Evidence for the Importance of Atmospheric Nitrogen Deposition to Eutrophic Lake Dianchi, China

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ABSTRACT: Elevated atmospheric nitrogen (N) deposition has significantly influenced aquatic ecosystems, especially with regard to their N budgets and phytoplankton growth potentials. Compared to a considerable number of studies on oligotrophic lakes and oceanic waters, little evidence for the importance of N deposition has been generated for eutrophic lakes, even though emphasis has been placed on reducing external N inputs to control eutrophication in these lakes. Our high-resolution observations of atmospheric depositions and riverine inputs of biologically reactive N species into eutrophic Lake Dianchi (the sixth largest freshwater lake in China) shed new light onto the contribution of N deposition to total N loads. Annual N deposition accounted for 15.7% to 16.6% of total N loads under variable precipitation conditions, 2-fold higher than previous estimates (7.6%) for the Lake Dianchi. The proportion of N deposition to total N loads further increased to 27−48% in May and June when toxic blooms of the ubiquitous non-N2 fixing cyanobacteria Microcystis spp. are initiated and proliferate. Our observations reveal that reduced N (59%) contributes a greater amount than oxidized N to total N deposition, reaching 56−83% from late spring to summer. Progress toward mitigating eutrophication in Lake Dianchi and other bloom-impacted eutrophic lakes will be difficult without reductions in ammonia emissions and subsequent N deposition.

INTRODUCTION

Human activities have significantly increased emissions of reactive nitrogen (N, including reduced and oxidized forms) to the atmosphere and their resultant deposition. Results of global atmospheric chemical transport models indicated that atmospheric deposition of N has increased by approximately 3-fold since preindustrial times, especially over the East and South Asia. The ecological effects of elevated atmospheric N deposition have been observed or modeled for oligotrophic lakes and oceanic waters, showing that such N inputs can stimulate phytoplankton growth. This also leads to an increase in the stoichiometric N/phosphorus (P) ratio, which can induce a shift from N-limited to P-limited phytoplankton. In contrast, the contribution made by atmospherically deposited N to the total N input for eutrophic lakes has rarely received attention because it was commonly considered to be much less than N inputs from watersheds to these lakes. However, recent evidence from the few field observations demonstrated that the contribution of N deposition cannot be neglected compared to riverine inputs for eutrophic lakes. For instance, Luo et al. measured wet N deposition in mesotrophic Lake Taihu, the third largest freshwater lake in China, indicating that atmospheric inputs of N accounted for approximately 14% of the total N input. Therefore, atmospheric N deposition should be considered in the management of eutrophic lakes.
of total N inputs. A similar percentage was reported for a mesotrophic lake in Venezuela\textsuperscript{11} (Lake Maracaibo, wet deposition, 19%). Furthermore, the percentage increased considerably when both dry and wet deposition (i.e., gaseous, particulate and dissolved N) were included.\textsuperscript{12} With long-term controls of point and nonpoint sources from watersheds taking place in developed countries, atmospheric N deposition is becoming an increasingly dominant N source to these waters (e.g., 15–42% of the total N inputs\textsuperscript{13} in the northeastern and mid-Atlantic regions of the United States).

However, the contribution made by atmospheric N deposition to total inputs remains challenging to accurately assess, especially at a high-resolution temporal scale. Multiple factors are associated with the difficulty in assessment, especially when quantifying dry deposition and N inputs from watersheds to receiving waters. First, gaseous N (i.e., ammonia [NH\textsubscript{3}], nitrogen dioxide [NO\textsubscript{2}], nitric acid and nitrous acid [HNO\textsubscript{2}/HNO\textsubscript{3}]) deposition have generally been ignored or are only partially considered by current observation networks globally (e.g., EANET,\textsuperscript{14} CASTNET,\textsuperscript{15} AMoN,\textsuperscript{15} IMPROVE NH\textsubscript{x},\textsuperscript{15} and EMEP\textsuperscript{16}) and in previous regional assessments.\textsuperscript{12} Likewise, wet or bulk deposition of inorganic N (NH\textsubscript{4}+–N + NO\textsubscript{3}−–N) has been measured systematically in China, but organic N was seldom analyzed.\textsuperscript{12} Those omissions may result in a considerable underestimation of direct atmospheric N deposition to these waters. Second, most measurements applying passive samplers yield ambient exposures on a monthly interval, mainly due to low concentrations of gaseous or particulate N.\textsuperscript{12} Although active samplers (e.g., denuder systems) can be deployed to determine hourly or daily concentrations of gaseous constituents and fine-fraction particulates,\textsuperscript{17} they are unsuitable for long-term field observations without a power supply.\textsuperscript{12} In addition, riverine N inputs (i.e., N concentration × river discharge) were generally determined by using process-based models that simulate the N available for transport to the waters from different watershed N sources (runoff from agriculture, urban areas and upland forests, point sources).\textsuperscript{13,18,19} A key limitation in such models is the large uncertainties arising from model structure and parameter choices.\textsuperscript{20} Such uncertainties can be reduced by performing observations of river discharge and N concentrations at high spatial and temporal resolutions.

The primary objective of this study is to test the hypothesis that atmospheric N deposition makes an important contribution to total N loads for eutrophic Lake Dianchi, the sixth largest freshwater lake in China, through high-resolution (daily or biweekly) systematic observations of N deposition and inputs. Dry and wet deposition fluxes of all N species and riverine N inputs were quantified and their annual and biweekly contributions to total N loads were determined. In addition, we considered the reliability of the fluxes as well as the importance of N deposition on lake eutrophication, and implications for lake ecosystem management.
MATERIALS AND METHODS

Study Area. Lake Dianchi is located in Southwest China (24°29′ to 25°28′N, 102°29′ to 103°01′E). It has a surface area of 310 km² at the elevation of 1887.5 m, a mean depth of 5.3 m, a shoreline of 150 km, and water retention time of 3.5 years.1 Watershed area for Lake Dianchi is 2920 km², of which 1088.6 km² is controlled by 10 large to medium-size reservoirs that have no outflows. The lake is artificially divided into two segments by an embankment and a dam, where the area of the northern part (Coahai) is ~12 km² surrounded by urban area of Kunming city and the southern part (Waahai, 298 km²) is bordered by an intensively managed farmland. Land use in the watershed is primarily forest (47%), cropland (20%), and urban areas (16%; Figure 1). The total population in the Lake Dianchi Watershed has increased steadily from 1.8 million in 1992 to 4.1 million in 2015. Consequently, the lake has suffered from severe eutrophication, despite the fact that approximately $2 billion RMB (equal to 7.7 billion in USD) has been invested to mitigate lake eutrophication by the Chinese Central Government during 1996–2015.2

Dry N Deposition Samples around the Lake. Sampling of dry N deposition, including gaseous and particulate N, was conducted from April 1, 2010 to March 31, 2011. Five deposition monitoring stations were evenly located around Lake Dianchi (Figure 1), including Kunming (KM), Baofeng (BF), Chenggong (CG), Jinning (JN), and Haikou (HK). For each station, Ogawa passive samplers (Ogawa & Co., FL, U.S.A.) were used to collect ammonia [NH₃] and nitrogen dioxide [NO₂] samples, which showed no significant differences from the continuous active samplers used in previous studies.3 Another passive sampler (USDA Forest Service, CA, U.S.A.) was used for collecting nitrous acid/nitric acid [HNO₂/HNO₃] samples. Previous comparative experiments showed that this sampler was reliable for determining wide ranges of ambient HNO₂/HNO₃.4 Detailed descriptions of these passive samplers can be found in Roadman et al.5 and Bytnerowicz et al.6 High volume aerosol samplers (Laoshan Elec. Inc., Qingdao, China) were applied to collect total suspended particulates (TSP) at a flow rate of 1.05 m³ min⁻¹. Three replicate samplers for dry N deposition were installed at a height of 3.5 m from the ground and at a distance of <50 m from the shoreline of Lake Dianchi. Passive samplers for gaseous N were exposed for 2 weeks, while active samplers for particulate N were operated 24 h every week on Monday, but was manually halted during rainfall events. The exposed samplers were placed in a resealable plastic bag, the bagged sampler was placed into a brown airtight container taken to laboratory for analysis. Detailed information for collection and measurements are provided in Text S1 of the Supporting Information (SI).

Wet N Deposition Samples around the Lake. At five deposition monitoring stations, rainfall samples were taken on an event basis using a stainless steel funnel and plexiglass bottle, along with precipitation automatically measured by tipping-bucket rain gauge (Guoxinhuaayuan Tech. Inc., Beijing, China). The collector was kept completely closed and was opened only at the beginning of a rainfall event by the staff from the Kunming Environmental Monitoring Center (KEMC). The collected samples were then immediately stored in acid-washed high density polyethylene (HDPE) bottles. To prevent biological utilization of deposited N, chloroform was used as a biocide in the wet bottle. A cap covered the “wet” bottle which, in combination with the inherently low rainwater pH of 4 to 5, minimized the potential for NH₃ volatilization. Wet deposition was sampled for all rainfall events having precipitation of >2 mm. All collected samples were frozen at ~18 °C at each site until delivery to the laboratory for analysis. Collectors were cleaned for each event with 1% HCl and Nanopure water.

River Discharge and Water Samples. Although river discharges data were not available except for Panlongjiang (No. 8) and Baoxiang (No. 10) Rivers, water levels in the other 17 rivers were continuously measured in this study (Figure 1). These 19 rivers contribute more than 95% of lake’s streamflow.7 The rating curves were developed by simultaneously measuring water levels and the current velocities for a cross-section of each river following the USGS guideline for the current-meter method,8 where the sizes of cross-section (i.e., width, depth, and area) were measured for each river (Table S1). To ensure rating curves over a wide range of flow rates, enhanced observations were conducted for three stormwater events (precipitation >30 mm day⁻¹). Each sampling procedure lasted more than 48 h per event. The 10-min water levels were continuously observed for each tributary by HOBO data logger (U20-001-01, Onset Computer Corporation, MA, U.S.A.), and the current velocities were measured by the Global Water flow probe (FP201) in each subsection of a channel cross-section. Water level was further converted into daily discharge based on the derived rating curves for each river. In addition, vertical water samplers were used to collect water samples of 19 rivers on biweekly interval. The procedure was referenced to the Technical Specifications Requirements for Monitoring of Surface Water and Waste Water in China (HJ/T91–2002). As all rivers have width of ≤50 m and depth of ≤5 m (Table S1), one sample was collected in the river centerline at 0.5 m below or half depth of the river. All water samples were rapidly transported to the KEMC for analysis.

Chemical Analysis. After the exposures, all filters for gaseous and particulate N were placed in polyethylene tubes into which 8 and 20 mL distilled/deionized water was added, respectively. The tubes were shaken on a shaker for 15 min at midrange speed. The extracted solutions of dry deposition, wet deposition samples, and water samples were then filtered with a 0.45 μm syringe filter. N quantities in the filtered solutions were analyzed following the Standard Methods for the Examination of Water and Wastewater9 in China. Ammonium nitrogen [NH₄⁺–N], nitrate nitrogen [NO₃⁻–N], total nitrogen (TN), and Kjeldahl nitrogen (KTN) were measured by Nessler’s reagent colorimetric method, ultraviolet spectrophotometric method, K₂S₂O₈ oxidation-ultraviolet spectrophotometric method, as well as Semi-Micro-Kjeldahl method, respectively. Organic N was determined by the difference between KTN and NH₄⁺–N. The detection limits for NH₄⁺–N, NO₃⁻–N, TN, and KTN are 0.025, 0.02, 0.05, and 0.025 mg N L⁻¹, respectively. For gaseous N deposition, ambient concentrations of NH₃, NO₂, and HNO₂/HNO₃ were calculated as product of their quantities collected in passive samplers and conversion coefficients divided by exposure time (see details at http://ogawaus.com/wp-content/uploads/2014/04/ConcentrationInAmbientAir.pdf ConcentrationInAmbientAir.pdf and Bytnerowicz et al.6). Additionally, for each batch of field exposed samples for gaseous and particulate N, three blank filters were also prepared to represent extracts from unexposed samplers that are prepared in the laboratory and transported between laboratory and monitoring sites. The blank samples were extracted and analyzed using the same methods as the exposed samples. The details of blank samples can be found in Text S1.
Estimation of Deposition Fluxes and Riverine Inputs. Biweekly flux of dry deposition (mg km⁻² s⁻¹) was calculated as a product of N concentration and deposition velocity (V_d, m s⁻¹) at each site. The concentrations of gaseous and particulate N were determined as described above. V_d of gaseous N and particulate N over lake surface was estimated by a well-tested deposition velocity model combined with the on-site hourly meteorological data. One of the national meteorological stations was located at the north end of Lake Dianchi (Figure 1), where meteorological variables used in the model included air temperature, relative humidity, air pressure, solar radiation, wind speed at 10-m height, precipitation, and dew point on hourly intervals. Prior to the estimate, V_d was converted into wind speed at 3.5-m height, v, according to a power law (i.e., v = v_10 × (3.5/10)°, where α was set as 0.35). Gas V_d over Lake Dianchi was calculated using the big-leaf resistance analogy model. In contrast to vegetation, roughness length was set as 0.0003 m based on the results for the East China Sea, and surface resistance was set as 0 s m⁻¹ for NH₃ and HNO₃ but 20,000 s m⁻¹ for NO₂. Particulate V_d was parametrized according to Slinn, where parameters related to surface resistance and gravitational settling velocity were determined according to Zhang et al. Details of these models were described in Text S2. In addition, the flux of wet deposition was calculated as a product of N concentration and rainfall amount for each rainfall event and monitoring site.

Daily riverine N inputs were calculated as a function of river discharge, by means of Load Estimator (LOADEST):  

\[ \ln(L) = a_0 + a_1 \ln Q + a_2 \ln Q^2 + a_3 \sin(2\pi d\text{time}) + a_4 \cos(2\pi d\text{time}) + a_5 \text{dtime}^2 + \epsilon \]  

where Q is daily river discharge, dtime is decimal time, a_0~ a_5 are the fitted coefficients in the multiple regression model, and \( \epsilon \) is estimate error. Note that the biweekly TN concentration was multiplied by the corresponding daily discharge so that eq 1 is a function of load \( (L) \) instead of concentration. Within LOADEST, the model to estimate N loads was set to be automatically selected from predefined regression models. To select the best one, LOADEST calculated model coefficients using each calibration data set (i.e., observed TN loads in few of days), and models with the lowest Akaike information criterion (AIC) values were selected for load estimations.

RESULTS

Dry N Deposition. The N flux of dry deposition over the Lake Dianchi was 42.1 ± 10.3 mg km⁻² s⁻¹ in 2010–2011 (1σ as the standard deviation of fluxes occurring in 5 sites; Figure 2), with 37.1 ± 10.5 mg km⁻² s⁻¹ for gaseous N and 5.0 ± 0.7 mg km⁻² s⁻¹ for particulate N. The major species, accounting for 99% of the dry deposition, was NH₃ (32.8 ± 10.1 mg km⁻² s⁻¹, 77.9%), HNO₂/HNO₃ (4.0 ± 0.6 mg km⁻² s⁻¹, 9.5%), particulate ON (2.9 ± 0.6 mg km⁻² s⁻¹, 6.8%), and particulate NO₃⁻N (2.0 ± 0.1 mg km⁻² s⁻¹, 4.7%), while the rest of 1% were NO₂ (0.7%) and particulate NH₄⁺N (0.3%). The corresponding concentrations and V_d of gases and TSP at 5 sites can be found in Figures S1 and S2. Regardless of three missing samples in early April–June, the dry deposition N flux showed strong seasonality (Figure 2b). High fluxes were observed from May to August, which were 1.7–13.3 times higher than the remaining periods. Dry deposition of NH₃ was the major source (>86.0%) of total dry deposition N flux from May to August, with biweekly coefficient of variation [CV] of 115%. The NO₂ flux was highest in autumn and early winter (0.4 ± 0.2 mg km⁻² s⁻¹) and remained low in early spring and summer (0.2 ± 0.1 mg km⁻² s⁻¹). The remaining N fluxes were negligible with a seasonal variability with CV of ~27%.

Wet N Deposition. Wet deposition over the Lake Dianchi yielded an N flux of 46.5 ± 13.2 mg km⁻² s⁻¹, 10.5% higher than dry deposition (Figure 2c). NO₃⁻N dominated the total flux of...
wet deposition \( (24.3 \pm 7.5 \text{ mg km}^{-2} \text{s}^{-1}, 52.2\%) \), followed by ON \( (11.9 \pm 5.4 \text{ mg km}^{-2} \text{s}^{-1}, 25.6\%) \) and \( \text{NH}_4^+ \text{-N} \) \( (10.3 \pm 3.9 \text{ mg km}^{-2} \text{s}^{-1}, 22.2\%) \). The corresponding concentrations of diverse N species at all 5 sites can be found in Figure S1. Peak N fluxes of wet deposition occurred in the wet season (i.e., from later July to early October; Figure 2d), following the dry deposition peaks. The N flux during this period were up to \( 172.9 \pm 72.2 \text{ mg km}^{-2} \text{s}^{-1} \), 6-fold greater than mean value of the rest of the year. N fluxes from January to March were only \( 4.4 \pm 7.7 \text{ mg km}^{-2} \text{s}^{-1} \), due to this being the period of lowest precipitation (Figure 2d). This
trend of wet deposition fluxes was highly consistent with precipitation patterns over all 5 sites ($r = 0.83, P < 0.001$; Figure S3), but it was not significantly correlated with N concentrations ($r = 0.32, P < 0.001$; Figure S3). Seasonal changes of NO$_3^−$–N and ON were analogous to the total flux of wet deposition, whereas the changes of NH$_4^+$–N matched up best with dry deposition.

In addition, Figure S4 showed that the ratio of dry to wet deposition of N was 0.91 for the entire period but increased by more than 1.40 in early spring and winter. The ratio of reduced N (NH$_4$ + NH$_4^+$–N) to oxidized N (NO$_3^−$ + NO$_2^−$–N + HNO$_2$/HNO$_3$) in total (wet plus dry) N deposition fluxes was more than 1.4 (Figure S4). The ratio increased to 1.7–3.8 in May, early July, and later August (Figure S4). This was slightly larger than national averages (1.2), but smaller than ratios reported for the United States (1.3–3.5).

**Riverine N Inputs.** Using the observed daily discharge as an independent variable, the LOADEST was able to reproduce observed N inputs from watersheds to Waihai, as shown by a coefficient of determination ($R^2$) of 0.89 and the AIC of 1.8 (Figure 3). In the case of riverine N inputs to Caohai, $R^2$ and AIC were 0.94 and 1.4 (Figure 3), respectively. Thus, the biweekly and ON were analogous to the total HNO$_3$ in total (wet plus dry) N deposition S3), but it was not significantly correlated with N concentrations over all 5 sites ($r = 0.83, P < 0.001$; Figure S3).

**Contributions from Atmospheric Deposition.** Combining dry and wet deposition with daily riverine inputs of N, we derived the mean proportion of atmospheric N deposition to total N loads (i.e., sum of atmospheric deposition and riverine inputs), which was 16.0 ± 0.8% for the period 2010–2011, during an extremely dry period with precipitation of 707 mm yr$^{-1}$ (Figure 4a). The standard deviation due to uncertainties from deposition velocities (assuming CV = 10%), sampling processes (assuming CV = 5%), and riverine N inputs was determined by a Monte Carlo ensemble simulation. More importantly, the contribution made by atmospheric N deposition increased to 25.5% (2015) and 29.2% (2020) accordingly (Figure 4d). The proportion of atmospheric N deposition to total N loads into Lake Dianchi increases to 25.5% (2015) and 29.2% (2020) accordingly (Figure 4d).

**DISCUSSION**

Reliable estimation of the contributions made by atmospheric N deposition is fundamental to our understanding of the lake N budget and is needed to enable policy makers to formulate a comprehensive N management plan. However, previous estimates of the proportion of atmospheric N deposition to total N loads showed large discrepancies using traditional approaches. This is mainly due to the lack of consideration of diverse N forms in atmospheric deposition and lack of direct observations of riverine inputs. Our high-resolution observations of N deposition, along with simultaneous measurements of riverine N inputs, permit new insight into the contribution of N deposition for a typical eutrophic lake.

Recent gaseous and particulate inorganic N have been measured systematically in the Nationwide Nitrogen Deposition Monitoring Network (NNDMN) in China, conducted by China Agricultural University. Two of 43 sites over vegetation were located close to Lake Dianchi and were operated from April 2009 to March 2010. Mean concentrations of NH$_3$, HNO$_2$/HNO$_3$, and NO$_2$ at the two sites of NNDMN (7.0, 6.3, and 0.75 μg N m$^{-3}$) were consistent with those of our study (7.6, 6.5, and 0.98 μg N m$^{-3}$), but slightly larger in particulate N (1.44 vs. 1.10 μg N m$^{-3}$; Figure S1). Accordingly, dry deposition of gaseous N and particulate inorganic N in our study (10.5 ± 0.3 and 0.67 ± 0.1 kg N ha$^{-1}$ yr$^{-1}$) were 44.7% and 46.0% lower than the mean values of the two sites in NNDMN (19.0 ± 0.7 and 1.2 ± 0.6 kg N ha$^{-1}$ yr$^{-1}$), respectively. Such a large discrepancy is primarily due to the difference in deposition velocities of NO$_2$ and HNO$_2$/HNO$_3$ (Figure S2) and additionally due to the difference in concentration of particulate N (Figure S1). For example, due to a larger surface resistance ($R_s$) over water compared to land use types, the mean value of $V_A$ for NO$_2$ in our study (0.0049 ± 0.0009 m s$^{-1}$) was much lower than that for vegetation ($0.18 ± 0.08$ m s$^{-1}$) but close to that over oceanic water ($0.0053 ± 0.003$ m s$^{-1}$). $V_A$ for HNO$_2$/HNO$_3$ over Lake Dianchi (0.39 ± 0.27 m s$^{-1}$) was only one-quarter of that over vegetation but approximately two-thirds of that over the coastal ocean ($0.48 ± 0.27$ m s$^{-1}$). Indeed, HNO$_2$/HNO$_3$ is prone to dissolve in water and its $R_s$ can be negligible, the lower $V_A$ for HNO$_2$/HNO$_3$ over water may be therefore due to the changing aerodynamic
resistance along the gradient of wind speed or roughness length. Lake Dianchi has a smaller roughness length compared to vegetation surface, meaning it has higher aerodynamic resistance. Relative to oceanic water, wind speed over Lake Dianchi is indeed lower and thus results in a smaller $V_d$ for $\text{HNO}_2/\text{HNO}_3$. In addition, annual mean fluxes of NH$_3$ over Lake Dianchi were greater than the national averages over the surface rather than waters ($26.0 \text{ mg km}^{-2} \text{s}^{-1}$) and mean values across United States and European countries, but less for the rest forms of dry deposition of N.

Our results also indicate that NH$_3$ dominated the fluxes of dry deposition over Lake Dianchi. The seasonality of NH$_3$ deposition, consistent with previous observations, was not surprising. First, a growing number of field experiments indicate that NH$_3$ volatilization from croplands increases with air temperature in a nonlinear manner. This temperature-dependent relationship implies that NH$_3$ deposition flux remained higher in warm season than cool season, even though $V_d$ over Lake Dianchi is relatively lower in summer and autumn (Figure S2). To further understand the peaks of NH$_3$ deposition flux, in-house surveys of farmers were conducted in the Lake Dianchi watershed. Three hundred representative farmers were selected for a face-to-face questionnaire-based household survey to collect information on fertilizer use in 13 towns surrounding Lake Dianchi (Figure S7). Figure S7 showed that high rates of N fertilizer use mainly occur in April, June, and July (367.7, 473.4, 568.7 kg N ha$^{-1}$ month$^{-1}$, respectively), partly explaining the large amount of NH$_3$ volatilization and hence abrupt increase of local deposition over the lake (<1 km) from late May onward. In contrast, the fluxes of N oxidation products decreased in summer compared to those in other seasons at 5 sites, in agreement with previous observations in South China. Such seasonal patterns could be explained by discrepancies in atmospheric mixing and photochemical reactions between cold and warm seasons.

Likewise, wet or bulk deposition of NH$_4$$^+$/NO$_3$$^-$ has been measured systematically in China. Inorganic N is therefore used in the following comparative analysis. Wet deposition of inorganic N in this study (10.9 kg N ha$^{-1}$ yr$^{-1}$) was larger than the precipitation-adjusted annual mean flux (7.7 kg N ha$^{-1}$ yr$^{-1}$, $n = 2$) at the two NNDMN sites but approximately one-third less than observation-based “diagnostic” value from bulk N deposition (14.5 kg N ha$^{-1}$ yr$^{-1}$ adjusted according to precipitation difference between 2000s and 2010–2011 in Lake Dianchi). These discrepancies between our current and previous estimates stem from both site-to-site variation and sampling method differences. In our study, wet deposition of inorganic N in KM and JN was comparable to that of the two sites in NMDMN but 25% less than that in the other 3 sites (BF, CG, HK) located in east and west portions of Lake Dianchi. Thus, the mean wet deposition of the 5 sites exceeded that of the two sites in NMDMN. In addition, bulk N deposition used in previous observations denotes the deposition flux as a combination of wet deposition and partial dry deposition (e.g.,
coarse experimental studies showed that the proportion of dry deposition accounts for approximately 25% of bulk N deposition. When excluding the fraction attributable to dry deposition, observation-based “diagnostic” N deposition was decreased to 10.8 kg N ha⁻¹ yr⁻¹, almost equal to our estimate of wet N deposition.

In addition to atmospheric deposition, the reliability of riverine N inputs strongly influences estimates of the relative contribution made by atmospheric N deposition. Fortunately, the state-of-the-art LOADEST has proven to be an effective tool to estimate daily loads from biweekly sampling, based on the empirical relationship with daily discharges. However, when using monthly concentrations and discharges from the routine monitoring program of local government (i.e., green solid circles in Figure 3), total N inputs from all rivers were changed to 5589 t N yr⁻¹, which was 22.4% greater than our observations. Consequently, the contribution made by N deposition decreased to 13% of the total N load. Ambient NH₃ has not been historically and routinely measured due to lack of specific regulatory requirements for its measurement. If we further exclude the dry deposition of NH₃, then the proportion of N deposition to total N loads declined to 8.9%, which is comparable to previous estimates (7.6%). This new finding suggests that the importance of atmospheric N deposition has likely been underestimated, especially for eutrophic lakes close to intensively managed farmland.

Our scenario-based predictions indicated that atmospheric N deposition is a quantitatively important source of biologically available N input in Lake Dianchi (Figure 4d). Furthermore, the role in atmospherically deposited N may increase if no effective controls on emissions are undertaken (Figure 4d). With regard to the lake’s cyanobacterial (e.g., Microcystis) bloom formation and proliferation, the relative contribution to total external N loading was highest (27 ± 5–48 ± 8% of total N loads) in late spring and early summer (Figure 4c). This period coincides with that of maximum phytoplankton growth (measured as chlorophyll-a in the period of 2009–2012, 18–39% month⁻¹; Figure 5b), and can in part support phytoplankton production, leading to maximal biomass accumulation in early autumn (152 µg L⁻¹; Figure 5c). Specifically, for the cyanobacterial bloom former Microcystis spp., which accounts for 84% of phytoplankton abundance, an initial spring proliferation (5.1 × 10⁴ to 6.7 × 10⁴ cell L⁻¹; Figure 5c) and maximum growth rate (35–90% month⁻¹; Figure 5b) overlap closely with the period of maximum atmospheric N deposition relative to total N inputs (Figure 5a). In addition, during the summer Microcystis blooms in Lake Dianchi, soluble reactive phosphorus (SRP) concentrations remained quite high in the water column, while dissolved inorganic N decreased rapidly (Figure 8f). N availability therefore controlled biomass production if having excess SRP in the lake. A similar scenario has been observed in eutrophic Lake Taihu, China, which like Lake Dianchi is impacted by summer Microcystis blooms. In situ nutrient enrichment bioassays have shown these blooms to be largely N-limited. This temporal linkage implies that atmospheric N deposition, as a highly significant N source, may support Microcystis growth during the critical initial proliferation period and for sustaining summer blooms in Lake Dianchi. It should be noted that this toxic bloom-forming genus is not a nitrogen (N₂) fixer and hence has a strong requirement for externally supplied N to support growth. This places even more weight on timely and quantitatively significant inputs of atmospheric N deposition during a period of maximum bloom potential and N demand by this genus. To confirm this scenario, in situ bioassays should be conducted in the future to determine seasonal patterns of N limitation and thresholds for cyanobacteria growth. To mitigate the heavily N-dependent Microcystis growth during this critical period, N emissions that dominates the flux of N deposition into Lake Dianchi should be effectively controlled in addition to the reduction of riverine N inputs. For example, cropland NH₃ volatilization in the watershed can be reduced through site-specific nitrogen management that aims to optimize the supply of fertilizers over time and space while maintaining or enhancing crop yields.

At present, the contribution made by atmospheric N deposition to the total N input contains some uncertainties and is influenced by factors related to data sets used in calculation and scale of sampling. V₄ of gaseous N over the water were calculated using deposition velocity model based on local meteorological data, which was not validated by direct observations. Chemical form of gaseous and particulate N also depends on meteorological conditions and atmospheric composition (e.g., humidity, temperature, oxygen radicals). Any change in these factors will result in differences in atmospheric N deposition over lake surface. In addition, runoff of atmospheric N deposition from watersheds (i.e., indirect deposition) to Lake Dianchi, was not measured or estimated in this study. However, NO₃⁻N, a highly soluble N in water, was a dominant N constituent in precipitation (Figure 2c). If assuming no differences in NO₃⁻N concentrations in precipitation between the sites around and away from the lake, then N deposition flowing into the lake from watershed runoff was at least 141.1 t N during 2010–2011, which was calculated as the product of mean NO₃⁻N concentration in wet samples from 5 monitoring sites and the fraction of streamflow generated from the excess rainfall during a major storm event (see details in Text S4 and Figure S9). Accordingly, the contribution of annual N deposition to total N loads increased from 16% (direct portion) to >18% (direct and indirect portions). Therefore, further efforts at making long-term measurements of dry deposition using relaxed eddy accumulation systems are needed. In addition, the contribution of indirect N deposition on watersheds needs to be determined. Lastly, groundwater discharge, as an additional N input to the lake, was also not included because no measurements were available. We estimated it using a lake water balance approach for the period of April 2010–March 2011 (see details in Text S5). Figure S10 indicated that groundwater discharge (3.6 × 10⁴ m³ yr⁻¹) was much lower than streamflow during study period (380.9 × 10⁴ m³ yr⁻¹). Although uncertainties exist in this estimate, N contributions from groundwater discharge were considered negligible.

In summary, atmospheric N deposition proved to be an important source of N to Lake Dianchi, especially during initial proliferation and periods of maximum Microcystis bloom formation. Decreases in riverine N inputs from watersheds are expected to continue into the future as China aims to lower the N inputs from reuse of reclaimed water and urban stormwater. Current projections of increasing NH₃ emissions, meanwhile, suggest that N deposition levels, especially reduced N, will increase and form a larger fraction of total N loading to the lake in the future. We note that reduced N (as NH₄⁺) is the preferred N source for bloom-forming cyanobacteria, including Microcystis spp. Compared to Lake Dianchi, the contribution made by atmospheric N depostions may be even greater in other eutrophic lakes in East and North China which are experiencing extremely poor air quality. Although China’s Central Govern-
ment has issued regulatory policies for cleaner air and water,5,6 local administrators still face challenges in coordinating the reductions of atmospheric deposition and riverine inputs, given limited financial resources. In addition to implementing a comprehensive assessment of watershed N loads (i.e., N deposition and inputs) in China, identifying effective approaches that yield positive benefits for both air and water quality remains a high priority for lake ecosystem management.

**ASSOCIATED CONTENT**

**SUPPORTING INFORMATION**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.0c00615.

Details of sampling design, deposition velocity models, details of atmospheric deposition at 5 sites, results of in-house surveys, and the other information (PDF)

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**Notes**

The authors declare no competing financial interest.

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