Ecological consequences of long-term browning in lakes

Craig E. Williamson1, Erin P. Overholt1, Rachel M. Pilla1, Taylor H. Leach1, Jennifer A. Brentrup1, Lesley B. Knoll2, Elizabeth M. Mette1 & Robert E. Moeller1

Increases in terrestrially-derived dissolved organic matter (DOM) have led to the browning of inland waters across regions of northeastern North America and Europe. Short-term experimental and comparative studies highlight the important ecological consequences of browning. These range from transparency-induced increases in thermal stratification and oxygen (O2) depletion to changes in pelagic food web structure and alteration of the important role of inland waters in the global carbon cycle. However, multi-decadal studies that document the net ecological consequences of long-term browning are lacking. Here we show that browning over a 27 year period in two lakes of differing transparency resulted in fundamental changes in vertical habitat gradients and food web structure, and that these responses were stronger in the more transparent lake. Surface water temperatures increased by 2–3 °C in both lakes in the absence of any changes in air temperature. Water transparency to ultraviolet (UV) radiation showed a fivefold decrease in the more transparent lake. The primary zooplankton grazers decreased, and in the more transparent lake were largely replaced by a two trophic level zooplankton community. These findings provide new insights into the net effects of the complex and contrasting mechanisms that underlie the ecosystem consequences of browning.

The causes of the browning of inland waters have been shown to be related to a combination of recovery from anthropogenic acidification1, increases in precipitation2,3, and other factors such as changes in land use and DOM-iron interactions4,5. In some regions where browning is prevalent such as northeastern North America, up to a 71% increase in very heavy precipitation events has been reported in the past 5 decades6. Browning results from increases in not only dissolved organic carbon (DOC) concentration1, but also DOC color (DOC-specific absorbance at 320 nm, a*320nm)7, and is often accompanied by increasing water pH1.

Prior short-term experimental, comparative, and modeling studies suggest that long-term browning will alter vertical habitat gradients in pelagic ecosystems with important ecological consequences8. Two contrasting mechanisms have been hypothesized to underlie these ecological consequences of browning: (1) the provision of ecosystem subsidies in the form of nutrients or fixed carbon, and (2) the effects of shading8,9. Terrestrial DOM can provide nutrient subsidies that enhance zooplankton production10, while the shading effects of DOM inhibit photosynthesis and consequently reduce food supply for higher trophic levels including zooplankton and fish11. Simultaneous increases in pH and decreases in UV exposure may also alter consumer-resource interactions during browning12. All of these mechanisms can lead to reductions in water transparency to UV and photosynthetically active radiation (PAR, 400–700 nm) that have the potential to influence vertical habitat gradients, food web structure, and ultimately ecosystem services. Accumulating evidence suggests that the effects of browning are context dependent and vary with the initial color of the lake. Clear-water oligotrophic lakes are hypothesized to be more sensitive to browning than are lakes with initially higher DOM13. Modeling and comparative studies even suggest a unimodal response wherein clear-water oligotrophic lakes respond to browning in the opposite way as do already brown lakes14. But do the results of these short-term experimental, comparative, and modeling studies accurately forecast the response of lakes to long-term browning? How do the ecological consequences differ in clear-water versus brown-water lakes? And are important water transparency related phenological events such the UV or PAR clear-water phases15 altered during browning? The multi-decadal observational studies that are necessary to resolve these questions and their implications for important ecosystem services are lacking. Here we present 27 years of data on two small lakes of differing transparency that demonstrate changes in the vertical habitat gradients and food web structure during browning that are especially strong in the clearer lake.

The two small study lakes are located in protected watersheds in northeastern Pennsylvania, USA, a region that has experienced both the decrease in acid deposition as well as the increase in precipitation characteristic of regions

1Department of Biology, Miami University, Oxford, OH 45056, USA. 2Lacawac Sanctuary, Lake Ariel, PA 18436, USA. Correspondence and requests for materials should be addressed to C.E.W. (email: craig.williamson@miamiOH.edu)
Figure 1. Interannual changes in meteorological and limnological variables during the study period. (a) Mean spring (March – mid-July date of sampling) and annual air temperature (°C). (b) Average spring (March – April) wind speed (m s⁻¹). (c) Annual precipitation (cm yr⁻¹). (d) Annual atmospheric deposition of NO₃⁻ and SO₄²⁻ (kg ha⁻¹ yr⁻¹). (e) DOC concentration (mg L⁻¹) in the surface waters. (f) DOC-specific absorbance at 320 nm (a₃₂₀ nm, m² (g C)⁻¹) in the surface waters, a measure of DOC color. (g) Surface water pH. (h) Surface water acid-neutralizing capacity (ANC, μEq L⁻¹). (i and j) Water transparency to UV radiation (320 nm) (i) and PAR (400–700 nm) (j) expressed as the depth (m) to which 1% of subsurface irradiance
where lakes are browning\(^2,3,6,16\). Lake Lacawac is a 21 ha dystrophic brown-water lake (DOC = 5–6 mg C L\(^{-1}\)) with a maximum depth of 13 m and a catchment located within a sanctuary preserve, while Lake Giles is a 48 ha oligotrophic clear-water lake (DOC = 1–2 mg C L\(^{-1}\)) with a maximum depth of 23 m and a catchment located within a privately owned and well-protected watershed. These lakes are near and below the global median DOC concentration of 5.7 mg C L\(^{-1}\) for lakes\(^17\). Both are seepage lakes with no tributary streams or rivers other than a small spring on the shore of each lake. There have been no changes in land use or land cover in the privately owned and well-protected catchments, thus leaving only atmospheric or meteorological factors as potential agents of external forcing in these lakes.

### Results

There was no significant trend in either mean annual air temperature or spring (March – mid-July date of sampling) air temperature in the region during the 1988–2014 study period (Fig. 1a). There was a significant increase in spring (March – April) wind speed (Fig. 1b). These two different “spring” periods were chosen as those most likely to influence water temperatures in the epilimnion (air temperature in the whole ice-free period before the sampling), and hypolimnion (wind speed in March-April following ice-out, but before thermal stratification is established). Annual precipitation also increased significantly during this period (Fig. 1c), and atmospheric deposition of both SO\(_4^{2-}\) and NO\(_3^-\) decreased (Fig. 1d).

In-lake changes in Giles were greater than, but not opposite those in Lacawac. Average DOC concentrations and color (\(a^{*}_{320}\)) increased in both lakes but increases were statistically significant only in Giles where DOC concentration doubled (Fig. 1e,f). The increase in pH showed over a ten-fold decrease in hydrogen ion concentration in Giles and a smaller but significant trend in Lacawac (Fig. 1g). Both lakes experienced significant increases in acid neutralizing capacity (ANC), with Giles recovering from a negative alkalinity in the 1990s (Fig. 1h). Browning in both lakes was evident from significant declines in water transparency to both UV and PAR penetration during the study period (Fig. 1i,j). In Giles the 1% UV attenuation depth declined from over 10 m to about 2 m (Fig. 1i), and the 1% PAR attenuation depth declined from 23 m to less than 15 m (Fig. 1j). Declines in water transparency were more modest but also statistically significant in Lacawac (Fig. 1j).

Water temperature and O\(_2\) showed complex, depth-specific changes during browning. Comparison of the average responses in the first versus last five years of the study period reveal that surface water temperatures have warmed by 2–3°C in both lakes, while deeper waters cooled or showed no significant change depending on the lake and depth (Fig. 2a). In Lacawac the waters at intermediate depths trended toward cooling while the deeper waters in Giles cooled by as much as 4°C (Fig. 2a). These depth specific changes in water temperature led to stronger thermal stratification in both lakes. Surface water temperatures were unrelated to mean air temperature during the months preceding sampling in both lakes (Fig. 3a), but were strongly related to water transparency in Lake Giles (Fig. 3b). Hypolimnetic temperatures were not related to spring wind speed in either lake, despite its increasing trend (Fig. 3c), but did show a strong decrease with decreasing water transparency in Lake Giles (Fig. 3d). Increases in wind speed would suggest the potential for greater wind-driven mixing and warmer hypolimnion temperatures; however, this relationship was not observed. Percent saturation of O\(_2\) declined at mid-water depths in Lacawac with the greatest decline at a depth of 6 m, while in Giles the decline was in deeper depths with the greatest decline at 21 m (Fig. 2b). The average (±S.E.M.) depth of O\(_2\) supersaturation (defined here by ≥110% saturation) in Lake Giles shallowed from 12.0 ± 0.61 m during 1988–1992 to 8.0 ± 0.42 m during 2010–2014. This led to opposite trends in O\(_2\) at intermediate depths between about 5–8 m in the two lakes wherein O\(_2\) declined in Lacawac and increased in Giles (Fig. 2b). These changes are likely related to the observed reductions in water transparency and a consequently shallower chlorophyll maximum.

These altered vertical habitat gradients were accompanied by pronounced changes in the pelagic food web, particularly in Giles. Though data are limited to only a few years, extensive efforts to trap planktivorous juvenile (young-of-year, YOY) fish in Lake Giles in 1991 led to no captures, while catch per unit effort (CPUE) in the epilimnion of Lake Giles in the past 3 years averaged 5.5 individuals per 24 h trap period with a CPUE of up to 26 fish in a single trap. A similar increase was observed in YOY fish in Lacawac with an average CPUE of 1.6 and 5.0 in 1991 and 2014 respectively. The two dominant zooplankton grazers in Giles both showed substantial declines. The average May–August abundance of the cladoceran Daphnia in Giles declined to only half of its original numbers, and the calanoid copepods declined by five-fold (Fig. 4a,b). In contrast, the abundance of cyclopid copepods and their small rotifer prey increased in Giles with the cyclopoids increasing by five-fold or more (Fig. 4c,d). In Lacawac only the calanoid copepods declined significantly and their decline was much less pronounced than in Giles (Fig. 4b). No significant trends in chlorophyll a concentration were observed in either Giles (\(\tau = 0.28, p = 0.11, n = 18\)) or Lacawac (\(\tau = −0.22, p = 0.26, n = 16\)) during the study period.
Monthly data on water transparency indicate strong changes in the phenology of both UV and PAR transparency in the two study lakes. Pronounced UV and PAR clear-water phases that were observed early in the study period in both lakes have disappeared in more recent years (Fig. 5). These clear-water phases were historically strongest in Giles where the measured 1% attenuation depth for potentially damaging 320 nm UV increased from 4 m in the spring to over 10 m by mid-summer, and the measured 10% PAR attenuation depth increased from 9 m to 14 m during this same period (Fig. 5a, b). Similar but less pronounced changes in these UV and PAR clear-water phases were observed in Lacawac (Fig. 5c, d).

Discussion
Our long-term observations are generally consistent with short-term experimental, comparative, and modeling studies, and with emerging evidence for the central importance of shading effects during browning9. These shading effects are particularly important in small lakes that are sheltered by the surrounding forest and thus minimally influenced by wind18,19, as are Lacawac and Giles. Evidence includes decreases in UV and PAR transparency, changes in thermal structure with no increase in regional air temperature, increases in O2 depletion, a shift in the O2 maximum to a shallower depth in Giles, and declines in the two primary zooplankton grazers. In contrast, increases in YOY fish and the addition of a trophic level in the zooplankton community are more consistent with either a resource subsidy hypothesis or the release of YOY fish from low pH or high UV exposure12. Regardless of the underlying mechanisms, these ecological consequences of browning have important implications for critical ecosystem services. The changes in food web structure suggest an increase in the trophic transfer to higher trophic levels such as fish and are relevant to fishery productivity14. The decreases in water transparency to UV and PAR decrease the potential for solar disinfection of parasites and pathogens20. The increase in DOM itself can increase the production of carcinogenic disinfection byproducts in drinking water as well as the cost of water purification for municipal use21. Browning may also influence the temperature and O2 dependent production of CO2 and CH4 from lake sediments and the role of inland waters as a globally important source of greenhouse gases22. Browning may either increase23 or decrease24 the sequestration of greenhouse gases depending on the balance between mineralization of DOC to carbon dioxide and burial of organic carbon in the sediments. Colder temperatures and anoxia in the hypolimnion may increase deposition of fixed carbon in the sediments by reducing decomposition rates, but also increase the production of methane, which is a much more powerful greenhouse gas than is the carbon dioxide produced under well oxygenated conditions. Even short-term extreme precipitation events can induce browning that reduces water clarity, shifts lake metabolism from autotrophy to heterotrophy25, alters vertical distribution of important zooplankton grazers26, and depletes O2, leading to hypoxia or anoxia27,28.

The stronger long-term trends in Giles versus Lacawac are consistent with the concept that less transparent, higher DOM lakes are more resistant to browning while clear-water lakes are more sensitive sentinels of ecosystem level responses to browning13. The stronger thermal stratification in both lakes is likely due to an increased absorption of sunlight in the browner surface waters, and a consequent reduction in heat energy transferred to deeper waters29. Mechanistic models have clearly demonstrated the potential for DOM-induced changes in transparency to alter thermal structure in lakes to the extent observed in our two study lakes including warmer surface water temperatures and cooler deeper waters, as well as stronger effects in more transparent lakes13. In very clear lakes like Giles, these effects are particularly pronounced due to the strong changes in water transparency in response to increases in DOM. PAR penetrated to the bottom of Giles (1% of surface irradiance was 23 m, Fig. 1j) early in the
study period, but it often penetrated less than 15 m later in the study period, thus severely reducing the potential for solar heating of deeper waters in more recent years.

The changes in the phenology of water transparency were not predicted by shorter-term studies of browning and signal important changes in lake ecosystem structure and function. Zooplankton grazers such as *Daphnia* play an important role in grazing down spring phytoplankton blooms, leading to a seasonal increase in water transparency. Thus the disappearance of the PAR clear-water phase during browning is consistent with the observed decline in zooplankton grazers. The disappearance of the UV clear-water phase likely signals a change in DOM processing15. Breakdown of DOM by sunlight has recently been demonstrated to be more important in high latitude lakes than previously appreciated24, and is known to be responsible for a majority of the seasonal UV clear-water phases observed in the past in Lake Giles15.

It is critical to note that we have examined only two lakes, and the responses of other lakes to browning are likely to vary as a function of not only transparency, but lake and catchment geomorphology, geology, local hydrology, atmospheric deposition, and other environmental conditions. However, small lakes like these study lakes dominate numerically on a global scale and are critical to the processing and fate of organic matter27. Browning may also have consequences for larger lakes and coastal ocean waters. In the Arctic thawing permafrost is releasing large amounts of isoluble and photolabile DOM to inland and coastal waters and altering underwater UV environments as well as

---

**Figure 3. Relationships between water temperatures and air temperature, wind speed, and water transparency.** The relationships between (a) July mean epilimnion water temperature and average spring (March – mid-July date of sampling) air temperature (Lacawac $\tau = 0.174, P = 0.21$; Giles $\tau = 0.117, P = 0.40$), and (b) July mean epilimnion water temperature and water transparency measured as $a_{4320nm}$ (Lacawac $\tau = 0.18, P = 0.28$; Giles $\tau = 0.40, P = 0.01$), and between (c) July mean hypolimnion water temperature and average spring (March-April) wind speed (Lacawac $\tau = -0.004, P = 1$; Giles $\tau = -0.27, P = 0.09$). Relationships between (d) July mean hypolimnion water temperature and water transparency measured as $a_{4320nm}$ (Lacawac $\tau = -0.242, P = 0.14$; Giles $\tau = -0.596, P < 0.001$). Lines represent linear regression models with bold indicating significant trends ($P < 0.05$), Kendall non-parametric tests.
greenhouse gas production. Mesocosm experiments in coastal ocean waters have demonstrated that O2 depletion can cause P regeneration from the sediments as well as generate increases in DOM, leading to positive feedbacks that are similar to those described in lakes. Novel UV-based optical approaches coupled with remote sensing can identify terrestrially derived lignin components of DOM and enable visualization of the influence of browning on regions as expansive as the Arctic Ocean. These kinds of advanced sensor approaches will be critical to resolving the extent to which the ecological consequences and threat to ecosystem services observed in browning lakes are also a threat to coastal ocean ecosystems and global greenhouse gas production. While the greater importance of wind-driven mixing and a greater buffering capacity will likely mask the effects of browning in many larger lakes and oceans, recovery from acidification as well as interactions with iron may enhance the browning and hence increase thermal stratification of smaller inland lakes.

Our data indicate that the widespread increases in surface water temperatures and thermal stratification that have been associated with climate warming in lakes and oceans may be augmented by increases in precipitation-induced transport of terrestrially-derived DOM. Thus in regions where lakes and coastal oceans are browning, the wetter as well as the warmer conditions associated with climate change may combine to provide a double-edged sword that more strongly increases the strength of thermal stratification and potential for associated threats to ecosystem services across these inland and coastal ecosystems.

Figure 4. Interannual changes in the zooplankton. (a) The cladoceran Daphnia, a strong grazer on phytoplankton. (b) Calanoid copepods, omnivorous grazers on phytoplankton. (c) Predatory cyclopoid copepods. (d) Rotifers, a preferred prey of cyclopoid copepods. Lines are linear regressions; bold indicates significant trends ($P < 0.05$): Giles Daphnia $\tau = -0.43$, $P = 0.022$, $n = 16$; Lacawac Daphnia $\tau = -0.19$, $P = 0.381$, $n = 14$; Giles calanoids $\tau = -0.57$, $P = 0.003$, $n = 16$; Lacawac calanoids $\tau = -0.43$, $P = 0.038$, $n = 14$; Giles cyclopoids $\tau = 0.38$, $P = 0.043$, $n = 16$; Lacawac cyclopoids $\tau = -0.03$, $P = 0.913$, $n = 14$; Giles rotifers $\tau = 0.41$, $P = 0.038$, $n = 15$; Lacawac rotifers $\tau = -0.30$, $P = 0.193$, $n = 12$. 

![Figure 4](image_url)
Methods
Water samples were collected from a single depth in the epilimnion using a Van Dorn bottle. All variables except zooplankton were measured from a single sample collected between mid-July and mid-August, during the period of maximum thermal stratification. Strength of stratification was calculated by subtracting the average hypolimnion temperature from the average epilimnion temperature. In both lakes, a period of peak stratification (±10%) typically extends over a period of 7–11 weeks from early to mid-July into late August or early September. Zooplankton densities are annual averages from May to August. For measurements of DOC and absorbance, samples were filtered through pre-combusted 0.7 μm glass-fiber filters shortly (generally within 48 h) after collection. For DOC, samples were analyzed through high temperature oxidation in a TOC analyzer. Decadal dissolved absorbance at 320 nm was measured using a spectrophotometer and converted to Naperian dissolved absorption coefficients (a_d) using the following relationship:

\[ a_d = \frac{2.303 \times D}{r} \]

Where D is the raw absorbance value at 320 nm and r is the path length (m) of the quartz cuvette. The a^*320 nm was calculated as the absorbance at 320 nm per unit DOC (mg L^-1). Raw water samples were analyzed for pH using an Orion pH meter equipped with a Ross epoxy-body combination electrode, calibrated at pH 7.00 and 4.00 using commercial pH buffers. Following pH measurements, ANC was determined through potentiometric titration with dilute HCl (1988–1993) or H_2SO_4 (0.01 N for Giles and 0.1 N for Lacawac). Titration points between pH 4.4 and 3.7 were plotted after Gran transformation, and ANC was calculated as μEq L^-1. For Chl a analysis, water samples were analyzed through high temperature oxidation in a TOC analyzer. Decadal dissolved absorbance at 320 nm was measured using a spectrophotometer and converted to Naperian dissolved absorption coefficients (a_d) using the following relationship:

\[ a_d = \frac{2.303 \times D}{r} \]

Where D is the raw absorbance value at 320 nm and r is the path length (m) of the quartz cuvette. The a^*320 nm was calculated as the absorbance at 320 nm per unit DOC (mg L^-1). Raw water samples were analyzed for pH using an Orion pH meter equipped with a Ross epoxy-body combination electrode, calibrated at pH 7.00 and 4.00 using commercial pH buffers. Following pH measurements, ANC was determined through potentiometric titration with dilute HCl (1988–1993) or H_2SO_4 (0.01 N for Giles and 0.1 N for Lacawac). Titration points between pH 4.4 and 3.7 were plotted after Gran transformation, and ANC was calculated as μEq L^-1. For Chl a analysis, water samples were analyzed through high temperature oxidation in a TOC analyzer. Decadal dissolved absorbance at 320 nm was measured using a spectrophotometer and converted to Naperian dissolved absorption coefficients (a_d) using the following relationship:

\[ a_d = \frac{2.303 \times D}{r} \]
two minutes) was also employed at the midpoint of a 48 h freezer extraction from 2010–2014\textsuperscript{35}. Following fluorometric analysis, samples were acidified with 0.03N HCl to differentiate Chl-a from phaeophytin. Direct inter-comparisons of the acetone and acetone/methanol solvents for Chl extraction and the opposite (non-significant) Chl a trends in the two lakes indicate no methodological artifacts in the Chl a data.

Temperature, O\textsubscript{2}, and light were taken in situ at a 1 meter depth or higher resolution. Temperature (1988–1992) and O\textsubscript{2} (all years) were measured at 1-meter intervals using a YSI oxygen meter. Temperature from 1993–2014 was measured using a subsmerisible radiometer. The difference in both temperature and O\textsubscript{2} (in percent saturation) at each meter depth was calculated using the five most recent years (2010–2014) and the five earliest years (1988–1992). UV and PAR light profiles and the measured depth to which 1% of surface irradiance penetrated were collected using one of three methods, including a Li-Cor sensor (PAR; 1988–1993), PUV radiometer (PAR and UV; 1993–2003) or a BIC radiometer with a deck cell (PAR and UV; 2003–2014).

Zooplankton densities are from integrated net tows taken through the whole water column. In Giles, the zooplankton community consists of Daphnia catawba, Cyclops scutifer, calanoid copepods (87% Leptodiaptomus minutus, 12% Aglaodiaptomus spatulocrenatus), and rotifers (29% Keratella spp., 27% Kellicottia spp., 26% Polyarthra spp., and 11% Gastropus spp.). In Lacawac, there are two species of Daphnia (72% D. ambiguus and 27% D. catawba), a single species of calanoid copepod (L. minutus), two species of cyclopoid copepods (64% C. scutifer and 36% Mesocyclops edax), and rotifers (35% Keratella spp., 30% Polyarthra spp., and 18% Kellicottia spp.). No significant trends in the relative abundance of the two cyclopoid copepod species in Lacawac were observed during the study period. While some calanoid copepods feed on rotifers\textsuperscript{36–37}, the dominant rotifer genera in these lakes (Keratella, Kellicottia, and Polyarthra) have effective defense or avoidance responses to Diaptomid predators and are not the preferred prey of the very small body sized dominant calanoid, L. minutus, in our two study lakes. From 1990–1993 and 2005–2008 Daphnia were collected with a 202-μm bongo net and counted in a Bogorov chamber, except in Lacawac in 2008 where a Wisconsin-style 48-μm net and Sedgewick Rafter chamber were employed. In 1997, 2001, 2009–2011, and 2014 Daphnia were collected in a 48-μm Wisconsin-style bongo net and counted in a Bogorov chamber. Intercomparison of the nets demonstrated similar high collection efficiencies for the two mesh sizes. For most years, both calanoid and cyclopoid copepods, as well as rotifers were collected with a 48-μm net and counted in a Sedgewick Rafter chamber. Exceptions include 1990–1993 when female cyclopoid copepods were collected with a 202-μm Wisconsin-style bongo net and counted in a Bogorov chamber; and 2012 and 2013, when all copepods and Daphnia were collected using a 153-μm Wisconsin-type bongo net and counted in a Bogorov chamber. Copepod analyses include males, females and copepodids. Daphnia counts include juveniles and adults (not neonates). Total rotifer abundance includes all individual rotifers with the exception of the genus Conochilus, which can be found both as a colony and individually, and was therefore excluded from analysis.

Data on YOY larval fish represent 24 h trapping efforts using clear plastic replicate (2 separate sides) minnow traps suspended at a depth of 1.5 m, deployed between 15:00–17:00 h, and expressed as C.PUE. The traps used in the 1990’s and the 2010’s were the same identical traps. YOY trapping in Lacawac was conducted on 31 May – 1 June, 13–14 July, and 25–26 August, 1991 for a total of 12 traps deployed over each 2 day period (6 traps per 24 hour period that were emptied and immediately redeployed, with a similar effort in Lake Giles). In 2012 trapping occurred in Lake Giles on three different dates (17 June, 3 and 20 July) with six traps used per date. In 2013 only two traps were used per date, but the trapping was done on 12 different dates from early July through mid-August (2, 3, 9, 12, 16, 19, 23, 25, 30 July and 2, 6 and 8 August). In 2014 trapping in both Giles and Lacawac occurred on a single date (3, 13 July, respectively) with two traps per lake. The species captured in Lake Giles from 2012–2014 were approximately 90% Micropterus salmoides on all trap dates with up to 10% of the individuals being Lepomis spp. Fish species in Lacawac during the 1991 trapping included Perca flavescens as the only species captured 31 May – 1 June, 50% Lepomis spp. and 50% P. flavescens during the 13–14 July trapping, and only Lepomis spp. on 25–26 August. In 2014, the species present in Lacawac were largely M. salmoides and M. dolomieu with approximately 15% of the individuals captured being Lepomis spp.

Meteorological data on minimum and maximum daily air temperature and precipitation were based on daily data obtained from the National Oceanic and Atmospheric Administration National Climate Data Center using the nearest station (<15 km to the two lakes) located in Hawley, Pennsylvania, USA (GHCND:USC00363758). Mean daily air temperature was calculated from averaging daily minimum and daily maximum air temperature. Annual mean air temperature was the previous 365-day average of the temperature profile, inclusive of the sampling day (roughly 16 July – 15 July of the following year). Spring mean air temperature was calculated from the average 01 March to day of sampling for each year. Data for annual wet deposition of SO\textsubscript{2}\textsuperscript{−2} and NO\textsubscript{3}\textsuperscript{−} were downloaded from the National Atmospheric Deposition Program monitoring site at the nearest station (40 km to Lacawac, 23 km to Giles) located in Milford, Pennsylvania, USA (PA72\textsuperscript{28}). Wet deposition data prior to 1994 were excluded from our analysis due to contamination in the samples leading to overestimation, and are not comparable to deposition data after 1994. Wind speed data (1993–2014) were collected from an on-lake buoy at Lacawac using an anemometer at 2 m height (maintained by B.R. Hargreaves, Lehigh University). Spring wind speed was calculated from the average 01 March – 30 April daily wind speed for each year.

Statistical Analysis

Mann-Kendall non-parametric trend tests were used for temporal trends of all variables\textsuperscript{36,40}. Significance of trends were considered at the P < 0.05 level. For temporal trend models, LOESS 20-moving averages were used for most variables. For zooplankton trends, linear models were used instead due to many years of missing data and the high variability (Mann-Kendall trend tests were still used to assess significance). Kendall univariate tests were run to assess relationships between spring air temperature and epilimnion temperature, spring wind speed and hypolimnion temperature, and ad\textsubscript{320}nm and both epilimnion and hypolimnion temperature (independently). All
statistical analyses and models were run under the "Kendall" and "wq" packages and were completed in R (version 3.1.1).

References
1. Monteith, D. T. et al. Dissolved organic carbon trends resulting from changes in atmospheric deposition chemistry. Nature 450, 537–541 (2007).
2. Brothers, S. et al. A feedback loop links brownification and anoxia in a temperate, shallow lake. Limnol. Oceanogr. 59, 1388–1398 (2014).
3. Couture, S., Houle, D. & Gagnon, C. Increases of dissolved organic carbon in temperate and boreal lakes in Quebec, Canada. Environ. Sci Pollut. R. 19, 361–371 (2012).
4. Clark, J. M. et al. The importance of the relationship between scale and process in understanding long-term DOC dynamics. Sci. Total Environ. 408, 2768–2775 (2010).
5. Weyhenmeyer, G. A., Prairie, Y. T. & Tranvik, L. J. Browning of boreal freshwater coupled to carbon-iron interactions along the aquatic continuum. PLoS One 9, e88104 (2014).
6. Melillo, J., Richmond, T. C. & Yohe, G. W. Climate Change Impacts in the United States: The third national climate assessment. (U.S. Global Change Research Program, 2014).
7. Williamson, C. E. et al. Lakes as sensors in the landscape: Optical metrics as scalable sentinel responses to climate change. Limnol. Oceanogr. 59, 840–850 (2014).
8. Solomon, C. T. et al. Ecosystem consequences of changing inputs of terrestrial dissolved organic matter to lakes: Current knowledge and future challenges. Ecosystems, 10.1007/s10021-015-9848-y (2015).
9. Seekell, D. et al. The influence of dissolved organic carbon on primary production in northern lakes. Limnol. Oceanogr. 60, 1276–1285 (2015).
10. Kissman, C. E., Williamson, C. E., Rose, K. C. & Saros, J. E. Response of phytoplankton in an alpine lake to inputs of dissolved organic matter through nutrient enrichment and trophic forcing. Limnol. Oceanogr. 58, 867–880 (2013).
11. Kelly, P. T., Solomon, C. T., Weidel, B. C. & Jones, S. E. Terrestrial carbon is a resource, but not a subsidy, for lake zooplankton. Ecology 95, 1236–1242 (2014).
12. Williamson, C. E., Hargreaves, B. R., Orr, P. S. & Lovera, P. A. Does UV play a role in changes in predation and zooplankton community structure in acidified lakes? Limnol. Oceanogr. 44, 774–783 (1999).
13. Read, J. & Rose, K. C. Physical responses of small temperate lakes to variation in dissolved organic carbon concentrations. Limnol. Oceanogr. 58, 921–931 (2013).
14. Finstad, A. G., Helland, I. P., Ugedal, O., Hesthagen, T. & Hessen, D. O. Unimodal response of fish yield to dissolved organic carbon. Ecol. Lett. 17, 36–43 (2014).
15. Williamson, C. E., De Lange, H. J. & Leech, D. M. Do zooplankton contribute to an ultraviolet clear-water phase in lakes? Limnol. Oceanogr. 52, 662–667 (2007).
16. Strock, K. E., Nelson, S. J., Kahl, J. S., Saros, J. E. & Mc Dowell, W. H. Decadal trends reveal recent acceleration in the rate of recovery from acidification in the northeastern U.S. Environ. Sci. Technol. 48, 6681–6689 (2014).
17. Sobek, S., Tranvik, L. J., Prairie, Y. T., Kortelainen, P. & Cole, J. J. Patterns and regulation of dissolved organic carbon: An analysis of 7,500 widely distributed lakes. Limnol. Oceanogr. 52, 1208–1219 (2007).
18. Markfort, C. D. et al. Wind sheltering of a lake by a tree canopy or bluff topography. Water Resour. Res. 46, 10.1029/2009wr007759 (2010).
19. Read, J. S. et al. Lake-size dependency of wind shear and convection as controls on gas exchange. Geophys. Res. Lett. 39, 10.1029/2012gl051886 (2012).
20. Overholt, E. P. et al. Solar radiation decreases parasitism in Daphnia. Ecol. Lett. 15, 47–54 (2012).
21. Ledesma, J. L. J., Köhler, S. J. & Futter, M. N. Long-term dynamics of dissolved organic carbon: Implications for drinking water supply. Sci. Total Environ. 432, 1–11 (2012).
22. Marotta, H. et al. Greenhouse gas production in low-latitude lake sediments responds strongly to warming. Nat. Clim. Change 4, 467–470 (2014).
23. Fortino, K., Whalen, S. & Johnson, C. Relationships between lake transparency, thermocline depth, and sediment oxygen demand in Arctic lakes. Inland Waters 4, 79–90 (2014).
24. Cory, R. M., Ward, C. P., Crump, B. C. & Kling, G. W. Sunlight controls water column processing of carbon in arctic fresh waters. Science 345, 925–928 (2014).
25. Sadro, S. & Melack, J. The effect of an extreme rain event on the biogeochemistry and ecosystem metabolism of an oligotrophic high-elevation lake. Arct. Antarct. Alp. Res. 45, 222–231 (2012).
26. Rose, K. C. et al. The role of UV and fish in regulating the vertical distribution of Daphnia. Limnol. Oceanogr. 57, 1867–1876 (2012).
27. Verpoorter, C., Kutser, T., Seekell, D. & Tranvik, L. A global inventory of lakes based on high-resolution satellite imagery. Geophys. Res. Lett. 41, 6396–6402 (2014).
28. Skoog, A. C. & Arias-Esaquiel, V. A. The effect of induced anoxia and reoxygenation on benthic fluxes of organic carbon, phosphate, iron, and manganese. Sci. Total Environ. 407, 6085–6092 (2009).
29. Fichot, C. G. et al. Pan-Arctic distributions of continental runoff in the Arctic Ocean. Sci. Rep. 3, 1–6 (2013).
30. Gao, K. S. et al. Rising CO2, and increased light exposure synergistically reduce marine primary productivity. Nat. Clim. Change 2, 519–523 (2012).
31. Kraemer, B. M. et al. Morphometry and average temperature affect lake stratification responses to climate change. Geophys. Res. Lett. 42, 4981–4988 (2015).
32. Schneider, P. & Hook, S. J. Spatial distribution of inland water bodies show rapid surface warming since 1985. Geophys. Res. Lett. 37, 1–5 (2010).
33. Winslow, L. A., Read, J. S., Hansen, G. J. A. & Hanson, P. C. Small lakes show muted climate change signal in deepwater temperatures. Geophys. Res. Lett. 42, 355–361 (2015).
34. Kirk, J. T. O. et al. Measurement of UV-B radiation in two freshwater lakes: an instrument intercomparison. Archiv. Hydrobiol. 43, 71–99 (1994).
35. Pechar, L. Use of an acetone:methanol mixture for the extraction and spectrophotometric determination of chlorophyll-a in phytoplankton. Archiv. Hydrobiol. Supplement 78, 99–117 (1987).
36. Williamson, C. E. Predator-prey interactions between omnivorous diatomid copepods and rotifers: The role of prey morphology and behavior. Limnol. Oceanogr. 32, 167–177 (1987).
37. Williamson, C. E. & Butler, N. M. Predation on rotifers by the suspension-feeding calanoid copepod Diaptomus pallidus. Limnol. Oceanogr. 31, 393–402 (1986).
38. NADP (National Atmospheric Deposition Program). National Trends Network http://nadp.isws.illinois.edu/ntn/ (Accessed 01 March 2015).
39. Mann, H. B. Non-parametric tests against trend. Econometrica 13, 163–171 (1945).
40. Kendall, M. G. Rank Correlation Methods. 4th edn, (Charles Griffin, 1975).
41. McLeod, A. I. Kendall: Kendall rank correlation and Mann–Kendall trend test. R package version 2.2 http://CRAN.R-project.org/package=Kendall (2011).
42. Jassby, A. D. & Cloern, J. E. Some tools for exploring water quality monitoring data. R package version 0.4-1. http://CRAN.R-project.org/package=wq (2014).
43. R. Core Team. R: A Language and Environment for Statistical Computing http://www.R-project.org/. (R Foundation for Statistical Computing, 2014).

Acknowledgements
This paper is dedicated to the late Robert E. Moeller. Bruce Hargreaves and Don Morris assisted with the collection of some of the long-term data. We thank the many students and technicians who have been involved in the collection of the long term data through the years, and the Lacawac Field Station and Blooming Grove Club for providing access to the lakes.

Author Contributions
C.W. designed and led the long-term study, sampled the lakes, provided the lake data, and wrote the manuscript. E.O. collected and analyzed data. R.P. helped collect and process data, conducted statistical analyses, and produced figures. T.L. provided larval fish data. J.B. helped collect data. L.B.K. helped collect data and contributed numerical analysis on dissolved oxygen. E.M. compiled data and contributed to data analysis. R.M. helped with data collection and collation. All authors except R.M. edited the manuscript.

Additional Information
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Williamson, C. E. et al. Ecological consequences of long-term browning in lakes. Sci. Rep. 5, 18666; doi: 10.1038/srep18666 (2015).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/