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Global deposition of total reactive nitrogen oxides from 1996 to 2014 constrained with satellite observations of NO₂ columns

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Abstract. Reactive nitrogen oxides (NO_{ν}) are a major constituent of the nitrogen deposited from the atmosphere, but observational constraints on their deposition are limited by poor or nonexistent measurement coverage in many parts of the world. Here we apply NO₂ observations from multiple satellite instruments (GOME, SCIAMACHY, and GOME-2) to constrain the global deposition of NO_{y} over the last 2 decades. We accomplish this by producing top-down estimates of NO_x emissions from inverse modeling of satellite NO₂ columns over 1996–2014, and including these emissions in the GEOS-Chem chemical transport model to simulate chemistry, transport, and deposition of NO_v. Our estimates of long-term mean wet nitrate (NO_3^-) deposition are highly consistent with available measurements in North America, Europe, and East Asia combined (r = 0.83, normalized mean bias = -7%, N = 136). Likewise, our calculated trends in wet NO₃⁻ deposition are largely consistent with the measurements, with 129 of the 136 gridded modeldata pairs sharing overlapping 95 % confidence intervals. We find that global mean NO_{ν} deposition over 1996–2014 is $56.0 \text{ Tg N yr}^{-1}$, with a minimum in 2006 of 50.5 Tg N and a maximum in 2012 of 60.8 Tg N. Regional trends are large, with opposing signs in different parts of the world. Over 1996 to 2014, NO_{ν} deposition decreased by up to 60 % in eastern North America, doubled in regions of East Asia, and declined by 20 % in parts of western Europe. About 40 % of the global NO_{ν} deposition occurs over oceans, with deposition to the North Atlantic Ocean declining and deposition to the northwestern Pacific Ocean increasing. Using the residual between NO_x emissions and NO_y deposition over specific land regions, we investigate how NO_x export via atmospheric transport has changed over the last 2 decades. Net export from the continental United States decreased substantially, from 2.9 Tg N yr⁻¹ in 1996 to 1.5 Tg N yr⁻¹ in 2014. Export from China more than tripled between 1996 and 2011 (from 1.0 to 3.5 Tg N yr⁻¹), before a striking decline to 2.5 Tg N yr⁻¹ by 2014. We find that declines in NO_x export from some western European countries have counteracted increases in emissions from neighboring countries to the east. A sensitivity study indicates that simulated NO_y deposition is robust to uncertainties in NH₃ emissions with a few exceptions. Our novel long-term study provides timely context on the rapid redistribution of atmospheric nitrogen transport and subsequent deposition to ecosystems around the world.

1 Introduction

The introduction of reactive nitrogen to the environment by anthropogenic activities (e.g., from fossil fuel combustion and the production of fertilizers for agriculture) has drastically altered the global nitrogen cycle with consequences throughout the Earth system (Galloway et al., 2004). Reactive nitrogen dominates the chemical production of tropospheric ozone and contributes to inorganic aerosol formation, with implications for air quality and climate. Deposition of nitrogen from the atmosphere has been linked to eutrophication and acidification (Bouwman et al., 2002), reductions in biodiversity (Bobbink et al., 2010; Hernández et al., 2016; Isbell et al., 2013; De Schrijver et al., 2011), and climate impacts through coupling with the carbon cycle and greenhouse gas emissions (Liu and Greaver, 2009; Reay et al., 2008; Templer et al., 2012). Despite its global importance, observational constraints on nitrogen deposition are lacking in many parts of the world due to poor or nonexistent measurement coverage (Vet et al., 2014).

Atmospheric transport is a dominant process for distributing reactive nitrogen around the world (Galloway et al., 2008). Some forms of reactive nitrogen can be transported over distances greater than 1000 km (Neuman et al., 2006; Sanderson et al., 2008), depositing across national boundaries and continents. For example, the US is estimated to export 30-40 % of its reactive nitrogen emissions (Dentener et al., 2006; Holland et al., 2005; Zhang et al., 2012), while transport from China accounts for up to 66-92 % of total nitrogen deposition to parts of the northwestern Pacific Ocean (Zhao et al., 2015). Fertilization of the open ocean due to atmospheric transport and deposition of anthropogenic nitrogen may be a considerable factor in marine productivity (Duce et al., 2008), prompting important questions about the fate and impact of deposition far downwind of sources where observations are limited.

Reactive nitrogen oxides (NO_v \equiv NO + NO₂ + HNO₃ + HONO + organic nitrate molecules + aerosol nitrate) contribute about half of the total nitrogen deposited worldwide (Dentener et al., 2006). NO_v deposition was estimated to be around $45-50 \text{ Tg N yr}^{-1}$ in the late 1990s and early 2000s, representing a three- to fourfold increase since the preindustrial era (Dentener et al., 2006; Kanakidou et al., 2016; Lamarque et al., 2013). A substantial range exists in the trajectory of global NO_v deposition beyond the year 2000, depending on the emission scenario. Galloway et al. (2004) projected that NO_v deposition could increase by > 70% by 2050, while Dentener et al. (2006) projected changes between -25 to +50% by 2030 for maximum feasible reduction and "pessimistic" scenarios, respectively. More recent multimodel projections by Lamarque et al. (2013) estimate NO_v deposition would change by -16 to +5% for 2030 and by -48 to -25% for 2100, depending on the representative concentration pathway (RCP) scenario. This range in projections highlights the need for global observational constraints on contemporary changes in nitrogen oxide emissions.

Sources of NO_x (\equiv NO + NO₂), whose oxidation is responsible for the formation of other reactive nitrogen oxides, include fossil fuel combustion, biomass burning, lightning, and biogenic emission from soil. The magnitude and spatial distribution of NO_x emissions have changed considerably over the past several decades, corresponding to patterns of human development and emission control measures. Tropospheric NO₂ columns derived from satellite remote sensing observations have been used extensively to constrain regional and global NO_x emissions (Streets et al., 2013). This top-down approach complements bottom-up inventories that are assembled using regionally specific emission factors and fuel combustion data for various source categories. In particular, satellite NO₂ observations can provide insight into otherwise poorly constrained sources (Beirle et al., 2010; Jaegle et al., 2005; Richter et al., 2004; Vinken et al., 2014), produce coherent long-term trends (van der A et al., 2008; Lu et al., 2015; Stavrakou et al., 2008; Zhang et al., 2007), document interannual variability (Castellanos and Boersma, 2012; Konovalov et al., 2010; Russell et al., 2012), offer information to evaluate and improve bottom-up inventories on the global scale (Martin, 2003; Miyazaki et al., 2017), and provide timely emission updates (Lamsal et al., 2011; Mijling et al., 2013; Souri et al., 2016).

Satellite observations of NO₂ began with GOME (1995-2003), were followed by SCIAMACHY (2002-2011), and continue today with OMI (2004-), GOME-2 (2007-), and TROPOMI (2017-), resulting in a record of global atmospheric NO₂ abundance over the past 20 years. These observations have been used previously to estimate the deposition of nitrogen species, either in combination with chemical transport modeling or with empirical approaches (Cheng et al., 2013; Jia et al., 2016; Lu et al., 2013; Nowlan et al., 2014). For example, Nowlan et al. (2014) demonstrated how satellite-inferred surface concentrations of NO2 can be combined with modeling to produce spatially continuous estimates of NO₂ dry deposition fluxes. They found that dry deposition of NO₂ contributes as much as 85 % of total NO_{ν} deposition in urban areas, but represents only 3 % of global NO_x emitted. The remaining 97 % of global NO_y deposition is made up of both wet and dry deposition of other reactive nitrogen oxide compounds that are not directly observed by satellite-based instruments.

In this study, we expand on the approach of Nowlan et al. (2014) by using the satellite observations of NO₂ columns to constrain total NO_y deposition, including other oxidized nitrogen species and wet deposition which contribute substantially to NO_y deposition. We accomplish this by constraining surface NO_x emissions using the satellite observations of NO₂ and simulating subsequent NO_y deposition with a global chemical transport model. Given the effective mass balance between NO_x emissions and deposition of reactive nitrogen oxides, observational constraints on NO_x emissions provide a powerful top-down constraint on deposition (which to our knowledge has not yet been exploited in this way).

We leverage the long-term coverage of GOME, SCIA-MACHY, and GOME-2 observations to produce a globally consistent and continuous record of NO_y deposition from 1996 to 2014. We highlight long-term trends in satellite-constrained NO_y deposition around the world and discuss changes in regional export of NO_x. Our satellite-constrained estimates of NO_y deposition are evaluated using measured wet nitrate (NO₃⁻) deposition from a variety of sources worldwide. We also explore the sensitivity of the NO_y deposition estimates to uncertainties in NH₃ emissions and discuss other considerations.

2 Methods

2.1 Satellite-based constraints on NO_y deposition

The application of satellite NO₂ column observations to constrain NO_y deposition requires a method to propagate the observational information across the entire NO_y system containing species with lifetimes of days or longer. The short NO_x lifetime of hours during daytime satellite observations implies that a direct assimilation for NO₂ column abundance would rapidly lose the observational information as the assimilation returns to its unperturbed state well before the next satellite observation days later. We therefore apply satellite NO₂ observations to constrain NO_x emissions in a simulation of NO_y deposition so the information propagates across the entire NO_y system.

We calculate top-down surface NO_x emissions from 1996 to 2014 using observations from GOME (1995-2003), SCIAMACHY (2002–2011) and GOME-2 (2007–). The similar overpass time of these three instruments (from about 09:30 to 10:30 LT, local time) facilitates their combination to provide consistent long-term coverage (Geddes et al., 2015; Hilboll et al., 2013; Konovalov et al., 2010; van der A et al., 2008). We achieve consistency across all three instruments despite their varying pixel sizes $(320 \text{ km} \times 40 \text{ km}, 60 \text{ km} \times 30 \text{ km}, \text{ and } 80 \text{ km} \times 40 \text{ km} \text{ for})$ GOME, SCIAMACHY, and GOME-2, respectively) by gridding the daily observations from each to a regular coarse grid of $2^{\circ} \times 2.5^{\circ}$ latitude by longitude. Tropospheric NO₂ vertical column densities are provided by KNMI at http://www.temis. nl/airpollution/. In all cases, NO2 column densities are retrieved by differential optical absorption spectroscopy using backscattered radiance in the 425-450 nm wavelength range according to the retrieval algorithm documented in Boersma et al. (2004). The error in individual satellite-derived tropospheric NO₂ column retrievals is estimated to be around 35-60 % for polluted scenes and greater than 100 % for clean regions (Boersma et al., 2004). Boersma et al. (2016) describe well the value of accounting for vertically resolved instrument sensitivity. We use the averaging kernels provided with the data to replace a priori NO₂ vertical profiles with those from GEOS-Chem model following Lamsal et al. (2010). We use daily nadir observations with a cloud radiance fraction of less than 0.5. We minimize errors associated with wintertime retrievals by using a solar zenith angle cut-off of 50°.

We use the GEOS-Chem chemical transport model (www. geos-chem.org) v9-02 to conduct the inversion of satellite observations of NO₂ and constrain global NO_y deposition. The simulation is described in Appendix A. Briefly, GEOS-Chem is driven by assimilated meteorology from the NASA Global Modeling and Assimilation Office and simulates detailed HO_x-NO_x-VOC-aerosol (VOC: volatile organic compound) chemistry (Bey et al., 2001; Park et al., 2004). Removal occurs through wet deposition (Amos et al., 2012; Liu et al., 2001) and dry deposition based on the widely

used resistance-in-series formulation (Wesely, 1989). Anthropogenic, biogenic, soil, lightning, and biomass burning emissions are included (see Appendix A). In the case of NO_x , the bottom-up emissions are used as prior estimates that we then overwrite with the top-down emissions.

We adopt a finite-difference mass-balance inversion (Cooper et al., 2017; Lamsal et al., 2011) for computational expedience given the 19-year period of interest. In two initial simulations, a perturbation (30%) to the a priori emissions, E, in a grid cell is used to find the relationship between the a priori NO₂ column, Ω , and the change in the column resulting from that perturbation:

$$\frac{\Delta E}{E} = \beta \times \frac{\Delta \Omega}{\Omega}.$$
(1)

The factor β in Eq. (1) accounts for nonlinear feedbacks between a change in NO_x emissions and NO_x chemistry in a grid cell leading to grid cell NO₂ column abundance.

We then use monthly-mean gridded satellite observations, Ω_{sat} , in combination with monthly β values for each grid box to derive new gridded annual emissions, $E_{topdown}$, from the prior emissions estimates, E_{prior} , by rewriting Equation 1 as:

$$E_{\text{topdown}} = E_{\text{prior}} \cdot \left[1 + \beta \frac{\Omega_{\text{sat}} - \Omega_{\text{prior}}}{\Omega_{\text{prior}}} \right].$$
(2)

We use GOME observations for years 1996 to 2002, SCIA-MACHY observations for years 2003 to 2006, and GOME-2 observations for years 2007 to 2014. In all cases, monthly mean simulated NO2 columns are calculated using days with coincident satellite observations. The simulated NO₂ vertical column is output daily for late morning. We calculate scaling factors for every month with available satellite observations, then calculate an annual mean scaling factor that is used to infer annual mean top-down emissions. Our top-down emissions retain the same seasonality as the prior emissions to mitigate concerns about seasonally missing data (such as from snow or monsoonal clouds). The top-down emissions are then used in a final simulation to model NO_{ν} deposition. For locations without any satellite observations, the a priori emissions are used. The resultant simulation of NO_{y} deposition is thus constrained by, and consistent with, the satellite NO2 observations (similar in essence to an assimilation system). We note that uncertainty in tropospheric NO_2 from lightning will propagate into the inversion (Travis et al., 2016), but there is no evidence of a significant trend in lightning NO_x over the long term (Murray et al., 2012). A constant bias is unlikely to affect the trend analyses presented here.

Table 1 shows the annual global top-down NO_x emissions we obtain from our calculations. We derive global mean satellite-constrained NO_x emissions from 1996 to 2014 of $55.6 \pm 3.4 \text{ Tg N yr}^{-1}$. Our top-down global NO_x emissions for 2001 of 52.3 Tg N are consistent with the mean \pm standard deviation from over 20 models used in the Coordi-

Table 1. Global top-down NO_x emissions calculated using the finite mass-balance inversion approach with observations from GOME, SCIAMACHY, and GOME-2.

Year	Global NO _x
	emissions
	$(TgNyr^{-1})^*$
1996	60.1
1997	58.4
1998	59.2
1999	59.6
2000	53.4
2001	52.3
2002	55.1
2003	50.1
2004	51.5
2005	51.2
2006	50.0
2007	54.7
2008	56.1
2009	55.9
2010	57.5
2011	58.9
2012	59.3
2013	58.5
2014	54.0
Mean	55.6

* Includes anthropogenic HNO₃ flux of 2.3 ± 0.1 Tg N yr⁻¹.

nated Model Studies Activities of the Task Force on Hemispheric Transport of Air Pollution (HTAP) for the same year of 46.6 ± 7.8 Tg N (Vet et al., 2014).

2.2 Measurements of wet deposition

We use a variety of regional and global measurements of wet nitrate (NO_3^-) deposition to evaluate our satellite-constrained simulation from 1996 to 2014.

To evaluate overall global performance we use data compiled by the World Data Centre for Precipitation Chemistry for two time periods: 2000–2002 and 2005–2007 (http: //www.wdcpc.org/). The use of these data ensures optimal and consistent quality standards across all stations, allowing for evaluation of global performance with careful consideration of sampling protocols and data completeness (Vet et al., 2014).

To evaluate the long-term means and trends from 1996 to 2014, we obtain observations of wet NO_3^- deposition from North America, Europe, and East Asia where continuous measurements are available throughout most of our study period. Observations come from the National Atmospheric Deposition Program in the United States (http://nadp.sws.uiuc. edu/, available 1996–2014), from the Canadian Air and Precipitation Monitoring Network in Canada (http://www.ec.gc.



Figure 1. Long-term (1996–2014) mean NO_y deposition derived from the GEOS-Chem simulation, constrained by satellite observations of NO₂ columns from the GOME, SCIAMACHY, and GOME-2 instruments (**a**). Mean ratio of simulated dry NO_y deposition to total NO_y deposition (**b**).

ca/rs-mn/, available 1996–2011), from the European Monitoring and Evaluation Programme in Europe (http://www. emep.int/, available 1996–2014) and from the Acid Deposition Monitoring Network in East Asia (http://www.eanet. asia, available 2000–2014). In the US and Canada, wet deposition is measured by wet-only samplers that are triggered at the onset of precipitation. Measurements in Europe are made by bulk- and wet-only sampling methods, and we used both in this analysis. Measurements across East Asia are reported as wet-only, although at some stations this may not be accomplished by strictly wet-only samplers (http://www.eanet. asia/product/manual/prev/techwet.pdf).

For our trend analysis, we only included stations which had quality controlled annual data for at least 15 of the 19 years in our study. This left 128 stations across the United States, 14 stations in Canada, 18 stations across Europe, and 14 stations across East Asia. For comparison with the GEOS-Chem model, if multiple stations are available within a single grid box we grid all measurements of annual wet deposition to the model horizontal resolution.

3 Satellite-constrained estimates of NO_y deposition

Here we summarize the overall patterns in long-term mean deposition resulting from our satellite-constrained simulation, followed by a discussion of the long-term trends, changes in regional export, and the sensitivity of the simulated NO_y deposition to potential uncertainties in NH₃ emissions and other considerations.

3.1 Long-term mean NO_y deposition

Figure 1 (top) shows our satellite-constrained long-term mean NO_y deposition from 1996 to 2014. We find that 32.2 Tg N yr⁻¹ is deposited on average over the continents (57% of the total), and 23.8 Tg N yr⁻¹ is deposited on average over the oceans (43% of the total). This is similar to the estimate by Galloway et al. (2004) that 46% of modern day NO_y deposition occurs over the oceans. Critical nitrogen deposition loads for various natural freshwater and terrestrial ecosystems lie in the range of 5–20 kg N ha⁻¹ yr⁻¹, depending on the ecosystem, soil conditions, and land history (World Health Organization, 2000). We estimate that mean deposition of oxidized nitrogen alone exceeds 5 kg N ha⁻¹ yr⁻¹ over a land area of approximately 12.7 × 10⁶ km² (or ~ 8% total land area).

In the Northern Hemisphere, high NO_{ν} deposition tends to be associated with regions that have high anthropogenic NO_x sources. We find that mean NO_y deposition in the eastern United States exceeds $10 \text{ kg N} \text{ ha}^{-1} \text{ yr}^{-1}$ (maxi $mum = 11.4 \text{ kg N} \text{ ha}^{-1} \text{ yr}^{-1}$) with elevated deposition extending into southeastern Canada and hundreds of kilometers into the Atlantic Ocean. This is similar to the multimodel ensemble results from ACCMIP and HTAP, predicting between 5 and $15 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ in this region (Lamarque et al., 2013; Vet et al., 2014). A prior GEOS-Chem analysis over North America for the years 2006-2008 also predicted NO_{y} deposition exceeding $10 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ in the eastern US (Zhang et al., 2012). Elsewhere in North America, we find high NO_v deposition along the western coast of California (up to $6 \text{ kg N} \text{ ha}^{-1} \text{ yr}^{-1}$) and in the vicinity of Mexico City (up to $10 \text{ kg N ha}^{-1} \text{ yr}^{-1}$).

We find mean NO_y deposition is also elevated throughout Europe, with a maximum of 8.5 kg N ha⁻¹ yr⁻¹ located in northern Italy near the Po Valley region. Again, our long-term estimate in this region is similar to the AC-CMIP and HTAP ensemble means, predicting NO_y deposition in the range of 5–10 kg N ha⁻¹ yr⁻¹ (Lamarque et al., 2013; Vet et al., 2014). The elevated deposition here is also spatially consistent with the results from Holland et al. (2005). We find high deposition extending into western Russia with a hotspot in the vicinity of Moscow approaching 5 kg N ha⁻¹ yr¹. Our observation-constrained estimate also has isolated regions of high deposition in the Middle East (around 4–5 kg N ha⁻¹ yr⁻¹ in the vicinity of Tehran and around the Persian Gulf).

We find that the highest mean deposition in the world occurs in China, exceeding $10 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ in many regions. High deposition extends into the midlatitude western Pacific Ocean off the coast of East Asia. NO_y deposition in the ACCMIP and HTAP ensemble means also exceeds $10 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ throughout eastern China. We find the highest long-term mean deposition (with a maximum close to $20 \text{ kg N ha}^{-1} \text{ yr}^{-1}$) occurs in the south, around the Pearl River Delta and in the vicinity of Guangzhou, although

deposition is also high in the regions just west of Beijing and Shanghai.

In the Southern Hemisphere, high NO_y deposition is associated with biomass burning and soil NO_x sources, in addition to anthropogenic sources. For example, we find NO_y deposition is between 3 and 5 kg N ha⁻¹ yr⁻¹ in central and southern Brazil as well as in the tropical rainforests and moist savannahs of Africa. Our estimates in these biomassburning- and soil-NO_x-dominated regions are also generally consistent with the ACCMIP and HTAP ensemble estimates (2–5 kg N ha⁻¹ yr⁻¹). We find NO_y deposition up to 10 kg N ha⁻¹ yr⁻¹ in the vicinity of São Paulo and Rio de Janeiro, and in the vicinity of Johannesburg and the industrialized Mpumalanga Highveld of South Africa (all dominated by anthropogenic NO_x emissions). Our constrained simulation also identifies hotspots of deposition in the vicinity of Melbourne and Sydney, Australia (~4 kg N ha⁻¹ yr⁻¹).

Figure 1 (bottom) shows the simulated long-term ratio of dry NO_v deposition to total (wet + dry) NO_v deposition. Globally, dry and wet deposition contribute roughly equally to total NO_v deposition (52 and 48 %, respectively). Dry deposition usually accounts for more than 50% of the total over the continents and directly offshore, whereas wet deposition dominates over the remote oceans. In the generally arid regions of the world (e.g., the southwestern US, the Sahara Desert, the Arabian Peninsula, and the Gobi Desert) dry deposition accounts for \sim 85 % or more of the total deposition. Elsewhere, dry deposition fractions tend to be highest (>60 %) nearest to major surface NO_x sources (e.g., the eastern US, western Europe, and near other major urban centers around the world in addition to the soil- and biomassburning-dominated source regions in South America and Africa). HNO₃ typically makes the dominant contribution to dry NO_v deposition, although NO₂ and HNO₃ can make almost equal contributions in certain high-NO $_x$ environments. Isoprene nitrates and peroxyacetyl nitrates comprise $\sim 10-$ 30% of dry NO_v deposition in some densely forested and high-latitude environments, respectively.

We evaluate our estimates of NO_{v} deposition with measured wet NO_3^- from several sources. Figure 2 shows measurements of annual wet NO₃⁻ deposition from the World Data Centre for Precipitation Chemistry, available for two time periods: 2000-2002 (N = 470) and 2005-2007(N = 484). In both we see the patterns of elevated deposition in eastern North America, western Europe, and parts of South and East Asia, with lower deposition in western North America, across high latitudes in the Northern Hemisphere, and in the available observations in Africa. High deposition in the Southern Hemisphere is observed between São Paulo and Rio de Janeiro, and just southeast of Johannesburg. Figure 2 also shows the wet NO_3^- deposition from our constrained simulation during the same two time periods (2000-2002 and 2005-2007), which exhibits similar patterns to those found in total NO_v deposition (Fig. 1).



Figure 2. Annual wet NO_3^- deposition from measurements available through the World Data Centre for Precipitation Chemistry and from the GEOS-Chem simulation constrained with satellite observations of NO_2 . Two time periods are represented: 2000–2002 and 2005–2007.



Figure 3. Scatter plot of the satellite-constrained simulated wet NO_3^- deposition vs. measurements available through the World Data Centre for Precipitation Chemistry for specific subsets of the data. The red lines show the result of a reduced major axis linear regression. In the right column, the blue line shows the fit across all data and the red line shows the fit excluding the two circled data points that are discussed in the text (reported statistics refer to the red line fit).

We find a high degree of consistency between our estimate and the observations for both 2000–2002 (N = 306 model– data pairs) and 2005–2007 (N = 310 model–data pairs); the normalized mean bias (NMB) is -14 and -16%, respectively. The vast majority of pairs (>80%) agree to within 50% of each other. Figure 3 shows scatter plots for specific subsets of the global data. The agreement for both time periods is strongest over North America (r = 0.92 for both 2000–2002 and 2005–2007, NMB = +1.0 and -5.0%, respectively). Robust model agreement with wet nitrate deposition observations over densely monitored North America is characteristic of other global studies (Dentener et al., 2006; Lamarque et al., 2013; Vet et al., 2014). Our agreement is also good in Europe (r = 0.69 and 0.66, and NMB = -31.0 and -29.8 %, respectively). The weaker correlation and low bias in this region is likewise characteristic of global studies, although our spatial correlation (r = 0.66-0.69) is on the high end of previously reported multimodel ensembles ($r \sim 0.4-0.6$, Dentener et al., 2006; Lamarque et al., 2013; Vet et al., 2014). The negative bias over Europe compared to North America has previously been attributed to poor modeling of precipitation and/or spatial representativeness of the measurements compared to model resolution. Throughout the rest of the world (encompassing observations mostly over Asia, but also over eastern Russia, and some locations in the Southern Hemisphere) the combined spatial coverage of the observations is very low (N = 53). Normalized mean bias in these estimates is also high compared to North America (NMB = -19.5 and -17.8% for 2000–2002 and 2005– 2007, respectively), and our spatial correlation with the measurements is weak (r = 0.35 and 0.42, respectively). We find that our poor agreement here is disproportionately driven by the two observations that also have the highest measured deposition in the world: near Port Blair on the South Andaman Island in the Bay of Bengal, and in the Arunachal Pradesh state in northeastern India. Agreement is considerably better with the rest of the data (r = 0.78 and 0.72, NMB = +0.01and -0.01 % for 2000-2002 and 2005-2007, respectively). Excluding these two points substantially improves the global agreement as well (from r = 0.57 to 0.75 and r = 0.59 to 0.75, respectively). Site representativeness, precipitation errors, or uncertainty in our satellite-constrained NO_x emissions may explain the discrepancy at these two specific sites.

In addition to global data for 2000-2002 and 2005-2007 from the World Data Centre for Precipitation Chemistry, we also evaluate our estimates of NO_{ν} deposition over the long term (1996-2014) using continuous observations provided by regional networks. Figure 4 shows measured wet NO_{3}^{-} deposition over North America, Europe, and East Asia for locations where at least 15 years of quality-controlled annual data are available. These long-term mean observations demonstrate many of the same spatial patterns as the time slices from 2000-2002 and 2005-2007. In North America, a relatively smooth gradient is observed from low deposition in the west to high deposition at sites in the east. In Europe, the highest measured long-term mean wet NO₃⁻ deposition occurs at a coastal site in southern Norway, at a site just east of Copenhagen, and at locations in northern Italy and in Switzerland. At higher-latitude sites, deposition is lower. Across the eastern Asia network, the measurements show the highest deposition at sites in Southeast Asia (e.g., at a location between Kuala Lumpur and Singapore, and another in the vicinity of Jakarta) and in Japan. The lowest long-term mean deposition occurs at high-latitude sites along the border of Russia and Mongolia, while moderate to high deposition is measured on the coast of eastern China.

In general, our satellite-constrained estimate reflects the spatial variability that is seen in the measurements. Globally, the correlation between measured NO₃⁻ deposition and our estimated wet NO₃⁻ deposition is excellent (r = 0.83, NMB = -7.7 %, N = 136 gridded model-data pairs). The vast majority of pairs (>85 %) agree to within 50 % of each other. For the individual regions, normalized mean bias in our estimate is the smallest over North America (NMB = +2.4 %), and higher over Europe and East Asia (NMB = -32 and -25 %, respectively). The spatial corre-

lation over each region is strong (r = 0.89, r = 0.87, and r = 0.69 for North America, Europe, and East Asia, respectively), but sample sizes over Europe (N = 16) and East Asia (N = 11) are small so we emphasize caution in the interpretation of the statistics for these two regions. The lack of continuous measurement coverage even in parts of the world with routine network observations highlights the imperative of using other novel observational constraints on deposition (such as the global satellite observations of NO₂ used here).

3.2 Trends in global NO_y deposition from 1996 to 2014

Our long-term satellite-constrained estimate of NO_y deposition facilitates a unique and up-to-date investigation of the changes in NO_y deposition around the world. We calculate linear trends in annual NO_y deposition using the nonparametric Sen's method (Sen, 1968) and test for significance with the nonparametric Mann–Kendall method (Kendall, 1975; Mann, 1945). We treat increasing or decreasing trends as significant if p < 0.01. Given that this is a test for linear trends, regions where shorter-term trends in deposition may have changed signs over the period of study could result in erroneous or insignificant trends. Below we discuss particular regions where this is the case.

Figure 5 shows the long-term annual and seasonal trends calculated from our satellite-constrained estimate of total NO_y deposition across 1996–2014 (hatching indicates statistical significance). Figure 6 highlights time series of total NO_y deposition over three specific regions covering parts of North America, western Europe, and East Asia (as outlined in dashed boxes in Fig. 5a).

Substantial decreases are seen throughout North America, extending over the Atlantic Ocean to remote regions. The time series for this region (Fig. 6, left) shows that NO_{ν} deposition decreased by almost 40 % from 6.4 Tg N yr⁻¹ in 1996–1998 to 3.9 Tg N yr^{-1} in 2012–2014. The steepest local decline in the world appears over the Ohio River Valley area, with a maximum near Pittsburgh where NO_v deposition decreased by $-0.6 \text{ kg N ha}^{-1} \text{ yr}^{-2}$. NO_v deposition near Pittsburgh decreased from consistently exceeding $15 \text{ kg N} \text{ ha}^{-1} \text{ yr}^{-1}$ during 1996–2000, to below $6 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ by 2014. The strong decrease in the northeast is consistent with other long-term observational studies for the US (Sickles and Shadwick, 2007, 2015). Studies of US NO_x emissions derived from satellite observations have also highlighted the remarkable success of emission controls (Duncan et al., 2013; Russell et al., 2012). Our constrained estimate has the steepest declines during the summer (Fig. 5; June, July, and August, JJA), restricted tightly to the source regions. This also agrees with long-term observations showing the strongest reductions in the summer (Sickles and Shadwick, 2015), consistent with the shorter lifetime of NO_x and efficient dry deposition of NO_{ν} over the forested eastern US. We find significant decreases far downwind over the Atlantic Ocean during the other months, when NO_{y} can be



Figure 4. Long-term (1996–2014) wet NO_3^- deposition from available regional network measurements (**a**: NADP and CAPMON; **b**: EMEP; **c**: EANet) and from the GEOS-Chem simulation, constrained with satellite observations of NO_2 .

transported farther. The steep change in NO_y deposition in the eastern US over the last 20 years may have important consequences on tree mortality rates in the region, which have been demonstrated to be very sensitive to NO₃⁻ deposition in the range of $5-15 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ (Dietze and Moorcroft, 2011). The steeply decreasing trends across the US in our satellite-derived NO_y also support the increasing dominance of reduced nitrogen in total nitrogen deposition, evidenced by observations (Li et al., 2016) and model predictions (Ellis et al., 2013).

We find a small but statistically significant positive trend in NO_y deposition (+0.06 kg N ha⁻¹ yr⁻²) in northern Alberta, Canada, dominated by the trend in JJA. The region is downwind of development in the Canadian oil sands, which has seen notable changes in NO₂ column abundance as observed from space (McLinden et al., 2012). We estimate that deposition of NO_y in this area was at a maximum of 3.4 kg N ha⁻¹ yr⁻¹ in 2011 (up from 1.3 kg N ha⁻¹ yr⁻¹ in 1996–1997), and has since declined to 1.6 kg N ha⁻¹ yr⁻¹ by 2014. Elsewhere in Canada we estimate that NO_y deposition has decreased in the southern and eastern parts of the country, consistent with observational analyses (Zbieranowski and Aherne, 2011).

Declines in NO_y deposition are also found across Europe, but statistical significance tends to be limited to western continental Europe and the United Kingdom (while changes in the southern, northern, and eastern countries tend to be insignificant). According to the time series

for this region (Fig. 6, middle), NOy deposition decreased by about 15% (from 2.5 Tg N yr^{-1} in 1996–1998 to 2.1 Tg N yr^{-1} in 2012–2014). We find the steepest local trends $(-0.1 \text{ kg N ha}^{-2} \text{ yr}^{-1})$ in eastern Germany and the southern UK, where NO_v deposition in 2012–2014 decreased by 20 % compared to 1996–1998. Previous satellite constraints on NO_x emissions established that NO_x emissions in France, Germany, Great Britain, and Poland have declined since 1996 while emissions in Greece, Italy, Spain, and the Ukraine for example have either stayed constant or increased (Konovalov et al., 2008). The local variability in emission trends leads to notable transboundary impacts. For example, our simulation predicts no net trend in NO_v deposition over the Ukraine; but we find this is a result of opposing trends in dry (increasing) and wet (decreasing) deposition. This would be explained by increasing local emissions but decreasing transport from upwind. Similarly, we find significant increases in dry deposition in parts of western Russia but no significant trend in wet deposition.

Large increases in NO_y deposition are found throughout Asia, concentrated especially in eastern China and parts of Southeast Asia. Figure 6 shows the time series of wet and dry NO_y deposition within the rectangular region outlined in Fig. 5 that encompasses eastern China and part of the adjacent ocean. We find that NO_y deposition in the region increased by 65 % from 5.2 Tg N yr^{-1} in 1996–1998 to 8.6 Tg N yr^{-1} in 2012–2014. The time series also shows that NO_y deposition decreased after peakJ. A. Geddes and R. V. Martin: Global deposition of total reactive nitrogen oxides



Figure 5. Long-term trend (1996–2014) in the satellite-constrained simulation of NO_y deposition. (a) Annual mean; (b) December–January–February; (c) March–April–May; (d) June–July–August; (e) September–October–November. Diagonal hatching represents trend significance (p < 0.01). Hatching from top-left to bottom-right indicates a decreasing trend; hatching from bottom-left to top-right indicates an increasing trend.

ing around 9.3 Tg N yr^{-1} in 2011–2012. We find that the steepest increasing local trends in the world appear in eastern China, and in the Pearl River Delta region (up to +0.6 kg N ha⁻¹ yr⁻²). In fact, deposition in the Pearl River Delta region is the highest in the world for most of our record, exceeding $20 \text{ kg N} \text{ ha}^{-1} \text{ yr}^{-1}$ every year from 2003 to 2014 (almost doubling from just over $11 \text{ kg N ha}^{-1} \text{ yr}^{-1}$ in 1996). The trends in deposition over China are largest in summer when the NO_x lifetime is short, with more obvious indications of increasing NO_{ν} transport in the spring months (Fig 5; March-April-May, MAM). The substantial increase in NO_x emissions throughout East Asia has been inferred from satellite instruments in several previous studies (Mijling et al., 2013; Richter et al., 2005). The timing and extent of the reversal in NO_{ν} deposition that we see is also consistent with observed NO2 columns over eastern China derived from OMI (de Foy et al., 2016; Krotkov et al., 2016). The ability of our satellite-constrained NO_v deposition estimate to capture this sudden dramatic decrease over China, in contrast with previous projections (e.g., the RCP2.6, RCP4.5, and RCP8.5 projections to 2030, Lamaque et al., 2013), emphasizes an attribute of the satellite constraint.

Small statistically significant decreasing trends are found over the biomass-burning-dominated source regions of Africa. The decrease in NO_y deposition of about $\sim 3 \% \text{ yr}^{-1}$

relative to the long-term mean in northern Africa is consistent with the most recent GFED4 inventory from 1997 to 2014 (http://www.globalfiredata.org/), which has fire NO_x emissions decreasing at a rate of about 3 % yr⁻¹ in this region. In contrast, we also estimate a similar decrease in southern Africa that is not represented in the recent GFED4 emission time series. A reduction in NO₂ column abundance in this region (observed by GOME and SCIAMACHY) was also reported by van der A (2008). They postulate that this decline could be a result of deforestation leading to less biomass burning, but changing NO_x emission factors from biomass burning could also potentially contribute to the trend.

Despite the large regional trends described above, we find that global deposition changed very little between 1996– 1998 (56.1 Tg N yr⁻¹) and 2012–2014 (58.5 Tg N yr⁻¹) due to the opposing changes in different regions. Total NO_y deposition was lowest in 2006 (50.5 Tg N), and peaked at 60.8 Tg N in 2012. Since then, it appears that global NO_y deposition may be on the decline. Future observations in the coming years will be needed to establish whether this most recent decline is robust or temporary.

Figure 7 shows the calculated long-term trends in the measured wet NO₃⁻ deposition for locations across North America, Europe, and East Asia where at least 15 years of qualitycontrolled annual data are available (the coverage of these observation is the same as in Fig. 4). The observations over North America show the gradient in the trend from negligible in the west to steeply and significantly decreasing in the northeastern US and southeastern Canada. The steepest observed statistically significant trend $(-0.18 \text{ kg N ha}^{-1} \text{ yr}^{-2})$ occurs east of Detroit in southwestern Ontario, Canada. In Europe, only one of the gridded observations has a statistically significant trend $(-0.07 \text{ kg N ha}^{-1} \text{ yr}^{-2})$, located near the border of Denmark and Germany. The other locations in Europe show statistically insignificant trends over the longterm. In East Asia, we also we find that most of the stations record statistically negligible trends over the long term (only 2 of the 11 gridded observations have significant trends). The steepest observed trend in this region $(+0.39 \text{ kg N ha}^{-1} \text{ yr}^{-2})$ is found near Kuala Lumpur and is statistically significant.

We compare the long-term trends in these measurements with our satellite-constrained trends in wet NO_3^- deposition. We find a similar spatial gradient in North America, and the same magnitude of declines through the northeastern US and southern Ontario (-0.12 to -0.16 kg N ha⁻¹ yr⁻²). Over Europe, our estimates have low statistical significance in the trends throughout much of this region, consistent with the observations. Where we do see statistical significance (the northern United Kingdom, southern Denmark, and in some central to eastern European countries), observations are not available over the long-term for evaluation. In East Asia, our satellite-constrained trend estimates show statistically significant increases throughout much of the region (in contrast to most of the available observations). The trend over Kuala



Figure 6. Time series of annually integrated dry and wet NO_y deposition over specific regions (North America, Europe, and East Asia) as defined by the dashed rectangles in Fig. 5.



Figure 7. Long-term (1996–2014) trends in wet NO_3^- deposition from available regional network measurements (as in Fig. 4) and from the GEOS-Chem simulation, constrained by satellite observations of NO₂. Closed circles around the measurements indicate significant trends (p < 0.01); hatching indicates statistical significance in the simulation.

Lumpur is significant and positive $(+0.24 \text{ kg N ha}^{-1} \text{ yr}^{-2})$, as expected from the available measurements.

We again emphasize the small sample size in Europe (N = 16) and East Asia (N = 11). Moreover, in many cases trends in one (or both) datasets are small and/or insignificant. For these reasons, we focus on comparing the confidence intervals of the measured and satellite-constrained trends. We find that for 129 of the 136 gridded pairs (>90% of the data), the 95% confidence intervals overlap; of the pairs for which the intervals do not overlap, 3 (out of 109) occur in North America, with 1 (out of 16) in Europe and 3 (out of 11) in East Asia. For a large majority of the data in all three regions we therefore conclude that the satellite-derived trends are not significantly different from the trends inferred with ground-based measurements. Continued long-term measurements with better spatial coverage are imperative to better

evaluate long-term estimates of global NO_y deposition, especially throughout Europe and East Asia (but also in other parts of the world where long-term coverage is not available at all).

3.3 Changes in continental export of NO_y

NO_y deposition is a transboundary, and even intercontinental, issue (HTAP, 2010). In a multimodel study, Sanderson et al. (2008) found that between 3 and 10 % of NO_x emissions from Europe, North America, South Asia, and East Asia are ultimately deposited over foreign regions. Long-range transport events of NO₂ alone can be systematically detected by satellite observations (Zien et al., 2014). Here we extend such studies using our satellite-constrained long-term estimates of annual NO_y deposition to evaluate how the amount of NO_x

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exported from specific regions (i.e., the net balance between emissions and deposition over a land area) has changed over the last 2 decades.

Decreases in NO_y export over the Atlantic Ocean from North America and increases in export over the western Pacific Ocean from East Asia are evident in Fig. 5. We find that net export of NO_x from North America via atmospheric transport has decreased by more than 40 % (from 2.5 in 1996, to 1.4 Tg N yr⁻¹ in 2014). In contrast, we find that export of NO_x from Asia increased by 40 %, from 3.3 Tg N in 1996 to a maximum of 4.7 Tg N in 2011, with a subsequent decrease to 3.8 Tg N by 2014. As a result of these opposing trends, total deposition to the global oceans has changed remarkably little (25.0 Tg N yr⁻¹ in 1996–1998 compared to 24.4 Tg N yr⁻¹ in 2012–2014), but has experienced substantial regional redistribution.

 NO_{ν} export from North America has received considerable attention. Urban plumes from the eastern US that are transported across the North Atlantic for several days could still contain 20-50 ppb of reactive nitrogen oxides (Neuman et al., 2006). A recent detailed GEOS-Chem study of nitrogen deposition over the US estimated that net annual export of NO_x was around 38 % of NO_x emissions (or 2.5 Tg N) for 2006–2008 (Zhang et al., 2012). We estimate a similar fraction of export from the continental US using our observationally constrained simulation $(34\% \pm 2\%$ from 1996–2014), with a small decreasing trend from 35 % in 1996-1998 to 32 % in 2012–2014. As a result of declining emissions we find that absolute export from the continental US decreased by 50 % from 2.9 in 1996 to 1.5 Tg N in 2014. We find declines in NO_y deposition across the Atlantic Ocean, with small though statistically significant declines as far downwind as southern Greenland. The decreases downwind of the continent are clearest and most significant in the winter, spring, and fall (Fig. 5b, c, and e), while the trends are more local in the summer (when the NO_x lifetime is short and when midlatitude wind speeds are weaker).

We similarly calculate the net imbalance between NO_x emissions and NO_y deposition over western European countries and find a decrease of almost 40 %, from 2.2 to 1.3 Tg N. We calculate mean export of NO_x emissions from western European countries to be $45 \% \pm 4 \%$, with a notable decreasing trend from 50 % in 1996–1998 to 40 % in 2012–2014. As a result, the decrease in net export is steeper than the decrease in emissions from the region. As alluded to in Sect. 3.2, the decrease in NO_x export from some western European countries has likely compensated for increases in emissions in some of the central to eastern European countries, as well as in western Russia, where we find dry deposition has significantly increased, but wet deposition has decreased or shows no significant net trend.

Reactive nitrogen transport from Asia has previously been shown to contribute to O_x production across the midlatitude Pacific, reaching as far as the western coast of North America (Walker et al., 2010; Zhang et al., 2008), and major NO_x transport events from China can be indirectly observed by NO₂ columns (Lee et al., 2014). Our satellite-constrained estimate predicts that export from China alone more than tripled from 1.0 Tg N in 1996 to a maximum of 3.5 Tg N in 2011, then decreased to 2.5 Tg N by 2014. We estimate that an average of $24\% \pm 4\%$ emissions from China are exported, varying over time from as little as 15 % of emissions in 1998 to a maximum of 31 % of emissions in 2011 (with an overall increasing trend). Zhao et al. (2017) used a higher resolution $(0.5^{\circ} \times 0.667^{\circ})$ GEOS-Chem simulation and estimated that 36 % of China's NO_x emissions over 2008–2012 are exported. We calculate an export fraction of around 27 %for the same time period. The discrepancy between the two estimates may be attributed to the coarser horizontal resolution of our simulation $(2^{\circ} \times 2.5^{\circ})$, pointing to important resolution-dependent effects in global simulations of deposition. Other factors may include the use of different NO_x emissions (our satellite-constrained emissions indicate rapid change over this period of time), and the treatment of adjacent oceans.

Zhao et al. (2015) used GEOS-Chem to explore nitrogen deposition to the northwestern Pacific Ocean off the coast of China from 2008–2010. They estimated total (wet + dry) NO_y deposition of 6.9 and 3.1 kg N ha⁻¹ yr⁻¹ to the Yellow Sea and the South China Sea, respectively. Our simulation predicts that NO_y deposition to the same regions of the Yellow Sea increased from 5.1 in 1996 to 9.5 kg N ha⁻¹ yr⁻¹ by 2012 and to the South China Sea from 2.8 in 1996 to 4.3 kg N ha⁻¹ yr⁻¹ in 2011. Subsequent declines in the following years will hopefully have encouraging implications for nitrogen availability and the incidence of algal blooms in these regions (Hu et al., 2010).

Export of pollution from China has been shown to influence deposition over Japan in particular (e.g., Lin et al., 2008). Using observations of wet nitrate deposition, Morino et al. (2011) report increases throughout Japan from 1989-2008, and attribute this trend largely to transport from China. Likewise, integrated NO_v deposition over Japan increased (p < 0.01) in our satellite-constrained estimate. In fact, we find that Japan transitioned from a net "exporter" of NO_{ν} over 1996-2006 (emissions exceeded local deposition by up to 24 %) to a net "importer" of NO_v over 2007–2014 (local deposition exceeded emissions by up to 20%). The increase in deposition was dominated by statistically significant increases in wet deposition in some parts of the country. We find the increase over Japan is most uniform during the spring (Fig. 5, MAM), consistent with transport from China being pronounced during the spring season (Tanimoto et al., 2005). Nevertheless, the impacts of local NO_x controls can also be important. Dry deposition dominates the decline in annual NO_v deposition just west of Tokyo. Declines are seen throughout the southern part of the country during both the summer and fall seasons (Fig. 5, JJA and SON, September, October, and November). These results demonstrate the indirect relationship between local emissions and local deposition of NO_y for regions influenced by atmospheric transport and also show how long-term trends can depend strongly on the season and process (wet or dry deposition).

3.4 Sensitivity of NO_y deposition to NH₃ emissions

The transport and ultimate deposition of oxidized nitrogen may be tightly coupled with the reduced nitrogen $(NH_x = NH_3 + NH_4^+)$ and sulfate systems, due to the formation of NH₄NO₃ aerosol that becomes favorable once all H₂SO₄ has been neutralized (i.e., if there is "excess" NH₃). Examples of the resulting nonlinearity between PM_{2.5} concentrations and precursor emissions have been noted in the literature (Banzhaf et al., 2013; Derwent et al., 2009; Fowler et al., 2005). The formation of NH₄NO₃ aerosol at the expense of HNO3 with changing excess ammonia could therefore conceivably change the atmospheric lifetime of NO_{y} at the surface; accumulation-mode aerosol may have a dry deposition lifetime of days, whereas HNO₃ tends to have a dry deposition lifetime of shorter than a day. As a result, the predicted footprint of source impacts is sensitive to NH₃ emissions (Lee et al., 2016).

Contemporary emissions of NH3 are highly uncertain (Reis et al., 2009), so we perform a sensitivity experiment by perturbing (increasing) all anthropogenic and natural NH₃ emissions in the model by 25 % for the year 2012. Predicted NO_{ν} deposition from this simulation is compared to the predicted NO_{y} deposition in the 2012 simulation where NH_{3} emissions were not perturbed. Since we have not altered the emissions of oxidized nitrogen, simple mass balance dictates that increases in deposition over some regions will be countered by decreases elsewhere. Our perturbation is therefore to be interpreted as an experiment that tests how accurately the spatial pattern in NO_{ν} deposition at our model resolution can be predicted, given some uncertainty in NH₃ emissions. Given the horizontal resolution of our simulation $(2.5^{\circ} \times 2.0^{\circ})$, we acknowledge that our estimates of the sensitivity of NO_{ν} deposition to perturbations in NH₃ emissions may underestimate the importance of those interactions on finer spatial scales.

Figure 8 shows the results of this experiment. The sensitivity of NO_y deposition to an increase in NH₃ emissions is positive or negative depending on the region, while net deposition over the global domain does not change (to within 1– 2%). Over the continents, the sensitivity in total (wet + dry) NO_y deposition to the 25% perturbation in NH₃ emissions tends to be less than ± 5 %, with a few exceptions. We find differences in NO_y deposition on the order of 10% over parts of high-latitude Russia, northwest and central Africa, eastern China, southern South America, and Australia. However, with the exception of China, these are also regions where deposition is relatively low. We conclude that for most regions of interest, our satellite-constrained estimates of NO_y deposition over the continents and their trends will not be severely impacted by uncertainty in the NH₃ inventories.



Figure 8. Sensitivity of simulated NO_y deposition in 2012 to a 25 % perturbation (increase) in ammonia emissions in all grid boxes (shown separately for total deposition, dry deposition, and wet deposition).

Notably, the difference exceeds +50% over Myanmar, suggesting that simulated NO_y deposition over this country is extremely sensitive to changes in NH₃ emissions. It is clear from Fig. 8 that this results from a high sensitivity in dry deposition (middle panel) instead of wet deposition (bottom panel). Myanmar has some of the lowest estimated NH₃ emissions in all of South and East Asia (at least an order of magnitude lower than surrounding India, China, and Thailand), so this sensitivity reflects changes in the upwind emissions and subsequent transport of NO_y. We find the opposite sensitivity in nearby Cambodia, where the sensitivity of dry NO_y deposition to a 25\% perturbation is NH₃ emissions is -50%.

Over the oceans, the sensitivity of NO_y deposition to the 25 % increase in NH₃ emissions is generally low ($<\pm 5$ %), with the expected exceptions in areas that are directly offshore from major continental source regions. In the North Atlantic Ocean east of Canada and Greenland, the North Pacific Ocean off the coasts of China, Japan, and the South China Sea, the sensitivity of NO_y deposition is between 5 and 20 %. Our predicted decrease in dry NO_y deposition to the Yellow Sea given an increase in NH₃ emissions is consistent with previous adjoint analyses showing increased NO_y dry deposition in this region with a decrease in Asian NH₃ emissions (Zhao et al., 2015). Likewise, the sensitivity of deposition to the Mediterranean Sea is between 10 and 20 %. The differences in NO_y deposition over the oceans results from sensitivity in both dry and wet deposition (although in the case of the Mediterranean it is dominated by dry deposition). We conclude that although changes (or uncertainties) in NH₃ emissions can impact the distance of transport and deposition to oceans downwind of the major NO_x sources, the absolute magnitude of deposition is low where the sensitivity of NO_y deposition to NH₃ is relatively high.

3.5 Other considerations

A number of other uncertainties are important in an inversion of satellite NO₂ columns to calculate surface NO_x emissions and simulate long-term NO_y deposition. These can depend on, for example, the choice of inversion approach, errors in the satellite retrieval, and uncertainties in model processes (e.g., emissions, boundary layer mixing, chemical NO_x sinks, meteorology, and dry deposition).

Cooper et al. (2017) found that the finite mass-balance inversion approach used here can be improved upon by using an iterative method that performs with similar accuracy as a four-dimensional variational data assimilation. Multiconstituent data assimilation also shows considerable promise for constraints on surface NO_x emissions (Myazaki et al., 2017). Satellite retrieval algorithms continue to develop with advances that will improve the accuracy of future estimates of satellite-constrained NO_y deposition.

Uncertainties in model processes are also of interest. For example, uncertainties in the chemical sink of NO_x alone (e.g., the rate of HNO₃ formation, heterogeneous loss of N₂O₅ onto aerosol) can have a substantial impact on topdown emissions estimates (Stavrakou et al., 2013), suggesting more fundamental work in constraining these processes is required. Lin et al. (2010) found that top-down NO_x emissions estimates over East Asia are sensitive to other model uncertainties, including planetary boundary layer mixing scheme, lightning emissions, diurnal profile of emissions, and a priori NO_x , CO, and VOC emissions. Uncertainties in model meteorology are also important. For example, the MERRA precipitation fields used in our study are known to correlate weakly with observational datasets (Rienecker et al., 2011), but improvements can be expected from MERRA-2 due to the inclusion of gauge- and satellite-based precipitation corrections (Reichle et al., 2017). Finally, dry deposition schemes are also highly variable among models (Flechard et al., 2011; Hardacre et al., 2015), and future work in dry deposition evaluation should be a priority.

Nonetheless, despite these uncertainties we find a high degree of consistency between observations and our predictions in the long-term changes to deposition. Evidence continues to emerge about potential biases in bottom-up inventories (e.g., Travis et al., 2016), and our observational constraint on NO_x emissions mitigates against such biases. We expect that continued advancements in inversion approaches, satellite retrieval algorithms, and fundamental atmospheric chemistry processes will allow for increasingly accurate satellite-based constraints on deposition.

4 Conclusions

 NO_{y} deposition represents about half of the total reactive nitrogen deposited to Earth's surface. Even in the US where nitrogen oxide emissions have decreased substantially, constituents of NO_{ν} remain major contributors to the nitrogen deposited in areas of concern (Lee et al., 2016; Li et al., 2016). We applied NO₂ observations from multiple satellites over 1996–2014 together with the GEOS-Chem chemical transport model to estimate long-term changes to reactive nitrogen oxide deposition around the world. Given the effective global mass balance between NO_x emissions and deposition of reactive nitrogen oxides, we show that satellite constraints on NO_x emissions can provide a powerful topdown constraint on deposition in order to evaluate long-term changes worldwide. Observations from the GOME, SCIA-MACHY, and GOME-2 satellite instruments have provided continuous global coverage over the last 20 years, allowing observational constraints on NO_{ν} deposition that enhance the poor spatial coverage of ground-based deposition measurements.

We find substantial variability in regional trends of NO_{ν} deposition. NO_{ν} deposition declined most steeply throughout the northeastern United States at a rate of up to -0.6 kg N ha⁻¹ yr⁻², but has also decreased significantly throughout most of the country and in southern Canada. In Europe, statistically significant declines at a rate of up to -0.1 kg N ha⁻¹ yr⁻² are seen over some western countries. NO_v deposition has increased substantially throughout East Asia, exceeding a rate of $+0.6 \text{ kg N} \text{ ha}^{-1} \text{ yr}^{-2}$ in some parts. Since reductions in deposition over some regions were counteracted by increases in others, global NO_v deposition did not change considerably over the long term. However, we find that global NO_{ν} deposition could now be on the decline overall, since deposition in Asia peaked around 2010-2012. The ability to resolve the striking recent decline in NO_{ν} deposition in China (despite prior projections of increasing NO_x emissions) demonstrates one of the attributes of using a satellite-based constraint. Future observations will be important in evaluating whether this trend persists.

We find that changes over the last 2 decades in the export of reactive nitrogen oxides via atmospheric transport have impacted countries downwind of source regions. Export from North America has decreased by at least 40 %, while export from Asia has increased by the same relative amount. We find evidence that decreases in NO_x export from some western European countries have counteracted increases in local emissions from some eastern and central European countries, resulting in negligible net change in NO_y deposition over the long term. Likewise, Japan is highly sensitive to changes in export from China, but this depends strongly on the season and whether wet and dry deposition are both considered. While uncertainty in NH₃ emissions can impact the footprint of NO_y export and deposition, we show that this sensitivity is small in most regions of concern.

Direct measurements of deposition are sparse, inhibiting evaluation. This is especially challenging for global simulations, where individual measurements may not necessarily be regionally representative. Nevertheless, we find that for the vast majority of locations our satellite-derived trends are largely consistent with the observed trends. Expanded coverage of ground-based observations over the long-term is needed to more comprehensively evaluate long-term estimates of global NO_y deposition. This need also motivates the value of using alternative observational constraints such as the satellite NO₂ columns, as presented here.

Forthcoming satellite observations of NO₂ at higher spatial resolution (e.g., TROPOMI; Veefkind et al., 2012) and with diurnally varying observations (e.g., TEMPO; Zoogman et al., 2017, Sentinel-4, and GEMS) will offer increasingly robust constraints on NO_x emissions that affect NO_y deposition. Satellite observations of NH₃ (e.g., Van Damme et al., 2014) may offer additional opportunities to constrain the reactive nitrogen budget. Higher resolution global modeling will also be an important development to accurately account for nonlinear NO₂ losses in global emission inversions (Valin et al., 2011).

Our satellite-constrained estimates of NO_y document interannual changes over the past 2 decades worldwide. We expect that this information will be useful in future research into the impacts of nitrogen deposition to important biodiversity hotspots, in regions dealing with excessive nitrogen inputs leading to algal blooms, or for estimating the changing impacts of nitrogen deposition on global carbon uptake. Data availability. Wet NO_3^- deposition data from the World Data Centre for Precipitation Chemistry for 2000-2002 and for 2005-2007 can be accessed at http://www.wdcpc.org/assessment. Wet NO₃⁻ deposition data from the National Atmospheric Deposition Program in the United States can be accessed at http: //nadp.sws.uiuc.edu/data/. Wet NO₃ deposition data from the Canadian Air Precipitation Monitoring Network in Canada can be accessed at http://www.ec.gc.ca/rs-mn/default.asp?lang=En&n= 6C8C66C5-1. Wet NO₃⁻ deposition data from the Acid Deposition Monitoring Network in East Asia can be accessed at http: //www.eanet.asia/product/index.html. Satellite-derived NO2 tropospheric column density observations from GOME, SCIAMACHY, and GOME-2 can be accessed at http://www.temis.nl/airpollution/ no2.html. Our gridded satellite-derived estimates of global annual NO_v deposition can be downloaded from http://fizz.phys.dal.ca/ ~atmos/martin/?page_id=1520, or obtained by contacting the principal investigator, Jeffrey Geddes (jgeddes@bu.edu).

Appendix A

We simulate atmospheric chemistry from 1996 to 2014 using the GEOS-Chem chemical transport model (www. geos-chem.org) v9-02. Our simulations are driven with the MERRA meteorological product at a global horizontal resolution of $2.5^{\circ} \times 2.0^{\circ}$ and 47 vertical layers. GEOS-Chem includes detailed HO_x-NO_x-VOC-O₃-aerosol chemistry (Bey et al., 2001; Park et al., 2004), with isoprene chemistry following Paulot et al. (2009a, b) and gas-aerosol partitioning for the sulfate-nitrate-ammonium system calculated according to the ISORROPIA II equilibrium model (Fountoukis and Nenes, 2007). Gas-aerosol phase coupling occurs via N₂O₅ uptake (Evans and Jacob, 2005) and HO₂ uptake (Mao et al., 2013) in addition to other heterogeneous chemistry (Jacob, 2000) and aerosol effects on photolysis frequencies (Martin et al., 2003). Our simulations use time steps of 15 min for transport and convection, and 30 min for emissions and chemistry.

Removal by wet deposition occurs through scavenging in moist convective updrafts, as well as in-cloud and belowcloud scavenging during large-scale precipitation for watersoluble aerosol and gases (Amos et al., 2012; Liu et al., 2001). Removal by dry deposition is calculated based on the widely used resistance-in-series formulation from Wesely (1989), described for GEOS-Chem in Wang et al. (1998) and Zhang et al. (2001) for aerosol. Dry deposition of NO_y over the United States was recently explored and evaluated in detail by Zhang et al. (2012).

Anthropogenic emissions are prescribed by the NEI 2005 inventory for the United States (http: //www.epa.gov/ttnchie1/trends/), the CAC inventory for Canada (www.ec.gc.ca/indicateurs-indicators/default.asp? lang=en&n=E79F4C12-1), the BRAVO inventory for Mexico (Kuhns et al., 2005), the EMEP inventory for Europe (http://www.emep.int/), and Zhang et al. (2009) for China and Southeast Asia. Elsewhere, the EDGAR v3 emission inventory is used for anthropogenic NO_x , CO, and SO_x (Olivier et al., 2005); the GEIA inventory for NH₃ (Bouwman et al., 1997); and the RETRO inventory for VOCs (Hu et al., 2015). Monthly scaling of NO_x emissions is included in North America (based on the VISTAS inventory), Europe (based on the EMEP inventory), and Asia (based on the Zhang et al., 2009 inventory). Monthly scaling of EDGAR emissions is based on the seasonality from the Global Emission Inventory Activity (Benkovitz et al., 1996). Aircraft emissions are from the AEIC inventory (Stettler et al., 2011). Scale factors based on energy statistics following van Donkelaar et al. (2008) are used to scale NO_x , CO, and SO_x emissions between 1996 and 2010 for years when the emissions are unavailable from the inventory. For other species and for emissions beyond 2010, the closest available year is used. Biogenic VOC emissions are calculated using the MEGAN model (Guenther et al., 2006). Biomass burning emissions are according to the GFED3 inventory (Mu et al., 2011). Soil NO_x is calculated using the Berkeley–Dalhousie parameterization (Hudman et al., 2012). Lightning NO_x is implemented according to Murray et al. (2012). These a priori surface NO_x emissions are overwritten by our satellite-derived top-down estimates in the assessment of NO_y deposition.

Competing interests. The authors declare that they have no conflict of interest.

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