Atmospheric deposition of reactive nitrogen to a deciduous forest in the southern Appalachian Mountains

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Received: 17 June 2022 – Discussion started: 23 June 2022
Revised: 29 November 2022 – Accepted: 8 December 2022 – Published: 9 March 2023

Abstract. Assessing nutrient critical load exceedances requires complete and accurate atmospheric deposition budgets for reactive nitrogen (N_r). The exceedance is the total amount of N_r deposited to the ecosystem in excess of the critical load, which is the amount of N_r input below which harmful effects do not occur. Total deposition includes all forms of N_r (i.e., organic and inorganic) deposited to the ecosystem by wet and dry pathways. Here we present results from the Southern Appalachian Nitrogen Deposition Study (SANDS), in which a combination of measurements and field-scale modeling was used to develop a complete annual N_r deposition budget for a deciduous forest at the Coweeta Hydrologic Laboratory. Wet deposition of ammonium, nitrate, nitrite, and bulk organic N were measured directly. The dry deposited N_r fraction was estimated using a bidirectional resistance-based model driven with speciated measurements of N_r air concentrations (e.g., ammonia, ammonium aerosol, nitric acid, nitrate aerosol, bulk organic N in aerosol, total alkyl nitrates, and total peroxy nitrates), micrometeorology, canopy structure, and biogeochemistry. Total annual deposition was ~6.7 kg N ha⁻¹ yr⁻¹, which is on the upper end of N_r critical load estimates recently developed for similar ecosystems in the nearby Great Smoky Mountains National Park. Of the total (wet + dry) budget, 51.1 % was contributed by reduced forms of N_r (NHₓ = ammonia + ammonium), with oxidized and organic forms contributing ~41.3 % and 7.6 %, respectively. Our results indicate that reductions in NHₓ deposition would be needed to achieve the lowest estimates (~3.0 kg N ha⁻¹ yr⁻¹) of N_r critical loads in southern Appalachian forests.

1 Introduction

Prior to the Industrial Revolution, Earth’s ecosystems received reactive nitrogen (N_r) deposition rates of ~0.5 kg ha⁻¹ yr⁻¹ (Holland et al., 1999). Since the 19th century, anthropogenic activities, both industrial and agricultural, have resulted in unprecedented quantities of N_r being released into the atmosphere, subsequently altering biogeochemical cycles (Neff et al., 2002a, b; Ollinger et al., 2002; Bragazza et al., 2006; Doney et al., 2007; Galloway et al., 2008; Boonstra et al., 2017). Excessive atmospheric deposition of N_r to terrestrial ecosystems may lead to soil and aquatic acidification, nutrient imbalance and enrichment,
plant damage, and microbial community changes, as well as loss of biodiversity (Bobbink et al., 1998; Lohse et al., 2008; Simkin et al., 2016). Nitrogen deposition rates in many areas, including North America, Europe, and Asia, exceed 10 kg ha⁻¹ yr⁻¹ and may double by the year 2050 in some regions (Galloway et al., 2008).

The amount of N deposition below which significant harmful effects do not occur is known as the critical load (Nilsson and Grennfelt, 1988). Critical loads can be quantified using empirical relationships between ecosystem N input and ecosystem response (Pardo et al., 2011; Root et al., 2015), or mass-balance-type biogeochemical models (Lynch et al., 2017; McNulty et al., 2007). For the southern Appalachian Mountains, simple mass balance approaches yield critical loads similar to those derived from empirical approaches for forest health and biogeochemical responses. In a recent study employing a mass balance model for the Great Smoky Mountains National Park, Pardo et al. (2018) quantified critical loads for spruce–fir, beech, and mixed deciduous forests in the range of 2.8 to 7 kg N ha⁻¹ yr⁻¹, with the highest value corresponding to a high-elevation spruce–fir site experiencing disturbance-induced regrowth. Accurate and complete deposition budgets (i.e., including all forms of N) are required to quantify the amount of N input to ecosystems in excess of the critical load (i.e., the critical load exceedance).

Estimates of N deposition for critical load assessments can be derived from gridded chemical transport models (CTMs) (Ellis et al., 2013; Lee et al., 2016; Simkin et al., 2016; Clark et al., 2018; Makar et al., 2018), measurement–model fusion (MMF) techniques that combine measurements with CTM output (Schwede and Lear, 2014; Nanus et al., 2017; McDonnell et al., 2018; U.S. EPA, 2019), or inferential modeling with site-specific measurements (Flechard et al., 2011; Li et al., 2016). While these approaches reflect the state of the science and are widely used, they collectively suffer from incompleteness of the N deposition budget (and are therefore biased low) (Walker et al., 2019a). Monitoring networks for wet deposition (NADP/NTN) and air concentrations of N₂O (CASTNET) focus only on inorganic species, excluding organic forms of N, which account for ~25% of total N in wet deposition on average (Jickels et al., 2013). Due to difficulties in sampling (Walker et al., 2012) and the inability to fully speculate the wide range of constituents (Neff et al., 2002a; Altieri et al., 2012, 2018; Cape et al., 2011; Samy et al., 2013; Chen et al., 2018), organic N is not routinely monitored. Hence, deposition of organic N remains uncertain, and thus N deposition budgets developed from network monitoring data and CTMs remain incomplete.

Current N deposition estimates also have a relatively high degree of uncertainty in the estimation of dry deposition. While wet deposition is routinely measured, direct measurements of dry N deposition (i.e., flux measurements) in North America are relatively few (Walker et al., 2020). Estimates of dry deposition for ecosystem assessments are therefore derived from models (Schwede and Lear, 2014; Li et al., 2016; Lee et al., 2016). Of the inorganic N species, NH₃ is the most important contributor to dry deposition in many areas (Walker et al., 2019b) but also the most uncertain (Flechard et al., 2011) due to the bidirectionality of surface–atmosphere exchange. A paucity of flux measurements (Walker et al., 2020) precludes bias correction of dry deposition in CTMs and MMF techniques, making dry deposition much more uncertain relative to wet deposition.

The Coweeta study site represents southern Appalachian Mountain forests, which are highly diverse and productive ecosystems that provide a variety of ecosystem services, including a source of surface drinking water (Caldwell et al., 2014). While deposition of oxidized N to forests in the southeastern US has declined in response to the Clean Air Act, montane ecosystems continue to receive high rates of deposition due to elevation-induced precipitation gradients (Weathers et al., 2006; Knoepp et al., 2008). Southern Appalachian forests continue to show signs of sensitivity to N deposition. For example, litterfall N fluxes and foliar N concentrations at Coweeta have steadily increased over the past 2 decades (Knoepp et al., 2018). Highly spatially variable meteorological patterns typical of complex terrain are difficult to model (Lehner and Rotach, 2018), leading to uncertainties in precipitation amounts and wet deposition (Zhang et al., 2018) as well as the micrometeorological processes that govern dry deposition (Cowan et al., 2022). Estimates of deposition from gridded CTMs in mountainous terrain therefore contain a higher degree of uncertainty relative to low-elevation ecosystems. For these reasons, a better understanding of total N deposition in southern Appalachian forests is needed.

This study investigates the N deposition budget in a remote montane forest in the southeastern US. We combine long-term (1978–2020) and seasonal intensive (2015–2016) measurements of wet deposition, speciated air concentrations of N₂O, micrometeorology, biogeochemistry, and forest canopy structure with in situ inferential dry deposition modeling to develop an annual, speciated, total N deposition budget, including net and component NH₃ fluxes as well as dry and wet organic N deposition. Seasonal and annual total N deposition fluxes are presented in the context of long-term trends in air concentrations and wet deposition of inorganic N species. Spatial representativeness is characterized using measurements of air concentrations of the primary inorganic N species and previous wet deposition measurements along an elevation gradient across the topographically complex forested basin.

2 Methods

2.1 Site description

The study was conducted at the USDA Forest Service Coweeta Hydrologic Laboratory, a 2185 ha experimental
forest in southwestern North Carolina, USA (35.0605° N, 83.4305° W), near the southern end of the Appalachian Mountain chain. Topography is complex, with elevations ranging from 681 to 1594 m within the Coweeta Basin. Mean annual temperature and precipitation are 12.9 °C and 1795 mm, respectively. Dominant overstory species are Liriodendron tulipifera, Quercus alba, Betula lenta, and Acer rubrum, which comprise 24%, 18%, 11%, and 8% of the basal area, respectively, in the low-elevation forests where the study was conducted (Oishi et al., 2018). The dominant understory woody shrub species is Rhododendron maximum (evergreen), which comprises 15% of the basal area (Oishi et al., 2018). Species composition in the vicinity of the eddy flux tower (EFT), further described below, is detailed in Table S1 in the Supplement. Canopy height surrounding the EFT is ~30 m.

The Coweeta Basin has been a long-term monitoring site for atmospheric chemistry and deposition since the late 1970s. Weekly wet deposition of ammonium (NH₄⁺) and nitrate (NO₃⁻), along with sulfate (SO₄²⁻), chloride, and base cations, has been measured as part of the NADP/NTN (Site NC25, https://nadp.slh.wisc.edu/networks/ national-trends-network/, last access: 1 November 2022) since 1978. Weekly integrated air concentrations of particulate NH₄⁺, NO₃⁻, SO₄²⁻, chloride, and base cations, as well as nitric acid (HNO₃) and sulfur dioxide (SO₂), have been measured by CASTNET (Site COW137, https://www.epa.gov/castnet, last access: 1 November 2022) since 1987. Since 2011, biweekly integrated air concentrations of ammonia (NH₃) have been measured by the NADP Ammonia Monitoring Network (AMoN Site NC25, https://nadp.slh.wisc.edu/networks/ammonia-monitoring-network/, last access: 2 November 2022). Here we use these datasets to place our study results into historical context, to supplement the more intensive atmospheric chemistry measurements described below, and to use as inputs for inferential modeling of dry deposition. The long-term NADP and CASTNET measurements are collected in the lower part of the basin, indicated as NC25/COW137 in Fig. 1.

### 2.2 Southern Appalachian Nitrogen Deposition Study

Building on the long-term NADP and CASTNET measurements described above, the Southern Appalachian Nitrogen Deposition Study (SANDS) was conducted in 2015 and 2016 to better understand the atmospheric chemistry and deposition of reactive nitrogen at Coweeta. Intensive measurement campaigns were conducted from 21 May–9 June 2015, 6–25 August 2015, 9–26 September 2015, 19 April–11 May 2016, and 13 July–3 August 2016. A sub-set of measurements was conducted continuously between February 2015 and August 2016. As described below, time-resolved and time-integrated measurement techniques were used to characterize organic N in the gas phase, in particulate matter, and in wet deposition; the temporal variability of air concentrations of gas-phase oxidized and reduced forms of N; and the spatial variability of atmospheric N concentrations across the Coweeta Basin. Vertical profiles of air concentrations were measured within the forest canopy to examine source–sink processes, and measurements of soil and vegetation chemistry were conducted to characterize NH₃ emission potentials. Measurements were combined with NADP and CASTNET data to develop seasonal and annual total N deposition budgets employing inferential modeling for the dry deposition component. Vertical concentration profile and biogeochemical measurements were used to inform the parameterization of NH₃ bidirectional exchange. Sampling locations are described in Fig. 1 and Table 1. Measurement details are summarized in Table 2.

#### 2.2.1 Wet deposition

Additional wet deposition measurements were conducted adjacent to the NTN NC25 sampler to quantify the contribution of bulk water-soluble organic N (WSON) to water-soluble total nitrogen (WSTN) in precipitation. Weekly precipitation samples were collected in a modified wet-only sampler with a borosilicate glass funnel and amber glass bottle (Walker et al., 2012), shielded from sunlight, and maintained in the field under continuous refrigeration to maintain the stability of ON until retrieval (Walker et al., 2012). Samples were sent to the NADP Central Analytical Laboratory on ice for analysis of NH₄⁺, NO₃⁻, NO₂⁻, and WSTN as described by Walker et al. (2012). WSON concentration was calculated as

\[
WSON = WSTN - (NH_4^+ + NO_3^- + NO_2^-).
\]
Table 1. Sampling locations and type of sampler deployed.

<table>
<thead>
<tr>
<th>Site code</th>
<th>Latitude (N)</th>
<th>Longitude (W)</th>
<th>Elevation (m)</th>
<th>Sampler type</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC25/COW137</td>
<td>35.0605</td>
<td>83.4305</td>
<td>686</td>
<td>AMoN (NC25), CASTNET (COW137), Tisch PM$_{2.5}$, DD-CL, TD-PC-CL, NTN (NC25), EPA precipitation</td>
</tr>
<tr>
<td>EFT$^a$</td>
<td>35.0591</td>
<td>83.4274</td>
<td>690</td>
<td>MARGA$^c$, URG$^c$, passive NH$_3$ and HNO$_3$, micrometeorology</td>
</tr>
<tr>
<td>WS18</td>
<td>35.0512</td>
<td>83.4337</td>
<td>806</td>
<td>Passive NH$_3$ and HNO$_3$</td>
</tr>
<tr>
<td>SK$^b$</td>
<td>35.0482</td>
<td>83.4542</td>
<td>986</td>
<td>Passive NH$_3$ and HNO$_3$, CASTNET (COW005)</td>
</tr>
<tr>
<td>CS28</td>
<td>35.0466</td>
<td>83.4650</td>
<td>1189</td>
<td>Passive NH$_3$ and HNO$_3$</td>
</tr>
<tr>
<td>CS77</td>
<td>35.0303</td>
<td>83.4604</td>
<td>1425</td>
<td>Passive NH$_3$ and HNO$_3$</td>
</tr>
</tbody>
</table>

$^a$ Eddy flux tower. $^b$ Screwdriver Knob. $^c$ Tisch PM$_{2.5}$, MARGA, and URG denuder and filter pack samplers were deployed only during intensive sampling periods.

Table 2. Details of intensive and long-term atmospheric measurements at Coweeta.

<table>
<thead>
<tr>
<th>Sampler name</th>
<th>Operating periods</th>
<th>Measured species</th>
<th>Resolution</th>
<th>Height (m)$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DD-CL, TD-PC-CL</td>
<td>August 2015–August 2016</td>
<td>HNO$_3$, NO$_y$, $\Sigma$AN$^b$, $\Sigma$PN$^c$</td>
<td>Hourly</td>
<td>8</td>
</tr>
<tr>
<td>MARGA</td>
<td>Spring, summer 2016 intensives</td>
<td>HNO$_3$, NH$_3$, NO$_2^-$, SO$_4^{2-}$, NH$_4^+$</td>
<td>Hourly</td>
<td>$\sim$ 40</td>
</tr>
<tr>
<td>URG denuder and filter</td>
<td>All intensives 2015–2016</td>
<td>NO$_2^-$, NO$_3^-$, SO$_4^{2-}$, NH$_4^+$, WSTN</td>
<td>3 or 4 h integrated</td>
<td>Multiple</td>
</tr>
<tr>
<td>Tisch PM$_{2.5}$</td>
<td>All intensives 2015–2016</td>
<td>HNO$_3$, NH$_3$, NO$_2^-$, NO$_3^-$, SO$_4^{2-}$, NH$_4^+$</td>
<td>24 h integrated</td>
<td>$\sim$ 1</td>
</tr>
<tr>
<td>CASTNET (COW137)</td>
<td>Long-term</td>
<td>HNO$_3$, NO$_2^-$, SO$_4^{2-}$, NH$_4^+$, Cl$^-$, base cations</td>
<td>Weekly integrated</td>
<td>10</td>
</tr>
<tr>
<td>AMoN (NC25)</td>
<td>Long-term</td>
<td>HNO$_3$, NH$_3$</td>
<td>Biweekly</td>
<td>2</td>
</tr>
<tr>
<td>Passive HNO$_3$, NH$_3$</td>
<td>2015</td>
<td>HNO$_3$, NH$_3$</td>
<td>Biweekly</td>
<td>10</td>
</tr>
<tr>
<td>CASTNET (COW005)</td>
<td>2015</td>
<td>HNO$_3$, NO$_2^-$, SO$_4^{2-}$, NH$_4^+$, Cl$^-$, base cations</td>
<td>Weekly integrated</td>
<td>10</td>
</tr>
<tr>
<td>NADP/NTTN</td>
<td>Long-term</td>
<td>NO$_3^-$, NH$_4^+$, SO$_4^{2-}$, Cl$^-$, H$^+$, base cations</td>
<td>Weekly accumulated</td>
<td>Ground</td>
</tr>
<tr>
<td>EPA precipitation</td>
<td>February 2015–August 2016</td>
<td>NO$_2^-$, NO$_3^-$, SO$_4^{2-}$, NH$_4^+$, WSTN</td>
<td>Weekly accumulated</td>
<td>Ground</td>
</tr>
</tbody>
</table>

$^a$ Above ground. $^b$ Total alkynitrates. $^c$ Total peroxynitrates.

The method detection limit for WSON is 10 µg N L$^{-1}$ (Walker et al., 2012). These measurements were collected continuously from February 2015 to August 2016.

During the spring of 2015, thymol was added to the precipitation collection bottle as a biocide to inhibit organic nitrogen loss in the sample should the refrigerated collector malfunction or lose power. Thymol negatively affected the precision of the total nitrogen measurement, and its use was discontinued in fall of 2015. Ultimately, there were no issues with the refrigerated collector, and the thymol-containing samples were excluded from the analysis presented herein. However, this data loss resulted in a gap from August 2015 to mid-October 2015 of the 12-month period for which the total deposition budget is developed. The data gap comprised eight weekly periods in which precipitation occurred. For this period, the NH$_4^+$ and NO$_3^-$ concentrations from the collocated NADP/NTTN NC25 sampler were used. Based on the SANDS measurements, the ON concentration during this period was estimated by assuming that NH$_4^+$ and NO$_3^-$ contribute 89% of total nitrogen in rainfall, with WSON representing the balance (11%). For the annual budget, weekly concentrations were combined with measured precipitation depth to calculate weekly deposition (kg N ha$^{-1}$). Comparison between SANDS and NTN concentrations of NH$_4^+$ and
NO$_3^-$ showed very good agreement (Sect. S1 in the Supplement, Fig. S1).

### 2.2.2 Air concentrations

Hourly concentrations of NO$_y$, HNO$_3$, total gas-phase peroxy nitrates (ΣPN), and total gas-phase alkyl nitrates (ΣAN) were measured continuously from August 2015 to August 2016 at the height of 8 m adjacent to the COW137 CASTNET tower (Fig. 1, Tables 1 and 2). NO$_y$ and HNO$_3$ were measured using a modified model 42S NO–NO$_2$–NO$_y$ analyzer; the NO$_y$ technique is described in detail by Williams et al. (1998). Briefly, total oxidized reactive nitrogen (NO$_x$) is converted to NO using a molybdenum catalyst heated to 325 $^\circ$C. On a second channel, a metal denuder coated with potassium chloride (KCl) is used to remove HNO$_3$ before passing through a second molybdenum converter heated to 325 $^\circ$C. The difference between the total NO$_x$ measurement and the HNO$_3$-scrubbed NO$_x$ measurement is interpreted as HNO$_3$. Here we refer to the method as denuder difference chemiluminescence (DD-CL).

Total peroxy nitrates (ΣPNs) and total alkynitrates (ΣANs) were measured using a modification of the technique described by Day et al. (2002), in which PN$s$ and AN$s$ are thermally decomposed to NO$_2$ followed by measurement of the incremental NO$_2$ above ambient background for each decomposition step. Day et al. (2002) quantified NO$_2$ via laser-induced fluorescence, while photolytic conversion to NO and quantification of the resulting NO by NO–O$_3$ chemiluminescence is used in the current study. Here we refer to the method as thermal-decomposition photolytic-conversion chemiluminescence (TD-PC-CL). A single chemiluminescence analyzer was used for NO$_2$, HNO$_3$, ΣPN, and ΣAN measurements. Additional detail on the instrument and associated QA/QC procedures is included in Sect. S2.

Hourly concentrations of NH$_3$ and HNO$_3$ were measured on the eddy flux tower (EFT, Fig. 1, Tables 1 and 2) at two heights above the canopy (34 and 37.5 m.a.g., above ground, during spring 2016: 34 and 43.5 m.a.g. during summer 2016) using the Monitor for Aerosols and Gases in Ambient Air (MARGA, Metrohm Applikon B.V., the Netherlands). Details and principles of the MARGA system have been previously described (Remsey and Walker, 2016; Chen et al., 2017). Briefly, the MARGA 2S consisted of two sampler boxes positioned on the tower and a detector box located in a climate-controlled enclosure at the base of the tower. Sample boxes comprised an inlet of 1.27 cm outer diameter 30 cm long perfuorolalkoxy Teflon tubing with no particle size selection, through which airflow was mass-controlled at $\sim$ 16.7 L min$^{-1}$, a wet rotating denuder (WRD) for collection of soluble gases, and a steam jet aerosol collector (SJAC). Liquid sample from the WRD and SJAC is continuously drawn from the sample boxes down the tower to the analytical box for analysis by ion chromatography (IC) on an hourly basis at the detector unit located in a climate-controlled enclosure at the base of the tower. At the beginning and end of the measurement intensive, multi-level liquid NO$_3^-$ and NH$_4^+$ standards were introduced at the WRD and SJAC, with airflow turned off, to assess the analytical accuracy of the NH$_3$ and HNO$_3$ measurement. MARGA measurements were conducted during the spring and summer 2016 intensives. Comparisons of continuous and time-integrated methods for HNO$_3$ and NH$_3$ are summarized in Sect. S1 (Figs. S2 and S3).

Concentrations of NH$_3$, HNO$_3$, SO$_2$, NH$_4^+$, NO$_3^-$, and SO$_4^{2-}$ in air were measured concurrently on the EFT at 10 heights from just above the forest floor (0.5 m.a.g.) to several meters above the canopy (upper height of 37 m.a.g. during spring 2016, 43.5 m.a.g. during summer 2016) using a glass annular denuder and filter pack (URG Corporation, Chapel Hill, NC) system. The sampling assembly included a 1% Na$_2$CO$_3$-coated denuder for collection of acid gases followed by a 1% H$_3$PO$_4$-coated denuder for collection of NH$_3$, a filter pack containing a primary Teflon filter for collection of aerosol, and a backup Nylon filter (47 mm, Pall Corp, Port Washington, NY) to collect HNO$_3$ liberated by dissociation of NH$_4$NO$_3$ on the primary filter. Inlets were Teflon-coated glass impactors with a nominal 2.5 µm aerodynamic diameter cut point (URG Corporation, Chapel Hill, NC). Sample durations were typically 3 or 4 h at a flow rate of $\sim$ 16.7 L min$^{-1}$. Flow rates were controlled by critical orifice and were verified before and after each sampling period with a NIST-traceable primary standard flowmeter (Bios DryCal DC-Lite flowmeter, Mesa Laboratories, Inc., Lakewood, CO).

Denuders and filters were extracted with 10 mL of deionized water and analyzed by ion chromatography (IC, Dionex model ICS-2100, Thermo Scientific, Waltham, MA). Extracts were analyzed for cations using Dionex IonPac 2 mm CG12 guard and CS12 analytical columns; separations were conducted using 20 mM methanesulfonic acid (MSA) as eluent at a flow rate of 0.25 mL min$^{-1}$. Anions were analyzed (IonPac 2 mm AG23 guard column, AS23 analytical columns) using an isocratic eluent mix of carbonate–bicarbonate (4.5:0.8 mM) at a flow rate of 0.25 mL min$^{-1}$. Multi-point (≥ 5) calibrations were conducted using a mixture prepared from individual inorganic standards (Inorganic Ventures, Christiansburg, VA). A midlevel accuracy check standard was prepared from certified standards mix (AccuStandard, New Haven, CT) for quality assurance and quality control. Profile measurements were conducted during each of the five SANDS intensives.

Bulk organic nitrogen in aerosols was measured using a high-volume (Hi-Vol) Tisch TE-1000 (Tisch Environmental, Cleves, OH) dual cyclone PM$_{2.5}$ sampler operated at a flow rate of 230 L min$^{-1}$. The unit was deployed at ground level adjacent to the COW137 CASTNET tower and collected 24 h (started at 07:00 local time) integrated samples on pre-baked (550 $^\circ$C for 12 h) quartz-fiber (QF) filters (90 mm, Pall Corp, Port Washington, NY). Field blanks were collected the same.
way except being loaded in the sampler without the pump switched on. A QF punch (1.5 cm²) from each sample was extracted with deionized water (18.2 MΩ cm, Milli-Q reference system, Millipore, Burlington, MA) in an ultrasonic bath for 45 min. The sample extract was filtered through a 0.2 µm pore size polytetrafluoroethylene membrane syringe filter (Iso-disc, Sigma Aldrich, St. Louis, MO) before subsequent analyses.

Water-soluble total N (WSTN) concentrations were measured using a high-temperature catalytic combustion and chemiluminescence method that included a total organic carbon analyzer (TOC-VCSH) combined with a total nitrogen module (TNM-1) (Shimadzu Scientific Instruments, Columbia, MD). Briefly, the TN module converts all nitrogen compounds to NO at 720 °C in a combustion chamber, and NO is quantified by NO₂ chemiluminescence through reaction with ozone. A five-point calibration was conducted with KNO₃ standard solution for each batch of samples. Before and after each batch of samples was analyzed, quality assurance checks and analyses, including lab deionized and accuracy check standards, were conducted to ensure accuracy and precision. Inorganic species (NH₄⁺, NO₃⁻, NO₂⁻) were analyzed by IC as described above, and WSON was calculated according to Eq. (1). Comparisons of Hi-Vol and CASTNET PM measurements are summarized in Sect. S1 (Fig. S4).

To evaluate the spatial distribution of gaseous N across the Coweeta Basin, additional passive sampling of HNO₃ and NH₃ was conducted across an elevation gradient for the full year of 2015 (Fig. 1, Tables 1 and 2). Samplers were deployed for 2-week periods at the height of 10 m a.g. on an aluminum tilt-down tower. NH₃ measurements followed AMoN methods. HNO₃ was collected on 47 mm Nylon filters (Nylasorb, Pall Corp, Port Washington, NY) as described by Bytnerowicz et al. (2005). NH₃ and HNO₃ sampler preparation and analysis was performed by the NADP Central Analytical Laboratory and CASTNET laboratories, respectively. Field calibration of the passive HNO₃ measurements (Fig. S5) was based on comparison with a collocated CASTNET sampler at the Screwdriver Knob site (Fig. 1, Tables 1 and 2), which also operated for the full year of 2015.

2.2.3 Micrometeorology

Site characteristics of micrometeorology and ecosystem fluxes of water and carbon dioxide have been previously described (Novick et al., 2013, 2014; Oishi et al., 2018). Three-dimensional wind components were measured by a sonic anemometer (model 81000, R.M. Young Company, Traverse City, MI) above the forest canopy on the EFT. Momentum and kinematic heat fluxes were determined by eddy covariance (EC) from sonic data. For EC calculations, raw 10 Hz sonic data were processed into hourly averages after block average detrending and 2D coordinate rotation (Novick et al., 2013). Air temperature and relative humidity (RH) were measured at the top of the tower (EC155, Campbell Scientific, Logan, UT) and 2/3 canopy height (HMP-45, Vaisala, Helsinki, Finland). Photosynthetically active radiation (PAR; LI-190, LI-COR Biosciences, Lincoln, NE) as well as upward and downward shortwave and longwave radiation (CNR 4, Kipp & Zonen, Delft, the Netherlands) were measured at the top of the tower. Surface wetness was measured in the canopy crown, in the understory, and at the ground using leaf wetness sensors (model 237, Campbell Scientific). Soil volumetric water content (VWC) averaged over 0–30 cm depth was measured in four locations around the tower using water content reflectometers (CS616, Campbell Scientific). Soil temperature was measured at four depths (5, 20, 35, and 55 cm) in two locations near the tower using thermistors. For missing data, linear interpolation was used to fill short gaps (1–4 h). Longer gaps were filled by substitution using the average hourly diel profile calculated for each month. Micrometeorological data were used for inferential dry deposition modeling as described below.

2.2.4 Biogeochemistry

Ammonia emission potentials (Γ_) and compensation points for live vegetation, leaf litter on the forest floor, and soil were estimated from measurements of NH₄⁺ and pH in the leaf and litter tissue and soil pore water (Massad et al., 2010). Green leaves were collected from 18 species (Table S1) within the flux footprint of the tower and other locations in the Coweeta Basin. Leaf litter was collected along transects extending ~100 m to the northeast and southwest (i.e., predominant wind directions) of the flux tower. Litter included a composite of intact leaves and leaf fragments and excluded the more decomposed material at the top of the organic soil layer. Approximately 5 g of leaf tissue was ground in liquid nitrogen using a mortar and pestle and small coffee grinder, then extracted with 20 mL of deionized water. pH was determined directly on the extracts (Oakton pH 2100 m, Mettler Toledo InLab Micro electrode). The [NH₄⁺] in the extracts, which reflects the bulk tissue concentration, was determined by ion chromatography as described above for denuder measurements either directly or, for samples with high organic content, after separation of the NH₄⁺ from the solution as NH₃ using headspace equilibration. For the headspace method, 5 mL of tissue extract was added to a 250 mL high-density polyethylene jar containing two ALPHA passive samplers (Center for Ecology and Hydrology; Tang et al., 2001), without the diffusion barrier, affixed to the interior of the lid. The jar was sealed, and 5 mL of 0.3 N NaOH was added to the extract via a septum. The NH₃ liberated from the liquid extract into the headspace was collected by the passive diffusion samplers over a period of 48 h, after which the passive sampler was extracted with 10 mL of deionized water. Extracts were then analyzed by ion chromatography as described above.

Emission potentials of the vegetation (Γ_E) and litter (Γ_L) were estimated from measured concentrations of [H⁺] (M)
and [NH$_4^+$] (µg g$^{-1}$ tissue fresh weight) in the bulk tissue as

$$\Gamma_{s,1} = \frac{[\text{NH}_4^+]}{[\text{H}^+]} \times (5.56 \times 10^{-5}) \times LD,$$

(2)

where LD is leaf density (kg L$^{-1}$ fresh tissue, equivalent to g cm$^{-3}$ fresh tissue). In this case, LD values for woody deciduous and woody evergreen species are 0.37 and 0.42 kg L$^{-1}$, respectively (Poorter et al., 2009). Emission potentials for litter, which consisted of a mix of intact or partial leaves and needles, assume an average density of 0.4 kg L$^{-1}$. The factor of 5.56 $\times$ 10$^{-5}$ in Eq. (2) is necessary to convert [NH$_4^+$] from µg NH$_4^+$ g$^{-1}$ tissue to mol NH$_4^+$ kg$^{-1}$ tissue.

Soil chemistry was measured in 20 m $\times$ 20 m plots located in the vicinity of the tower. During 2010, soil NH$_4^+$ was determined on soil samples collected with polyvinyl chloride cores (5 cm diameter and 10 cm deep) in four locations (replicates) within each of four plots. Samples were collected bimonthly during the growing season. NH$_4^+$ was extracted within 2 h of collection using 5 g of sieved (< 5 mm) soil in 20 mL of 2 M potassium chloride followed by colorimetric analysis (Astoria2 autoanalyzer, Astoria-Pacific International; Coweeta Hydrologic Laboratory, 2016). Soil pH was measured on 0–10 cm samples collected in three plots in winter of 2013 (each sample representing a composite of 20–25 2.5 cm diameter soil cores). Soil pH was determined directly after mixing 5 g soil with 10 mL 0.01 M calcium chloride (Coweeta Hydrologic Laboratory, 2016). Soil emission potential (\(\Gamma_{\text{soil}}\)) (unitless) was estimated directly from measured molar concentrations of [H$^+$] and [NH$_4^+$] as

$$\Gamma_{\text{soil}} = \frac{[\text{NH}_4^+]}{[\text{H}^+]}.$$  

(3)

### 2.2.5 Above-canopy flux measurements

Above-canopy fluxes of NH$_3$ and HNO$_3$ were quantified from measurements of vertical concentration gradients conducted during the final (summer 2016) intensive when the tower was at maximum height and the greatest vertical separation of concentration measurements was achieved. Fluxes were determined using the modified Bowen ratio (MBR) method (Meyers et al., 1996) as

$$F = -K_c(z) \frac{dC}{dz},$$

(4)

where $K_c$ and $dC/dz$ are the eddy diffusivity and vertical concentration gradient of the chemical species of interest. The value of $K_c$ for trace gases was assumed to be equivalent to the eddy diffusivity of heat ($K_t$), calculated as

$$K_c = K_t = -\bar{u} T \frac{\Delta z}{\Delta t},$$

(5)

where $\bar{u} T$ is the kinematic surface heat flux measured by eddy covariance above the canopy (43.5 m a.g.l.), $\Delta t$ is the air temperature difference between the levels at 43 and 35 m a.g.l., and $\Delta z$ is the height interval of air temperature measurements. Air temperature was measured using aspirated thermocouples, and $\Delta t$ was corrected for a small bias between the sensors determined by collocated comparison. Concentration gradients were determined from URG annular denuder measurements at 34.6 and 43 m a.g.l., as described above in Sect. 2.2.2. Given the complexity of the topography, no attempt was made to correct for potential roughness sublayer effects on either the eddy diffusivity or concentration gradients, which should be acknowledged as a source of uncertainty in the calculated HNO$_3$ and NH$_3$ fluxes. Additional detail on the gradient measurements is included in Sect. S3.

### 2.2.6 Seasonal and annual deposition budget

Speciated seasonal and annual total nitrogen deposition budgets were developed for the period August 2015 to August 2016. The wet-deposited components, including NH$_4^+$, NO$_3^-$, and total WSON, were directly measured. Speciated dry deposition was estimated by combining measured air concentrations, micrometeorology, biogeochemistry, and canopy physical characteristics within a box version of the Surface Tiled Aerosol and Gaseous Exchange (STAGE) model, which is an option in the Community Multiscale Air Quality Model (CMAQ) version 5.3 (Appel et al., 2021).

The STAGE model treats the deposition of gases and particles separately. The air–surface exchange of gases is parameterized as a gradient process and is used for both bidirectional exchange and dry deposition following the widely used resistance model of Nemitz et al. (2001) and Massad et al. (2010):

$$F = -f_{\text{veg}} \frac{\chi_a(z) - \chi_{0g}}{R_a} - (1 - f_{\text{veg}}) \frac{\chi_a(z) - \chi_g}{R_a + R_g},$$

(6)

where $F$ is the net flux above the canopy (a negative value represents a net deposition flux and a positive value represents a net emission flux), which is the sum of the component cuticular ($F_{\text{cut}}$), stomatal ($F_s$), and ground ($F_g$) fluxes. The quantity $\chi_a(z)$ is the ambient concentration at a reference height ($z$), $\chi_{0g}$ is the concentration at height $d$ (displacement height) + $z_0$ (roughness length), $\chi_g$ is the ground layer compensation point, $R_a$ is the aerodynamic resistance between $z$ and $d + z_0$, $R_g$ is the total ground resistance including in-canopy aerodynamic resistance ($R_{\text{inc}}$), ground boundary layer resistance ($R_{\text{bg}}$), and soil resistance ($R_{\text{soil}}$ ($R_g = R_{\text{inc}} + R_{\text{bg}} + R_{\text{soil}}$)), and $f_{\text{veg}}$ is the vegetation coverage fraction. The ground is fully covered by vegetation at our forested site, and $f_{\text{veg}}$ is therefore set to 1.

The quantity $\chi_{0g}$ is related to the canopy ($\chi_c$) and ground compensation points ($\chi_g$) according to

$$\chi_{0g} = \frac{\chi_c(z) + \chi_g}{\frac{1}{R_a} + \frac{1}{R_{\text{inc}}} + \frac{1}{R_{\text{soil}}}},$$

(7)
where $R_{bd}$ is the leaf boundary layer resistance. $\chi_c$ follows Nemitz et al. (2001) but is modified to account for a cuticular compensation point ($\chi_c$):

$$\chi_c = \frac{161512}{T} 10^{-4.507.11} \Gamma_{s, cut, g}.$$

(9)

$\Gamma_{cut}$ is set to 0 in this study, and thus there is only deposition to leaf cuticles. For unidirectional exchange of gases other than NH$_3$, $\Gamma_s$ and $\Gamma_g$ are also set to 0. In the case of NH$_3$, the foliage and ground layers may act as a source or sink depending on the ratio of the ambient concentration to the respective compartment compensation point (Husted and Schjoerring, 1995). Here values of $\Gamma$ for NH$_3$ are derived from measurements of live vegetation, litter, and soil chemistry as described above. Values used in the base model simulation are described in Sect. 3.6, and the sensitivity of modeled NH$_3$ fluxes to $\Gamma$ is discussed in Sect. 5.5.

Formulas for each resistance component are summarized in Table S2. The resistances are largely estimated following Massad et al. (2010) with the following exceptions. The value of $R_s$ is based on the Noah (Chen and Dudhia, 2001) or P-X land surface scheme (Pleim and Xiu, 1995) in the Weather Research and Forecasting (WRF) model, and in this study, the P-X scheme is used. Deposition to wetted cuticular and ground surfaces considers the bulk accommodation coefficient, following Fahey et al. (2017), and can be a limiting factor for highly soluble compounds. The parameterization of $R_{inc}$ follows Shuttleworth and Wallace (1985) as do Massad et al. (2010) but here uses the canopy momentum attenuation parameterization from Yi (2008) and in-canopy eddy diffusivity following Harman and Finnigan (2007). This parameterization is similar to Bash et al. (2010), and detailed descriptions of model resistances can be found in the references mentioned above.

Dry deposition ($F$) of aerosol nitrogen (NH$_4^+$ and NO$_3^-$) is estimated as the product of the measured concentration ($C$) and the STAGE-modeled dry deposition velocity ($V_d$):

$$F = -V_d(z) \times C(z).$$

(10)

Aerosol dry deposition processes include gravitational settling, Brownian diffusion, surface impaction, and rebound. Similar to gases, STAGE calculates the averaged $V_d$ for particles by summing the $V_d$ over vegetative or non-vegetative surfaces, weighted by vegetation cover fraction, which is 1 (full coverage) at Coweeta. $V_d$ for a particle with aerodynamic diameter $D_p$ is calculated as

$$V_d(D_p) = \frac{V_g}{1 - \exp[-V_g (R_a + R_{bp})]},$$

(11)

where $R_{bp}$ is the boundary layer resistance for particles, and the gravitational settling velocity ($V_g$) is calculated as

$$V_g = \rho D_p^2 g \frac{C}{18\mu},$$

(12)

where $\rho$ is the density of the aerosol, $g$ is the acceleration of gravity, $\mu$ is the air dynamic viscosity, and $C$ is the Cunningham slip correction factor. The turbulent transport processes are considered to be similar for gas and aerosol, and $R_a$ can be formulated based on the similarity theory relationships. Unlike deposition of gases, the boundary layer resistance usually dominates the aerosol deposition process as Brownian diffusion is much slower than molecular diffusion is for gases (Pleim and Ran, 2011). Thus, $R_{bp}$ depends on the collection efficiency of the surface and can be determined following Shu et al. (2022):

$$R_{bp} = F_t u_s \left( Sc^{-\frac{2}{3}} + E_{im} \right)^{-1},$$

(13)

where $u_s$ is the friction velocity, and $Sc$ is the Schmidt number for particles. The quantity $E_{im}$ represents the collection efficiency by impaction and follows Slinn (1982) for vegetative canopies and Giorgi (1986) for smooth (non-vegetative) surfaces. The quantity $F_t$ is an empirical correction factor to account for increased deposition during convective conditions, parameterized as

$$F_t = V_{fac} \left( 1 + 0.24 \frac{w^2}{u_s^2} \right),$$

(14)

where $V_{fac}$ is an empirical constant representing the enhanced effects over vegetation canopies. For vegetative canopies, $V_{fac}$ is equal to the one-sided leaf area index (LAI) with a minimum value of 1, and for non-vegetative surface, a value of 1 is used. The quantity $w_s$ is the convective velocity scale (Deardorff velocity), defined as

$$w_s = \left( \frac{g}{T_v} \frac{z_i}{w^*/T^*} \right)^{\frac{1}{2}},$$

(15)

where $T_v$ is virtual air temperature, $z_i$ is average depth of the mixed layer, and $w^*/T^*$ is the measured kinematic surface heat flux.

A bulk $V_d$ for PM$_{2.5}$ is obtained by integrating size-resolved $V_d$ according to the particle size distribution. The
size distribution profiles for NH$_4^+$ and NO$_3^-$ are from measurements at eight Canadian rural forest sites (Zhang et al., 2008), and the size distribution for particulate organic nitrogen is estimated as an average of that for NH$_4^+$ and NO$_3^-$. Model sensitivities of particle nitrogen fluxes to assumed size distributions are discussed in Sect. S5.

The STAGE model is extracted from CMAQ v5.3 and executed in a one-dimensional mode. The prescribed surface parameters (e.g., $z_0$, $d$) were modified according to the site conditions. Continuous LAI data were extracted from the Moderate resolution Imaging Spectroradiometer (MODIS) global LAI product (MCD15A2H), which is generated daily at a 500 m spatial resolution, and each data point covers an 8 d period. The MODIS LAI (Fig. S6) was adjusted using in situ canopy measurements as described in Sect. S4. Hourly meteorological measurements, including air temperature, relative humidity, $u_{*}$, atmospheric pressure, precipitation rate, global radiation, and soil temperature and moisture, were used to drive STAGE. The Obukhov length, which is defined as

$$L = -rac{u_{*}^3 T_o}{(kgw/m^3)},$$

where $k$ is the von Karman constant, was also calculated from micrometeorological measurements.

2.2.7 Air concentrations for dry deposition modeling

Air concentration data used for dry deposition modeling are summarized in Table 3. Hourly measurements of HNO$_3$ by DD_CL as well as ΣAN and ΣPN by TD-PC-CL were conducted for a full year and were therefore used directly for modeling. Over the 12-month sampling period, 18%, 22%, and 22% of hourly HNO$_3$, ΣAN, and ΣPN concentrations were missing or invalid, respectively. Missing data were replaced with the corresponding hour from the median diel profile comprised of days with > 75% completeness. Surrogate formulas of nitrooxy-butanol (C$_4$H$_8$NO$_4$) and PAN (C$_5$H$_7$NO$_3$) were assumed for ΣAN and ΣPN, respectively.

Continuous NH$_3$ concentrations were only measured during the last two intensives. Biweekly AMoN NH$_3$ measurements, with corrections for travel blanks, were used to establish a continuous 12-month time series of air concentration for annual deposition modeling. Ammonia concentrations are known to exhibit pronounced diel patterns, even in remote areas (Wentworth et al., 2016). Variability in air concentration interacts with diel cycles in surface wetness, turbulence, and other factors to influence diel patterns in air-surface exchange rates. To incorporate this interaction, the diel concentration pattern determined during spring and summer 2016 by MARGA NH$_3$ measurements (Fig. S7) was imposed on the biweekly AMoN NH$_3$ concentration. The hourly profile of NH$_3$ concentrations was normalized by the corresponding overall mean concentration to produce a normalized mean diel concentration profile. This profile was then applied to each biweekly AMoN air concentration, temporally scaling the NH$_3$ concentration by time of day while maintaining the measured biweekly AMoN concentration. Gap filling of AMoN data was not required. Comparisons of NH$_3$ measurements are briefly discussed in Sect. S1, Fig. S2.

Hi-Vol measurements of speciated particulate N were only conducted during intensive periods to assess the relative contributions of inorganic and organic fractions to total water-soluble N. The CASTNET particulate NH$_4^+$ and NO$_3^-$ were used to provide a continuous 12-month time series of air concentration for annual deposition modeling. Concentrations of Hi-Vol and CASTNET measurements were shown to be comparable (Sect. S1, Fig. S4). For the annual time series, particulate organic nitrogen (PON) concentration was estimated based on speciated measurements during intensives, which showed that inorganic N accounts for ~ 88% of WSTN on average. Weekly average PON concentration was estimated from the weekly CASTNET measurements assuming that NH$_4^+$ + NO$_3^-$ represents 88% of total particulate nitrogen and PON represents the balance (12%). Weekly concentrations were then expressed at the hourly timescale for modeling. Gap filling of CASTNET data was not required.

Components of the atmospheric reactive N budget that are not routinely measured at Coweeta and were not directly measured during SANDS include NO, NO$_2$, HONO, and N$_2$O$_5$. At the continental scale, regional model simulations suggest that NO, HONO, and N$_2$O$_5$ make minor contributions to the total dry deposition of reactive N (~ 2%), though the contribution of NO$_2$ is larger (~ 6%) (Walker et al., 2020). While NO, HONO, and N$_2$O$_5$ have been excluded from our modeling analysis, we have included an estimate of NO$_2$ concentration, from which dry deposition is estimated. The “other” fraction of NO$_y$ (i.e., NO$_x$-HNO$_3$-ΣPN-ΣAN) measured at Coweeta represents between 47% (summer) and 76% (winter) of total NO$_x$ on a seasonal basis. This “other” fraction includes NO, NO$_2$, HONO, N$_2$O$_5$, and some NO$_3^-$ but is likely dominated by NO$_2$. The measured diel profile of “other” NO$_y$ (Fig. S8) concentration shows the typical pattern indicative of morning and evening modes related to mobile NO$_x$ emissions. Winds at Coweeta are from the east-northeast during the morning, which is the direction of local residences, the town of Otto, NC, and US Highway 23. Winds are from the west-southwest during the evening, which is the direction of the Nantahala National Forest. Consistent with the diel profile of “other” NO$_y$, a much larger morning peak in NO$_2$ is therefore expected. To estimate the concentration of NO$_2$ from the measured “other” NO$_y$, we examined the ratio of NO$_2$ to the quantity NO$_x$-HNO$_3$-PANS-NTR (e.g., “other” NO$_y$) simulated by CMAQ (V5.2.1) for the Coweeta site over the year 2015: “PANS” represents total peroxy nitrates, and NTR represents other organic nitrates. Relative to the measured NO$_x$ species, PANS and NTR are assumed to represent ΣPN and ΣAN, respectively. The ratio of CMAQ-estimated NO$_2$ to “other” NO$_y$ ranges from 0.51 during summer to 0.60 during winter.

https://doi.org/10.5194/bg-20-971-2023

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These seasonal factors were applied to the measured “other” NO\textsubscript{x} to estimate the hourly NO\textsubscript{x} concentration. Gap filling procedures for hourly “other” NO\textsubscript{x} follow those for HNO\textsubscript{3}, ΣPN, and ΣAN described above. Details of CMAQ V5.2.1 can be found in Table S3.

Regarding the use of measurements from different towers (Table 3) for inferential modeling, we acknowledge that differences in tower position on the landscape (i.e., within the forest –EFT – versus adjacent clearing – COW137) and the height of the measurement above the surface will be sources of variability in air concentrations. Given the complexity of the topography, no attempt was made to correct air concentrations for differences in measurement heights.

3 Results and discussion

3.1 Long-term trends in atmospheric N at Coweeta

Emissions of oxidized nitrogen (NO\textsubscript{x}) and sulfur (SO\textsubscript{4}) have declined significantly in the eastern US in response to the 1990 Clean Air Act Amendments. Trend data from the US EPA's National Emissions Inventory (NEI) indicate a nationwide decline of 74% and 46% for SO\textsubscript{4} and NO\textsubscript{x} emissions from the early 1990s to 2010s, respectively, comparing 1990–1994 to 2010–2014 annual averages (U.S. EPA, 2014). By contrast, NH\textsubscript{3} emissions have been reported as relatively unchanged or slightly increasing for the same periods (Ellis et al., 2013; Paulot and Jacob, 2014; Xing et al., 2013), depending on the location and region of the US. Declining NO\textsubscript{x} and SO\textsubscript{4} emissions resulted in decreasing trends in air concentrations of HNO\textsubscript{3} and SO\textsubscript{4} at Coweeta between the 1990s and 2010s (Fig. 2). Concentrations noticeably began to decline in 2008, the timeline of which likely indicates the effect of full implementation of the 2006 Tier 2 Gasoline Sulfur Program, as well as the enactment of the Clean Air Interstate Rule (CAIR), both of which aimed to further reduce NO\textsubscript{x} and SO\textsubscript{4} emissions (Sickles and Shadwick, 2015; LaCount et al., 2021). Compared to other species, NH\textsubscript{3} concentrations have only been measured at Coweeta for a relatively short period of time.

Atmospheric NH\textsubscript{3} reacts with acidic sulfate to form ammonium sulfate ((NH\textsubscript{4})\textsubscript{2}SO\textsubscript{4}) or bisulfate ((NH\textsubscript{4})HSO\textsubscript{4}) aerosol. Under favorable thermodynamic conditions (low temperature and high RH), NH\textsubscript{3} in excess of acidic sulfate will react with HNO\textsubscript{3} to form ammonium nitrate aerosol (NH\textsubscript{4}NO\textsubscript{3}). Concentrations of SO\textsubscript{4}– at Coweeta have tracked SO\textsubscript{2}, and subsequently NH\textsubscript{4}+ concentrations have declined substantially relative to early 1990s levels (Fig. 2). However, concentrations of NO\textsubscript{3}– aerosol, which are relatively low at Coweeta, have not followed trends in SO\textsubscript{4}2– and NH\textsubscript{4}+ (Fig. 2). Previous studies at other US sites have also reported non-proportional changes in PM\textsubscript{2.5} mass in response to SO\textsubscript{2} and NO\textsubscript{x} control strategies (Blanchard and Hidy, 2005; Sickles and Shadwick, 2015). Nonlinear reductions or increases in particulate NO\textsubscript{3}– with coincident SO\textsubscript{2} and NO\textsubscript{x} emission reductions relate to the thermodynamic equilibrium of the SO\textsubscript{4}2––NO\textsubscript{3}––NH\textsubscript{4}+–HNO\textsubscript{3}–NH\textsubscript{3} aerosol system. As ambient SO\textsubscript{4}2– concentrations decline, the capacity for NH\textsubscript{4}+ formation (i.e., neutralization) also decreases, leaving additional NH\textsubscript{3} in the gas phase. Amounts of NH\textsubscript{3} in excess of acidic SO\textsubscript{4}2– can subsequently react with HNO\textsubscript{3} to form NH\textsubscript{4}NO\textsubscript{3}, confounding the relationship between NO\textsubscript{x} emission reductions and atmospheric NO\textsubscript{3}– concentrations.

The long-term trend in NO\textsubscript{3}– wet deposition at Coweeta (Fig. 3) has tracked the downward trend in ambient HNO\textsubscript{3} concentration. Wet deposition of NH\textsubscript{4}+, however, shows no
Figure 2. Long-term CASTNET (weekly, site COW137) and AMoN (biweekly, site NC25) air concentrations (as N) of NO$_3^-$ (a), HNO$_3$ (b), NH$_4^+$ (c), NH$_3$ (d), SO$_4^{2-}$ (e), and SO$_2$ (f).

apparent trend, in contrast to the decline in NH$_4^+$ aerosol concentration. This pattern may relate to the combined effects of changes in regional NH$_3$ emissions, aerosol chemistry, and dry deposition rates on the long-term trend in atmospheric NH$_3$ concentrations. As noted above, declines in SO$_2$ emissions and SO$_4^{2-}$ aerosol result in less conversion of NH$_3$ to NH$_4^+$ aerosol, leaving more NH$_3$ in the gas phase. Furthermore, reduced air concentrations of acidic species such as SO$_2$ and HNO$_3$ result in lower dry deposition rates and subsequently less acidic deposition surfaces, which in turn reduces the deposition velocity (i.e., increases the atmospheric lifetime) of NH$_3$ (Sutton et al., 2003). In addition to changes in emissions, these two processes are thought to be at least partly responsible for the increases in NH$_3$ air concentrations that have been observed in some locations across the US (Butler et al., 2016; Yu et al., 2018; Yao and Zhang, 2019). While there is no discernable trend in NH$_3$ air concentrations over the relatively short period of record at Coweeta, a decline in wet deposition of NH$_3$ aerosol may have been offset to some extent by increased wet deposition of NH$_3$ gas (Asman, 1995), which is highly soluble, resulting in an overall lack of trend in NH$_3^+$ wet deposition at Coweeta over time. Similar to other areas in the US (Li et al., 2016), the downward trend in NO$_3^-$ wet deposition has led to an increase in the relative contribution of reduced forms of N (i.e., NH$_x = NH_3 + NH_4^+$) to inorganic wet N deposition at Coweeta (NH$_4^+$ : NO$_3^-$, Fig. 3).

### 3.2 Wet deposition

Of the various forms of N in precipitation, ammonium was the most abundant inorganic species, contributing 47.0 % of WSTN in weekly samples ($N = 52$) on average, followed by NO$_3^-$ (41.7 %, Fig. 4, Table S4). The contribution of NO$_2^-$ was negligible. Organic compounds (WSON) contributed 11 % of WSTN on average, which is within the range of values (3 % to 33 %) reported for other locations in the US (Scudlark et al., 1998; Whitall and Paerl, 2001; Keene et al., 2002; Beem et al., 2010; Walker et al., 2012; Benedict et al., 2013). While concentrations of N compounds were generally higher during warm months, a seasonal pattern in the percent contribution of WSON to WSTN was not apparent. In a previous study at Coweeta (1994–1996), Knoepp et al. (2008) found that organic nitrogen contributed 21 % of total nitrogen in bulk (wet + dry) deposition samples. Differences between Knoepp et al. (2008) and SANDS results may be related to interannual variability or trends in rainfall composition over the intervening 2 decades (e.g., Fig. 3), differences in collection method (wet only versus bulk deposition), or analytical techniques used for total N analysis (persulfate–UV diges-
3.3 Air concentrations of oxidized N

The oxidized fraction of reactive nitrogen (NO\textsubscript{x}) comprises a mixture of gaseous and particulate inorganic (NO, NO\textsubscript{2}, N\textsubscript{2}O\textsubscript{5}, HONO, HNO\textsubscript{3}, NO\textsubscript{3}\textsuperscript{-}) as well as organic compounds. Owing to its large deposition velocity and typical atmospheric concentration, HNO\textsubscript{3} is the primary contributor to dry deposition of inorganic oxidized N (Walker et al., 2020). Much less is known about the dry deposition of oxidized organic nitrogen compounds (Walker et al., 2020). Peroxy nitrates (PNs) and alkyl and multifunctional nitrates (ANs) are formed during the photochemical oxidation of volatile organic compounds (VOCs) in the presence of NO\textsubscript{x} (NO\textsubscript{x} = NO + NO\textsubscript{2}). While PNs exist in the gas phase, ANs can exist in the gas or particle phase and can be the dominant chemical sink for NO\textsubscript{x} in high-BVOC (biogenic VOC), low-NO\textsubscript{x} environments (Farmer and Cohen, 2008; Browne and Cohen, 2012; Paulot et al., 2012; Browne et al., 2013). Unlike PNs, ANs can also form at night via nitrate radical-induced oxidation of VOCs. Further, PNs and ANs have been shown to contribute significantly to the total NO\textsubscript{x} budget in geographically diverse rural and forested environments (e.g., Trainer et al., 1993; Nouaime et al., 1998; Farmer et al., 2008; Browne et al., 2013; Toma et al., 2019). Flux measurements at Blodgett Forest, CA, showed that PN dry deposition contributed 4%–19% of total N deposition at the site (Wolfe et al., 2009). Chemical transport modeling with current representation of the atmospheric oxidized nitrogen system suggests that PNs and ANs together contribute ~7% of total N deposition and ~11% of dry N deposition at the US continental scale compared to ~21% and 34% for HNO\textsubscript{3} and ~6% and 9% for particulate NO\textsubscript{3}\textsuperscript{-} (Walker et al., 2020).

The annual average concentration of NO\textsubscript{x} was 1.00 ppb (0.55 µg N m\textsuperscript{-3}), with the highest seasonal average concentration in the winter (1.32 ppb, 0.75 µg N m\textsuperscript{-3}) and lowest in the summer (0.64 ppb, 0.34 µg N m\textsuperscript{-3}) (Fig. 5, Table S5). The nearest rural NO\textsubscript{x} monitoring site is 85 km to the northwest at Look Rock in the Great Smoky Mountains National Park, where the annual concentration was 1.5 ppb over the same period (NPS, 2020). Similar to Coweeta, NO\textsubscript{x} concentrations at Look Rock are typically lowest during summer and highest in winter, though the seasonal cycle exhibits some interannual variability. The annual mean concentrations of HNO\textsubscript{3}, XPN, and ΣAN determined by TD-PC-CL were 0.14 (0.08), 0.1 (0.06), and 0.09 (0.05) ppb (µg N m\textsuperscript{-3}), respec-
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Reference

Figure 5. Seasonal and annual percent contribution of HNO$_3$, ΣPN, ΣAN, and other compounds to total NO$_x$ (a); seasonal and annual box plots of NO$_3^-$, HNO$_3$, ΣPN, and ΣAN – solid and dashed lines inside the box represent the median and mean, respectively; the top and bottom of the box represent 75th and 25th percentiles; whiskers represent 10th and 90th percentiles, and dots represent 5th and 95th percentiles. (b) Diurnal profiles of HNO$_3$, ΣPN, and ΣAN – observations represent median hourly concentration, and bars represent interquartile range (IQR). (c) “Other NO$_x$” is calculated as NO$_3^-$+HNO$_3$–ΣPN–ΣAN, which, while primarily comprised of NO$_x$, includes N$_2$O$_5$, HONO, NO$_2^-$, and possibly other organics.

4.4 Air concentrations of reduced N

Reduced forms of nitrogen (NH$_4^+$) represent another important component of the inorganic dry N deposition budget. At the continental scale, NH$_3$ dry deposition contributes $\sim 20 \%$ of total N deposition and $\sim 32 \%$ of dry N deposition, whereas the contributions from NH$_4^+$ aerosol are $\sim 4 \%$ and $\sim 6 \%$, respectively (Walker et al., 2020). Similar to oxidized forms of N, the partitioning of mass between the gas (NH$_3$) and particulate (NH$_4^+$) phases affects the dry deposition rate of NH$_2^+$, given the larger deposition velocity of NH$_3$ relative to NH$_4^+$.

During 2015–2020, with 2020 being the most recent full year of AMoN data, average concentrations of NH$_3$ and NH$_4^+$ were similar (Fig. 7). Both species displayed a seasonal pattern of lowest concentrations in the winter and higher concentrations during warm months. NH$_3$ concentrations peaked in summer, reflecting the temperature dependence of reactive nitrogen deposition.
Regional agricultural and biogenic emissions. NH$_4^+$ concentrations followed the seasonal cycle in SO$_4^{2-}$ concentrations, which were also similar in spring and summer and minimum in winter at Coweeta. The seasonal cycle of NH$_3$–NH$_4^+$ partitioning was driven more by NH$_3$ than NH$_4^+$, with the former exhibiting more seasonal variability. Hourly measurements conducted during spring and summer 2016 showed that NH$_3$ also displayed significant diel variability (Fig. S7), reaching a maximum around midday and a minimum overnight. Seasonal mean concentrations during the SANDS period fall within the IQR of the 6-year period between 2015 and 2020, with SANDS annual and 6-year averages being very similar. Concentrations of NH$_4^+$ were higher relative to NTO$_3$ during SANDS and over the longer term at Coweeta (Figs. 6 and 7).

### 3.5 Aerosol N composition

Ammonium was the most abundant inorganic species, contributing 86.8 % of WSTN (N = 103) on average (Fig. 8, Table S7). Organic compounds (WSON) contributed 11.5 % of WSTN, which is very similar to precipitation. The contributions of NO$_3^-$ and NO$_2^-$ were minor. Our study-wide average of percent WSON is slightly lower than measurements at other North American forest sites, including Duke Forest, North Carolina (33 %, Lin et al., 2010), and Rocky Mountain National Park (14 %–21 %) (Benedict, 2012), but is within the global range of 10 %–39 % (Cape et al., 2011). Similar to precipitation chemistry, there was no seasonal pattern in the percent contribution of WSON to WSTN in PM$_{2.5}$. Hi-Vol measurements of inorganic PM components compared well overall with collocated CASTNET measurements (Sect. S1, Fig. S4).

### 3.6 Biogeochemistry

Estimates of NH$_3$ emission potentials ($\Gamma$) for the ground and vegetation are needed to calculate compensation points ($\chi$) and fluxes in STAGE. Measurements of pH, NH$_4^+$, and corresponding $\Gamma$ of the leaves ($\Gamma_s$) and litter ($\Gamma_l$) are summarized in Fig. 9 and Tables S8 and S9. Measurements of $\Gamma_s$ are divided into green leaves collected during the growing season (spring and summer) and senescent leaves collected in October. NH$_3$ emission potentials ($\Gamma$) for green leaves ($\Gamma_s$) ranged from zero to 4070 with a median value (35.8) (Table S8) corresponding to a compensation point of $\chi_s = 0.25 \mu g \text{NH}_3 \text{m}^{-3}$ at 25 $^\circ$C. Large intra-species variability of tissue pH and NH$_4^+$ was observed (Table S9), and separating by crown versus understory species did not reveal distinct differences between groups. Given the variability of the observations, the median $\Gamma_s$ was used for STAGE simulations. Senescence marks the translocation of N in leaves to storage tissues (Schneider et al., 1996). Along with a decline in photosynthetic activity, degradation of chlorophyll, and other metabolic changes, glutamine synthetase (GS) activity also declines (Pearson et al., 2002). Glutamine synthetase catalyzes assimilation of NH$_4^+$ into glutamine and is therefore important in regulating the pool of NH$_4^+$ available for exchange as NH$_3$ between the leaf and atmosphere as well as remobilizing organic N for storage during senescence. A decline in GS activity can thus result in increased leaf NH$_4^+$ concentrations (Pearson et al., 2002; Wang et al., 2011). Senescent leaves were similar to green leaves with respect to median tissue pH but showed higher concentrations of tissue NH$_4^+$ (Mattsson et al., 2002; Mattsson and Schjoerring, 2002). Absolute differences between $\Gamma_s$ were correspondingly higher (113), equivalent to $\chi_s = 0.8 \mu g \text{NH}_3 \text{m}^{-3}$ at 25 $^\circ$C. For STAGE modeling, the median $\Gamma$ for senescent leaves was used for $\Gamma_s$ during the fall.

Regardless our method for estimating $\Gamma_s$, the fundamental assumption is that the chemistry of the bulk leaf tissue is representative of the apoplast. While a number of studies have shown positive correlations between bulk tissue chemistry, apoplastic chemistry, and independently quantified compensation points (David et al., 2009; Hill et al., 2002; Massad et al., 2010; Mattsson and Schjoerring, 2002; Mattsson et al., 2009), absolute differences between $\Gamma_s$ derived from bulk tissue versus apoplast measurements can be large. For example, Sutton et al. (2009) and Personne et al. (2015) both show that ratios derived from bulk tissue chemistry exceed those derived from apoplast chemistry. We did not perform experiments to validate the use of bulk tissue chemistry as a proxy for apoplast chemistry and acknowledge this source of uncertainty. However, our estimates of $\Gamma_s$ appear reasonable in the context of the range of existing observations (cited above).
Summary of AMoN NH$_3$ and CASTNET NH$_4^+$ concentrations (as N) from 2015–2020 during winter, spring, summer, and fall. Solid and dashed lines inside the box represent the median and mean, respectively. The top and bottom of the box represent 75th and 25th percentiles. Whiskers represent 10th and 90th percentiles, and dots represent 5th and 95th percentiles. “6-year” represents the statistics for the entire 6-year period. Squares represent the seasonal and annual mean concentration for the August 2015–August 2016 modeling period. Pie charts represent the average percent contribution of NH$_3$ and NH$_4^+$ to total NH$_x$ (NH$_3$ + NH$_4^+$) expressed as nitrogen. AMoN concentrations were adjusted by subtracting the mean travel blank for the 6-year period.

Contributions of N aerosol species to WSTN in 24 h Hi-Vol PM$_{2.5}$ samples during seasonal SANDS intensives (a) along with the percentage contribution of WSON to WSTN (b).

Box plots of pH (a), NH$_4^+$ (b) concentration (µg g$^{-1}$ fresh tissue), and equivalent emission potential ($\Gamma'$) (c) in tissue of green leaves, senescent leaves, and litter on the forest floor. The solid line inside the box represents the median. The top and bottom of the box represent 75th and 25th percentiles. Whiskers represent 10th and 90th percentiles, and dots represent 5th and 95th percentiles.

Leaf litter on the soil surface has been shown to be a source of NH$_3$ to the atmosphere in both natural and agricultural ecosystems (Nemitz et al., 2000b; David et al., 2009; Hansen et al., 2013). As litter decomposes, mineralization of organic N is a source of NH$_4^+$, some of which is lost to the overlying air as NH$_3$. Litter NH$_4^+$ concentrations were similar to green leaves but lower than senescent leaves (Fig. 9). However, the pH was higher than both green and senescent leaves. The resulting median $\Gamma'$ (69.3) was larger than green leaves but smaller than senescent leaves, equivalent to $\chi_l = 0.49$ µg NH$_3$ m$^{-3}$ at 25 °C. Litter $\Gamma'$ was much larger and summarized by Massad et al., 2010) and the general relationship between [NH$_4^+$]$_{\text{bulk}}$ and $\Gamma'$, put forth by Massad et al. (2010, Eq. 6). Measurements on bulk tissue are less labor-intensive and therefore more tempting than measurements on the apoplast. However, more studies comparing the two procedures are needed to extend the meta-analysis of Massad et al. (2010) to a wider range of natural ecosystems, particularly deciduous forests.
than that of the underlying soil. Average (0–10 cm soil depth) soil pH (4.18) and NH$_4^+$ (1.21 mg N kg$^{-1}$) correspond to $\Gamma_{\text{soil}} \sim 10$ at a soil mass wetness of 0.1 g g$^{-1}$, equivalent to $X_{\text{soil}} \sim 0.1$ µg NH$_3$ m$^{-3}$ at 25 °C. This very low $\Gamma_{\text{soil}}$ results from the low pH of the shallow soil.

Vertical profiles of air concentrations within and above the canopy were measured to investigate patterns of air–surface exchange with specific ecosystem compartments (i.e., canopy crown, understory, and ground). A detailed analysis of bidirectional N fluxes is forthcoming; thus, we limit the discussion of these data to NH$_3$ in the context of interpreting patterns observed in the biogeochemical emission potentials and their prescription in the STAGE model. Nitric acid, NH$_4^+$, and NO$_3^-$ showed expected decreasing concentrations from above the canopy to the forest floor, indicative of deposition. While NH$_3$ profiles showed patterns of deposition to the crown and understory, concentrations near the forest floor indicated both emissions and deposition (Fig. 10). Of the 76 daytime profiles measured, 40% showed decreases down to the forest floor, and 60% showed an increasing concentration from approximately the lower understory (∼ 5 m a.g.) to the forest floor. The former pattern is interpreted as deposition to the forest floor, and the latter is interpreted as emission. Thus, the mean profile suggests a source of NH$_3$ at the ground. The very low $\Gamma_{\text{soil}}$ suggests that emission from the soil is unlikely given such a low pH. The leaf litter layer, which indicates a much higher emission potential ($\Gamma_1$) than the soil, is a more likely source of NH$_3$. This hypothesis is consistent with Hansen et al. (2013, 2017), in which emissions of NH$_3$ from a beech (Fagus sylvatica) forest after leaf fall were attributed to the decomposition of new litter. Similar to our site, the underlying soil also had low pH (4–5). Given our observations, we used $\Gamma_1$ (median = 69.3, Table S8) rather than $\Gamma_{\text{soil}}$ as the ground emission potential ($\Gamma_\text{g}$) in STAGE.

Figure 10. Vertical concentration profiles of NH$_3$. The mean (filled circle) and standard deviation (bars) of concentrations are shown for $N = 76$ daytime profiles.

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3.7 N deposition budget

Total annual N deposition for the period August 2015–August 2016 was 6.7 kg N ha$^{-1}$, which is broken down by percent contribution of individual species to wet and dry deposition in Fig. 11. Over this period, wet deposition contributed 60.5% of total N deposition, of which NH$_4^+$ was the primary component (29.5%). Wet deposition of organic N contributed 5.4% of the total N deposition budget. Dry deposition accounted for 39.5% of total deposition, of which NH$_3$ was the primary contributor (20.5%). Reduced forms of inorganic N were the largest contributor to the budget (51.1%), with oxidized inorganic and organic N contributing 41.3% and 7.6% of total N deposition, respectively. Dry deposition of organic N made a small contribution (2.2%) to the total deposition budget.
Ammonia is the most important contributor to the dry deposition budget (52.1 %) and differs from the other species in that it is exchanged bidirectionally between the ground, canopy, and atmosphere.

Seasonal net canopy-scale and component fluxes are shown in Fig. S9. The mean net flux \( (F_{\text{net}}) \) is downward (i.e., deposition) during all seasons, generally following the seasonal pattern of the atmospheric NH\(_3\) concentration. The cuticular flux \( (F_{\text{cut}}) \), which is unidirectional in STAGE, is the dominant deposition pathway and ranged from \(-97.7 \text{ ng N m}^{-2} \text{ s}^{-1}\) (deposition) to near zero. The contribution of \( F_{\text{cut}} \) to the total net flux ranged from 84.7 % in spring to \( \sim 100 \% \) during fall. The stomatal flux \( (F_{\text{s}}) \) is bidirectional, ranging from \(-4.5 \text{ ng N m}^{-2} \text{ s}^{-1}\) (deposition) to 2.3 ng N m\(^{-2}\) s\(^{-1}\) (emission), with the largest fluxes occurring during warmer periods of the growing season when the stomatal resistance is lowest. The stomatal flux is smaller than \( F_{\text{cut}} \) for several reasons. First, \( R_{\text{s}} \) is generally larger than \( R_{\text{cut}} \). Also, in the current model formulation the cuticular compensation is \( X_{\text{cut}} = 0 \). Thus, the NH\(_3\) concentration gradient between air above the leaf (\( X_{\text{leaf}} \)) and \( X_{\text{s}} \) is smaller than for \( X_{\text{cut}} \). Finally, low LAI and large \( R_{\text{s}} \) in winter and fall as well as offsetting bidirectional fluxes in spring and summer result in a relatively small mean stomatal deposition flux \( (F_{\text{s}}) \) across seasons. \( F_{\text{g}} \) is also bidirectional, ranging from \(-3.9 \text{ ng N m}^{-2} \text{ s}^{-1}\) (deposition) to 2.5 ng N m\(^{-2}\) s\(^{-1}\) (emission). Fluxes are largest during spring as atmospheric NH\(_3\) begins to increase with warmer temperatures but before peak LAI is reached, after which the denser canopy increases the in-canopy aerodynamic \( (R_{\text{inc}}) \) and air-side ground boundary layer resistances \( (R_{\text{bg}}) \) (Table S2), thereby decreasing \( F_{\text{g}} \). On an annual scale, \( F_{\text{g}} \) and \( F_{\text{s}} \) make similar contributions to \( \sim 30 \% \) \( F_{\text{net}} \).

Nitric acid was the second-largest component of dry deposition, contributing 36.0 % of the total. While HNO\(_3\) deposits more rapidly than NH\(_3\) (Table S10), the overall importance to the dry N budget is constrained by relatively low air concentrations at this remote forest site. Particulate species made much smaller contributions to the budget due to much lower deposition velocities \( (V_{\text{d}} = \text{flux} \times \text{air concentration}) \) relative to their gaseous counterparts (Table S10). For example, while NH\(_4^+\) and NH\(_3\) contributed approximately equally to the atmospheric NH\(_3\) load (Fig. 7), the NH\(_4^+\) flux budget was regulated by the much more rapid exchange of NH\(_3\) between the forest and atmosphere relative to NH\(_4^+\). A similar example was observed for oxidized N. While NO\(_2\) represents an important fraction of the oxidized N concentration budget via its contribution to “other NO\(_x\)”, NO\(_2\) deposits much less rapidly than HNO\(_3\) (Table S10), thereby contributing a relatively small fraction (3.4 %) of the dry N flux. Of the organic N species, AN contributed slightly more (2.9 %) to dry N deposition than PN (2.2 %) owing to a higher deposition velocity (Table S10). Similar to particulate NH\(_4^+\) and NO\(_3^-\), PON made a small contribution to dry N deposition (0.4 %) due to its low \( V_{\text{d}} \) (Table S10). Reduced forms of N accounted for the majority of dry N deposition (54.9 %), with oxidized inorganic and organic forms of N contributing 39.6 % and 5.5 %, respectively.

Total N deposition peaked during the summer (2.5 kg N ha\(^{-1}\)) and reached a minimum in the fall (1.0 kg N ha\(^{-1}\)) (Fig. 12). Wet deposition exceeded dry deposition during all seasons. Seasonal variability in wet deposition was primarily driven by precipitation amount, whereas dry deposition was influenced by seasonality in air concentrations of the primary N\(_f\) species (Figs. 6 and 7), LAI (Fig. S6), turbulence, and other surface characteristics. Ammonia fluxes followed the seasonal pattern of air concentration, peaking in the summer and reaching a minimum in winter. Concentrations and fluxes of HNO\(_3\) peaked in the spring and reached a minimum in the fall. Deposition velocities, which can be thought of as the concentration-normalized flux, peaked during the summer and reached a minimum during winter for most N species. This pattern likely reflects the combined effect of seasonal cycles in LAI and turbulence characteristics. The seasonal pattern of \( V_{\text{d}} \) for HNO\(_3\) differed slightly from the other species, peaking in spring and reaching a minimum in fall. In contrast to other N species, HNO\(_3\) deposition is limited by turbulent transfer, with the canopy (surface) resistance assumed to be zero. The pattern of HNO\(_3\) \( V_{\text{d}} \) thus partially reflects seasonal patterns in wind speed and the degree of turbulent mixing above the canopy.

### 3.8 Evaluation of the dry deposition model

While total uncertainty in the dry deposition budget cannot be rigorously quantified (Walker et al., 2019a), the sensitivity of the model to parameterizations and key inputs can elucidate important aspects of model uncertainty and inform a potential range of dry deposition estimates. Here we undertake such an exercise by evaluating several alternative modeling scenarios to assess the sensitivity of fluxes and total dry deposition to assumptions regarding LAI, NH\(_3\) emission,

https://doi.org/10.5194/bg-20-971-2023

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but lower than $V_d$ showed MBR
As shown in Fig. S11, fluxes measured during summer 2016 and NH$_3$ concentrations of NO$_3$ become, for example, by equilibrium of HNO$_3$ and NO$_3$ on leaf surfaces (Nemitz et al., 2004a). This pattern may also reflect the influence of flux divergence related to NH$_4$NO$_3$ evaporation in the canopy crown, which would reduce the magnitude of the downward vertical gradients, and therefore the measured $V_d$, of HNO$_3$ and NH$_3$ (Nemitz et al., 2004b). In this analysis, concentrations of NO$_3$ (mean 0.08 µg m$^{-3}$) were much lower than HNO$_3$ (mean 0.47 µg m$^{-3}$), and NO$_3$ gradients were therefore difficult to resolve, precluding a definitive explanation of HNO$_3$ $V_d < V_{dmax}$. Ignoring potentially significant uncertainties related to the measurement of chemical and temperature gradients within the roughness sublayer, our results suggest that periods of NH$_3$ emission during the day, particularly at higher air temperature and lower humidity, may be underestimated. Our results also reinforce the need for temporally extensive measurements of concentrations and fluxes of HNO$_3$, NH$_3$, and NO$_3$ to examine exchange processes and uncertainties related to chemical flux divergence.

3.9 Spatial and temporal representativeness of the deposition budget

The complexity of atmospheric flows in mountainous terrain influences the spatial variability of wet and dry deposition processes (Lehner and Rotach, 2018). As the deposition budget presented above is based on measurements from the lowest elevation portion of the Coweeta Basin, the degree to which the budget is spatially representative must be considered. Potential effects on dry deposition were assessed by characterizing the magnitude and spatial variability of HNO$_3$ and NH$_3$ concentrations along an elevation gradient (Fig. 1, Table 1) from the lower to upper portions of the Coweeta Basin. It should be noted that HNO$_3$ concentrations at NC25 were measured by CASTNET, while HNO$_3$ passive samplers were used at the other locations. Concentrations are summarized in Fig. 13, in which the sites are ordered left to right from lowest to highest elevation. Nitric acid and NH$_3$ concentrations increase slightly with elevation, an explanation for which is not obvious. Nitric acid concentrations are highest at Screwdriver Knob, which is distinct from the other sites in that the measurement tower was situated on a relatively narrow exposed ridge. The measurements are therefore higher above the surrounding vegetation than at the other sites. Overall, variability of air concentrations across sites, even including SK, is sufficiently small such that spatial variability of dry deposition across the basin would likely be driven more by variability in meteorology than air concentrations.

A quantitative assessment of the effects of airflow on dry deposition across the basin is not possible, but the work of Hicks (2008) illustrates the relevant effects in the context of the resistance analogy for $V_d$. Over flat homogeneous terrain, flux to vegetation is driven by turbulent diffusion in the vertical direction above the canopy and horizontal advection is assumed to be zero. In the extreme case of airflow approaching a steep forested slope, horizontal flow penetrates the canopy and the transfer of material (deposition) to the canopy elements becomes dominated by horizontal advection and filtration rather than vertical diffusion. In the context of $V_d$, this situation is analogous to the aerodynamic resistance ($R_a$) approaching zero. Taking HNO$_3$ as an example under the typical assumption that the canopy resistance is $R_c = 0$, $V_d$ becomes limited by the quasi-laminar boundary layer resistance at the vegetation surfaces ($R_b$). Following the analysis of Hicks (2008), $V_d$ for HNO$_3$ could be enhanced by a factor of $[1 + (R_d/R_b)]^{1/2}$ (Hicks, 2008). Using median values of $R_d$ and $R_b$ from our modeling period, this would increase $V_d$ for HNO$_3$ by a factor of $\sim 1.4$. For gases that have a signif-
Concentrations (as N) of HNO$_3$ and NH$_3$ measured at different elevations, increasing from left to right (see Fig. 1 and Table 1), across the Coweeta Basin. Solid and dashed lines inside the box represent the median and mean, respectively. The top and bottom of the box represent 75th and 25th percentiles. Whiskers represent 90th and 10th percentiles, and dots represent 95th and 5th percentiles.

Regarding the temporal representativeness of the deposition budget calculated here, wet deposition of inorganic N (NO$_3^−$ + NH$_4^+$) during our 12-month model period (3.69 kg N ha$^{-1}$) agrees well with the mean annual deposition rate measured at NTN site NC25 (3.72 kg N ha$^{-1}$) over the period 2015–2020, with 2020 being the most recent full year of observations reported at the time of this analysis. Air concentrations of NO$_3^−$ and HNO$_3$ (Fig. 6) as well as NH$_3$ and NH$_4^+$ (Fig. 7) during our model period are also similar to the 6-year (2015–2020) mean concentrations measured by CASTNET and AMoN. In this context, our results are deemed temporally representative of the most recently available complete years of monitoring data.

4 Conclusions

Due to the success of the Clean Air Act, air concentrations and wet deposition of reactive N at Coweeta are the lowest observed since the beginning of routine monitoring in the late 1970s. However, even at historically low levels, our results show that NH$_3$ deposition remains highly ecologically relevant in the context of critical loads. Our estimate of total N$_r$ deposition of 6.7 kg N ha$^{-1}$ yr$^{-1}$ is near the upper-end estimate of mass-balance-derived critical loads (2.8 to 7 kg N ha$^{-1}$ yr$^{-1}$) recently reported for spruce–fir, beech, and mixed deciduous forests by Pardo et al. (2018) in the nearby Great Smoky Mountains National Park. Our result also falls within the range of empirical critical loads of N for combined tree health and biogeochemical responses (3–8 kg N ha$^{-1}$ yr$^{-1}$) as well as changes in mycorrhizal fungi spore abundance, community structure, and community composition (5–12 kg N ha$^{-1}$ yr$^{-1}$) in eastern temperate forests (Pardo et al., 2011).

A key feature of the deposition budget derived for Coweeta is the predominance of reduced forms (NH$_3$) of inorganic nitrogen (51.1%) over oxidized inorganic N (41.3%). Reductions in deposition of NH$_3$ will be needed to achieve the lower-end estimates of critical N loads (3 kg N ha$^{-1}$ yr$^{-1}$) for southern Appalachian forests. This presents a challenge, as emissions and air concentrations of NH$_3$ remain unregulated. Our results also show that organic forms of N make a nontrivial contribution (7.6%) to total N deposition, primarily via wet deposition. It is noted, however, that the gas-phase dry component of deposition only considers oxidized forms as alkyl and peroxy nitrates, excluding contributions from reduced (i.e., NH) organic compounds. While our results represent an advancement in accounting for organic dry N$_r$ deposition in total N$_r$ deposition, the application of new measurement technologies (Walker et al., 2019b) for broader chemical speciation of organic forms of dry N$_r$ deposition is needed.

Our results underscore the need for long-term measurements of reactive chemical fluxes, as well as the coupling of atmospheric and biogeochemical measurements, to im-
prove air–surface exchange models. Novel measurements that more directly elucidate the role of cuticular exchange of NH₃ and more temporally extensive measurements of leaf NH₃ emission potentials are particularly needed. For forest ecosystems, a physically representative parameterization for resistance to NH₃ diffusion through the leaf litter layer and more temporally extensive measurements of the litter NH₃ emission potential combined with more thorough understanding of litter decay dynamics are also needed. For sensitive ecosystems located in mountainous and other topographically complex landscapes, which includes much of the Class I wilderness area in the US, identification of locations suitable for micrometeorological flux measurements will be challenging. Novel flux measurement methods and application of in situ models, including translation of measurements from more ideal to complex locations, will likely be needed. Furthermore, long-term flux datasets are needed to assess the interactive effects of changing air quality and climate on both atmosphere–biosphere exchange and ecosystem response to deposition (e.g., Van Houtven et al., 2019).


Data availability. NADP, AMoN, and CASTNET data are available at the websites referenced herein (see Sect. 2.1 for more information). Other data appearing in figures and tables are available at https://catalog.data.gov/dataset/epa-sciencehub (EPA ScienceHub, 2023).

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/bg-20-971-2023-supplement.

Author contributions. JTW: conceptualization, formal analysis, methodology, funding acquisition, project administration, validation, visualization, writing. XC: formal analysis, investigation, methodology, validation, writing. ZW: formal analysis, investigation, methodology, software validation, writing. DS: investigation, formal analysis, validation. AD: data curation, investigation, methodology, resources. ACO: conceptualization, formal analysis, methodology, validation. EE: data curation, funding acquisition, formal analysis, methodology, validation, resources. JB: formal analysis, methodology, software. JK: data curation, investigation. MP: conceptualization, funding acquisition, resources. JI: formal analysis, investigation, writing. CFM: conceptualization, funding acquisition, resources, writing.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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Acknowledgements. We gratefully acknowledge field and laboratory support from USDA Forest Service staff at the Coweeta Hydrologic Laboratory, including Christine Sobek, Patsy Clinton, Chuck Marshall, and Cindi Brown. David Kirchgeissner (retired, U.S. EPA) tirelessly supported field and laboratory activities during SANDS intensives. Lee Nanny (former U.S. EPA) and Mark Barnes (U.S. EPA) supported field intensives and logistics. We also appreciate the support of Kevin Mishoe (Wood, Inc.) and Christopher Rogers (Wood, Inc.) for support of CASTNET field activities and data management, respectively.

Financial support. This research has been supported by the U.S. Environmental Protection Agency (Intramural) and the USDA Forest Service, Southern Research Station, Coweeta Hydrologic Lab.

Review statement. This paper was edited by Ivonne Trebs and reviewed by Chris Flechard and one anonymous referee.

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