Wet Inorganic Nitrogen Deposition at the Daheitin Reservoir in North China: Temporal Variation, Sources, and Biomass Burning Influences

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Received: 27 October 2020; Accepted: 17 November 2020; Published: 23 November 2020

Abstract: Atmospheric nitrogen deposition is of great concern to both air quality and the ecosystem, particularly in northern China, which covers one-quarter of China’s cultivated land and has many heavily air polluted cities. To understand the characteristics of wet N deposition at rural sites in northern China, one-year wet deposition samples were collected in the Daheitin reservoir region. Due to the intense emissions of gaseous nitrogen compounds from heating activities during cold seasons and distinct dilution effects under different rainfall intensities and frequencies, the volume weighted mean concentrations of wet N deposition showed higher levels in dry seasons but lower levels in wet seasons. In contrast, the wet N deposition rates varied consistently with precipitation, i.e., high during the wet season and lower during the dry season. The annual wet deposition rate of total inorganic ions (the sum of NO$_3^-$–N and NH$_4^+$–N) at the rural site in North China from July 2019 to June 2020 was observed at 18.9 kg N ha$^{-1}$ yr$^{-1}$, still remained at a relatively high level. In addition, biomass burning activities are ubiquitous in China, especially in northern China; however, studies on its impact on wet N deposition are limited. Non-sea salt potassium ion (nss-K$^+$) was employed as a molecular tracer to investigate the characteristics of biomass burning activities as well as their impact on the chemical properties of wet N deposition. Three precipitation events with high nss-K$^+$ levels were captured during the harvest season (June to July). The variations in the patterns of nss-K$^+$, deposited N species, and ratios of nss-K$^+$ to nitrogen species as well as their relationships all indicated that biomass burning emissions contributed remarkably to NO$_3^-$–N but had a minor influence on NH$_4^+$–N.

Keywords: precipitation; nitrogen deposition; potassium; biomass burning; air pollution

1. Introduction

With the rapid development of agriculture, industry, and the expansion of the urban population over the last several decades, anthropogenic activities, such as fossil fuel combustion, chemical fertilizer, etc., have greatly increased the levels of reactive nitrogen (N) in the atmosphere [1–4]. Reactive N in the atmosphere not only leads to air quality degradation by contributing to the production of ozone and PM$_{2.5}$ [5–11] but also disturbs the N cycle through excess deposition of N, leading to aquatic eutrophication, soil acidification, and impacting on ecosystem functions [2,12–15].
Nitrogen deposition rates decreased in the USA and Europe several decades ago [3,12,16–18]. In China, however, NH$_3$ and NO$_x$ have both continuously increased due to the high loading emissions from agricultural and industrial activities under the rapid economic development over the past few decades [19,20]. The annual total reactive nitrogen deposition (27.2 ± 6.7 kg N ha$^{-1}$) significantly increased at a rate of 10% from 2011 to 2018 [21]. SO$_4^{2-}$ has previously been reported for many sites to be the most dominant inorganic PM$_{2.5}$ component in China, followed by NO$_3^-$ and NH$_4^+$ [22,23]. Under rising NO$_x$ emissions, a result of industrialization and increase in vehicle numbers, NO$_3^-$ concentrations have been observed to exceed those of SO$_4^{2-}$ in parts of China [10,24–26]. Meanwhile, NH$_3$ emissions have increased over the several past decades [27]; for example, the annual concentration of NH$_3$ in 2018 was 14.2 µg N m$^{-3}$, which is twice that in 2011 (7.0 µg N m$^{-3}$), in Zhengzhou, China [28]. High emissions of NH$_3$ and NO$_x$, combined with high loading of NO$_3^-$ and NH$_4^+$ in ambient particulates subsequently contributed to the high deposition of atmospheric N in China [2,29–32]. In spite of the site-to-site variability in the data, total inorganic N deposition increased significantly with an average annual increase of 0.41 kg of nitrogen per hectare (kg N yr$^{-1}$) between 1980 and 2010 [2]. To address severe air pollution issues, the Chinese government implemented the Air Pollution Prevention and Control Action Plan (Action Plan) in 2013. As a result, China’s air quality has improved through the substantial reduction of several pollutants, in particular, SO$_2$ and particulates [19,20]. Under these air pollutant control measures, a subsequent decrease in atmospheric N deposition was expected. For example, the average wet total inorganic N deposition rate in Beijing during 1998 to 2004 was 30.6 kg N ha$^{-1}$ yr$^{-1}$ [33], while it decreased to 8.0 kg N ha$^{-1}$ yr$^{-1}$ at a rural site in Beijing during 2017–2018 [34]. The North China Plain (NCP) is one of the most polluted regions in China and also greatly affects the air quality status of downwind regions [35–37]. Scattered among the vast areas of farmlands in the NCP, which comprises one-quarter of China’s cultivated land [26,38], are many industries of (such as power, steel, chemical industry, etc.) heavily populated large cities. High loadings of ambient particles and gas phase pollutants (e.g., NH$_3$, NO$_2$, and SO$_2$) in the NCP are transferred from the atmosphere to the Earth’s surface and input into ecosystems increasing the N deposition rate [39]. The annual deposition of inorganic N in northern China was in the range of 15 to 50 kg N ha$^{-1}$ yr$^{-1}$ over 2003 to 2006 [40] and 16.3 to 28.2 kg N ha$^{-1}$ yr$^{-1}$ over 2007 to 2010 [32]. This deposition can lead to significant and serious ecological degradation in terrestrial and aquatic systems. Previous works evaluating N deposition fluxes as well as the influence of air pollutant control measures were mostly limited to urban areas [28,34,41], while researches in rural areas of North China were rarely reported.

Additionally, previous works have mostly focused on the agricultural or industrial sources of N deposition in the atmosphere, while the influence of biomass burning on N deposition has rarely been reported [29,42–44]. Biomass burning activities are ubiquitous in China, especially in North China, where there is an enormous rural population and winter heating custom. Based on previous studies, two intensive biomass burning episodes are typically observed in China: one occurring in the late spring and early summer (mostly in June), featuring open-field burning of agricultural residue after wheat straw harvest, and the other one in the late fall (between October to November), caused by agricultural residue burning after wheat or corn harvest [45,46]. Thus, the objectives of this study were to quantify: (1) the amount and temporal variation of wet N deposition at a rural site in North China after the clean air policy in China was implemented; (2) the sources of information as well as the influences of biomass burning for wet N deposition at the rural site in North China.

2. Experimental Section

2.1. Sample Collection

This study was carried out at the area of Daheitin Reservoir (DHT) (40.17° N, 118.33° E), which is an important drinking water source for the city of Tianjin and Tangshan (two megacities in North China) (Supplementary Materials, Figure S1). The sampling station was located on top of a two-story
building, approximately 3.0 m above ground, surrounded by a small village with approximately 500 inhabitants, roughly 1.0 km to the DHT Reservoir. Precipitation samples were collected using an automatic wet deposition collector that opens its lid automatically to collect a rainwater sample when it rains and closes automatically when the rain stops. The collected rainwater samples were transferred to pre-cleaned brown polyethylene bottles (200 mL) and stored in a refrigerator at −20 °C until chemical analysis. A total of 37 precipitation samples were collected from June 2019 to June 2020, covering the whole year, including the wet season (June–September) as well as the dry season (October–May).

2.2. Chemical Analysis

The rain samples were filtered with 0.45 µm filters to separate insoluble fractions before chemical analysis of the ions. The water-soluble inorganic ions in precipitation sample, including three anions (i.e., \( \text{SO}_4^{2-} \), \( \text{NO}_3^- \), \( \text{Cl}^- \)) and five cations (i.e., \( \text{NH}_4^+ \), \( \text{Ca}^{2+} \), \( \text{Na}^+ \), \( \text{K}^+ \), and \( \text{Mg}^{2+} \)) were analyzed by a Dionex ICS-3000 ion chromatograph (Thermo Fisher Scientific. Ltd., Waltham, MA, USA). The potassium concentrations presented here were corrected by a sea salt indicator (i.e., sodium; \( C_{\text{K}_{\text{sea}}} = C_{\text{K}} - 0.0355 \times C_{\text{Na}} \)) that has been widely used for various field observations [47–49]. More details about the water-soluble inorganic ion analysis method can be found elsewhere [50].

2.3. Calculations

2.3.1. The Annual Volume-Weighted Mean Concentrations (\( C_{\text{vwm}, \text{mg N L}^{-1}} \))

\[
C_{\text{vwm}} = \frac{\sum_{i=1}^{n} (C_i \times P_i)}{\sum_{i=1}^{n} P_i} \quad (1)
\]

The annual volume-weighted mean (VWM) concentrations of inorganic nitrogen species (\( C_{\text{vwm}, \text{mg N L}^{-1}} \)) were calculated using Equation (1). \( C_i \) is the mass concentration of inorganic nitrogen species in the \( i \)th sample (mg N L\(^{-1}\)), \( P_i \) is the precipitation amount collected within the \( i \)th sampling period (mm), and \( n \) was the total number of precipitation samples during the whole year.

2.3.2. Flux of Wet N Deposition (\( F_{\text{wd}, \text{kg N ha}^{-1}} \))

\[
F = 0.01 \times \sum_{i=1}^{n} C_i \times P_i \quad (2)
\]

\( F_{\text{wd}} \) represents the monthly, seasonal, and annual wet deposition N flux, calculated by Equation (2). \( C_i \) is the mass concentration of \( \text{NH}_4^+\)-N and \( \text{NO}_3^-\)-N (mg N L\(^{-1}\)), and \( P_i \) (in mm) is the total amount of precipitation over a month, season or year.

3. Results and Discussion

3.1. Concentrations of Inorganic Nitrogen Species in Rainwater

The total amount of precipitation for the whole year in the DHT reservoir area from July 2019 to June 2020 was 595 mm, similar to what was observed at other sites in North China [34] but obviously lower than in South China [29], East China [51], and Southeast China [42,52]. \( \text{NO}_3^- \), \( \text{SO}_4^{2-} \), and \( \text{NH}_4^+ \) were the three most abundant ions in the rainwater samples with annual VWM concentration of 4.69 mg L\(^{-1}\), 4.03 mg L\(^{-1}\), and 2.71 mg L\(^{-1}\), respectively. With the rapid urbanization and industrialization, vehicle emission sources have become increasingly prevalent in China, even in rural areas, leading to \( \text{NO}_3^- \) concentrations exceeding those of \( \text{SO}_4^{2-} \) at the DHT site with an annual average mass ratio (\( \text{NO}_3^- / \text{SO}_4^{2-} \)) of 1.20, which is higher than previous aerosol observations in North China (<1.0) [22–24]. \( \text{Ca}^{2+} \) was the second most abundant cation with an annual VWM concentration of 1.54 mg L\(^{-1}\), exhibiting a non-negligible contribution of soil dust sources in the atmosphere at the rural site in North China (Table 1).
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Table 1. The statistical summary of water-soluble ions in precipitation and their volume-weighted mean (VWM) concentrations from July 2019 to June 2020. nss = Non-sea salt.

| Parameter         | Unit | Average | SD   | Minimum | Maximum | Median | C_vwm |
|-------------------|------|---------|------|---------|---------|--------|-------|
| nss-K⁺            | mg L⁻¹ | 0.35    | 0.67 | 0.01    | 3.59    | 0.14   | 0.21  |
| Na⁺               | mg L⁻¹ | 1.30    | 3.56 | 0.04    | 15.3    | 0.19   | 0.74  |
| Ca²⁺              | mg L⁻¹ | 2.21    | 2.60 | 0.17    | 12.6    | 1.39   | 1.54  |
| Mg²⁺              | mg L⁻¹ | 0.42    | 0.49 | 0.07    | 2.48    | 0.28   | 0.30  |
| NH₄⁺              | mg L⁻¹ | 3.00    | 1.83 | 0.34    | 8.64    | 2.27   | 2.71  |
| NO₃⁻              | mg L⁻¹ | 6.24    | 5.27 | 0.51    | 32.7    | 4.06   | 4.69  |
| SO₄²⁻              | mg L⁻¹ | 5.04    | 4.16 | 0.59    | 25.5    | 3.90   | 4.03  |
| F⁻                 | mg L⁻¹ | 0.09    | 0.09 | 0.01    | 0.43    | 0.06   | 0.07  |
| NH₄⁺–N            | mg L⁻¹ | 0.96    | 1.58 | 0.10    | 9.04    | 0.55   | 0.59  |
| NO₃⁻–N            | mg L⁻¹ | 2.33    | 1.42 | 0.27    | 6.72    | 1.77   | 2.28  |
| Ratio of NH₄⁺–N/NO₃⁻–N |           | 1.97    | 0.64 | 0.26    | 3.41    | 1.92   | 1.97  |
| Mass ratio of NO₃⁻/SO₄²⁻ |           | 1.20    | 0.30 | 0.66    | 1.75    | 1.21   | -     |

Most of the rain events occurred during the summer from June–August (Figure 1; Table 2). During the entire sampling period, precipitation in the DHT region was concentrated in the summer season (416 mm), 70% of the whole year’s precipitation (595 mm). The annual VWM concentrations of deposition of inorganic N species, i.e., NH₄⁺–N, NO₃⁻–N, and TIN (the total sum VWM concentrations of NH₄⁺–N and NO₃⁻–N) were 2.11 mg N L⁻¹, 1.06 mg N L⁻¹, and 3.17 mg N L⁻¹, respectively. Notably, seasonal VWM concentrations of NH₄⁺–N and NO₃⁻–N showed higher levels in autumn and winter, and the lowest in summer (Figure 1). In late autumn and winter, more gaseous nitrogen compounds (i.e., NOₓ) are accumulated in the atmosphere due to the increased emissions from coal combustion for heating and biomass combustion activities, as well as rare precipitation wash out, subsequently resulting in higher concentrations of N deposition. In contrast, there was no coal combustion for heating over the summer season, the emission of precursor gases were relatively lower, and coupled with the dilution of frequent rainfall events. The VWM concentration of NO₃⁻–N was observed at its lowest level in summer (Figure 1) and showed a negative relationship with precipitation amount (R² = 0.22, p < 0.05, Figure S2).
Figure 1. Monthly patterns of (a) precipitation amounts and VWM concentrations (mg N L\(^{-1}\)) of inorganic nitrogen species (i.e., NH\(_4^+\)–N and NO\(_3^-\)–N); (b) proportion of NH\(_4^+\)–N and NO\(_3^-\)–N to TIN in wet deposition at the Daheitin (DHT) reservoir, a rural site in North China from July 2019 to June 2020. There was no precipitation during December 2019 and January 2020.

Table 2. Annual and seasonal variations in rainfall (mm), NH\(_4^+\)–N/NO\(_3^-\)–N ratios, annual VWM concentrations (mg N L\(^{-1}\)) of inorganic nitrogen (NH\(_4^+\)–N, NO\(_3^-\)–N, and TIN), and wet N deposition rates (kg N ha\(^{-1}\)) at the DHT reservoir, a rural site in North China from July 2019 to June 2020.

| Seasons                     | Rainfall (mm) | N Concentration in Rainwater (mg N L\(^{-1}\)) | Wet N Deposition Rate (kg N ha\(^{-1}\)) |
|-----------------------------|---------------|-----------------------------------------------|----------------------------------------|
|                             |               | NH\(_4^+\)–N NO\(_3^-\)–N TIN NH\(_4^+\)/NO\(_3^-\) Ratio | NH\(_4^+\)–N NO\(_3^-\)–N DIN          |
| Spring (March–May)          | 122           | 2.35 1.43 3.78 1.81                           | 2.87 1.75 4.62                         |
| Summer (June–August)       | 416           | 1.94 0.84 2.78 2.15                           | 8.06 3.51 11.6                         |
| Autumn (September–November)| 51.3          | 2.70 1.76 4.47 1.35                           | 1.39 0.90 2.29                         |
| Winter (December–January)  | 5.80          | 4.20 2.55 6.75 1.65                           | 0.24 0.15 0.39                         |
| Annual                      | 595           | 2.11 1.06 3.17 1.97                           | 12.6* 6.31* 18.9*                      |

* The unit is kg N ha\(^{-1}\) yr\(^{-1}\). DIN: deposition rate of wet inorganic nitrogen.

Compared to summer months, the gap between NH\(_4^+\)–N and NO\(_3^-\)–N was smaller in the autumn and winter months (Figure 1a). This may have been due to the large amount of NO\(_3^-\) emissions during heating-related coal combustion in November to March in northern China and coupled with the decreased NH\(_3\) emissions from agriculturally associated sources; the gap between NH\(_4^+\)–N and NO\(_3^-\)–N was reduced during the cold winter season. Accordingly, the proportion of NO\(_3^-\)–N to TIN in wet deposition showed a similar variation pattern, with lower proportions in the summer months, i.e., June (31.3%) and August (31.0%) and higher ones in September (48.2%) and November (46.2%) as shown in Figure 1b. The annual VWM concentrations of NH\(_4^+\)–N and NO\(_3^-\)–N observed in this study (2.11 mg N L\(^{-1}\) and 1.06 mg N L\(^{-1}\), respectively) were lower than those measured in North China during the years before the Action Plan was implemented in 2013, for example, during 1999 to 2004 (2.22 mg N L\(^{-1}\) for NO\(_3^-\)–N and 4.76 mg N L\(^{-1}\) for NH\(_4^+\)–N) [33]. However, this level was still higher than those observed in South China over 2016–2017 (0.5–0.9 mg N L\(^{-1}\) for NO\(_3^-\)–N and 0.7–1.3 mg N L\(^{-1}\) for NH\(_4^+\)–N) [29], a karst catchment from 2016–2017 (0.3 mg N L\(^{-1}\) for NO\(_3^-\)–N and 0.7–0.9 mg N L\(^{-1}\) for NH\(_4^+\)–N) [42], and rural sites in the central Sichuan Basin of China from 2011–2013 (1.05 ± 0.91 mg N L\(^{-1}\) for NO\(_3^-\)–N and 1.48 ± 1.05 mg N L\(^{-1}\) for NH\(_4^+\)–N) [52].

3.2. Wet Inorganic Nitrogen Species Deposition Flux

The monthly variations in the amounts of precipitation, wet deposition rates of N species, as well as their proportions of the total deposition rate of wet inorganic nitrogen (DIN, sum of wet deposition rates of NH\(_4^+\)–N and NO\(_3^-\)–N) are shown in Figure 2. The deposition rates of NH\(_4^+\)–N, NO\(_3^-\)–N, and DIN were generally higher in the wet season than those in the dry season. The wet deposition flux of NH\(_4^+\)–N, NO\(_3^-\)–N, and DIN during summer were 8.06 kg N ha\(^{-1}\) yr\(^{-1}\), 3.51 kg N ha\(^{-1}\) yr\(^{-1}\), and 11.6 kg N ha\(^{-1}\) yr\(^{-1}\), respectively, which were 33.6 times, 23.4 times and 29.7 times of those at winter (0.24 kg N ha\(^{-1}\) yr\(^{-1}\), 0.15 kg N ha\(^{-1}\) yr\(^{-1}\) and 0.39 kg N ha\(^{-1}\) yr\(^{-1}\), respectively) (Table 2). Wet NH\(_4^+\)–N and NO\(_3^-\)–N deposition accounted for 29.2–48.2% and 51.8–70.8% to DIN deposition, respectively (Figure 2b). The monthly varied patterns of wet N species deposition rates were consistent with that of precipitation as shown in Figure 2. Furthermore, the relationship between monthly deposition rates of NH\(_4^+\)–N and NO\(_3^-\)–N were positively correlated with the amount of monthly rainfall (R\(^2\) = 0.89, p < 0.01; R\(^2\) = 0.92, p < 0.01, respectively, Figure S3) indicating that the amount of precipitation is crucial in controlling wet N deposition.
Moreover, this level was also obviously higher than the wet DIN deposition rate in the USA (0.5–3.5 kg N ha\(^{-1}\) yr\(^{-1}\)) (Table S1) \cite{3,16,53}. This indicates that although the wet N deposition flux at the rural site in North China from July 2019 to June 2020. There was no precipitation during December 2019 and January 2020.

![Figure 2. Monthly patterns of (a) wet deposition rates (kg N ha\(^{-1}\)) of NO\(_3^−\)-N, NH\(_4^+\)-N, and (b) their proportion of the total deposition rate of wet inorganic nitrogen (DIN) at the DHT reservoir, a rural site in North China from July 2019 to June 2020. There was no precipitation during December 2019 and January 2020.](image)

The annual total wet N deposition flux in this study was 18.9 kg N ha\(^{-1}\) yr\(^{-1}\); a comparison to previous studies is shown in Table S1. The level of annual wet deposition flux in this study was lower prior to the implementation of the Action Plan, when anthropogenic impact was even stronger, for example, in northern China from 2007–2010 (16.3–28.2 kg N ha\(^{-1}\) yr\(^{-1}\)) \cite{32}, urban sites in East China from 2003–2005 (26.8 kg N ha\(^{-1}\) yr\(^{-1}\)) \cite{43}, and urban sites, for example, Zhengzhou, China, during 2011 (33.3 kg N ha\(^{-1}\)) \cite{28}. However, these results were still higher than those observed at a rural site of Beijing from 2017–2018 (8.0 kg N ha\(^{-1}\) yr\(^{-1}\)) \cite{34}, the Three Gorges Reservoir Region in the Southwest China in 2015 (7.1–16.8 kg N ha\(^{-1}\) yr\(^{-1}\)) \cite{44}, and rural areas in South China from 2017–2018 (13.2) \cite{29}. Moreover, this level was also obviously higher than the wet DIN deposition rate in the USA (0.5–3.5 kg N ha\(^{-1}\) yr\(^{-1}\)) (Table S1) \cite{3,16,53}. This indicates that although the wet N deposition flux at the rural site in North China decreased due to the Action Plan’s implementation; it still remained at a relatively high level.

3.3. Sources of Wet Inorganic N Deposition

3.3.1. NH\(_4^+\)-N/NO\(_3^−\)-N Ratio

In general, NH\(_4^+\) in rainwater is formed from the precursor gas NH\(_3\), the main anthropogenic source of which is the volatilization of fertilizer and livestock manure; while NO\(_3^−\) in rainwater is produced by the precursor gases NO\(_x\) (i.e., NO and NO\(_2\)), which are mainly emitted from fossil fuel combustion, such as from power plants, vehicles, and heating activities. Moreover, NH\(_3\) is readily converted to NH\(_4^+\), having a relatively shorter transportation distance, indicating NH\(_4^+\) in rainwater
is subjected to local sources [42]. Thus, the \( \text{NH}_4^+/\text{NO}_3^- \) ratio is a good indicator of the relative contributions of agricultural sources (i.e., fertilizer and livestock manure) and non-agricultural sources (e.g., industrial fossil fuel combustion and vehicle emissions) [42,43,54,55]. Generally, in areas with advanced industrialization, the \( \text{NH}_4^+/\text{NO}_3^- \) ratio is usually smaller than 1.0, e.g., at an urban site in Nanjing (0.94), China [43], Pearl River Delta (PRD) (0.33–4.0, averaged at 1.1), China [56], and New York (0.76–0.99) [57]. On the contrary, areas with intensive agriculture, such as rural or agricultural sites, are usually characterized by wet N deposition with \( \text{NH}_4^+/\text{NO}_3^- \) ratios greater than 1.0. In China, annual average \( \text{NH}_4^+/\text{NO}_3^- \) ratios were reported to be 1.76 for an agricultural site and 1.14 for a rural site in South China [29], 2.2–2.5 for a rural site in the Three Gorges Reservoir Region, Southwest China [44,58], 3.6 for the grasslands at Qinghai Lake [59], 1.9–2.0 for the North China Plain [40,54], and 2.05 for an agricultural site in Northeast China [60].

The monthly variation of \( \text{NH}_4^+/\text{NO}_3^- \) ratio at the DHT reservoir is shown in Figure 3. In this study, the annual average \( \text{NH}_4^+/\text{NO}_3^- \) ratio was 1.97 with all monthly average values exceeding 1.0 (in the range of 1.08 to 2.26). Thus, compared to \( \text{NO}_3^- \) from fossil fuel combustion in industry and transportation, \( \text{NH}_4^+ \) deposition from agriculture and excrement from humans and animals still occupied the larger portion in the DHT region. This was mainly because the sampling site was located in a rural area, surrounded by scattered rice paddy fields and villages, where N fertilization had a great effect on \( \text{NH}_4^+ \) deposition. \( \text{NH}_4^- \) was the dominant form of N deposition; however, the contribution from \( \text{NO}_3^- \) significantly increased over the past decades in China [2]. The ratio of \( \text{NH}_4^- \) to \( \text{NO}_3^- \) in wet precipitation decreased significantly by approximately five to two over time from 1980 to 2010 in China [2]. Similarly, the \( \text{NH}_4^-/\text{NO}_3^- \) ratio in wet deposition sharply decreased from 5.8 to 1.2 in Yangtze River Delta Region from 1980 to 2005 [43]. This indicates a great enhancement in \( \text{NO}_3^- \) emissions over the past few decades, corresponding to the rapid increase in fossil fuel consumption in the industry and transportation sectors.

Figure 3. A box chart of the monthly patterns of \( \text{NH}_4^+/\text{NO}_3^- \) ratios at the DHT reservoir, a rural site in North China from July 2019 to June 2020. There was no precipitation during December 2019 and January 2020. (×: The minimum and maximum values of the \( \text{NH}_4^+/\text{NO}_3^- \) ratio; the average value; upper and lower edge mean the 25% and 75% value; lines in the box indicate the median value of the \( \text{NH}_4^+/\text{NO}_3^- \) ratio).

The \( \text{NH}_4^+/\text{NO}_3^- \) ratios also varied with season, peaking in the summer months (i.e., June (2.23), July (2.02) and August (2.26)) and reaching lower values in the dry seasons, e.g., September (1.08), November (1.38), and March (1.43). This seasonal pattern is similar to those reported by other studies in
China, e.g., high values, particularly, in the wet season and low values in the dry season were observed in South China [29,56]. At the same, the annual average NH$_4^+$–N/NO$_3^-$–N ratio was 2.3 in the western part of South Korea from 2007–2008, but the lowest NH$_4^+$–N/NO$_3^-$–N ratio was observed during the winter season (0.63) [14]. This seasonal variation of wet N deposition is in accordance with the nitrogen-fertilizing period during the crop growing season in China. Meanwhile, temperatures over the wet season were higher than in the winter season, and hot temperature conditions were much easier for agriculturally associated NH$_3$ volatilization from fertilizer (urea, di-ammonium phosphate, etc.) and livestock manure applied in agricultural soils. The contribution of agricultural activity to atmospheric N deposition was substantially weakened during the winter season. Moreover, when entering into late autumn, the weather becomes cold, and the coal and biomass burning combustion activities for heating become abundant in North China, resulting in obvious elevated concentrations of NO$_3^-$ in ambient particles, and thus decreased the NH$_4^+$–N/NO$_3^-$–N ratio during winter season.

3.3.2. Biomass Burning Influences on Wet Inorganic N Deposition

Potassium is normally regarded as a tracer of biomass burning [5,47,61]. Non-sea-salt potassium concentrations were used to exclude the influence of potassium derived from sea-salt. Therefore, the ratio of non-sea salt (nss)-K$^+$ to NH$_4^+$–N and NO$_3^-$–N in the atmosphere can applied to estimate the effect of biomass burning to wet N deposition. Three precipitation events occurred during the harvest season on 6 June, 29 June and 30 July, 2019 (Figure 4). The nss-K$^+$ concentrations in those rainwater samples were significantly higher than in other precipitation samples, indicating the field biomass combustion was active during these periods. Fire pixel counts observed from MODIS (Moderate Resolution Imaging Spectroradiometer) Terra and Aqua also identified many fire spots in North China during June and July 2019, when it is time to harvest wheat and corn along with the common practice of straw burning (Figure 5a,b).

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Temporal variations of (a) concentrations of NH$_4^+$–N and NO$_3^-$–N, (b) nss-K$^+$ and Cl$^-$, (c) the ratios of nss-K$^+$/NO$_3^-$–N and nss-K$^+$/NH$_4^+$–N, (d) the ratios of NO$_3^-$–N/TIN and NH$_4^+$–N/TIN, and (e) rainfall in wet depositions in the DHT reservoir region from 6 June 2019 to 30 June 2020. The gray shades in the biomass burning’s impact (precipitation events at 6 June 2019, 29 June and 30 July 2019), and the pink shades in the heating start time with coal combustion and waste incineration (precipitation event at 21 November 2019).
wet N deposition. Several studies speculate that biomass burning might be a source of water-soluble organic N in wet deposition [29]. In this study, we provided evidence for the fact that biomass burning contributed to the polluted areas of North China with dense fire spots (Figure S4a–c). Abundant combustion aerosols were transported from these source areas to the DHT region, associated with significantly increased levels of biomass burning tracers, i.e., nss-K⁺, as well as Cl⁻ (Figure 4). Cl⁻ is regarded as another common tracer for biomass burning [63], which synchronously changed with nss-K⁺ in the wet deposition over the entire year, except for the precipitation event on 21 November 2019. Based on fuel combustion studies, Cl⁻ in aerosols can also be contributed by coal combustion [64,65] and chlorinated plastics waste burning [66]. Entering into middle November, the weather became cold, thus, coal combustion activities commenced in rural areas and central heating systems started operating. Subsequently, the level of Cl⁻ sharply increased during the precipitation event on 21 November 2019 at the start of the heating period. There was almost no influence on NH₃ emissions from biomass burning; hence, the ratio of nss-K⁺/NH₄⁺–N in the atmosphere was expected to be higher when biomass burning occurred. Correspondently, the ratio of nss-K⁺/NH₄⁺–N obviously increased during the summer harvest biomass burning episode but displayed no variation over the winter heating season (Figure 4c), which further confirmed that the high levels of Cl⁻ during the precipitation event on 21 November 2019 was emitted from coal combustion or chlorinated plastics waste burning but not biomass burning.

The concentration of nss-K⁺ was positively correlated with NO₃⁻–N (R² = 0.98, p < 0.01), but revealed no relationship to NH₄⁺–N (p > 0.5) (Figure 6). In general, NH₄⁺–N/TIN ratios were higher than NO₃⁻–N/TIN ratios throughout the observations. However, due to the substantial contribution of biomass burning to atmospheric NOₓ emissions during the harvest season or intense fossil fuel combustion during the start of the heating season, the ratios of NO₃⁻–N/TIN were obviously elevated, and surpassed the ratios of NH₄⁺–N/TIN (Figure 4d) during those intense air pollution periods. The change in the N species in the atmosphere exhibited the influences of air pollution on wet N deposition. Several studies speculate that biomass burning might be a source of water-soluble organic N in wet deposition [29]. In this study, we provided evidence for the fact that biomass burning

![Figure 5](image-url)
evidently contributed to the wet deposition of some inorganic N (e.g., NO$_3^-$–N); however, it had a minor influence on others (such as NH$_4^+$–N).

![Figure 6](image_url)

**Figure 6.** The relationships between the concentrations of NO$_3^-$–N, NH$_4^+$–N, and nss-K$^+$ in wet depositions in the DHT reservoir region from 6 June 2019 to 30 June 2020. Statistical analysis of the data was conducted with the linear fitting method.

**4. Conclusions**

One-year wet deposition samples were collected at the Daheitin reservoir in North China. The annual VWM concentrations of NH$_4^+$–N and NO$_3^-$–N were 2.11 mg N L$^{-1}$ and 1.06 mg N L$^{-1}$, respectively. Due to the intense emissions of more gaseous nitrogen compounds (i.e., NO$_x$ and NH$_3$) in heating activities in the cold seasons and the dilution effect by different rainfall intensity and frequency, the seasonal VWM concentrations of NH$_4^+$–N and NO$_3^-$–N showed higher levels during the dry season (i.e., autumn and winter) and lower levels during the wet season, i.e., summer. In contrast, the wet N deposition rates varied consistently with precipitation (i.e., high during the wet season and low during the dry season), indicating that the amount of precipitation was crucial in controlling wet N deposition. Compared to the years prior to the implementation of the Action Plan in 2013, the wet inorganic nitrogen deposition rate at the rural site in North China decreased to 18.9 kg N ha$^{-1}$ yr$^{-1}$ but still maintained a relatively high level.

Non-sea salt K$^+$ was employed as a molecular tracer of biomass burning emission, and three precipitation events with high nss-K$^+$ levels were captured during the harvest season (June to July). Based on the variations in the patterns of nss-K$^+$ and deposition of N species, the ratios of nss-K$^+$ to N species as well as their relationships, it was indicated that biomass burning emissions contributed remarkably to NO$_3^-$–N but had a minor influence on NH$_4^+$–N. These results highlight that biomass burning activity had a substantial influence on the composition of wet nitrogen deposition at the rural site in North China. Understanding the variations in trends and the influences of wet nitrogen deposition is important for effective air pollution control and for gaining insight into the atmospheric nitrogen cycle.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/2073-4433/11/11/1260/s1, Table S1: Wet N deposition rates (kg N ha$^{-1}$ yr$^{-1}$) in DHT region compared to those reported in other studies, Figure S1: Location of Daheittin Reservoir station (red star) and the surrounding cities, Figure S2: Relationship between the concentrations of NO$_3^-$–N in rainwater and rainfall amounts during each precipitation event at the
DHT reservoir, a rural site in North China from July 2019 to June 2020. Statistical analysis was conducted with logarithmic fitting method, Figure S3: Relationship between monthly deposition rates of NH$_4^+$.N and NO$_3^-$-N and rainfall amounts at the DHT reservoir, a rural site in North China from July 2019 to June 2020. Statistical analysis was conducted with linear fitting method, Figure S4: Air mass origins to DHT region, shown by clusters of 72 h backward trajectories arriving at 500 m aboveground level in (a) 6 June, 2019, (b) 29 June, 2019, (c) 30 July, 2019, (d) 21 November, 2019.

**Author Contributions:** Conceptualization, L.W., L.L., and X.L.; Data curation, L.W., K.L., W.X., and G.Z.; Formal analysis, W.X.; Funding acquisition, L.L., W.X., L.L., K.L., and W.H.; Investigation, P.Z., X.Z., and C.L.; Methodology, K.L. and L.L.; Project administration, L.L.; Supervision, L.L.; Writing—original draft preparation, L.W. and L.L.; Writing—review and editing, L.W., L.L., K.L., W.X., and W.H. All authors have read and agreed to the published version of the manuscript.

**Funding:** This work was supported by the Open Research Fund of the State Key Laboratory of Simulation and Regulation of Water Cycle in River Basin, China Institute of Water Resources and Hydropower Research (No. 201913), the National Natural Science Foundation of China (No. 51879279), Beijing Natural Science Foundation (No. 8192055), and the National Key Research and Development Program of China (No. 2016YFC0401701; 2017YFC0405804).

**Acknowledgments:** We are grateful to Hongbing Cheng and Yusi Liu for their help with the analysis of water-soluble ions. The authors would like to acknowledge Chengbiao Zhao for their help with precipitation sample collection.

**Conflicts of Interest:** The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**References**

1. Galloway, J.N.; Dentener, F.J.; Capone, D.G.; Boyer, E.W.; Howarth, R.W.; Seitzinger, S.P.; Asner, G.P.; Cleveland, C.C.; Green, P.A.; Holland, E.A.; et al. Nitrogen Cycles: Past, Present, and Future. *Biogeochemistry* 2004, 70, 153–226.

2. Liu, X.; Zhang, Y.; Han, W.; Tang, A.; Shen, J.; Cui, Z.; Vitousek, P.; Erismann, J.W.; Goulding, K.; Christie, P.; et al. Enhanced nitrogen deposition over China. *Nature* 2013, 494, 459–462. [PubMed]

3. Du, E.; de Vries, W.; Galloway, J.N.; Hu, X.; Fang, J. Changes in wet nitrogen deposition in the United States between 1983 and 2012. *Environ. Res. Lett.* 2014, 9, 095004.

4. Asner, G.P.; Townsend, A.R.; Riley, W.J.; Matson, P.A.; Neff, J.C.; Cleveland, C.C. Physical and biogeochemical controls over terrestrial ecosystem responses to nitrogen deposition. *Biogeochemistry* 2001, 54, 1–39.

5. Liang, L.; Engling, G.; Zhang, X.; Sun, J.; Zhang, Y.; Xu, W.; Liu, C.; Zhang, G.; Liu, X.; Ma, Q. Chemical characteristics of PM 2.5 during summer at a background site of the Yangtze River Delta in China. *Atmos. Res.* 2017, 198, 163–172.

6. Kulkarni, P.S.; Dasari, H.P.; Sharma, A.; Bortoli, D.; Salgado, R.; Silva, A.M. Nocturnal surface ozone enhancement over Portugal during winter: Influence of different atmospheric conditions. *Atmos. Environ.* 2016, 147, 109–120.

7. Aksoyoglu, S.; Ciarelli, G.; El-Haddad, I.; Baltensperger, U.; Prévôt, A.S.H. Secondary inorganic aerosols in Europe: Sources and the significant influence of biogenic VOC emissions, especially on ammonium nitrate. *Atmos. Chem. Phys.* 2017, 17, 7757–7773.

8. Pye, H.O.T.; Zuend, A.; Fry, J.L.; Isaacman-VanWertz, G.; Capps, S.L.; Appel, K.W.; Foroutan, H.; Xu, L.; Ng, N.J.; Goldstein, A.H. Coupling of organic and inorganic aerosol systems and the effect on gas-particle partitioning in the southeastern US. *Atmos. Chem. Phys.* 2018, 18, 357–370.

9. Zheng, B.; Zhang, Q.; Zhang, Y.; He, K.B.; Wang, K.; Zheng, G.J.; Duan, F.K.; Ma, Y.L.; Kimoto, T. Heterogeneous chemistry: A mechanism missing in current models to explain secondary inorganic aerosol formation during the January 2013 haze episode in North China. *Atmos. Chem. Phys.* 2015, 15, 2031–2049.

10. Zhang, R.; Sun, X.; Shi, A.; Huang, Y.; Yan, J.; Nie, T.; Yan, X.; Li, X. Secondary inorganic aerosols formation during haze episodes at an urban site in Beijing, China. *Atmos. Environ.* 2018, 177, 275–282.

11. Yao, X.; Ge, B.; Yang, W.; Li, J.; Xu, D.; Wang, W.; Zheng, H.; Wang, Z. Affinity zone identification approach for joint control of PM2.5 pollution over China. *Environ. Pollut.* 2020, 265 (Pt B), 115086. [PubMed]

12. Bobbink, R.; Hicks, K.; Galloway, J.; Spranger, T.; Alkemade, R.; Ashmore, M.; Bustamante, M.; Cinderby, S.; Davidson, E.; Dentener, F.; et al. Global assessment of nitrogen deposition effects on terrestrial plant diversity: A synthesis. *Ecol. Appl.* 2010, 20, 30–59. [PubMed]
13. Liu, X.; Xu, W.; Duan, L.; Du, E.; Pan, Y.; Lu, X.; Zhang, L.; Wu, Z.; Wang, X.; Zhang, Y.; et al. Atmospheric Nitrogen Emission, Deposition, and Air Quality Impacts in China: An Overview. *Curr. Pollut. Rep.* 2017, 3, 65–77.

14. Lee, K.-S.; Lee, D.-S.; Lim, S.-S.; Kwak, J.-H.; Jeon, B.-J.; Lee, S.-L.; Lee, S.-M.; Choi, W.-J. Nitrogen isotope ratios of dissolved organic nitrogen in wet precipitation in a metropolis surrounded by agricultural areas in southern Korea. *Agric. Ecosyst. Environ.* 2012, 159, 161–169.

15. Xie, Y.; Xiong, Z.; Xing, G.; Yan, X.; Shi, S.; Sun, G.; Zhu, Z. Source of nitrogen in wet deposition to a rice agroecosystem at Tai lake region. *Atmos. Environ.* 2008, 42, 5182–5192.

16. Benedict, K.B.; Carrico, C.M.; Kreidenweis, S.M.; Schichtel, B.; Malm, W.C.; Collett, J.L., Jr. A seasonal nitrogen deposition budget for Rocky Mountain National Park. *Ecol. Appl.* 2013, 23, 1156–1169.

17. Ellermann, T.; Nygaard, J.; Christensen, J.; Løfstrøm, P.; Geels, C.; Nielsen, I.; Poulsen, M.; Monies, C.; Gyldenkærne, S.; Brandt, J.; et al. Nitrogen Deposition on Danish Nature. *Atmosphere* 2018, 9, 447.

18. Houlton, B.Z.; Boyer, E.; Finzi, A.; Galloway, J.; Leach, A.; Liptzin, D.; Melillo, J.; Rosenstock, T.S.; Sobota, D.; Townsend, A.R. Intentional versus unintentional nitrogen use in the United States: Trends, efficiency and implications. *Biogeochimistry* 2012, 114, 11–23.

19. Zhang, Q.; Zheng, Y.; Tong, D.; Shao, M.; Wang, S.; Zhang, Y.; Xu, X.; Wang, J.; He, H.; Liu, W.; et al. Drivers of improved PM2.5 air quality in China from 2013 to 2017. *Proc. Natl. Acad. Sci. USA* 2019, 116, 24463–24469.

20. Zheng, B.; Tong, D.; Li, M.; Liu, F.; Hong, C.; Geng, G.; Li, H.; Li, X.; Peng, L.; Qi, J.; et al. Trends in China’s anthropogenic emissions since 2010 as the consequence of clean air actions. *Atmos. Chem. Phys.* 2018, 18, 14095–14111.

21. Fu, Y.; Xu, W.; Wen, Z.; Han, M.; Sun, J.; Tang, A.; Liu, X. Enhanced atmospheric nitrogen deposition at a rural site in northwest China from 2011 to 2018. *Atmos. Res.* 2020, 245, 105071.

22. He, K.; Zhao, Q.; Ma, Y.; Duan, F.; Yang, F.; Shi, Z.; Chen, G. Spatial and seasonal variability of PM$_{2.5}$ acidity at two Chinese megacities: Insights into the formation of secondary inorganic aerosols. *Atmos. Chem. Phys.* 2012, 12, 1377–1395.

23. Yang, F.; Huang, L.; Duan, F.; Zhang, W.; He, K.; Ma, Y.; Brook, J.R.; Tan, J.; Zhao, Q.; Cheng, Y. Carbonaceous species in PM$_{2.5}$ at a pair of rural/urban sites in Beijing, 2005–2008. *Atmos. Chem. Phys.* 2011, 11, 7893–7903.

24. Tan, J.; Duan, J.; Zhen, N.; He, K.; Hao, J. Chemical characteristics and source of size-fractionated atmospheric particle in haze episode in Beijing. *Atmos. Res.* 2016, 167, 24–33.

25. Sun, Y.L.; Wang, Z.F.; Fu, P.Q.; Yang, T.; Jiang, Q.; Dong, H.B.; Li, J.; Jia, J.J. Aerosol composition, sources and processes during wintertime in Beijing, China. *Atmos. Chem. Phys.* 2013, 13, 4577–4592.

26. Xu, D.; Ge, B.; Wang, Z.; Sun, Y.; Chen, Y.; Ji, D.; Yang, T.; Ma, Z.; Cheng, N.; Hao, J.; et al. Below-cloud wet scavenging of soluble inorganic ions by rain in Beijing during the summer of 2014. *Environ. Pollut.* 2017, 230, 963–973.

27. Kang, Y.; Liu, M.; Song, Y.; Huang, X.; Yao, H.; Cai, X.; Zhang, H.; Kang, L.; Liu, X.; Yan, X.; et al. High-resolution ammonia emissions inventories in China from 1980 to 2012. *Atmos. Chem. Phys.* 2016, 16, 2043–2058.

28. Luo, X.; Liu, X.; Pan, Y.; Wen, Z.; Xu, W.; Zhang, L.; Kou, C.; Lv, J.; Goulding, K. Atmospheric reactive nitrogen concentration and deposition trends from 2011 to 2018 at an urban site in north China. *Atmos. Environ.* 2020, 224, 117298.

29. Yu, X.; Pan, Y.; Song, W.; Li, S.; Li, D.; Zhu, M.; Zhou, H.; Zhang, Y.; Li, D.; Yu, J.; et al. Wet and Dry Nitrogen Depositions in the Pearl River Delta, South China: Observations at Three Typical Sites With an Emphasis on Water-Soluble Organic Nitrogen. *J. Geophys. Res. Atmos.* 2020, 125, e2019JD030983.

30. Li, Z.; Walters, W.W.; Hastings, M.G.; Zhang, Y.; Song, L.; Liu, D.; Zhang, W.; Pan, Y.; Fu, P.; Fang, Y. Nitrate Isotopic Composition in Precipitation at a Chinese Megacity: Seasonal Variations, Atmospheric Processes, and Implications for Sources. *Earth Space Sci.* 2019, 6, 2200–2213.

31. Xu, W.; Liu, L.; Cheng, M.; Zhao, Y.; Zhang, L.; Pan, Y.; Zhang, X.; Gu, B.; Li, Y.; Zhang, X.; et al. Spatial–temporal patterns of inorganic nitrogen air concentrations and deposition in eastern China. *Atmos. Chem. Phys.* 2018, 18, 10931–10954.

32. Pan, Y.P.; Wang, Y.S.; Tang, G.Q.; Wu, D. Wet and dry deposition of atmospheric nitrogen at ten sites in Northern China. *Atmos. Chem. Phys.* 2012, 12, 6515–6535.

33. Liu, X.; Ju, X.; Zhang, Y.; He, C.; Kopsch, J.; Fusuo, Z. Nitrogen deposition in agroecosystems in the Beijing area. *Agric. Ecosyst. Environ.* 2006, 113, 370–377.
34. Xu, W.; Wen, Z.; Shang, B.; Dore, A.J.; Tang, A.; Xia, X.; Zheng, A.; Han, M.; Zhang, L.; Zhao, Y.; et al. Precipitation chemistry and atmospheric nitrogen deposition at a rural site in Beijing, China. Atmos. Environ. 2020, 223, 117253.

35. Sahu, L.K.; Kondo, Y.; Miyazaki, Y.; Kuwata, M.; Koike, M.; Takegawa, N.; Tanimoto, H.; Matsueda, H.; Yoon, S.C.; Kim, Y.J. Anthropogenic aerosols observed in Asian continental outflow at Jeju Island, Korea, in spring 2005. J. Geophys. Res. 2009, 114. [CrossRef]

36. Zeng, J.; Yue, F.-J.; Li, S.-L.; Wang, Z.-J.; Qin, C.-Q.; Wu, Q.-X.; Xu, S. Agriculture driven nitrogen wet deposition in a karst catchment in southwest China. Agric. Ecosyst. Environ. 2020, 294, 106883.

37. Zhao, M.; Huang, Z.; Qiao, T.; Zhang, Y.; Xiu, G.; Yu, J. Chemical characterization, the transport pathways and potential sources of PM2.5 in Shanghai: Seasonal variations. Atmos. Res. 2015, 158–159, 66–78.

38. Liang, L.; Engling, G.; Cheng, Y.; Tian, M.; Chen, Y.; Qiao, B.; Zhang, L.; Yang, F.; Zhang, L.; Luo, Q. Wet deposition and sources of inorganic nitrogen in the Three Gorges Reservoir Region, China. Environ. Pollut. 2018, 233, 520–528. [PubMed]

39. Liang, L.; Engling, G.; Du, Z.; Cheng, Y.; Duan, F.; Liu, X.; He, K. Seasonal variations and source estimation of saccharides in atmospheric particulate matter in Beijing, China. Chemosphere 2016, 150, 365–377. [PubMed]

40. Zhang, Z.; Gao, J.; Engling, G.; Tao, J.; Chai, F.; Zhang, L.; Zhang, R.; Sang, X.; Chen, C.-Y.; Lin, Z.; et al. Characteristics and applications of size-segregated biomass burning tracers in China’s Pearl River Delta region. Atmos. Environ. 2015, 102, 290–301.

41. Zhang, Z.; Gao, J.; Engling, G.; Tao, J.; Chai, F.; Zhang, L.; Zhang, R.; Sang, X.; Chen, C.-Y.; Lin, Z.; et al. Characteristic of 2-methyltetrols in ambient aerosol in Beijing, China. Atmos. Environ. 2012, 59, 376–381.

42. Chen, Z.; Huang, T.; Huang, X.; Han, X.; Yang, H.; Cai, Z.; Yao, L.; Han, X.; Zhang, M.; Huang, C. Characteristics, sources and environmental implications of atmospheric wet nitrogen and sulfur deposition in Yangtze River Delta. Atmos. Environ. 2019, 219, 116904.

43. Kuang, F.; Liu, X.; Zhu, B.; Shen, J.; Pan, Y.; Su, M.; Goulding, K. Wet and dry nitrogen deposition in the central Sichuan Basin of China. Atmos. Environ. 2016, 143, 39–50.

44. Fenn, M.E.; Ross, C.S.; Schilling, S.L.; Baccus, W.D.; Larrabee, M.A.; Lofgren, R.A. Atmospheric deposition of nitrogen and sulfur and preferential canopy consumption of nitrate in forests of the Pacific Northwest, USA. For. Ecol. Manag. 2013, 302, 240–253.
54. Zhang, C.-H.; Guo, H.-R.; Huang, H.; Ma, T.-Y.; Song, W.; Chen, C.-J.; Liu, X.-Y. Atmospheric nitrogen deposition and its responses to anthropogenic emissions in a global hotspot region. *Atmos. Res.* 2021, 248, 105137.

55. Calvo-Fernández, J.; Marcos, E.; Calvo, L. Bulk deposition of atmospheric inorganic nitrogen in mountainous heathland ecosystems in North-Western Spain. *Atmos. Res.* 2017, 183, 237–244.

56. Zheng, L.; Chen, W.; Jia, S.; Wu, L.; Zhong, B.; Liao, W.; Chang, M.; Wang, W.; Wang, X. Temporal and spatial patterns of nitrogen wet deposition in different weather types in the Pearl River Delta (PRD), China. *Sci. Total Environ.* 2020, 740, 139936.

57. Fahey, T.J.; Williams, C.J.; Rooney-Varga, J.N.; Cleveland, C.C.; Postek, K.M.; Smith, S.D.; Bouldin, D.R. Nitrogen Deposition In and Around an Intensive Agricultural District in Central New York. *J. Environ. Qual.* 1999, 28, 1585–1600.

58. Zhang, L.; Tian, M.; Peng, C.; Fu, C.; Li, T.; Chen, Y.; Qiu, Y.; Huang, Y.; Wang, H.; Li, Z.; et al. Nitrogen wet deposition in the Three Gorges Reservoir area: Characteristics, fluxes, and contributions to the aquatic environment. *Sci. Total Environ.* 2020, 738, 140309.

59. Zhang, X.; Lin, C.; Zhou, X.; Lei, K.; Guo, B.; Cao, Y.; Lu, S.; Liu, X.; He, M. Concentrations, fluxes, and potential sources of nitrogen and phosphorus species in atmospheric wet deposition of the Lake Qinghai Watershed, China. *Sci. Total Environ.* 2019, 682, 523–531.

60. Yu, W.T.; Jiang, C.M.; Ma, Q.; Xu, Y.G.; Zou, H.; Zhang, S.C. Observation of the nitrogen deposition in the lower Liaohe River Plain, Northeast China and assessing its ecological risk. *Atmos. Res.* 2011, 101, 46–468.

61. Yu, S.; Zhang, C.; Yuan, C.; Xu, H.; Ma, L.; Fang, Q.; Chen, G. Investigation on the influence of sulfur and chlorine on the initial deposition/fouling characteristics of a high-alkali coal. *Fuel Process. Technol.* 2020, 198, 106234.

62. NASA, Fire Information for Resource Management System. Available online: https://firms.modaps.eosdis.nasa.gov/map. (accessed on 20 October 2020).

63. Meng, X.; Zhou, W.; Rokni, E.; Chen, G.; Sun, R.; Levendis, Y.A. Release of Alkalis and Chlorine from Combustion of Waste Pinewood in a Fixed Bed. *Energy Fuels* 2019, 33, 1256–1266.

64. Yu, S.; Zhang, C.; Yuan, C.; Xu, H.; Ma, L.; Fang, Q.; Chen, G. Investigation on the influence of sulfur and chlorine on the initial deposition/fouling characteristics of a high-alkali coal. *Fuel Process. Technol.* 2020, 198, 106234.

65. Wilczyńska-Michalik, W.; Dańko, J.; Michalik, M. Characteristics of Particulate Matter Emitted from a Coal-Fired Power Plant. *Pol. J. Environ. Stud.* 2020, 29, 1411–1420.

66. Zhao, K.; Hu, Y.; Tian, Y.; Chen, D.; Feng, Y. Chlorine removal from MSWI fly ash by thermal treatment: Effects of iron/aluminum additives. *J. Environ. Sci.* 2020, 88, 112–121.

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