscillation in Lake Annie)\(^5\). The lakes included in our analysis represent a wide range of surface area (0.02 to 68,800 km\(^2\)), maximum depth (2.3 to 1,642 m), latitude (60°S to 69°N) and elevation (–212 to 1,987 m above sea level) (see Supplementary Table 1 for more information).

Temperature data. In total, we used more than 32 million lake temperature measurements for our analyses. The number of observations per lake ranged from 368 (Lake Stensjon) to 7,636,767 (Lake Superior) with approximately 232,000 observations per lake on average. Temperature data from each lake came from in situ temperature profiles\(^4\)\(^\text{–}\)\(^6\) for lakes smaller than 169 km\(^2\) and from a combination of in situ temperature profiles and remotely sensed surface water temperatures for 21 larger lakes. Remote sensing data were used in recognition that temperature and warming rates can vary substantially across altitude and longitude for large lakes\(^7\). The mean length of the temperature time series was 36 years with a range from 15 to 101 years. All lakes had temperature data which started in the year 2000 or earlier and ended in 2000 or later. Lakes had on average 29 temperature profiles per year (inner quartile range: 7–26). In situ temperature data were measured using a wide variety of temperature sensors. Data collection methods included regularly collected discrete temperature profiles, high-resolution thermistor chains and other commonly accepted tools for measuring aquatic temperature. The in situ data are publicly available through the environmental data initiative\(^8\).

Remotely sensed lake surface temperatures were measured using the Advanced Very High-Resolution Radiometer (AVHRR) and processed by the Group for High Resolution Sea Surface Temperature (GHRSST) project\(^9\). AVHRR data have been validated against buoy data from the North American Great Lakes and found to have a root mean squared error of 0.55°C compared with in situ measurements\(^2\). AVHRR temperature data were included to capture horizontal variability in temperature and warming in 21 of the 139 lakes that would not be captured by temperature profiles from a single central location. AVHRR data were pooled with in situ data for temperature interpolation.

Temperature interpolation. Temperature data were spatially and temporally interpolated for each lake. All temperature profile data were first linearly interpolated across depth because temperature variability with depth is highly constrained by lake physics and typically allows for robust interpolations. The largest lake gap over depth in each lake was interpolated using a 0.1 × mean depth of each lake. Following interpolation across depth, data were interpolated across time using standard spline interpolation models with a Kalman filter\(^10\). The model output was used to fill data gaps to produce a continuous, daily time series over the day of the year range for which temperature profiles had been regularly measured. Some times of the year were excluded from specific lakes because they lacked regular measurements throughout the length of the long-term time series. Thus, the same starting and ending day of the year was used for each lake throughout its time series, and was often shorter than the full annual cycle (Supplementary Table 1). The largest gap in time over which interpolation occurred was 30 days and this included extrapolations for lakes with missing data at the beginning or end of seasonal coverage in a specific year. Years with longer gaps were omitted from the analysis and the length of the seasonal coverage was optimized to minimize the number of years that needed to be removed. For large lakes with many sampling points (for example, Baikal, Superior, Victoria), temperature data were divided into 1,000 km\(^2\)/latitude–longitude bins and interpolated across depth and across time separately for each bin. The mean seasonal coverage of the interpolated lake time series was 245 days per year with a minimum of 17 days per year and a maximum of 365 days per year.

The interpolated temperature output had a daily temporal resolution and a depth resolution which varied continuously over depth. At the lake surface, we interpolated temperatures every 0.1 m (for example, 0, 0.1 m, 0.2 m), to every 1 m starting at a depth of 10 m (for example, 10 m, 11 m, 12 m) and every 100 m starting at a depth of 1,000 m (for example, 1,000 m, 1,100 m, 1,200 m). These depth increments were used because they consistently gave good coverage over all major lake strata, regardless of each lake’s morphometric characteristics, while minimizing computational intensity by eliminating redundancy within lake strata.

Thermal habitat non-overlap calculations. After interpolating the temperature data across depth and season for each lake, we bisected it into an early part (part \(a\)) and a later part (part \(b\)). Parts \(a\) and \(b\) were iteratively delineated at three points positioned serially along the time series—at the 30th, 50th and 70th quantiles. We averaged the final non-overlap values across these three delineations for each lake so that the results depended less on the somewhat arbitrary decision of where to split the time series. For each lake we determined 10,000 temperature values from each of parts \(a\) and \(b\). This was repeated ten times resulting in a total of 300,000 temperature values across all three time series delineations and all ten repetitions for each lake (10,000 × 3 × 10). The sampling probability for temperature values in each comparison was weighted by the volume increment associated with each temperature value (depth increment \((L)\times\text{cross-sectional area at each depth (}C\)) \(L\), was calculated as the difference between the depth of the sampled temperature value and the next depth in the depth resolution of the interpolated temperatures. \(C\) at each depth for each lake was calculated using standard, three-parameter models for estimating lake cross-sectional area based on surface area, maximum depth and mean depth\(^8\). For large lakes with temperature data at multiple locations across latitude and longitude, \(C\) was divided by the number of latitude–longitude bins used for each lake. Temperature values from large lakes were sampled regardless of their associated latitude–longitude bins. As a result of the volume-weighting procedure, temperature measurements were sampled in proportion to the volume of water represented by each value, with temperatures representing larger volumes being sampled more often. As a consequence of this volume-weighting procedure, the resulting temperature distributions were robust to moderate changes in the depths used for the temperature interpolation (Supplementary Fig. 1).

We defined thermal non-overlap (TNO) as the symmetric difference (\(|\setminus\)) between the kernel density estimations of temperature values from parts \(a\) and \(b\) of the time series as a proportion of the union (\(\cup\)) of both kernel density estimations, following an established method\(^6\). Conversely, we defined the thermal habitat overlap (as opposed to non-overlap) as the intersection (\(\cap\)) of the kernel density estimations as a proportion of the union (\(\cup\)) of both distributions. All values were converted to percentages by multiplying by 100.

\[
\text{\normalfont TNO} = 100 \times \frac{T_{\text{recent}} \setminus T_{\text{baseline}}}{T_{\text{recent}} \cup T_{\text{baseline}}} = 100 \times \frac{1 - \frac{1}{T_{\text{recent}} \cup T_{\text{baseline}}}}{1 - \frac{1}{T_{\text{recent}} \cup T_{\text{baseline}}}} \tag{1}
\]

We used simulations to test the sensitivity of TNO to changes in mean and s.d. of temperature. We primed these simulations with three baseline temperature distributions all with a mean of 15°C but with varying s.d. (4, 6, 8°C). We simulated a range of additional temperature distributions by increasing and decreasing the mean and s.d. of the baseline temperature distributions and then calculated the corresponding values of TNO. The simulated change in both mean and s.d. was varied from −3 to +3°C. We found that TNO was sensitive to changes in mean and s.d. but was slightly more sensitive to reductions in s.d. compared with increases. TNO values also depended on the baseline s.d., such that lower starting s.d. elevates values of non-overlap given an equivalent change in temperature (Extended Data Fig. 1).

We also quantified null values of thermal non-overlap (TNO\(_0\)) by repeating the thermal non-overlap calculations but where parts \(a\) and \(b\) were defined by randomly dividing the individual years of data into two separate groups as opposed to sequentially dividing them along the time series.

\[
\text{\normalfont TNO}_0 = 100 \times \frac{T_{\text{random a}} \setminus T_{\text{random b}}}{T_{\text{random a}} \cup T_{\text{random b}}} \tag{2}
\]

To calculate standardized thermal non-overlap (TNO\(_s\)), we subtracted TNO\(_0\) from TNO thereby setting the null expectation to zero.

\[
\text{\normalfont TNO}_s = \frac{\text{\normalfont TNO} - \text{\normalfont TNO}_0}{\text{\normalfont TNO}_0} \tag{3}
\]

In this case, if the temperature distributions in the recent and baseline time periods were identical, the TNO\(_s\) would equal approximately zero. Values different from zero reflect a combination of random noise and long-term temperature change. All non-overlap values described in the main text and shown in Figs. 2–6 reflect values of TNO\(_s\). A comparison between raw values of TNO and TNO\(_s\) can be found in Extended Data Fig. 5. Thermal non-overlap values and the null values were calculated using the ‘overlap’ function from the ‘overlapping’ package\(^6\) in the R environment for statistical computing and visualization. In the function, we set the number of equally spaced points at which the overlapping kernel density estimation is evaluated to 100 for all comparisons because it minimized the values of TNO\(_0\), (we considered a range of values from 5 to 10,000).
To assess the effect of seasonal habitat restrictions ($S_{\text{season}}$) and volumetric habitat restrictions ($V_{\text{limit}}$), we modified equations (1)-(3) by comparing temperature values only from a specified range of depths and/or days of the year. We considered a range of habitat restrictions scaled from 0 to 0.95 instead of in units of days or m$^3$, so that habitat restrictions could be more readily compared across lakes. Comparing a $V_{\text{limit}}$ value of 0.8 across lakes of different sizes assumes that a habitat restriction of 2 m$^2$ in a 10 m$^3$ lake would be comparable to a 20 m$^2$ habitat delineation in a 200 m$^3$ lake. The actual size of the seasonal habitat restrictions for each lake in units of days were calculated using the value of $S_{\text{season}}$ as follows:

$$S = (\text{day}_{\text{max}} - \text{day}_{\text{min}}) \times (1 - S_{\text{season}})$$

where $S$ is the seasonal habitat restriction in units of days, $\text{day}_{\text{max}}$ is the maximum day of the year of the lakes' seasonal coverage, $\text{day}_{\text{min}}$ is the minimum day of the year of the lakes' seasonal coverage and $S_{\text{season}}$ is the seasonal habitat restriction scaled from 0 to 0.95. For example, in a lake with a seasonal coverage from day of the year 1 to day of the year 365, with an $S_{\text{season}}$ value of 0.75, we compared randomly selected temperature values from time periods $a$ and $b$ separately for four seasonal bins (days of the year 1–91, 92–183, 184–273 and 274–365). Simiarly, the actual size of the volumetric habitat restrictions ($V$) for each lake in units of m$^3$ were calculated using the value of $V_{\text{limit}}$ as follows:

$$V = (\text{volume}) \times (1 - V_{\text{limit}})$$

where $V$ is the volumetric habitat restriction in units of m$^3$, $\text{volume}$ is the lake's total volume and $V_{\text{limit}}$ is the volumetric habitat restriction value scaled from 0 to 0.95. For example, if a lake with a volume of 100 m$^3$ had a $V_{\text{limit}}$ value of 0.75, we randomly selected temperature values from time periods $a$ and $b$ which were within four 25 m$^3$ (100 m$^3 \times (1 - 0.8))$ bins. Volume bins were subsequently translated into sequential depth bins for the purpose of temperature value selection, making them functionally depth limits, and they are presented as such in the main text.

We factorioly combined a discrete series of values for $S_{\text{season}}$ and $V_{\text{limit}}$ (0, 1/2, 2/3, 5/6, 8/9, 12/13 and 19/20) to test a range of combined seasonal and volumetric habitat restrictions. We assessed the statistical significance of these lake-specific thermal non-overlap values estimated with the Wilcoxon signed-rank test. In the test, we compared $TN_0$ values to $TN_0$ values separately for each combination of time series delineation, seasonal habitat restriction and depth habitat restriction ($n = 108$). The average $P$ values from these tests for each lake are shown in Supplementary Table 1.

We compared thermal non-overlap values to a more widely used metric of whole-lake thermal change—whole-lake temperature trends. Whole-lake temperature trends were calculated based on the annual averages of all temperature values sampled for the pairwise thermal non-overlap calculations to maximize the comparability of the resulting temperature trends and thermal non-overlap values. Due to the temperature sampling probability being volume-weighted, the temperature trend was also indirectly volume-weighted. Temperature trends were calculated separately for each lake and for each time series set. For each lake, whole-lake temperature trends and thermal non-overlap values were not strongly correlated ($r = 0.10$, Kendall's rank correlation coefficient; Extended Data Fig. 4). All statistics and graphics were produced in the R statistical computing environment$^{17}$.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**
The remote sensing lake temperature data used can be found at https://doi.org/10.5067/GHAO-4BC02 and through the Earthdata website (https://earthdata.nasa.gov/). The in situ lake temperature data used available through the Environmental Data Initiative (EDI) data portal (https://portal.edirepository.org/nis/home.jsp) at https://doi.org/10.6073/pasta/f037d682eace2f46764e24e26068ee15. Geospatial and morphometric data for each lake are available from the HydroLAKES database at https://doi.org/10.1038/ncomms13603 and can be found at http://www.hydrosheds.org. The country vector data shown in Fig. 3 were sourced from Natural Earth Data (https://www.naturalearthdata.com) under a public domain licence that allows full and unrestricted use. The lake-specific panels in Figs. 2 and 4 have been repeated for all 139 lakes and are available at https://doi.org/10.5281/zenodo.4686874.

**Code availability**
All code is available under the identifier https://doi.org/10.5281/zenodo.4688292.

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Author contributions

B.M.K. conceived of the work, led the study, completed the analysis and wrote the manuscript with input from all co-authors. All authors edited and revised the manuscript. R.A., P.R.L., R.M.P. and R.I.W. provided additional feedback, edits and revisions. Data contributions were made by O.A., S.B., W.C.-M., S.P.D., M.T.D., E.E.G., K.D.H., D.O.H., S.N.H., K.D.J., W.K., L.B.K., B.M.K., P.R.L., F.L., M.S.L., S.C.M., D.C.M.-N., D.C.P., R.M.P., D.C.R., M.R., A.M.P., J.A.R., S.S., N.S., E.A.S., D.S., M.A.T., W.T., P.V., R.S., G.A.W. and R.A. R.M.P. and B.M.K. compiled, organized and quality checked the data across all lakes.

Competing interests

The authors declare no competing interests.

Additional information

Extended data are available for this paper at https://doi.org/10.1038/s41558-021-01060-3. Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41558-021-01060-3. Correspondence and requests for materials should be addressed to B.M.K.

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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | Sensitivity of thermal habitat change values to changes in mean and standard deviation of three hypothetical temperature distributions. The three hypothetical baseline temperature distributions considered here had a mean of 15 °C and standard deviations of 4, 6, and 8 °C (a). We simulated the effects of temperature changes (increases and decreases in both the mean and the standard deviation of the baseline temperature distributions) on resulting values of thermal habitat change (% non-overlap). The effects on thermal habitat change were simulated for each of the three baseline conditions (b-d). Values of thermal habitat change (% non-overlap) were sensitive to changes in mean and standard deviation of temperature as well as to the standard deviation of the baseline temperature distribution. Higher standard deviation of the baseline distribution lead to lower values of thermal habitat change.
Extended Data Fig. 2 | Partial dependency plots for each of the 12 predictors in the boosted regression trees. Colored lines are the raw partial dependence plot values from each of the 100 different BRTs which were fit to random subsets (70%) of the 139 lakes included in this analysis and the black lines are the locally weighted scatterplot smoothed (LOESS) lines across all BRTs. The panels are ordered by the relative importance of each predictor in the model (a-l) with the most important predictors in the upper left (a) and the least important in the bottom right (l). Note that the scales on the x- and y-axes vary from plot to plot.
Extended Data Fig. 3 | Relative importance of each of the 12 predictors in the boosted regression trees. The relative importance of each predictor variable in the model is a function of the frequency with which it was included in the BRT’s individual regression trees and the improvement to the model that resulted from its inclusion. Relative importance values are scaled to sum to 100%. The individual colored points represent 100 different BRTs fit to random subsets (70%) of the 139 lakes included in this analysis and they are jittered to aid their visualization. The colored violins are the kernel density estimations of the distribution and the black boxes show the first, second and third quartiles of the distributions.
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Comparison of thermal non-overlap and whole-lake warming rates across all 139 lakes. The density plot (a) shows the distribution of volume-weighted whole-lake warming rates across the 139 lakes included in the study. Thermal non-overlap and the absolute value of whole-lake warming rates were not strongly correlated ($r = 0.10, p = 0.081$, Kendall’s rank correlation; b).
Extended Data Fig. 5 | Comparison of raw thermal non-overlap values with null non-overlap values prior to removing the effects of time series characteristics. Raw thermal non-overlap values were calculated by comparing the first and second parts of each lake’s time series whereas the null non-overlap values were calculated by comparing two groups of randomly selected years. Values shown here are for the case with no seasonal or depth habitat restrictions. The non-overlap values in the main text are ‘standardized’ by subtracting the null non-overlap values from the raw non-overlap values shown here.
### Extended Data Fig. 6 | Model performance statistics from the cross-validation for the boosted regression trees analysis

Summary statistics are described in the ‘Summary statistic’ column and the values of each statistic are reported in the adjacent ‘value column’.

| Summary statistic                                                                 | Value                                      |
|----------------------------------------------------------------------------------|--------------------------------------------|
| Training data correlation between predicted and observed values where correlations were averaged across all BRTs | Kendall = 0.96  
Spearman = 0.99 |
| Training data correlation between predicted and observed values where predicted values were averaged across all BRTs | Kendall = 0.98  
Spearman = 0.99 |
| Test data correlation between predicted and observed values where correlations were averaged across all BRTs | Kendall = 0.56  
Spearman = 0.75 |
| Test data correlation between predicted and observed values where predicted values were averaged across all BRTs | Kendall = 0.61  
Spearman = 0.81 |
| Overall correlation between predicted and observed values where correlations were averaged across all BRTs and individual correlations were calculated from pooled training and test datasets | Kendall = 0.92  
Spearman = 0.93 |
| Overall correlation between predicted and observed values where predicted values were averaged across all BRTs from pooled training and test datasets | Kendall = 0.90  
Spearman = 0.99 |
| Standard deviation (SD) of standardized thermal non-overlap values ($TNO_L$) prior to cross validation | 16.24 % non-overlap |
| Training data SD of model residuals where SDs were averaged across all BRTs | 0.99 % non-overlap |
| Training data SD of model residuals where predicted values were averaged across all BRTs | 0.59 % non-overlap |
| Test data SD of model residuals (PRESS) where SDs were averaged across all BRTs | 10.44 % non-overlap |
| Test data SD of model residuals (PRESS) where predicted values were averaged across all BRTs | 9.36 % non-overlap |
| Overall SD between predicted and observed values where SDs were averaged across all BRTs and individual SDs were calculated from pooled training and test datasets | 5.84 % non-overlap |
| Overall SD between predicted and observed values where predicted values were averaged across all BRTs from pooled training and test datasets | 2.93 % non-overlap |
| Median absolute deviation (MAD) of standardized thermal non-overlap values ($TNO_L$) prior to cross validation | 11.34 % non-overlap |
Extended Data Fig. 7 | Analysis of model residuals calculated as the difference between observed values and the average predicted values across all 100 BRTs. The distribution of thermal habitat change (% non-overlap) values compared to the distribution of model residuals (a). Model residuals show minimal patterning when compared to each predictor in the BRT analysis (b–m). Colors represent the counts of observations in 2-dimensional bins where the color scale has been log₁₀-transformed.
Reporting Summary

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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

No software was used to collect the lake temperature data.

Data analysis

The R statistical computing environment was used for the analysis of all data and for the production of all figures in this manuscript. All code can be found under the identifier, DOI: 10.5281/zenodo.3894908

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Study description
We use long-term time series of lake temperature profiles to determine the magnitude of thermal habitat change in 139 widely distributed lakes. Time series were interpolated across depth and across seasons to generate data with consistent resolutions across lakes. To assess temperature change, we used a metric, “thermal novelty”, based on standardized Euclidean distances applied to the full length of each lake’s interpolated time series. We calculated the metric for a range of plausible seasonal and depth shift capacities for aquatic species in the face of climate change. We used a boosted regression tree (BRT) to explain variability across lakes in their thermal habitat novelty using lake characteristics (shore development index, residence time, mean depth, surface area, latitude, longitude, and elevation) while controlling for the characteristics of the time series for each lake (starting day of the year, ending day of the year, starting year, and ending year) and the shift limit (seasonal shift limit and depth shift limit).

Research sample
In situ temperature measurements were recorded digitally or in writing at the time of measurement.

Sampling strategy
Long-term lake water temperature data were assembled from as many monitoring stations around the world as possible.

Data collection
The data reported here reflect the collective efforts of myriad dedicated field crews, laboratory staff, data management and quality control staff, analysts and many others from a wide variety of nations, states, tribes, agencies, universities, and other organizations.

Timing and spatial scale
The timing and spatial scale of temperature measurements are complex and vary from lake to lake. The duration and seasonal coverage for each the temperature profile time series from each lake are summarized in the supplementary material.

Data exclusions
No data were excluded.

Reproducibility
There are no experimental findings in our study.

Randomization
The lake water temperature data reported here were collected on an opportunistic basis from as many lakes as possible. The length and seasonal coverage of lake water temperature time series varied from lake to lake and this variation was accounted for using boosted regression trees.

Blinding
Blinding was not relevant for this observational study based on long-term monitoring data which does not include experiments.

Did the study involve field work?  Yes  No

Field work, collection and transport

Field conditions
Lake water temperature measurements were collected across 139 lakes under a wide variety of field conditions over several decades.

Location
All sampling locations can be found in the supplementary material for the manuscript.

Access & import/export
All temperature data were recorded digitally or in writing upon measurement and no physical samples were imported or exported for the purpose of this study.

Disturbance
The measurement of lake water temperature is generally not an intrusive process.

Reporting for specific materials, systems and methods

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