Dear Dr. Zhang,

Thank you very much for your constructive comments. According to the comments, we revised the manuscript carefully. The detailed responses to the comments are shown as below (editor's comments in black, authors' responses in blue). The marked-up manuscript version was combined at the end of this response letter. Please note that the line numbers in the following responses indicate the line numbers in the revised version of the manuscript, but NOT the marked-up manuscript version.

Again, we are grateful for your patience, guidance, and hard work on our manuscript.

Kind regards,

Yongwen Liu, Xu-Ri, Yuesi Wang, Yuepeng Pan, and Shilong Piao

Responses to Editor's Comments:

EC – Editor's Comments, AR – Authors' Responses.

EC: Dear Authors,

EC: Please proofread the paper to correct grammar issues and loosely structured sentences. Simplify discussions wherever you can since there are many repetitions. A few examples are provided here, as well as some minor comments.

AR: We corrected grammar issues and loosely structured sentences, and simplified discussions wherever we can. Please see the detailed responses to the specific comments as below.

EC: Lines 32-35, modify the sentence to "Results from the present study suggest that earlier estimations based on chemical transport model simulations and/or limited field measurements likely overestimated substantially the regional inorganic N wet deposition in the TP."

AR: Corrected (Lines 32-35).

EC: Line 157, use "precipitation"

AR: Corrected (Line 155).

EC: Line 162, use "analyzing"

AR: Corrected (Line 161).

EC: Line 206, use "sources"

AR: Corrected (Line 205).

EC: Line 221, change "using of the" to "using the"

AR: Corrected (Line 221).

EC: Line 233 and 235, use "ranging from"

AR: Corrected (Lines 233 and 235).

EC: Line 257, use "patterns"

AR: Corrected (Line 257).

EC: Line 287, use "3rd"

AR: Corrected (Line 287).

EC: Line 288, use "different from"

AR: Corrected (Line 288).

EC: The whole page of 16: Differences in wet deposition were largely explained by differences in precipitation amount. Does this mean the air concentrations are similar? Is there any information available on air concentration from literature?

AR: Indeed, based on our *in situ* measurements, differences in inorganic N wet deposition across the sites were mainly explained by differences in precipitation amount. As far as we know, no regional scale air concentrations in the TP was reported. It remains unclear whether the air concentrations are similar over the TP. Thus, we did not discuss this point in the revised manuscript.

EC: Line 356 and below, could be simplified substantially.

AR: We agree. The two paragraphs after this line was combined and simplified as follows (Lines 357-380):

We applied backward trajectory analysis to identify the long range transport of atmospheric inorganic N wet deposition at the five sites in the TP (Fig. 5). There is large spatial heterogeneity of air mass transport pathways across the five sites. At Muztagh Ata Station, wet deposition was mainly transported from Central Asia and Middle East (Fig. 5a). This is probably because Muztagh Ata Station is located in the northwestern TP, where is almost completely controlled by westerlies rather than Indian Monsoon (Yao et al., 2013). Thus, anthropogenic activities in Central Asia and Middle East are the principal source of the inorganic N wet deposition in the northwestern TP. Except for Muztagh Ata Station, inorganic N wet deposition at the other four sites was probably transported by India Monsoon (Figs. 5b-5e). At Ngari Station, 90.0% of wet deposition was transported from Nepal and North India through Indian Monsoon, and 10.0% of wet deposition came from Central Asia and Qaidam Basin through westerlies (Fig. 5b). At Qomolangma Station and Nam Co Station, inorganic N wet deposition was mainly influenced by the anthropogenic activities in the northeastern India and Bangladesh (Figs. 5c and 5d). At Southeast Tibet Station, 90.6% of wet deposition was transported from India, Bangladesh and Myanmar by India Monsoon, and the other 9.4% came from the western TP and Middle East (Fig. 5e). Therefore, inorganic N wet deposition at these four stations principally was influenced by the anthropogenic N emissions in South Asia (e.g. India). Actually, after China and USA, India has been the third largest producer and consumer of fertilizers due to intensification of agriculture, resulting in high anthropogenic N emissions (Aneja et al., 2012). For instance, ammonia (NH3) emissions from livestock and fertilizer applications in India in 2003 was estimated as 1705 Gg yr⁻¹ and 1697 Gg yr⁻¹, respectively (Aneja et al., 2012). Moreover, in India, field burning of crop residue (FBCR) is another critical anthropogenic activity leading to N emissions. In 2010, 6300 Gg of dry biomass are estimated to be subjected to FBCR in India, resulting in 350 Gg N emissions (Sahai et al., 2011). Besides Indian monsoon, biomass-burning emissions in South Asia could be across the Himalayas and transported to the TP by the mountain/valley wind (Cong et al., 2015).

EC: Line 407 and other similar places, use "Firstly".

AR: Corrected (Line 399).

Wet deposition of atmospheric inorganic nitrogen at five remote sites on in the Tibetan Plateau

- 3 Y. W. Liu^{1,3}, Xu-Ri^{1,2}, Y. S. Wang⁴, Y. P. Pan⁴, and S. L. Piao^{1,2,3}
- ⁴ ¹Key Laboratory of Alpine Ecology and Biodiversity, Institute of Tibetan Plateau Research, Chinese
- 5 Academy of Sciences, Beijing 100101, China
- 6 ²CAS Center for Excellence in Tibetan Plateau Earth Sciences, Beijing 100101, China
- ⁷ ³Sino-French Institute for Earth System Science, College of Urban and Environmental Sciences,
- 8 Peking University, Beijing 100871, China
- ⁹ ⁴State Key Laboratory of Atmospheric Boundary Layer Physics and Atmospheric Chemistry (LAPC),
- 10 Institute of Atmospheric Physics, Chinese Academy of Sciences, Beijing 100029, China
- 11 Correspondence to: Xu-Ri (xu-ri@itpcas.ac.cn)

12 Abstract

13 Since the mid-20th century, nitrogen (N) deposition has shown an increasing trend on-in the Tibetan Plateau (TP), where alpine ecosystems are sensitive to elevated N deposition. However, the 14 15 quantitative characterization of N deposition in on the TP remains unclear, due in most part to the lack of in situ measurement. Using the Tibetan Observation and Research Platform network, we conducted 16 short-term *in situ* measurements of major ions (NO₃⁻, Cl⁻, SO₄²⁻, NH₄⁺, Na⁺, K⁺, Ca²⁺, and Mg²⁺) wet 17 deposition at five remote sites on the TPin the TP during 2011-2013. At Southeast Tibet Station, Nam 18 19 Co Station, Qomolangma Station, Ngari Station, and Muztagh Ata Station, the NH4⁺-N wet deposition was 0.63, 0.68, 0.92, 0.36 and 1.25 kg N ha⁻¹ yr⁻¹, respectively; the NO₃⁻⁻N wet deposition was 0.28, 20

21	0.24, 0.03, 0.08 and 0.30 kg N ha ⁻¹ yr ⁻¹ , respectively; and the inorganic N <u>wet</u> deposition was 0.91,
22	0.92, 0.94, 0.44 and 1.55 kg N ha ^{-1} yr ^{-1} , respectively. The inorganic N wet deposition mainly occurred
23	as the form of NH4 ⁺ -N during the summer season at all stationssites. Results of enrichment factor
24	analysis and principal component analysis demonstrated that both NH_4^+ -N and NO_3^- -N wet deposition
25	in on-the TP were mainly influenced by anthropogenic activities. Backward trajectory analysis showed
26	that the inorganic N deposition at Muztagh Ata Station was mainly transported from Central Asia and
27	Middle East through westerlies. At Southeast Tibet Station, Nam Co Station, Qomolangma Station and
28	Ngari Station, the inorganic N deposition was mainly contributed by anthropogenic sources in South
29	Asia, and was mainly transported by Indian monsoon. Combining site-scale in situ measurements of
30	inorganic N wet deposition in this and previous studies, the average wet deposition of atmospheric
31	NH4 ⁺ -N, NO ₃ ⁻ -N, and inorganic N on the TPin the TP was estimated to be 1.09, 0.57 and 1.66 kg N
32	ha^{-1} yr ⁻¹ , respectively. The average NH ₄ ⁺ -N:NO ₃ ⁻ -N ratio in precipitation on the TP in the TP was
33	approximately 2:1. Results from the present study suggest that earlier estimations based on chemical
34	transport model simulations and/or limited field measurements likely overestimated substantially the
35	regional inorganic N wet deposition in the TP. Compared to the present study, previous regional-scale
36	estimation of inorganic N wet deposition for the entire TP, either through atmospheric chemistry
37	transport model simulations or interpolations based on limited field observations, has been highly
38	overestimated. To clarify the total N deposition on the TPin the TP more clearly, it is essential to
39	conduct long-term monitoring of both wet and dry deposition of atmospheric N in various climate
40	zones on the TP in the future.

1 Introduction

42 The global nitrogen (N) cycle has been disturbed by elevated reactive N emissions from anthropogenic 43 activities since the mid-19th century (Canfield et al., 2010; Galloway et al., 2008; Gruber and Galloway, 44 2008). Accumulated reactive N in the environment has led to a series of effects on climate change and ecosystems, e.g. air pollution, stratospheric ozone depletion, the potential alteration of global 45 temperature, drinking water contamination, freshwater eutrophication, biodiversity loss, grassland 46 seed bank depletion, and dead zones in coastal ecosystems (Basto et al., 2015; Erisman et al., 2011; 47 48 Erisman et al., 2013; Lan et al., 2015; Pinder et al., 2012; Shi et al., 2015; Zaehle et al., 2010). To 49 examine the actual amount of N inputted into ecosystems, several monitoring networks have been 50 established at national or continent scales, e.g. the National Atmospheric Deposition Program National 51 Trends Network (NADP/NTN, United States) (Lehmann et al., 2005), the Canadian Air and Precipitation Monitoring Network (CAPMoN, Canada) (Zbieranowski and Aherne, 2011), the 52 European Monitoring and Evaluation Programme (EMEP, Europe) (Fagerli and Aas, 2008), the 53 54 Austrian Precipitation Sampling Network (Austria) (Puxbaum et al., 2002), and the Japanese Acid 55 Deposition Survey (JADS, Japan) (Morino et al., 2011).

56 Besides Europe and North America, East Asia has become another high N deposition region, due to rapid economic growth in recent decades (Dentener et al., 2006). Across China, inorganic N wet 57 58 deposition has increased since the mid-20th century, albeit with inconsistent estimations of the change: 8 kg N ha⁻¹ yr⁻¹ (from 13.2 kg N ha⁻¹ yr⁻¹ in the 1980s to 21.1 kg N ha⁻¹ yr⁻¹ in the 2000s) (X. J. Liu 59 et al., 2013);), 2.8 kg N ha⁻¹ yr⁻¹ (from 11.11 kg N ha⁻¹ yr⁻¹ in the 1980s to 13.87 kg N ha⁻¹ yr⁻¹ in 60 the 2000s) (Jia et al., 2014);), and 7.4 kg N ha⁻¹ yr⁻¹ (from 12.64 kg N ha⁻¹ yr⁻¹ in the 1960s to 20.07 61 kg N ha⁻¹ yr⁻¹ in the 2000s) (Lu and Tian, 2014). Enhanced N deposition has changed the structure 62 and function of terrestrial, aquatic and coastal ecosystems in China (Liu et al., 2011). To accurately 63

estimate the N deposition in China, several monitoring networks have been established at the regional
scale, e.g. in northern China (Pan et al., 2012), in forest ecosystems along the North–South Transect
of Eastern China (NSTEC; based on the ChinaFLUX network) (Sheng et al., 2013), and in subtropical
forest ecosystems in South China (Chen and Mulder, 2007). However, there are few observation sites
distributed in western China, particularly on-in the Tibetan Plateau (TP), resulting in uncertainty
regarding the N deposition for China as a whole (Jia et al., 2014; X. J. Liu et al., 2013; Lu and Tian,
2014).

The TP covers an area of about 2.57 million km², occupying approximately 1/4 of the land area of 71 72 China (Zhang et al., 2002). On the TPOver the TP, alpine ecosystems are widely distributed and are 73 sensitive to elevated N deposition. Multi-level N fertilization experiments have shown that alpine 74 grassland ecosystems are N limited and have potential capacity to absorb increased N deposition (Y. 75 W. Liu et al., 2013; Xu et al., 2014). However, long-term N addition can decrease the species richness 76 of both vegetation and soil seed banks in alpine meadow ecosystems on the TP in the TP (Ma et al., 77 2014). Ice core records show that the inorganic N deposition on the TP in the TP has increased during 78 recent decades (Hou et al., 2003; Kang et al., 2002a, b; Thompson et al., 2000; Zhao et al., 2011; Zheng 79 et al., 2010), and. tThis trend is also apparent in sediment cores of alpine lakes in the western and 80 southeastern TP (Choudhary et al., 2013; Hu et al., 2014). To recognize the characteristics of ion 81 deposition on the TP in the TP, a number of observations of precipitation chemistry have been carried 82 out in the eastern TP in recent years (Jia, 2008; Tang et al., 2000; Zhang et al., 2003; N. N. Zhang et al., 2012). Nevertheless, in the central and western TP, observation sites are scarce, indicating that the 83 84 situation in terms of N deposition across the entire TP remains unclear.

To quantitatively estimate the inorganic N wet deposition on the TPin the TP, we investigated the precipitation chemistry characteristics at five remote sites, situated mainly on in the central and western TP. The sites are part of the Tibetan Observation and Research Platform (TORP) network (Ma et al., 2008). Specifically, our aims were to (1) clarify the <u>characteristics amount</u> of inorganic N wet deposition on <u>in</u> the central and western TP, and (2) <u>quantitatively</u> assess the <u>amount of</u>-inorganic N wet deposition for <u>in</u> the entire TP by combining site-scale *in situ* measurements <u>in this with and those</u> in-previous studies.

92 **2 Materials and methods**

93 **2.1 Precipitation sampling and chemical analysis**

Using the Tibetan Observation and Research Platform (TORP) network (Ma et al., 2008), precipitation
chemistry observations were conducted at five sampling sites: Southeast Tibet Station, Nam Co Station,
Qomolangma Station, Ngari Station, and Muztagh Ata Station (Fig. 1), situated from the eastern to
western TP and covering various climatic zones and vegetation types. A brief description of the five
sites is shown in Table1.

99 During 2011–2013, we collected precipitation samples at each site, lasting at least one year. 100 Precipitation samples were collected following each precipitation event, using an inner removable 101 high-density polyethylene (HDPE) plastic bag in a pre-cleaned HDPE bucket. The HDPE bucket was 102 placed 1.5 m above the ground. We opened the plastic bag at the beginning of the precipitation event, 103 and collected precipitation samples at the end of the precipitation process. Then, the samples were 104 transferred into pre-cleaned HDPE bottles (50 mL). Snowfall samples were melted at room 105 temperature before being transferred into the HDPE bottles. All samples were kept frozen at the station and during transport until analysis in the laboratory. A total of 259 precipitation samples were collected,
among which eight samples were abandoned due to breakage during transportation or the samples
volume being less than 10 mL.

109 We analyzed the chemical composition of all precipitation samplings at the State Key Laboratory of 110 Environmental Aquatic Chemistry, Research Center for Eco-Environmental Sciences, Chinese Academy of Sciences. Analyzed ions included NO₃⁻, Cl⁻, SO₄²⁻, NH₄⁺, Na⁺, K⁺, Ca²⁺-, and Mg²⁺. All 111 ions were analyzed by the Ion Chromatography of Dionex-ICS2100. Samples for cation analysis were 112 eluted on a Dionex 4-mm CS12A separatory column using 20 mM Methanesulfoni acid solution for 113 114 an eluent pumped with a flow rate of 1.0 mL min⁻¹. Suppression was provided by a Dionex CSRS300 115 suppressor in recycle mode. For anion analysis, an IonPac AS19-HC column, 25 mM NaOH eluent, and ASRS300 suppresser were used. The analytical detection limit was 2 ng g^{-1} for all ions. 116

117 **2.2 Data quality control**

Previously documented methods (Rodhe and Granat, 1984;Safai et al., 2004) were used for quality assurance and quality control purposes., resulting in sSix (2.4%) samples, which fell outside the range $(m-3\delta, m+3\delta)$, and therefore were, being excluded. Here, *m* is the mean value and δ is the standard deviation. The Pearson correlation between Σ_{anions} and $\Sigma_{cations}$ was 0.82 (*p*-*P* < 0.001), suggesting credible data quality. The ratio of total anions to total cations was calculated following Eq. (1),

123
$$\frac{\sum \text{anions}}{\sum \text{cations}} = \frac{\sum_{k=1}^{n} (NO_{3}^{+} + Cl^{-} + SO_{4}^{2-})}{\sum_{k=1}^{n} (NH_{4}^{+} + Na^{+} + K^{+} + Ca^{2+} + Mg^{2+})},$$
(1)

where *n* is the number of samples. The ratio of $(\Sigma_{anions}/\Sigma_{cations})$ -was 0.26, indicating that at least one major anion was not measured (C. Li et al., 2007). Considering that pH is-was alkaline in both 126 precipitation and the surface soil layer (Ding et al., 2004; Y. H. Yang et al., 2012), the unmeasured 127 anion was likely HCO_3^- (C. Li et al., 2007).

128 **2.3 Statistical Analysis**

For each site, consecutive samples in one <u>year-year-round sampling period</u> were selected to analyze the annual mean values of ions. The sampling times of the samples used at the five sites were as follows: Southeast Tibet Station, November 2011 to October 2012; Nam Co Station, August 2011 to July 2012; Qomolangma Station, April 2011 to March 2012; Ngari Station, January 2013 to December 2013; Muztagh Ata Station, January 2011 to December 2011. A total of 168 precipitation samples were selected, among which the number of samples for Southeast Tibet Station, Nam Co Station, Qomolangma Station, Ngari Station, and Muztagh Ata Station was 53, 27, 30, 39, and 19, respectively.

136 The annual average ion concentration was calculated as the volume-weighted mean (VWM) following137 Eq. (2),

138
$$C = \frac{\sum_{i=1}^{n} (C_i \times P_i)}{\sum_{i=1}^{n} P_i},$$
 (2)

where *C* is the annual average ion concentration ($\mu eq L^{-1}$), *C_i* is the ion concentration of an individual sample *i* ($\mu eq L^{-1}$), and *P_i* is the precipitation amount corresponding to the sample *i* (mm).

141 Wet deposition of atmospheric N was calculated following Eq. (3),

142
$$N_{\text{wet}} = 0.00014 \times C_{\text{N}} \times P_{\text{annual}},\tag{3}$$

where N_{wet} is the annual wet deposition of atmospheric inorganic N (NH₄⁺-N or NO₃⁻-N, kg N ha⁻¹ yr⁻¹),-); C_N is the annual average equivalent concentration of N in precipitation (NH₄⁺-N or NO₃⁻-N, $\mu eq L^{-1}, \underline{}; P_{annual} \text{ is annual precipitation (mm yr}^{-1}), \underline{}; \text{ and } 0.00014 \text{ is the shift coefficient for the unit}$ $146 \quad \text{of } \mu eq L^{-1} \times \text{mm yr}^{-1} \text{ to the unit of } \text{kg N ha}^{-1} \text{ yr}^{-1}. \text{ Here, } 1 \mu eq \text{ NH}_4^+ \text{-N or NO}_3^- \text{-N contains } 1 \mu \text{mol N},$ $147 \quad \text{and the weight of } 1 \mu \text{mol N is } 14 \times 10^{-9} \text{ kg}. \text{ Thus, } \mu eq L^{-1} \times \text{mm yr}^{-1} = 14 \times 10^{-9} \text{ kg N} \times 10^3 \text{ m}^{-3} \times$ $148 \quad 10^{-3} \text{ m yr}^{-1} = 14 \times 10^{-9} \text{ kg N} \times 10^4 \text{ ha}^{-1} \text{ yr}^{-1} = 0.00014 \text{ kg N ha}^{-1} \text{ yr}^{-1}.$

149 **2.4 Source assessment of ion wet deposition**

150 **2.4.1 Enrichment factor**

151 Enrichment factor (EF) has been widely used to examine the source contributions of major ions wet 152 deposition in previous studies (Cao et al., 2009; Chabas and Lefevre, 2000; Kulshrestha et al., 1996; 153 Lu et al., 2011; Okay et al., 2002; Shen et al., 2013; Xiao et al., 2013; Zhang et al., 2007). Commonly, 154 Na is considered as the best reference element for seawater, due to its almost purely marine origin 155 (Keene et al., 1986; Kulshrestha et al., 2003). Another element, Ca is normally used as a reference element for continental crust, because Ca is a typical lithophile element and its composition in soil 156 157 barely changes (Zhang et al., 2007). In this study, Na and Ca were also-used as reference element for 158 seawater and continental crust, respectively.

On the TPIn the TP, multiple lines of evidence demonstrate that Na⁺ in precipitation on the TP mainly comes from oceans. Balestrini et al. (2014) monitored the chemical and isotopic compositions of precipitations at the Pyramid International Laboratory (5050_m a.s.l.) on the southern slope of the Himalayas, and data analysis suggested that Na⁺ and Cl⁻ were derived from the long-range transport of marine aerosols. Ice records in the central Himalayas show that Cl⁻/Na⁺ was positively related with the monsoon rainfall in northeast India, and there was a teleconnection between the Na⁺ and Cl⁻ concentrations and the North Atlantic Oscillation₄₅ indicating that Na⁺ in the ice core mainly came from oceans (Wang et al., 2002). Na⁺ has been used as a marine tracer when analyzinge the source
contributions of ions wet deposition on-in the northeastern TP (Li et al., 2015), the southeastern TP (B.
Liu et al., 2013) and the southern slope of central Himalayas (Tripathee et al., 2014).

169 On-Over the TP, sandy desertification land covers about 3.1×10^5 km², accounting for 14% of the whole 170 plateau, of which moderate sandy desertification land occupies 55.44% (Liu et al., 2005). The TP was 171 is regarded as an important dust source region (Fang et al., 2004; Han et al., 2009; Han et al., 2008). 172 The TP dust sources contribute 69% of dust at the surface and 40% of dust in the lower troposphere 173 on-over the TP (Mao et al., 2013). Moreover, arid regions are widely distributed surrounding the TP, 174 e.g. central Asia, and the deserts of in western China. The dust on over the TP partly comes from the 175 adjacent dust source regions, e.g. Taklimakan Desert in western China (Huang et al., 2007; Xia et al., 176 2008). Atmospheric dust aerosols over the TP are strongly impacted by local sources and enriched with 177 Ca (Zhang et al., 2001). These dust aerosols in the atmosphere can interact with clouds and precipitation (Huang et al., 2014), and deposit on the surface with precipitation. Thus, Ca^{2+} is 178 179 commonly used as a proxy of dust for in ice core studies in the TP (Kang et al., 2002a; Kang et al., 2010; Kaspari et al., 2007; Wang et al., 2008). As a dust proxy, Ca²⁺ record in an ice core from the 180 181 central TP even-was significantly related regional zonal wind (westerlies) trends, and reflected the 182 long-term control of regional atmospheric circulation strength over atmospheric dust concentrations (Grigholm et al., 2015). In addition, Ca also has been used as a reference element for continental crust 183 184 when assessing sources of ion wet deposition in precipitation on in the northern TP (Li et al., 2015).

In this study, the EF of an ion in precipitation relative to the ion in sea was estimated using Na as areference element following Eq. (4),

187
$$EF_{sea} = \frac{[X/Na^+]_{rain}}{[X/Na^+]_{sea}},$$
(4)

where EF_{sea} is the EF of an ion in precipitation relative to the ion in sea, X is an ion in precipitation, [X/Na⁺]_{rain} is the ratio from precipitation composition ($\mu eq X / \mu eq Na^+$), +); and [X/Na⁺]_{sea} is the ratio from sea composition (Keene et al., 1986; Turekian, 1968) ($\mu eq X / \mu eq Na^+$).

191 The EF of an element in precipitation relative to the element in soil was estimated using Ca as a192 reference element following Eq. (5),

193
$$\text{EF}_{\text{soil}} = \frac{[X/\text{Ca}^{2+}]_{\text{rain}}}{[X/\text{Ca}^{2+}]_{\text{soil}}},$$
 (5)

where EF_{soil} is the EF of an element in precipitation relative to the element in soil, X is an ion in precipitation, $(X/Ca^{2+})_{soil}$ is the ratio from precipitation composition ($\mu g X / \mu g Ca^{2+}$), +); and $[X/Ca^{2+}]_{soil}$ is the ratio from soil composition (Taylor, 1964) ($\mu g X / \mu g Ca^{2+}$).

197 To estimate fractions of marine, crustal and anthropogenic contributed to ions in precipitation, we 198 calculated the sources of ionic components in precipitation using equations from previous studies (Cao 199 et al., 2009; Lu et al., 2011; Zhang et al., 2007) as follows:

200 SSF (%) =
$$\frac{[X/Na^+]_{sea}}{[X/Na^+]_{rain}} \times 100,$$
 (6)

201 CF (%) =
$$\frac{[X/Ca^{2+}]_{soil}}{[X/Ca^{2+}]_{rain}} \times 100,$$
 (7)

202 AF (%) =
$$100 - SSF - CF$$
, (8)

where SSF is sea salt fraction, <u>CF</u> is crust fraction, <u>ind</u> AF is anthropogenic fraction. Note that if
SSF is greater than 1, SSF is recalculated as the difference between 1 and CF, and if CF is greater than
1, CF is recalculated as the difference between 1 and SSF.

206 **2.4.2 Principal component analysis**

Principal component analysis has been widely used in precipitation chemical studies to determine the effect of natural and anthropogenic sources on chemical composition of precipitation (Balasubramanian et al., 2001; Cao et al., 2009; Migliavacca et al., 2005; Zhang et al., 2007). In this study, principal component analysis was also used to examine the various sources of major ions in precipitation at the five remote sites on the TPin the TP. Varimax-rotated principal component analysis was performed using "principal" function in package "psych" of R 3.2.0 (R Core Team, 2015; http://www.R-project.org, last visited October 6, 2015).

214 **2.4.3 Backward trajectory analysis**

215 To identify the long range transport of water-soluble ions in precipitation, seven-day backward 216 trajectories arriving at the sampling sites for each individual precipitation event were calculated. 217 Backward trajectories calculated using TrajStat (version 1.4.4R4, were http://www.meteothinker.com/TrajStatProduct.html, last visited October 6, 2015), which is a 218 219 Geographic Information System based software, including a trajectory calculation module of 220 HYSPLIT (Hybrid Single Particle Lagrangian Integrated Trajectory Model; 221 http://www.arl.noaa.gov/ready/hysplit4.html, last visited October 6, 2015) (Wang et al., 2009). The 222 input meteorological data was the Global Data Assimilation System (GDAS) meteorological data 223 archives of the Air Resource Laboratory, National Oceanic and Atmospheric Administration (NOAA)

224	(ftp://arlftp.arlhq.noaa.gov/pub/archives/gdas1, last visited October 6, 2015). All backward trajectories
225	were calculated at 6-h interval (00:00, 06:00, 12:00, 18:00 UTC) at each sampling day, with an arrival
226	height of 500 m above the ground. Then, cluster analysis was performed using of the trajectories during
227	the one year-round sampling period at each site using TrajStat (version 1.4.4R4).

228 3 Results

229 **3.1 Chemical composition of atmospheric precipitation**

230 Figure 2 shows the seasonal dynamics of ion concentrations in precipitation at the five remote sites on 231 the TP in the TP. Wet deposition of all ions mainly occurs in summer season at all sites. Compared to 232 the sites with relatively higher precipitation amounts, e.g. Southeast Tibet Station and Nam Co 233 StationNgari Station and Muztagh Ata Station, the sites with relatively lower precipitation amounts 234 had relatively higher ion concentrations, e.g. Ngari Station and Muztagh Ata Station Southeast Tibet 235 Station and Nam Co Station (Fig. 2, Table 2). Ca²⁺ had the highest annual VWM concentration in 236 precipitation at most sites (except for Nam Co Station), with the highest proportion accounting for 237 measured ions of 54.6% at Southeast Tibet Station (Fig. 2 and 3). At Nam Co Station, NH4⁺ in 238 precipitation had the highest proportion accounting for measured ions of 39.5%, higher than those at 239 the other sites (ranging from 12.9% at Southeast Tibet Station to 18.9% at Muztagh Ata Station) (Fig. 240 3). Compared to NH₄⁺, NO₃⁻ had much lower proportion accounting for measured ions in precipitation, 241 ranging from 0.6% at Qomolangma Station to 14% at Nam Co Station (Fig. 3). The order of the average annual VWM of ion deposition at the five stations sites was: $Ca^{2+} > NH_4^+ > SO_4^{2-} > Cl^- > Na^+ > Mg^{2+} > Cl^- > Na^+ > Mg^{2+} > NH_4^+ > SO_4^{2-} > Cl^- > Na^+ > Mg^{2+} > SO_4^{2-} > Cl^- > SO_4^{2-} > Cl^- > SO_4^{2-} > Cl^- > SO_4^{2-} > SO_4^$ 242 243 $NO_3^- > K^+$ (Table 2). All major ion concentrations in precipitation on the TP in the TP were much lower 244 than those in northern and southern China (Table 2).

245 **3.2 Wet deposition of atmospheric inorganic N**

246 At Southeast Tibet Station, Nam Co Station, Qomolangma Station, Ngari Station, and Muztagh Ata Station, the NH₄⁺-N wet deposition was 0.63, 0.68, 0.92, 0.36 and 1.25 kg N ha⁻¹ yr⁻¹, respectively; 247 the NO₃⁻-N wet deposition was 0.28, 0.24, 0.03, 0.08 and 0.3 kg N ha⁻¹ yr⁻¹, respectively; and the 248 249 inorganic N wet deposition was 0.91, 0.92, 0.94, 0.44 and 1.55 kg N ha⁻¹ yr⁻¹, respectively (Table 3). 250 Besides the above five sites of the TORP network, previous site-scale in situ measurements of 251 inorganic N wet deposition at other sites on the TP in the TP were also collected, e.g. at Waliguan (Tang 252 et al., 2000), Wudaoliang (Yang et al., 1991), Lhasa (Zhang et al., 2003), Naidong (Jia, 2008), Biru 253 (Jia, 2008), Jiangda (Jia, 2008), and Lijiang (N. N. Zhang et al., 2012). Combining the site-scale in situ 254 measurements in our study with and those in previous studies, the average wet deposition of 255 atmospheric NH₄⁺-N, NO₃⁻-N and inorganic N on the TP in the TP were was estimated to be 1.09, 0.57 and 1.66 kg N ha⁻¹ yr⁻¹, respectively, and the estimated NH₄⁺-N:NO₃⁻-N ratio in precipitation on the 256 257 **TP**in the TP was approximately 2:1. Both NH₄⁺-N and NO₃⁻-N wet deposition on the TPin the TP were 258 much lower than those in northern and southern China (Table 3).

3.3 Seasonal dynamics of inorganic N wet deposition

The inorganic N wet deposition mainly occurred as the form of NH_4^+ -N during the summer season at all sites (Fig. 4). Both concentrations of NH_4^+ and NO_3^- did not exhibited any clear seasonal pattern (Fig. 2). The seasonal dynamics inorganic N wet deposition at most stations appeared the shape of single peak type (Fig. 4). The seasonal patterns of inorganic N wet deposition were similar to the seasonal patterns of precipitation, rather than that of NH_4^+ or NO_3^- concentration (Fig. 2).

3.4 Source assessment of wet deposition of inorganic N and other ions

266 **3.4.1 Enrichment factors**

Table 4 shows the EFs of precipitation constituents at the five sites relative to seawater and soil. If the EF value of an ion in precipitation is much higher (lower) than 1, the ion is considered to be enriched (diluted) relative to the reference source. Among the five sites, Cl⁻ had a relatively lower EF_{sea} value, ranging from 0.50 (Nam Co Station) to 0.90 (Qomolangma Station), but a relatively higher EF_{soil} value, ranging from 42.4 (Muztagh Ata Station) to 286 (Qomolangma Station). Different from Cl⁻, NH₄⁺ in precipitation was enriched relative to both marine origin and soil reference source at all sites, because its EF_{sea} values ranged from 11629 to 80684, and its EF_{soil} ranged from 350 to 2378. Similar to NH₄⁺,

 NO_3^- also had a relatively high value of both EF_{sea} and EF_{soil} at all five sites.

Table 5 shows the source contributions for major ions in precipitation of the five remote sites in this study. Almost all Cl⁻ and Na⁺ in precipitation on the TPin the TP appeared to be of marine origin, with SSF value above 95% at the five sites. Nearly all Ca²⁺ in precipitation came from crust at the five sites, with the CF value being above 90%. Among Across the five sites, anthropogenic sources contributed at least 99% of NH₄⁺ in precipitation, and NO₃⁻ in precipitation was also mainly influenced by anthropogenic activities, with AF values ranging from 95.3% to 99.9%.

3.4.2 Principal component analysis

Table 6 shows the 1st, 2nd and 3rd component of principal component analysis, which account<u>ed</u> for at least 85% of the total variance across the five sites. Na⁺ and Cl⁻ were mainly explained by the same component at all sites. Principal component analysis shows that the variances of Ca²⁺ and Na⁺ were represented by different components at four of five sites (except Southeast Tibet Station) (Table 6). The common variance of Ca²⁺, Mg²⁺ and SO₄²⁻ as 1st component represents the largest proportion of

287 the total specie variation at the 3 sites (Nam Co Station, Ngari Station, and Muztagh Ata Station) in 288 the central and western TP (Table 6). At Qomolangma Station, Na⁺, Cl⁻, K⁺ and NH₄⁺ as 1st component 289 represents the largest proportion of the total specie variation (Table 6). Except for Qomolangma Station, 290 at the other four sites, the variances of NH_4^+ were mainly represented by 3rd component (Table 6). At Southeast Tibet Station, both Ca²⁺ and Na⁺ variances were mostly represented by the 1st component, 291 292 but NO₃⁻ variances were mainly represented by the 2nd component (Table 6). At Nam Co Station, 293 Qomolangma Station, and Ngari Station, NO₃⁻ variances were mainly represented by the 3rnd 294 component, which were different with from that of both Ca^{2+} and Na^+ (Table 6). However, NO_3^- 295 variances were mainly represented by the 1st component at Muztagh Ata Station (Table 6).

3.4.3 Backward trajectory analysis

297 Fig. 5 shows the seven-day backward trajectories of air mass arriving at the five remote sites at the 298 sampling days. The transport pathways of air masses were various with the different sites (Fig. 5). The 299 cluster trajectory results showed that at Muztagh Ata Station, nearly all air masses at sampling days 300 were transported from Central Asia and Middle East (Fig. 5a). Different to from Muztagh Ata Station, 301 almost all air masses at Nam Co Station were transported from South Asia (Fig. 5d). For Ngari Station, 302 Qomolangma Station, and Southeast Tibet Station, the air masses at sampling days were mainly 303 transported from South Asia, with the proportion of 90%, 79.8%, and 90.6%, respectively (Figs. 5b-304 5e). Besides South Asia, Central Asia, Qaidam Basin, and Middle East was the second source of air 305 masses at sampling days for Ngari Station, Qomolangma Station, and Southeast Tibet Station, 306 respectively (Figs. 5b-5e).

307 4 Discussion

4.1 Wet deposition of atmospheric inorganic N on in the TP

309 According to our field observations, wet deposition of atmospheric inorganic N on-in the western TP 310 was lower than that on-in the eastern TP. For example, the rates of inorganic N wet deposition at Ngari Station and Muztagh Ata Station, located on the western TP, were 0.44 and 1.55 kg N ha⁻¹ yr⁻¹, B11 312 respectively. These inorganic N wet deposition in the western TP, which were much lower than those at the stations sites on in the eastern TP, e.g. Jiangda (1.91 kg N ha⁻¹ yr⁻¹), Lijiang (1.89 kg N ha⁻¹ yr⁻¹) 313 and Waliguan (2.92 kg N ha^{-1} yr⁻¹) (Table 3). However, the concentrations of inorganic N in 314 315 precipitation at the sites on in the western TP were comparable to those at the sites on in the eastern 316 TP. For instance, the annual average concentrations of NH4⁺ in precipitation at Ngari Station and 317 Muztagh Ata Station were 20.5 and 42.0 μ eq L⁻¹, respectively, which were even higher than those at 318 stations on-in the eastern TP, e.g. Lijiang (11.4 μ eq L⁻¹) and Lhasa (14.3 μ eq L⁻¹) (Table 2). Meanwhile, 319 compared to Lijiang and Lhasa, Ngari Station and Muztagh Ata had lower annual precipitation rates of 124.6 and 213.6 mm yr⁻¹ (Table 2). Therefore, compared to the eastern TP, the western TP had 320 321 relatively lower inorganic N deposition, probably due to its lower precipitation amount rather than its 322 comparable inorganic N concentration in precipitation.

Wet deposition of inorganic N for the entire TP was much lower than that in northern and southern China (Table 3). The average wet deposition of atmospheric inorganic N (sum of NH_4^+ -N and NO_3^+ -N) for the TP was estimated to be 1.66 kg N ha⁻¹ yr⁻¹. This was much lower than the inorganic N wet deposition <u>in-at</u> the cities <u>of-in</u> both northern and southern China, e.g. Beijing, Tianjin, Tangshan, Dalian, Nanjing, Hangzhou, Ningbo, Shanghai, Shenzhen, and Guiyang (Table 3). Moreover, the inorganic N wet deposition <u>on the TPin the TP</u> was also lower than that in the forest ecosystems of eastern China, e.g. TieShanPing, LiuChongGuan, LeiGongShan, CaiJiaTang, and XiLiuHe (Table 3). 330 Overall, compared to eastern China, the TP had relatively lower inorganic N wet deposition, probably 331 due to following two reasons. Firstly, except for Southeast Tibet Station and Lijiang, most N 332 observation sites on the TP in the TP are located in typical arid and semi-arid regions, with annual precipitation ranging from 124.6 mm yr⁻¹ at Ngari Station to 582 mm yr⁻¹ at Biru. Compared to this, 333 334 annual precipitation rates at sites in eastern China are much higher, particularly in southern China, where annual precipitation ranges from 825.5 mm yr^{-1} at Shanghai to 1769 mm yr^{-1} at Shenzhen (Table 335 336 3). Secondly, the average annual concentration of inorganic N (NH₄⁺-N or/and, NO₃⁻-N) in 837 precipitation on the TP in the TP was much lower than that in eastern China, especially at cities in 338 northern China (Table 2). This is probably because the effects of anthropogenic activities in eastern 339 China are much more intense than that on the TP in the TP, which is referred to as "The Third Pole" 340 and has an average altitude exceeding 4,000 meters above sea level and is referred to as "The Third 841 Pole" –(Qiu, 2008; Yao et al., 2012).

4.2 Source assessment of atmospheric inorganic N wet deposition on the TP in the TP

To analyze the source contributions of major ions wet deposition, EF was applied using Na and Ca as 343 344 reference element for seawater and continental crust, respectively. Here, Na and Ca in precipitation on 345 the TPin the TP was hypothesized mainly coming seawater and continental crust, respectively. This 346 assumption was partly confirmed by the results of principal component analysis in this study (Table 6). Principal component analysis shows that the variances of Ca^{2+} and Na^{+} were represented by 347 348 different components at four of five sites (except Southeast Tibet Station), indicating different source 349 of Ca^{2+} and Na^{+} in precipitation on the TP in the TP (Table 6). Moreover, Na^{+} and Cl^{-} were mainly 350 explained by the same component at all sites. This indicates that Na⁺ and Cl⁻ were likely contributed by the same source: sea-salt (Table 6). This assumption was also confirmed by the relatively high 351

Pearson correlation between Na⁺ and Cl⁻ at all five sites (Table S1 in the supplementary material). At Southeast Tibet Station, both Ca²⁺ and Na⁺ variances were mostly represented by the 1st component (Table 6). This probably because South Asia is also an important source of dust aerosols in the southeastern TP during the during the monsoon period (Zhao et al., 2013).

356 EF analysis results showed that at all the five sites, both NH₄⁺ and NO₃⁻ in precipitation were mainly 357 contributed by anthropogenic sources (Table 5). This was also confirmed by principal component analysis. Different with Ca²⁺ and Na⁺, NH₄⁺ variances were mainly represented by 3rd component at 358 four in five sites (except for Qomolangma Station) (Table 6). Except for Muztagh Ata Station, at the 359 360 other four stations, NO₃⁻ variances were also represented by different component with that of Ca²⁺ and 361 Na⁺ variances (Table 6). This indicates that the source of inorganic N wet deposition was probably 362 different with the sources of Ca^{2+} or Na^+ wet deposition. Meanwhile, at all five sites, Na and Ca mainly came from seawater and continental crust, respectively. Therefore, inorganic N wet deposition at the 363 364 five sites on the TP in the TP was mainly influenced by anthropogenic activities.

365 TP is mainly controlled by westerlies and Indian monsoon (Yao et al., 2013). The northern TP is mainly 366 influenced by westerlies, and the southern TP is mainly controlled by the Indian monsoon in summer seasons, but by the westerlies during non-monsoon seasons (Yao et al., 2013). N emissions from 367 368 anthropogenic activities in Central Asia might be transported to the northern TP by westerlies. N 369 emissions from anthropogenic activities in South Asia (e.g. India) might be transported to the southern 370 TP by Indian monsoon. After China and USA, India has been the third largest producer and consumer 371 of fertilizers due to intensification of agriculture, resulting in high anthropogenic N emissions (Aneja 372 et al., 2012). At present, reactive N emissions from crop and livestock farming in India were second 373 only to that in China (Aneja et al., 2012). For instance, ammonia (NH₃) emissions from livestock and fertilizer applications in India in 2003 was estimated as 1705 Gg yr⁺ and 1697 Gg yr⁺, respectively (Aneja et al., 2012). Moreover, in India, field burning of crop residue (FBCR) is another critical anthropogenic activity leading to N emissions. N emissions in India due to FBCR showed increase trend in the past 3 decades (Sahai et al., 2011). In 2010, 6300 Gg of dry biomass are estimated to be subjected to FBCR in India, resulting in 350 Gg N emissions (Sahai et al., 2011). Besides Indian monsoon, biomass burning emissions in South Asia could be across the Himalayas and transported to the TP by the mountain/valley wind (Cong et al., 2015).

381 We applied backward trajectory analysis to identify the long range transport of atmospheric inorganic 382 N wet deposition at the five sites on the TP in the TP (Fig. 5). There is large spatial heterogeneity of air 383 mass transport pathways across the five sites. At Muztagh Ata Station, wet deposition was mainly 384 transported from Central Asia and Middle East (Fig. 5a). It seems that air mass from South Asia has 385 no influence on the wet deposition at Muztagh Ata Station. This is probably because Muztagh Ata 386 Station is located on-in the northwestern TP, where is almost completely controlled by westerlies τ 387 rather than Indian Monsoon (Yao et al., 2013). Thus, anthropogenic activities in Central Asia and 388 Middle East are the principal source of the inorganic N wet deposition at Muztagh Ata Station in the 389 northwestern TP. Except for Muztagh Ata Station, inorganic N wet deposition at the other four sites 390 was mainly influenced by anthropogenic activities in South Asia and was probably transported by India 391 Monsoon (Figs. 5b-5e). At Ngari Station, 90.0% of wet deposition was transported from Nepal and 392 North India through Indian Monsoon, and 10.0% of wet deposition was transported came from Central 393 Asia and Qaidam Basin through westerlies (Fig. 5b). At Qomolangma Station and Nam Co Station, 394 inorganic N wet deposition was mainly influenced by the anthropogenic activities in the northeastern 395 India and Bangladesh (Figs. 5c and 5d). At Southeast Tibet Station, 90.6% of wet deposition was

396	transported from India, Bangladesh and Myanmar by India Monsoon, and the other 9.4% came from
397	the western TP and Middle East (Fig. 5e). Therefore, inorganic N wet deposition at these four stations
398	principally was influenced by the anthropogenic N emissions in South Asia (e.g. India). Actually, after
399	China and USA, India has been the third largest producer and consumer of fertilizers due to
400	intensification of agriculture, resulting in high anthropogenic N emissions (Aneja et al., 2012). For
401	instance, ammonia (NH ₃) emissions from livestock and fertilizer applications in India in 2003 was
402	estimated as 1705 Gg yr ⁻¹ and 1697 Gg yr ⁻¹ , respectively (Aneja et al., 2012). Moreover, in India, field
403	burning of crop residue (FBCR) is another critical anthropogenic activity leading to N emissions. In
404	2010, 6300 Gg of dry biomass are estimated to be subjected to FBCR in India, resulting in 350 Gg N
405	emissions (Sahai et al., 2011). Besides Indian monsoon, biomass-burning emissions in South Asia
406	could be across the Himalayas and transported to the TP by the mountain/valley wind (Cong et al.,
407	<u>2015).</u>

408 **4.3 Comparison of inorganic N wet deposition on the TP** in the TP with previous 409 estimations

Long-term dataset series of N deposition have been established based on observations (Lu and Tian, 2007, 2014, 2015) or model simulations (Dentener et al., 2006). These datasets have been used to estimate global or regional N deposition (Dentener et al., 2006; Lu and Tian, 2007) and drive ecosystem models to examine the ecological effects of elevated N deposition (Lu and Tian, 2013). Thus, reliable N deposition datasets are <u>a</u>-prerequisite<u>s</u> for N deposition estimation or driving ecosystem models. Here, the estimation of N wet deposition <u>on the TPin the TP</u> based on our field observations is compared with previous estimations via limited observations or simulations.

Lu and Tian (2007) estimated the inorganic N wet deposition as ranging from 4.16 kg N ha⁻¹ yr⁻¹ in 417 418 Tibet Autonomous Region (on thein the western TP) to 4.76 kg N ha⁻¹ yr⁻¹ in Qinghai Province (on 419 thein the eastern TP). Recently, Jia et al. (2014) estimated the inorganic N wet deposition during the 2000s as ranging from 6.11 kg N ha⁻¹ yr⁻¹ in Tibet Autonomous Region to 7.87 kg N ha⁻¹ yr⁻¹ in 420 421 Qinghai Province. Those estimations were even much higher even than the highest record of inorganic 422 N wet deposition observations on the TP in the TP (3.08 kg N ha⁻¹ yr⁻¹ at Biru during 2006-2007) (Table 423 3). In this study, combing *in situ* measurements at 5 sites in this study and 7 sites in previous studies 424 (Table 2), the average wet deposition of atmospheric NH4⁺-N, NO3⁺-N, and inorganic N on the TP in 425 <u>the TP</u> were estimated to be 1.09, 0.57, and 1.66 kg N ha⁻¹ yr⁻¹, respectively. According to our study, 426 both Lu and Tian (2007) and Jia et al. (2014) highly overestimated inorganic N wet deposition on the 427 TP in the TP, probably likely due to following two reasons. Firstly, compared to our study, the 428 observations used in previous regional-scale estimations were used much fewerlimited, with fewer in 429 *situ* measurement observation sites than used in the present study. For example, there were only four 430 sites in Tibet Autonomous Region and one site in Qinghai Province used in the estimation of Jia et al. 431 (2014). Such limited field observations probably lead to large uncertainty in the conclusions drawn 432 regarding inorganic N wet deposition on their the entire TP. Secondly, the Kriging interpolation 433 technique was used in both Lu and Tian (2007) and Jia et al. (2014) to estimate the spatial pattern of 434 inorganic N wet deposition in China. However, observation sites are sparsely distributed on the TPin 435 the TP, and the estimation of inorganic N wet deposition is largely influenced by N deposition 436 observations in the surrounding regions of much lower altitude. The average altitude of the TP is above 437 4000 m, where both the climate and anthropogenic activities are substantially different with those in lower altitude areas. For example, the average inorganic N wet deposition was 1.66 kg N ha⁻¹ yr⁻¹, 438 which was much lower than that in northern and southern China (Table 3). The interpolations at the 439

440 national scale in Lu and Tian (2007) and Jia et al. (2014) probably-likely overestimated the regional 441 inorganic N wet deposition on the TP in the TP. In addition, we also estimated the inorganic N wet 442 deposition for the entire TP using Kriging interpolation, but only based on the site-scale in situ 443 measurements on the TP in the TP (12 sites, including: 5 sites in this study and 7 sites in previous field 444 observations), rather than the observations in the surrounding regions of much lower altitude. The 445 inorganic N wet deposition for the entire TP estimation based on the Kriging interpolation in our study 446 is 1.76 kg N ha⁻⁻¹ yr⁻⁻¹ (Fig. S1 and spatial data as a NetCDF file in the supplementary material), which 447 is much lower than that in previous interpolation studies (Lu and Tian, 2007; Jia et al., 2014), but is 448 comparable with the averaged inorganic N wet deposition among the 12 sites (1.66 kg N ha⁻¹ yr⁻¹) 449 (Table 2).

450 Atmospheric chemistry transport models are commonly used to calculate current and future N 451 deposition. Dentener et al. (2006) used 23 atmospheric chemistry transport models to assess both global and regional N deposition. Compared to observation records, Dentener et al. (2006) 452 underestimated inorganic N wet deposition over the whole of China (Lu and Tian, 2007), but 453 overestimated it over for the TP. According to Dentener et al. (2006), the NH₄⁺, NO₃⁻, and inorganic 454 455 N wet deposition on the TP in the TP are 1.97, 0.99, and 2.96 kg N ha⁻¹ yr⁻¹, respectively—nearly 456 double that of N deposition estimated in our study. Based on site-scale in situ measurements, we 457 provide a more accurate regional-scale estimation of inorganic N wet deposition on the TP in the TP, 458 which can be used as background information in research studies focusing on the responses of alpine ecosystems to elevated N deposition. Besides assessment of N deposition, N deposition simulated by 459 460 atmospheric chemistry transport models is usually used to drive large-scale ecosystem models for integrated ecosystem assessment (Xu-Ri et al., 2012; Zaehle, 2013). The ecological effects of N 461

addition are probably not only influenced by the quantity of N deposition, but also by the proportions 462 463 of each component, e.g. the NH₄⁺-N:NO₃⁻-N ratio. For example, in African savannas, plants 464 demonstrate N uptake preference, which is likely influenced by the NH4⁺-N:NO3⁻-N ratio in their native habitats (Wang and Macko, 2011). However, in most current N fertilization experiments, the N 465 466 forms of fertilizer are NH₄NO₃, NH₄⁺-N or NO₃⁻-N (Liu and Greaver, 2009), with the NH₄⁺-N:NO₃⁻-467 N ratio of N wet deposition at experimental sites not considered. Our work shows that the estimated 468 NH4⁺-N:NO₃⁻-N ratio of inorganic N wet deposition on the TP is approximately 2:1, which 469 is consistent with the modelled estimation of Dentener et al. (2006), but lower than the NH4⁺-N:NO3⁻-470 N ratio of 2.5 in forest ecosystems in eastern China (Du et al., 2014). This NH₄⁺-N:NO₃⁻-N ratio (2:1) 471 is recommended to be considered when N fertilization experiments are conducted in alpine ecosystems 472 on the TP in the TP.

473 **4.4 Uncertainty and recommendations**

474 Combining our *in situ* measurements at five remote sites and previous site-scale field observations, the 475 inorganic N wet deposition on the TP in the TP was quantitatively assessed in this study. The 476 assessment is conducive to accurately estimating N wet deposition for the entire nation of China, and 477 provides background information of N wet deposition for the studies focusing on the alpine ecological 478 effects of elevated N deposition. Despite this, there are uncertainties in the estimation of N deposition 479 on the TP in the TP due to following reasons. Firstly, total N deposition comprises wet deposition (in 480 the form of precipitation) and dry deposition (in the form of gases and particles). Considering the 481 whole of China, dry deposition contributes 30% to total inorganic N deposition (Lu and Tian, 2007, 482 2014, 2015). In northern China, this ratio is much higher, at 60% (Pan et al., 2012). However, in this 483 study, we only estimated the inorganic N wet deposition on the TP in the TP, with the situation 484 regarding dry deposition remaining unclear. Thus, investigation of N dry deposition is critical for 485 assessing total N deposition on the TP in the TP. Secondly, the TP covers an area of about 2.57 million 486 km², occupying approximately 1/4 of the land area of China (Zhang et al., 2002). Precipitation on the 487 **TP**in the TP is influenced by both the Indian monsoon and westerlies, leading to spatial variation in 488 the origins of N wet deposition. Therefore, it is necessary to establish N wet deposition observation 489 sites in different climatic zones. Thirdly, besides spatial heterogeneity, N deposition on the TP in the 490 TP also possesses temporal heterogeneity. Inorganic N wet deposition on the TP has increased 491 during recent decades, as recorded in ice cores (Hou et al., 2003;Kang et al., 2002a, b;Thompson et al., 492 2000;Zhao et al., 2011;Zheng et al., 2010) and sediment cores of alpine lakes (Choudhary et al., 493 2013; Hu et al., 2014). The long-term trend and inter-annual variability of inorganic N wet deposition 494 on the TP in the TP can't be quantitatively characterized by the short-term *in situ* measurements in this 495 study. Overall, critical questions remain open regarding the quantitative understanding of N deposition 496 on the TP in the TP. Therefore, t To deepen our understanding N deposition on the TP in the TP, it is 497 essential to perform long-term in situ measurements of N wet and dry deposition in various climate 498 zones in the future.

499 **5 Conclusion**

Alpine ecosystems on the TPin the TP are sensitive to elevated N deposition, and the inorganic N deposition has been increasing since the mid-20th century. However, the amount of inorganic N wet deposition on in the Tibetan TP remains unclear, due to a paucity of *in situ* measurement. In this study, using stations in the TORP network, we conducted *in situ* measurements of major ions wet deposition at five remote sites, situated mainly on thein the central and western TP. Among the five sites, both NH_4^+ -N and NO_3^- -N were mainly contributed by anthropogenic sources. Combining site-scale *in situ*

506 measurements in our and previous studies, the average wet deposition of atmospheric NH₄⁺-N, NO₃⁻-=N, and inorganic N on the TP in the TP are estimated to be 1.09, 0.57, and 1.66 kg N ha⁻¹ yr⁻¹, 507 508 respectively. Considering the entire TP, according to our results, previous regional-scale assessment have highly overestimated inorganic N wet deposition, either through simulations with atmospheric 509 510 chemistry transport models (Dentener et al., 2006) or interpolations based on limited field observations 511 for the whole of China (Jia et al., 2014; Lu and Tian, 2007). The NH₄⁺-N:NO₃⁻-N ratio in precipitation 512 on the TP in the TP was found to be approximately 2:1, which is consistent with model simulations (Dentener et al., 2006). To clarify the total N deposition on the TP in the TP more clearly, we 513 514 recommend conducting long-term monitoring of both wet and dry deposition of N in various climate 515 zones in the future work.

516 Acknowledgements

517 We are grateful to the staff at the Southeast Tibet Observation and Research Station for the Alpine 518 Environment, Chinese Academy of Sciences (Southeast Tibet Station), Nam Co Monitoring and 519 Research Station for Multisphere Interactions, Chinese Academy of Sciences (Nam Co Station), 520 Qomolangma Atmospheric and Environmental Observation and Research Station, Chinese Academy 521 of Sciences (Qomolangma Station), Ngari Desert Observation and Research Station (Ngari Station), 522 and Muztagh Ata Westerly Observation and Research Station (Muztagh Ata Station) for their 523 assistance in collecting the samples and providing the precipitation data. The authors also acknowledge 524 thank-Da Wei, Dongxue Dai, Xiaodong Geng, Tenzin Tarchen, and Shan Lu for their contributions to 525 the field work. The authors acknowledge the constructive comments of the two anonymous referees, 526 whose helpful feedback resulted in a greatly improved manuscript. This work was supported by the Strategic Priority Research Program-Climate Change: Carbon Budget and Related Issues, of the 527

528 Chinese Academy of Sciences (XDA05050404-3-2, XDA05020402), and the National Natural 529 Science Foundation of China (40605032, 40975096, 41175128).

530 References

- 531 Aneja, V. P., Schlesinger, W. H., Erisman, J. W., Behera, S. N., Sharma, M., and Battye, W.: Reactive
- nitrogen emissions from crop and livestock farming in India, Atmos. Environ., 47, 92-103, doi:
 10.1016/j.atmosenv.2011.11.026, 2012.
- Balasubramanian, R., Victor, T., and Chun, N.: Chemical and statistical analysis of precipitation in
 Singapore, Water. Air Soil Poll., 130, 451-456, doi: 10.1023/A:1013801805621, 2001.
- 536 Balestrini, R., Polesello, S., and Sacchi, E.: Chemistry and isotopic composition of precipitation and
- 537 surface waters in Khumbu valley (Nepal Himalaya): N dynamics of high elevation basins, Sci.
 538 Total. Environ., 485, 681-692, doi: 10.1016/j.scitotenv.2014.03.096, 2014.
- 539 Basto, S., Thompson, K., Phoenix, G., Sloan, V., Leake, J., and Rees, M.: Long-term nitrogen
- 540 deposition depletes grassland seed banks, Nat. Commun., 6, 6185, doi: 10.1038/ncomms7185,
 541 2015.
- 542 Canfield, D. E., Glazer, A. N., and Falkowski, P. G.: The evolution and future of Earth's nitrogen cycle,
 543 Science, 330, 192-196, doi: 10.1126/science.1186120, 2010.
- Cao, Y. Z., Wang, S. Y., Zhang, G., Luo, J. Y., and Lu, S. Y.: Chemical characteristics of wet
 precipitation at an urban site of Guangzhou, South China, Atmos. Res., 94, 462-469, doi:
 10.1016/j.atmosres.2009.07.004, 2009.
- 547 Cong, Z. Y., Kawamura, K., Kang, S. C., and Fu, P. Q.: Penetration of biomass-burning emissions from
- 548 South Asia through the Himalayas: new insights from atmospheric organic acids, Sci. Rep.-Uk,
- 549 5, 9580, doi: 10.1038/srep09580, 2015.

550	Chabas, A., and Lefevre, R. A.: Chemistry and microscopy of atmospheric particulates at Delos
551	(Cyclades-Greece), Atmos. Environ., 34, 225-238, doi: 10.1016/S1352-2310(99)00255-1, 2000.
552	Chen, X. Y., and Mulder, J.: Atmospheric deposition of nitrogen at five subtropical forested sites in
553	South China, Sci. Total. Environ., 378, 317-330, doi:10.1016/j.scitotenv.2007.02.028, 2007.
554	Choudhary, P., Routh, J., and Chakrapani, G. J.: A 100-year record of changes in organic matter
555	characteristics and productivity in Lake Bhimtal in the Kumaon Himalaya, NW India, J.
556	Paleolimnol., 49, 129-143, doi: 10.1007/s10933-012-9647-9, 2013.
557	Dentener, F., Drevet, J., Lamarque, J. F., Bey, I., Eickhout, B., Fiore, A. M., Hauglustaine, D., Horowitz,
558	L. W., Krol, M., Kulshrestha, U. C., Lawrence, M., Galy-Lacaux, C., Rast, S., Shindell, D.,
559	Stevenson, D., Van Noije, T., Atherton, C., Bell, N., Bergman, D., Butler, T., Cofala, J., Collins,
560	B., Doherty, R., Ellingsen, K., Galloway, J., Gauss, M., Montanaro, V., Muller, J. F., Pitari, G.,
561	Rodriguez, J., Sanderson, M., Solmon, F., Strahan, S., Schultz, M., Sudo, K., Szopa, S., and Wild,
562	O.: Nitrogen and sulfur deposition on regional and global scales: A multimodel evaluation, Global
563	Biogeochem. Cycles, 20, GB4003, doi: 10.1029/2005GB002672, 2006.
564	Ding, G. A., Xu, X. B., and Wang, S. F.: Database from the acid rain network of China meteorological
565	administration and it's preliminary analyses, Journal of Applied Meteorological Science, 15, 85-
566	94, 2004 (in Chinese with English abstract).
567	Ding, M., Yao, F., Chen, J., Wang, X., and Yang, S.: Chemical characteristics of acidic precipitation in
568	Tiantong, Zhejiang Province, Acta Scientiae Circumstantiae, 32, 2245-2252, 2012 (in Chinese
569	with English abstract).
570	Du, E., Jiang, Y., Fang, J. Y., and de Vries, W.: Inorganic nitrogen deposition in China's forests: Status

571 and characteristics, Atmos. Environ., 98, 474-482, doi: 10.1016/j.atmosenv.2014.09.005, 2014.

572 Erisman, J. W., Galloway, J., Seitzinger, S., Bleeker, A., and Butterbach-Bahl, K.: Reactive nitrogen

- in the environment and its effect on climate change, Curr. Opin. Env. Sust., 3, 281-290, doi:
 10.1016/j.cosust.2011.08.012, 2011.
- 575 Erisman, J. W., Galloway, J. N., Seitzinger, S., Bleeker, A., Dise, N. B., Petrescu, A. M. R., Leach, A.
- 576 M., and de Vries, W.: Consequences of human modification of the global nitrogen cycle, Philos.
- 577 T. R. Soc. B, 368, 20130116, doi: 10.1098/rstb.2013.0116, 2013.
- 578 Fagerli, H., and Aas, W.: Trends of nitrogen in air and precipitation: Model results and observations at
- 579 EMEP sites in Europe, 1980-2003, Environ. Pollut., 154, 448-461, doi:
 580 10.1016/j.envpol.2008.01.024, 2008.
- Fang, X. M., Han, Y. X., Ma, J. H., Song, L. C., Yang, S. L., and Zhang, X. Y.: Dust storms and loess
 accumulation on the Tibetan Plateau: A case study of dust event on 4 March 2003 in Lhasa,
 Chinese Sci. Bull., 49, 953-960, doi: 10.1360/03wd0180, 2004.
- 584 Galloway, J. N., Townsend, A. R., Erisman, J. W., Bekunda, M., Cai, Z. C., Freney, J. R., Martinelli,
- 585 L. A., Seitzinger, S. P., and Sutton, M. A.: Transformation of the nitrogen cycle: Recent trends,
- questions, and potential solutions, Science, 320, 889-892, doi: 10.1126/science.1136674, 2008.
- 587 Gao, J. G., Zhang, Y. L., Liu, L. S., and Wang, Z. F.: Climate change as the major driver of alpine

grasslands expansion and contraction: A case study in the Mt. Qomolangma (Everest) National

- 589 Nature Preserve, southern Tibetan Plateau, Quatern. Int., 336, 108-116, doi:
 590 10.1016/j.quaint.2013.09.035, 2014.
- 591 Grigholm, B., Mayewski, P. A., Kang, S., Zhang, Y., Morgenstern, U., Schwikowski, M., Kaspari, S.,
- 592 Aizen, V., Aizen, E., Takeuchi, N., Maasch, K. A., Birkel, S., Handley, M., and Sneed, S.:
- 593 Twentieth century dust lows and the weakening of the westerly winds over the Tibetan Plateau,
- 594 Geophys. Res. Lett., 42, 2434-2441, doi: 10.1002/2015gl063217, 2015

588

595 Gruber, N., and Galloway, J. N.: An Earth-system perspective of the global nitrogen cycle, Nature, 451,

- 596 293-296, doi: 10.1038/nature06592, 2008.
- Han, Y. X., Fang, X. M., Kang, S. C., Wang, H. J., and Kang, F. Q.: Shifts of dust source regions over
- 598 central Asia and the Tibetan Plateau: Connections with the Arctic oscillation and the westerly jet,
- 599 Atmos. Environ., 42, 2358-2368, doi: 10.1016/j.atmosenv.2007.12.025, 2008.
- Han, Y. X., Fang, M., Zhao, T. L., Bai, H. Z., Kang, S. C., and Song, L. C.: Suppression of precipitation
- by dust particles originated in the Tibetan Plateau, Atmos. Environ., 43, 568-574, doi:
 10.1016/j.atmosenv.2008.10.018, 2009.
- Hou, S.: Chemical Characteristics of Precipitation at the Headwaters of the Ürümqi River in the
 Tianshan Mountains, Journal of Glaciology and Geocryology, 23, 80-84, 2001 (in Chinese with
 English abstract).
- Hou, S. G., Qin, D. H., Zhang, D. Q., Kang, S. C., Mayewski, P. A., and Wake, C. P.: A 154a highresolution ammonium record from the Rongbuk Glacier, north slope of Mt. Qomolangma
 (Everest), Tibet-Himal region, Atmos. Environ., 37, 721-729, doi:10.1016/S13522310(02)00582-4, 2003.
- Hu, Z. J., Anderson, N. J., Yang, X. D., and McGowan, S.: Catchment-mediated atmospheric nitrogen
- deposition drives ecological change in two alpine lakes in SE Tibet, Glob. Change Biol., 20, 16141628, doi: 10.1111/Gcb.12435, 2014.
- Huang, J. P., Minnis, P., Yi, Y. H., Tang, Q., Wang, X., Hu, Y. X., Liu, Z. Y., Ayers, K., Trepte, C., and
- 614 Winker, D.: Summer dust aerosols detected from CALIPSO over the Tibetan Plateau, Geophys.
- 615 Res. Lett., 34, L18805, doi: 10.1029/2007gl029938, 2007.
- Huang, J. P., Wang, T. H., Wang, W. C., Li, Z. Q., and Yan, H. R.: Climate effects of dust aerosols over
- 617 East Asian arid and semiarid regions, J. Geophys. Res.-Atmos., 119, 11398-11416, doi:
- 618 10.1002/2014jd021796, 2014.

619	Huang, K., Zhuang, G. S., Xu, C., Wang, Y., and Tang, A. H.: The chemistry of the severe acidic
620	precipitation in Shanghai, China, Atmos. Res., 89, 149-160, doi: 10.1016/j.atmosres.2008.01.006,
621	2008.

- Huang, Y. L., Wang, Y. L., and Zhang, L. P.: Long-term trend of chemical composition of wet
- atmospheric precipitation during 1986-2006 at Shenzhen City, China, Atmos. Environ., 42, 3740-
- 624 3750, doi: 10.1016/j.atmosenv.2007.12.063, 2008.
- Jia, J.: Study of atmospheric wet deposition of nitrogen in Tibetan Plateau, Master, Tibet University,
 2008 (in Chinese with English abstract).
- Jia, Y., Yu, G., He, N., Zhan, X., Fang, H., Sheng, W., Zuo, Y., Zhang, D., and Wang, Q.: Spatial and
- decadal variations in inorganic nitrogen wet deposition in China induced by human activity, Sci.
 Rep.-UK, 4, 3763, doi: 10.1038/srep03763, 2014.
- 630 Kang, S., Mayewski, P. A., Qin, D., Yan, Y., Hou, S., Zhang, D., Ren, J., and Kruetz, K.:
- 631 Glaciochemical records from a Mt. Everest ice core: relationship to atmospheric circulation over
- 632 Asia, Atmos. Environ., 36, 3351-3361, doi:10.1016/S1352-2310(02)00325-4, 2002a.
- 633 Kang, S. C., Mayewski, P. A., Qin, D. H., Yan, Y. P., Zhang, D. Q., Hou, S. G., and Ren, J. W.:
- 634 Twentieth century increase of atmospheric ammonia recorded in Mount Everest ice core, J.
 635 Geophys. Res.-Atmos., 107, 4595, doi: 10.1029/2001jd001413, 2002b.
- Kang, S. C., Zhang, Y. L., Zhang, Y. J., Grigholm, B., Kaspari, S., Qin, D. H., Ren, J. W., and Mayewski,
- P.: Variability of atmospheric dust loading over the central Tibetan Plateau based on ice core
 glaciochemistry, Atmos. Environ., 44, 2980-2989, doi: 10.1016/j.atmosenv.2010.05.014, 2010.
- 639 Kaspari, S., Mayewski, P., Kang, S., Sneed, S., Hou, S., Hooke, R., Kreutz, K., Introne, D., Handley,
- 640 M., Maasch, K., Qin, D., and Ren, J.: Reduction in northward incursions of the South Asian
- monsoon since approximate to 1400 AD inferred from a Mt. Everest ice core, Geophys. Res. Lett.,

- 642 34, L16701, doi: 10.1029/2007gl030440, 2007.
- Keene, W. C., Pszenny, A. A. P., Galloway, J. N., and Hawley, M. E.: Sea-salt corrections and
 interpretation of constituent ratios in marine precipitation, J. Geophys. Res.-Atmos., 91, 66476658, doi: 10.1029/Jd091id06p06647, 1986.
- 646 Kulshrestha, U. C., Sarkar, A. K., Srivastava, S. S., and Parashar, D. C.: Investigation into atmospheric
- deposition through precipitation studies at New Delhi (India), Atmos. Environ., 30, 4149-4154,
 doi: 10.1016/1352-2310(96)00034-9, 1996.
- 649 Kulshrestha, U. C., Kulshrestha, M. J., Sekar, R., Sastry, G. S. R., and Vairamani, M.: Chemical
- 650 characteristics of rainwater at an urban site of south-central India, Atmos. Environ., 37, 3019-
- 651 3026, doi: 10.1016/S1352-2310(03)00266-8, 2003.
- Lan, Z., Jenerette, G. D., Zhan, S., Li, W., Zheng, S., and Bai, Y.: Testing the scaling effects and
 mechanisms of N-induced biodiversity loss: Evidence from a decade-long grassland experiment,
- 654 J. Ecol., doi: 10.1111/1365-2745.12395, 2015.
- Lehmann, C. M. B., Bowersox, V. C., and Larson, S. M.: Spatial and temporal trends of precipitation
 chemistry in the United States, 1985-2002, Environ. Pollut., 135, 347-361, doi:
 10.1016/j.envpol.2004.11.016, 2005.
- Li, C., Kang, S. C., Zhang, Q. G., and Kaspari, S.: Major ionic composition of precipitation in the Nam
- 659 Co region, Central Tibetan Plateau, Atmos. Res., 85, 351-360,
 660 doi:10.1016/j.atmosres.2007.02.006, 2007.
- Li, M., Ma, Y., Ishikawa, H., Ma, W., Sun, F., Wang, Y., and Zhu, Z.: Characteristics of
 micrometeorological elements near surface and soil on the northern slope of Mt. Qomolangma
 area, Plateau Meteorology, 26, 1263-1268, 2007 (in Chinese with English abstract).
- 664 Li, Z. J., Li, Z. X., Wang, T. T., Gao, Y., Cheng, A. F., Guo, X. Y., Guo, R., Jia, B., Song, Y. X., Han,

665	C. T., and Theakstone, W.: Composition of wet deposition in the central Qilian Mountains, China,
666	Environ. Earth Sci., 73, 7315-7328, doi: 10.1007/s12665-014-3907-0, 2015.
667	Liu, B., Kang, S. C., Sun, J. M., Zhang, Y. L., Xu, R., Wang, Y. J., Liu, Y. W., and Cong, Z. Y.: Wet
668	precipitation chemistry at a high-altitude site (3,326 m a.s.l.) in the southeastern Tibetan Plateau,
669	Environ. Sci. Pollut. R., 20, 5013-5027, doi: 10.1007/s11356-012-1379-x, 2013.
670	Liu, L. L., and Greaver, T. L.: A review of nitrogen enrichment effects on three biogenic GHGs: the
671	CO ₂ sink may be largely offset by stimulated N ₂ O and CH ₄ emission, Ecol. Lett., 12, 1103-1117,
672	doi: 10.1111/j.1461-0248.2009.01351.x, 2009.
673	Liu, X. J., Duan, L., Mo, J. M., Du, E. Z., Shen, J. L., Lu, X. K., Zhang, Y., Zhou, X. B., He, C. N.,
674	and Zhang, F. S.: Nitrogen deposition and its ecological impact in China: An overview, Environ.
675	Pollut., 159, 2251-2264, doi: 10.1016/j.envpol.2010.08.002, 2011.
676	Liu, X. J., Zhang, Y., Han, W. X., Tang, A. H., Shen, J. L., Cui, Z. L., Vitousek, P., Erisman, J. W.,
677	Goulding, K., Christie, P., Fangmeier, A., and Zhang, F. S.: Enhanced nitrogen deposition over
678	China, Nature, 494, 459-462, doi: 10.1038/nature11917, 2013.
679	Liu, Y. H., Dong, G. R., Li, S., and Dong, Y. X.: Status, causes and combating suggestions of sandy
680	desertification in Qinghai-Tibet Plateau, Chinese Geogr. Sci., 15, 289-296, doi: 10.1007/s11769-
681	005-0015-9, 2005.
682	Liu, Y. W., Xu-Ri, Xu, X. L., Wei, D., Wang, Y. H., and Wang, Y. S.: Plant and soil responses of an
683	alpine steppe on the Tibetan Plateau to multi-level nitrogen addition, Plant Soil, 373, 515-529,
684	doi: 10.1007/s11104-013-1814-x, 2013.

- Lu, C. Q., and Tian, H. Q.: Spatial and temporal patterns of nitrogen deposition in China: Synthesis of
 observational data, J. Geophys. Res.-Atmos., 112, D22S05, doi:10.1029/2006JD007990, 2007.
- 687 Lu, C. Q., and Tian, H. Q.: Net greenhouse gas balance in response to nitrogen enrichment:

perspectives from a coupled biogeochemical model, Glob. Change Biol., 19, 571-588, doi:
10.1111/gcb.12049, 2013.

690 Lu, C. Q., and Tian, H. Q.: Half-century nitrogen deposition increase across China: A gridded time-

691 series data set for regional environmental assessments, Atmos. Environ., 97, 68-74, doi:

692 10.1016/j.atmosenv.2014.07.061, 2014.

- 693 Lu, C. Q., and Tian, H. Q.: Reply to "Comments on 'Half-century nitrogen deposition increase across
- China: A gridded time-series dataset for regional environmental assessments'", Atmos. Environ.,
 101, 352-353, doi:10.1016/j.atmosenv.2014.11.032, 2015.
- Lu, X. W., Li, L. Y., Li, N., Yang, G., Luo, D. C., and Chen, J. H.: Chemical characteristics of spring
- 697 rainwater of Xi'an city, NW China, Atmos. Environ., 45, 5058-5063, doi:
 698 10.1016/j.atmosenv.2011.06.026, 2011.
- Ma, Y. M., Kang, S. C., Zhu, L. P., Xu, B. Q., Tian, L. D., and Yao, T. D.: Tibetan Observation and
- Research Platform–atmosphere–land interaction over a heterogeneous landscape, B. Am.
 Meteorol. Soc., 89, 1487-1492, doi: 10.1175/2008bams2545.1, 2008.
- 702 Ma, Z., Ma, M. J., Baskin, J. M., Baskin, C. C., Li, J. Y., and Du, G. Z.: Responses of alpine meadow
- seed bank and vegetation to nine consecutive years of soil fertilization, Ecol. Eng., 70, 92-101,
 doi: 10.1016/j.ecoleng.2014.04.009, 2014.
- Mao, R., Gong, D. Y., Shao, Y. P., Wu, G. J., and Bao, J. D.: Numerical analysis for contribution of the
- Tibetan Plateau to dust aerosols in the atmosphere over the East Asia, Sci. China Earth Sci., 56,
 301-310, doi: 10.1007/s11430-012-4460-x, 2013.
- 708 Migliavacca, D., Teixeira, E. C., Wiegand, F., Machado, A. C. M., and Sanchez, J.: Atmospheric
- precipitation and chemical composition of an urban site, Guaiba hydrographic basin, Brazil,
- 710 Atmos. Environ., 39, 1829-1844, doi: 10.1016/j.atmosenv.2004.12.005, 2005.

- 711 Morino, Y., Ohara, T., Kurokawa, J., Kuribayashi, M., Uno, I., and Hara, H.: Temporal variations of
- 712 nitrogen wet deposition across Japan from 1989 to 2008, J. Geophys. Res.-Atmos., 116, D06307,
 713 doi: 10.1029/2010jd015205, 2011.
- 714 Okay, C., Akkoyunlu, B. O., and Tayanc, M.: Composition of wet deposition in Kaynarca, Turkey,
- 715 Environ. Pollut., 118, 401-410, doi: 10.1016/S0269-7491(01)00292-5, 2002.
- Pan, Y. P., Wang, Y. S., Tang, G. Q., and Wu, D.: Wet and dry deposition of atmospheric nitrogen at
 ten sites in Northern China, Atmos. Chem. Phys., 12, 6515-6535, doi: 10.5194/acp-12-6515-2012,
 2012.
- 719 Pinder, R. W., Davidson, E. A., Goodale, C. L., Greaver, T. L., Herrick, J. D., and Liu, L. L.: Climate
- change impacts of US reactive nitrogen, P. Natl. Acad. Sci. USA, 109, 7671-7675, doi:
 10.1073/pnas.1114243109, 2012.
- R Core Team. R: A language and environment for statistical computing. R Foundation for Statistical
 Computing, Vienna, Austria, 2015.
- Puxbaum, H., Simeonov, V., Kalina, M., Tsakovski, S., Loffler, H., Heimburger, G., Biebl, P., Weber,
- A., and Damm, A.: Long-term assessment of the wet precipitation chemistry in Austria (1984-
- 726 1999), Chemosphere, 48, 733-747, doi: 10.1016/S0045-6535(02)00125-X, 2002.
- 727 Qiu, J.: The third pole, Nature, 454, 393-396, doi: 10.1038/454393a, 2008.
- Rodhe, H., and Granat, L.: An evaluation of sulfate in European precipitation 1955-1982, Atmos.
- 729 Environ., 18, 2627-2639, doi: 10.1016/0004-6981(84)90327-5, 1984.
- 730 Safai, P. D., Rao, P. S. P., Mornin, G. A., All, K., Chate, D. M., and Praveen, P. S.: Chemical
- composition of precipitation during 1984-2002 at Pune, India, Atmos. Environ., 38, 1705-1714,
- 732 doi: 10.1016/j.atmosenv.2003.12.016, 2004.
- 733 Sahai, S., Sharma, C., Singh, S. K., and Gupta, P. K.: Assessment of trace gases, carbon and nitrogen

emissions from field burning of agricultural residues in India, Nutr. Cycl. Agroecosys., 89, 143-

735 157, doi: 10.1007/s10705-010-9384-2, 2011.

- 736 Shen, W. J., Ren, H. L., Jenerette, G. D., Hui, D. F., and Ren, H.: Atmospheric deposition and canopy
- exchange of anions and cations in two plantation forests under acid rain influence, Atmos.
- 738 Environ., 64, 242-250, doi: 10.1016/j.atmosenv.2012.10.015, 2013.
- 739 Sheng, W. P., Yu, G. R., Jiang, C. M., Yan, J. H., Liu, Y. F., Wang, S. L., Wang, B., Zhang, J. H., Wang,
- 740 C. K., Zhou, M., and Jia, B. R.: Monitoring nitrogen deposition in typical forest ecosystems along
- 741 a large transect in China, Environ. Monit. Assess., 185, 833-844, doi: 10.1007/s10661-012-2594-
- 742 0, 2013.
- Shi, Y. L., Cui, S. H., Ju, X. T., Cai, Z. C., and Zhu, Y. G.: Impacts of reactive nitrogen on climate
 change in China, Sci. Rep.-UK, 5, 8118, doi: 10.1038/Srep08118, 2015.
- 745 Tang, J., Xue, H., Yu, X., Cheng, H., Xu, X., Zhang, X., and Ji, J.: The preliminary study on chemical
- characteristics of precipitation at Mt. Waliguan, Acta Scientiae Circumstantiae, 20, 420-425, 2000
 (in Chinese with English abstract).
- Taylor, S. R.: Abundance of chemical elements in the continental crust a new table, Geochim.
 Cosmochim. Ac., 28, 1273-1285, doi: 10.1016/0016-7037(64)90129-2, 1964.
- 750 Thompson, L. G., Yao, T., Mosley-Thompson, E., Davis, M. E., Henderson, K. A., and Lin, P. N.: A
- high-resolution millennial record of the South Asian Monsoon from Himalayan ice cores, Science,
- 752 289, 1916-1919, doi: 10.1126/science.289.5486.1916, 2000.
- 753 Tripathee, L., Kang, S. C., Huang, J., Sillanpaa, M., Sharma, C. M., Luthi, Z. L., Guo, J. M., and
- 754 Paudyal, R.: Ionic composition of wet precipitation over the southern slope of central Himalayas,
- 755 Nepal, Environ. Sci. Pollut. R., 21, 2677-2687, doi: 10.1007/s11356-013-2197-5, 2014.
- Tu, J., Wang, H. S., Zhang, Z. F., Jin, X., and Li, W. Q.: Trends in chemical composition of precipitation

- 757 in Nanjing, China, during 1992-2003, Atmos. Res., 73, 283-298, doi:
 758 10.1016/j.atmosres.2004.11.002, 2005.
- 759 Turekian, K. K.: Oceans, Prentice-Hall, New Jersey, United States, 1968.
- 760 Wang, L. X., and Macko, S. A.: Constrained preferences in nitrogen uptake across plant species and
- 761 environments, Plant Cell Environ., 34, 525-534, doi: 10.1111/j.1365-3040.2010.02260.x, 2011.
- Wang, P. L., Yao, T. D., Tian, L. D., Wu, G. J., Li, Z., and Yang, W.: Recent high-resolution
 glaciochemical record from a Dasuopu firn core of middle Himalayas, Chinese Sci. Bull., 53,
- 764 418-425, doi: 10.1007/s11434-008-0098-7, 2008.
- Wang, Y., Ma, Y., Zhu, Z., and Li, M.: Variation characteristics of meteorological elements in near
 surface layer over the Lulang valley of southeastern Tibetan Plateau, Plateau Meteorology, 29,
 63-69, 2010 (in Chinese with English abstract).
- Wang, Y. Q., Zhang, X. Y., and Draxler, R. R.: TrajStat: GIS-based software that uses various trajectory
- statistical analysis methods to identify potential sources from long-term air pollution
 measurement data, Environ. Model. Soft., 24, 938-939, doi: 10.1016/j.envsoft.2009.01.004, 2009.
- Xia, X. G., Wang, P. C., Wang, Y. S., Li, Z. Q., Xin, J. Y., Liu, J., and Chen, H. B.: Aerosol optical
- depth over the Tibetan Plateau and its relation to aerosols over the Taklimakan Desert, Geophys
 Res Lett, 35, L16804, doi: 10.1029/2008gl034981, 2008.
- Xiao, H. W., Xiao, H. Y., Long, A. M., Wang, Y. L., and Liu, C. Q.: Chemical composition and source
 apportionment of rainwater at Guiyang, SW China, J. Atmos. Chem., 70, 269-281, doi:
 10.1007/s10874-013-9268-3, 2013.
- 777 Xu, H., Bi, X. H., Feng, Y. C., Lin, F. M., Jiao, L., Hong, S. M., Liu, W. G., and Zhang, X. Y.: Chemical
- composition of precipitation and its sources in Hangzhou, China, Environ. Monit. Assess., 183,
- 581-592, doi: 10.1007/s10661-011-1963-4, 2011.

- Xu-Ri, Prentice, I. C., Spahni, R., and Niu, H. S.: Modelling terrestrial nitrous oxide emissions and
 implications for climate feedback, New. Phytol., 196, 472-488, doi: 10.1111/j.14698137.2012.04269.x, 2012.
- Xu, X. L., Wanek, W., Zhou, C. P., Richter, A., Song, M. H., Cao, G. M., Ouyang, H., and Kuzyakov,
- Y.: Nutrient limitation of alpine plants: Implications from leaf N : P stoichiometry and leaf δ¹⁵N,
 J. Plant Nutr. Soil Sc., 177, 378-387, doi: 10.1002/jpln.201200061, 2014.
- 786 Yang, F., Tan, J., Shi, Z. B., Cai, Y., He, K., Ma, Y., Duan, F., Okuda, T., Tanaka, S., and Chen, G.:
- Five-year record of atmospheric precipitation chemistry in urban Beijing, China, Atmos. Chem.
- 788 Phys., 12, 2025-2035, doi: 10.5194/acp-12-2025-2012, 2012.
- Yang, L., Ren, Y., and Jia, L.: Preliminary study of chemical composition of precipitation at
 Wudaoliang, Qinghai Province, Plateau Meteorology, 10, 209-216, 1991 (in Chinese with English
 abstract).
- Yang, Y. H., Ji, C. J., Ma, W. H., Wang, S. F., Wang, S. P., Han, W. X., Mohammat, A., Robinson, D.,
- and Smith, P.: Significant soil acidification across northern China's grasslands during 1980s2000s, Glob. Change Biol., 18, 2292-2300, doi: 10.1111/j.1365-2486.2012.02694.x, 2012.
- 795 Yang, Z., Ou Yang, H., Xu, X., and Yang, W.: Spatial heterogeneity of soil moisture and vegetation
- coverage of alpine grassland in permafrost area of the Qinghai-Tibet Plateau, Journal of Natural
 Resources, 25, 426-434, doi: 10.11849/zrzyxb.2010.03.008, 2010 (in Chinese with English
- abstract).
- Yao, T. D., Masson-Delmotte, V., Gao, J., Yu, W. S., Yang, X. X., Risi, C., Sturm, C., Werner, M., Zhao,
- 800 H. B., He, Y., Ren, W., Tian, L. D., Shi, C. M., and Hou, S. G.: A review of climatic controls on
- 801 δ^{18} O in precipitation over the Tibetan Plateau: observations and simulations, Rev. Geophys., 51,
- 802 525–548, doi: 10.1002/rog.20023, 2013.

803	Yao, T., Thompson, L. G., Mosbrugger, V., Zhang, F., Ma, Y., Luo, T., Xu, B., Yang, X., Joswiak, D.
804	R., Wang, W., Joswiak, M. E., Devkota, L. P., Tayal, S., Jilani, R., and Fayziev, R.: Third Pole
805	Environment (TPE), 3, 52-64, doi:10.1016/j.envdev.2012.04.002, 2012.
806	Zaehle, S., Friedlingstein, P., and Friend, A. D.: Terrestrial nitrogen feedbacks may accelerate future
807	climate change, Geophys. Res. Lett., 37, 20130125, doi: 10.1029/2009gl041345, 2010.
808	Zaehle, S., and Friend, A. D.: Carbon and nitrogen cycle dynamics in the O-CN land surface model: 1.
809	Model description, site-scale evaluation, and sensitivity to parameter estimates, Global
810	Biogeochem. Cycles, 24, GB1005, doi: 10.1029/2009gb003521, 2010.
811	Zaehle, S.: Terrestrial nitrogen - carbon cycle interactions at the global scale, Philos. T. R. Soc. B, 368,

- 812 L01401, doi: 10.1098/rstb.2013.0125, 2013.
- Zbieranowski, A. L., and Aherne, J.: Long-term trends in atmospheric reactive nitrogen across Canada:
 1988-2007, Atmos. Environ., 45, 5853-5862, doi: 10.1016/j.atmosenv.2011.06.080, 2011.
- Zhang, D. D., Peart, M., Jim, C. Y., He, Y. Q., Li, B. S., and Chen, J. A.: Precipitation chemistry of
- 816 Lhasa and other remote towns, Tibet, Atmos. Environ., 37, 231-240, doi:10.1016/S1352817 2310(02)00835-X, 2003.
- 818 Zhang, M., Wang, S., Wu, F., Yuan, X., and Zhang, Y.: Chemical compositions of wet precipitation
- and anthropogenic influences at a developing urban site in southeastern China, Atmos. Res., 84,
- 820 311-322, doi: 10.1016/j.atmosres.2006.09.003, 2007.
- 821 Zhang, N. N., He, Y. Q., Cao, J. J., Ho, K. F., and Shen, Z. X.: Long-term trends in chemical
- 822 composition of precipitation at Lijiang, southeast Tibetan Plateau, southwestern China, Atmos.
- 823 Res., 106, 50-60, doi: 10.1016/j.atmosres.2011.11.006, 2012.
- 824 Zhang, X. Y., Arimoto, R., Cao, J. J., An, Z. S., and Wang, D.: Atmospheric dust aerosol over the
- Tibetan Plateau, J. Geophys. Res.-Atmos., 106, 18471-18476, doi: 10.1029/2000jd900672, 2001.

- Zhang, X. Y., Jiang, H., Zhang, Q. X., and Zhang, X.: Chemical characteristics of rainwater in northeast
 China, a case study of Dalian, Atmos. Res., 116, 151-159, doi: 10.1016/j.atmosres.2012.03.014,
 2012.
- 829 Zhang, Y. J., Kang, S. C., You, Q. L., and Xu, Y. W.: Climate in the Nam Co basin, in: Modern
- 830 environmental processes and changes in the Nam Co basin, Tibetan Plateau, edited by: Kang, S.
- C., Yang, Y. P., Zhu, L. P., and Ma, Y. M., China Meteorological Press, Beijing, 15-24, 2011 (in
 Chinese).
- Zhang, Y. L., Li, B. Y., and Zheng, D.: A discussion on the boundary and area of the Tibetan Plateau
 in China, Geographical Research, 21, 1-8, 2002 (in Chinese with English abstract).
- 835 Zhao, H. B., Xu, B. Q., Yao, T. D., Tian, L. D., and Li, Z.: Records of sulfate and nitrate in an ice core
- from Mount Muztagata, central Asia, J. Geophys. Res.-Atmos., 116, D13304, doi:
 10.1029/2011jd015735, 2011.
- 838 Zhao, Z. Z., Cao, J. J., Shen, Z. X., Xu, B. Q., Zhu, C. S., Chen, L. W. A., Su, X. L., Liu, S. X., Han,
- Y. M., Wang, G. H., and Ho, K. F.: Aerosol particles at a high-altitude site on the Southeast Tibetan
- 840 Plateau, China: Implications for pollution transport from South Asia, J. Geophys. Res-Atmos.,
- 841 118, 11360-11375, doi: 10.1002/jgrd.50599, 2013.
- Zheng, W., Yao, T. D., Joswiak, D. R., Xu, B. Q., Wang, N. L., and Zhao, H. B.: Major ions composition
- records from a shallow ice core on Mt. Tanggula in the central Qinghai-Tibetan Plateau, Atmos.
- Res., 97, 70-79, doi: 10.1016/j.atmosres.2010.03.008, 2010.

Station name	Station name expanded	Latitude	Longitude	Altitude	Annual mean	Annual	Vegetation type	References
					temperature	precipitation		
				m a.s.l.	°C	mm yr^{-1}		
Southeast Tibet Station	Southeast Tibet Observation and Research	29°46′N	94°44′E	3326	5.6	800-1000	Subalpine coniferous forest	Wang et al.,
	Station for the Alpine Environment,						and temperate deciduous	(2010)
	Chinese Academy of Sciences						conifer mixed forest	
Nam Co Station	Nam Co Monitoring and Research Station	30°47′N	90°58′E	4730	-0.6	414.6	Alpine meadow and alpine	Zhang et al.,
	for Multisphere Interactions, Chinese						steppe	(2011)
	Academy of Sciences							
Qomolangma Station	Qomolangma Atmospheric and	28°13′N	86°34′E	4300	3.9	402.8	Alpine meadow and alpine	Gao et al.,
	Environmental Observation and Research						steppe	(2014) and M.
	Station, Chinese Academy of Sciences							Li et al.,
								(2007)
Ngari Station	Ngari Desert Observation and Research	33°24′N	79°43′E	4264		124.6	Desert steppe	This study
	Station							
Muztagh Ata Station	Muztagh Ata Westerly Observation and	38°17′N	75°1′E	3650		213.6	Alpine steppe	This study
	Research Station							

Table 1. Descriptions of the five precipitation sampling sites on the TP<u>in the TP</u>.

<u> </u>			or proorproor			(-)							
				Precipitati										
Area	Sites	Represents	Periods	on	$NH_4{}^+$	Na^+	\mathbf{K}^+	Mg^{2+}	Ca ²⁺	NO_3^-	Cl^{-}	${\rm SO_4^{2-}}$	Data type	Reference
Tibetan	Southeast Tibet	Remote site	2011-2012	914.6	4.9	3.8	1.0	0.9	20.8	2.2	2.5	2.1	VWM	This study
Plateau	Nam Co	Remote site	2011-2012	382.5	12.7	1.9	0.4	0.9	7.9	4.5	1.1	2.9	VWM	This study
	Qomolangma	Remote site	2011-2012	258	25.4	26.0	4.5	1.7	53.4	0.8	27.1	3.2	VWM	This study
	Ngari	Remote site	2013	124.6	20.5	12.4	1.8	4.8	50.9	4.8	11.9	11.6	VWM	This study
	Muztagh Ata	Remote site	2011	213.6	42.0	10.1	3.0	11.3	119.4	9.9	8.9	17.4	VWM	This study
	Waliguan	Remote site	1997	388	45.5	8.7	3.8	12.1	34.0	8.3	6.1	24.0	Mean	Tang et al. (2000)
	Wudaoliang	Remote site	Aug. 1989	266.5ª	27.1	21.7	6.2	=	Ξ	13.2	25.6	29.2	Mean	Yang et al. (1991)
	Lhasa	Remote city	1998-2000	250-500	14.3	11.2	5.1	10.9	197.4	6.9	9.7	5.2	Mean	Zhang et al. (2003)
	Lijiang	City	1989–2006	900	11.4	2.5	=	7.7	50.2	3.6	11.6	32.6	Mean	Zhang et al. (2012a)
				Average	22.6	10.9	3.2	6.3	66.8	6.0	11.6	14.3		
Northern	Beijing	City	2001-2005	441	236.0	22.5	13.8	48.4	209.0	106.0	34.9	314.0	VWM	F. Yang et al. (2012)
China	Dalian	City	2007	602	107.8	36.2	6.87	25.29	78.92	51.38	59.83	168.0	VWM	Zhang et al. (2012e)
	Nanjing	City	1992-2003	648-1242	193.2	23.0	12.1	31.7	295.4	39.6	142.6	241.8	VWM	Tu et al. (2005)
	Tianshan Mountain	Remote site	1995–1996	=	=	55.7	14.9	15.8	78.0	22.3	40.9	88.1	Mean	Hou (2001)
Southern	Hangzhou	City	2006-2008	1435	79.9	12.2	4.2	7.1	51.9	38.4	13.9	110.0	VWM	Xu et al. (2011)
China	NingBo	City	2010-2011	1374.7	46.2	22.4	7.0	9.3	31.5	38.7	31.0	72.6	VWM	Ding et al. (2012)
	Shanghai	City	2005	825.5	80.7	50.1	14.9	29.6	204.0	49.8	58.3	199.6	VWM	K. Huang et al. (2008)
	Shenzhen	City	1986–2006	1769	35.2	40.3	7.2	9.7	77.7	22.1	37.9	74.3	Mean	Y. L. Huang et al. (2008)
	Guiyang	City	2008-2009	1171	112.8	13.9	9.6	10.5	182.9	7.3	20.7	265.6	VMW	Xiao et al. (2013)

Table 2. Annual mean concentrations of major ions ($\mu eq L^{-1}$) in precipitation at five remote sites on the TP in the TP and other sites in China.

850 Unit of precipitation is mm yr⁻¹ Units of precipitation are millimeters (mm yr⁻¹). VWM indicates volume-weighted mean.

851 ^a Precipitation amount data was obtained from Yang et al. (2010).

Table 3. Annual inorganic nitrogen wet deposition (kg N ha⁻¹ yr⁻¹) at five remote sites on the

Figure 4.5.3 TPin the TP, as well as other sites in China. Units of precipitation are is millimeters (mm yr⁻¹). DIN

854	means inorganic	e nitrogen (sum	of NH4 ⁺ -N	and NO ₃ N).	VWM indicates	volume-weighted mean
-----	-----------------	-----------------	------------------------	-------------------------	---------------	----------------------

Area	Sites	Represents	Periods	Precipitation	NH4 ⁺ -N	NO ₃ -N	DIN	Data type	Reference
Tibetan	Southeast	Remote site	2011-2012	914.6	0.63	0.28	0.91	VWM	This study
Plateau	Nam Co	Remote site	2011-2012	382.5	0.68	0.24	0.92	VWM	This study
	Qomolangma	Remote site	2011-2012	258	0.92	0.03	0.94	VWM	This study
	Ngari	Remote site	2013	124.6	0.36	0.08	0.44	VWM	This study
	Muztagh Ata	Remote site	2011	213.6	1.25	0.30	1.55	VWM	This study
	Waliguan	Remote site	1997	388	2.47	0.45	2.92	Mean	Tang et al. (2000)
	Wudaoliang	Remote site	Aug. 1989	266.5 ^a	1.55	1.11	2.66	Mean	Yang et al. (1991)
	Lhasa	Remote city	1998–2000	250-500	0.54	0.45	0.99	Mean	Zhang et al. (2003)
	Naidong	Remote city	2006-2007	451	0.91	0.82	1.72	VWM	Jia (2008)
	Biru	Remote city	2006-2007	582	1.22	1.86	3.08	VWM	Jia (2008)
	Jiangda	Remote city	2006-2007	547	1.11	0.80	1.91	VWM	Jia (2008)
	Lijiang	Remote city	1989–2006	900	1.43	0.46	1.89	Mean	N. N. Zhang et al. (2012)
				Average	1.09	0.57	1.66		
Northern	Beijing	City	2001-2005	441	14.57	6.54	21.12	VWM	F. Yang et al. (2012)
China	Dalian	City	2007	602	9.08	4.33	13.41	VWM	X. Y. Zhang et al. (2012)
	Nanjing	City	1992-2003	648-1242	25.56	5.23	30.79	VWM	Tu et al. (2005)
	Beijing	City	2008-2010	572	=	=	27.9	VWM	Pan et al. (2012)
	Tianjin	City	2008-2010	544	=	=	18.1	VWM	Pan et al. (2012)
	Baoding	Industrial	2008-2010	513	=	=	23.1	VWM	Pan et al. (2012)
	Tanggu	Industrial	2008-2010	566	=	=	28.2	VWM	Pan et al. (2012)
	Tangshan	Industrial	2008-2010	610	=	=	21.6	VWM	Pan et al. (2012)
	Yangfang	Suburban	2008-2010	404	=	=	20.7	VWM	Pan et al. (2012)
	Cangzhou	Suburban	2008-2010	605	=	=	22.6	VWM	Pan et al. (2012)
	Luancheng	Agricultural	2008-2010	517	=	=	22.2	VWM	Pan et al. (2012)
	Yucheng	Agricultural	2008-2010	566	=	=	24.8	VWM	Pan et al. (2012)
	Xinglong	Rural	2008-2010	512	=	=	16.3	VWM	Pan et al. (2012)
Southorn	TiaShanDing	Domoto sito	1000 2004	1000	25 50	0.80	25 20	VIXINA	Chan and Muldar (2007)
China	LinChang	Remote site	1999–2004	1228	25.50	9.80	55.50 2.70		Chen and Mulder (2007)
China	LiuChongGuan	Remote site	1999–2004	854	2.40	1.50	5.70		Chen and Mulder (2007)
	CeilieTere	Remote site	1999–2004	1/14	5.70 21.10	2.00	0.30		Chen and Mulder (2007)
		Remote site	1999–2004	1232	21.10	12.70	33.80		Chen and Mulder (2007)
	LiuxiHe	Remote site	1999–2004	1620	4.30	7.50	11.80	VWM	Chen and Mulder (2007)
	Hangzhou	City	2006-2008	1435	16.1	7.7	23.77	VWM	Xu et al. (2011)
	Ningbo	City	2010–2011	13/4.7	8.9	7.4	16.34	VWM	Ding et al. (2012)
	Shanghai	City	2005	825.5	9.3	5.8	15.08	VWM	K. Huang et al. (2008)
	Shenzhen	City	1986-2006	1769	8.7	5.5	14.19	Mean	Y. L. Huang et al. (2008)
	Guiyang	City	2008–2009	1171	18.5	1.2	19.69	VMW	X1ao et al. (2013)

855 Notes: Tang et al. (2000), Yang et al. (1991), Zhang et al. (2003), N. N. Zhang et al. (2012), F. Yang et al. (2012), X. Y.

856 Zhang et al. (2012), Xu et al. (2011), Ding et al. (2012), K. Huang et al. (2008), Y. L. Huang et al. (2008) and Xiao et al.

857 (2013) reported the concentrations of NH₄⁺-N and NO₃⁻-N in precipitation, but did not calculate nitrogen wet deposition.

858 For these previous studies, we recalculated the annual inorganic nitrogen wet deposition according to the reported

- 859 concentrations of NH₄⁺-N and NO₃⁻-N in precipitation and annual precipitation. ^a Precipitation amount data was obtained
- 860 from Yang et al. (2010).

Table 4. Enrichment factors (EFs) relative to seawater and soil for precipitation constituents of

863 **five remote sites on the TP**in the TP.

	Southeast Tibet Station		Nam Co Station		Qomolangma Station		Ngari Station		Muztagh Ata Station		[X/Na ⁺] _{sea}	$[X/Ca^{2+}]_{soil}$
	EFsea	$\mathrm{EF}_{\mathrm{soil}}$	EF _{sea}	$\mathrm{EF}_{\mathrm{soil}}$	$\mathrm{EF}_{\mathrm{sea}}$	$\mathrm{EF}_{\mathrm{soil}}$	$\mathrm{EF}_{\mathrm{sea}}$	EF _{soil}	EF _{sea}	$\mathrm{EF}_{\mathrm{soil}}$		
Na^+	1.0	0.36	1.0	0.48	1.0	0.99	1.0	0.5	1.0	0.17	1.0000	0.5690
$NH_{4}{}^{+}$	15707	350	80684	2378	11629	701	19706	594	49751	519	0.0001ª	0.0006°
\mathbf{K}^+	11.5	0.18	8.5	0.17	7.83	0.33	6.4	0.13	13.5	0.10	0.0220	0.5040
Mg^{2+}	1.1	0.05	2.0	0.12	0.29	0.03	1.7	0.10	4.9	0.10	0.2270	0.5610
Ca^{2+}	126.2	1.0	95.3	1.0	46.6	1.0	93.1	1.0	269.5	1.0	0.0440	1.0000
Cl^-	0.57	67.3	0.50	77.8	0.90	286	0.8	132	0.76	42.2	1.1600	0.0031
NO_3^-	23842	154	98242	840	1237	21.7	15869	139	40619	123	0.0000^{b}	0.0021 ^d
${\rm SO_4^{2-}}$	4.7	13.0	12.7	46.7	1.03	7.76	7.7	29.2	14.3	18.6	0.1210	0.0188 ^e

^a Marine nitrogen ions <u>were</u> regarded as entire NH₄⁺.

^bMarine nitrogen ions<u>were</u> regarded as entire NO₃⁻.

866 ° Soil nitrogen was regarded as entire range of NH_3 compounds.

^dSoil nitrogen <u>was</u> regarded as entire range of NO₃ compounds.

^eSoil sulfur <u>was</u> regarded as entire range of SO₄ compounds.

Table 5. Source contributions (%) for major ions in precipitation of five remote sites on the

TPin the TP. SSF indicates sea salt fraction; CF indicates crust fraction; AF indicates anthropogenic

	South	east Ti	bet	Nam Co			Qomo	olangm	a	Ngari			Muztagh Ata			
	Station			Station			Statio	Station			n		Station			
	SSF	CF	AF	SSF	CF	AF	SSF	CF	AF	SSF	CF	AF	SSF	CF	AF	
$\mathrm{NH_4^+}$	0.0	0.3	99.7	0.0	0.0	100	0.0	0.1	99.8	0.0	0.2	99.8	0.0	0.2	99.8	
NO_3^-	0.0	0.6	99.3	0.0	0.1	99.9	0.1	4.6	95.3	0.0	0.7	99.3	0.0	0.8	99.2	
${\rm SO_4^{2-}}$	21.3	7.7	71.0	7.9	2.1	90.0	87.1	12.9		12.9	3.4	83.7	7.0	5.4	87.6	
Ca^{2+}	0.8	99.2		1.0	99.0		2.1	97.9		1.1	98.9		0.4	99.6		
\mathbf{K}^+	8.7	91.3		11.8	88.2		12.8	87.2		15.5	84.5		7.4	92.6		
Mg^{2+}	92.9	7.1		49.0	51.0		0	100		59.2	40.8		20.2	79.8		
\mathbf{Cl}^{-}	98.5	1.5		98.7	1.3		99.7	0.3		99.2	0.8		97.6	2.4		
Na ⁺	100			100			100			100			100			

Table 6: Varimax-rotated principal component analysis of major ions in precipitation at five remote sites on the TPin the TP. PC1, PC2 and PC3 indicates the 1st, 2nd
and 3rd component, respectively. CT means communality. *N* indicates the number of precipitation samples at each site. Boldfaced values are the largest value among
the 3 components for each ion at each site.

	Southeast Tibet				Nam Co				Qomolangma				Ngari				Muztagh Ata			
	(<i>N</i> = 53)				(<i>N</i> = 27)			(<i>N</i> = 30)				(<i>N</i> = 39)				(<i>N</i> = 19)				
	RC1PC1	<u>P</u> ₽C2	<u>P</u> ₽C3	СТ	<u>₽</u> ₽C1	<u>P</u> ₽C2	PRC3	СТ	<u>P</u> ₽C1	<u>P</u> ₽C2	<u>₽</u> ₽C3	СТ	<u>P</u> ₽C1	PRC2	<u>P</u> ₽C3	СТ	<u>P</u> ₽C1	<u>P</u> ₽C2	<u>₽</u> ₽C3	СТ
Na ⁺	0.94	0.02	0.24	0.94	0.62	0.73	0.00	0.93	0.91	0.28	0.10	0.92	0.62	0.73	0.06	0.92	0.22	0.96	0.08	0.98
$\mathbf{NH_4}^+$	0.25	0.22	0.93	0.98	0.36	0.13	0.89	0.94	0.69	-0.15	0.24	0.56	-0.11	0.57	0.75	0.90	-0.19	0.14	0.96	0.98
K^+	0.88	0.10	0.35	0.91	0.11	0.93	0.07	0.89	0.84	0.35	0.27	0.89	0.15	0.84	0.40	0.89	0.47	0.79	-0.05	0.85
Mg^{2+}	0.77	0.45	0.29	0.89	0.89	0.19	0.38	0.97	0.39	0.88	0.18	0.97	0.67	0.56	0.09	0.78	0.90	0.38	-0.16	0.99
Ca ²⁺	0.66	0.46	0.21	0.70	0.76	0.11	0.57	0.92	-0.02	0.96	0.03	0.92	0.85	0.17	0.15	0.77	0.85	0.35	-0.25	0.90
Cl^{-}	0.91	0.10	-0.06	0.85	0.05	0.95	0.20	0.95	0.92	0.25	0.11	0.92	0.37	0.83	0.00	0.82	0.25	0.92	0.17	0.94
NO_3^-	0.03	0.95	0.09	0.91	0.32	0.13	0.92	0.96	0.26	0.16	0.94	0.98	0.36	-0.01	0.84	0.84	0.88	0.12	-0.18	0.83
${\rm SO_4}^{2^-}$	0.24	0.89	0.18	0.89	0.80	0.10	0.52	0.92	0.64	0.64	0.31	0.90	0.91	0.22	0.15	0.90	0.87	0.31	0.11	0.87
Variance (%)	46	27	15		33	30	30		43	30	15		34	33	19		43	35	14	
Cumulative (%)	46	73	88		33	63	93		43	74	88		34	66	85		43	78	92	

878 Figure captions

879 Figure 1. Map of the precipitation inorganic nitrogen N wet deposition sampling sites on the TP in 880 the TP. The red points indicate the five remote sampling stations sites of this study. The black points 881 indicate the sampling sites from previous records. Southeast Tibet Station is short for Southeast Tibet 882 Observation and Research Station for the Alpine Environment, Chinese Academy of Sciences; Nam Co Station is short for Nam Co Monitoring and Research Station for Multisphere Interactions, Chinese 883 884 Academy of Sciences; Qomolangma Station is short for Qomolangma Atmospheric and Environmental 885 Observation and Research Station, Chinese Academy of Sciences; Ngari Station is short for Ngari 886 Desert Observation and Research Station; and Muztagh Ata Station is short for Muztagh Ata Westerly 887 Observation and Research Station. Figure 2. Seasonal dynamics of ion concentrations (unit: $\mu eq L^{-1}$) and precipitation (unit: mm) 888

at five remote sites on the TPin the TP. The sampling times of the five stations sites were as follows:
Southeast Tibet Station, November 2011 to October 2012; Nam Co Station, August 2011 to July 2012;
Qomolangma Station, April 2011 to March 2012; Ngari Station, January 2013 to December 2013;
Muztagh Ata Station, January 2011 to December 2011.

Figure 3. Annual average volume-weighted concentration percentages of measured ions in precipitation (unit: $\mu eq L^{-1}/\mu eq L^{-1}$) at five remote sites on the TP in the TP.

Figure 4. Seasonal dynamics of inorganic N wet deposition at five remote sites on the TPin the
TP. The sampling time windows of those stations sites are same to Fig. 2.

Figure 5. Seven-day backward trajectories at five remote sites on the TP in the TP. Black lines

show the backward trajectories calculated at 6-h interval (00:00, 06:00, 12:00, 18:00 UTC) at sampling

days, with an arrival height of 500 m above the ground. Red lines show the clustering trajectories.

Figure 1.





Figure 3.



Figure 4.



Figure 5.

