Rapid deposition of oxidized biogenic compounds to a temperate forest

Tran B. Nguyen1,2, John D. Crounse1, Alex P. Tenga, Jason M. St. Clair2, Fabien Paulotb,c, Glenn M. Wolfea,d,e, and Paul O. Wennberga,f

Divisions of 1Geological and Planetary Sciences and 2Engineering and Applied Science, California Institute of Technology, Pasadena, CA 91125; 3Geophysical Fluid Dynamics Laboratory, National Oceanic and Atmospheric Administration, Princeton, NJ 08540; 4Atmospheric and Oceanic Sciences, Princeton University, Princeton, NJ 08544; 5Atmospheric Chemistry and Dynamics Lab, NASA Goddard Space Flight Center, Greenbelt, MD 20771; and 6Joint Center for Earth Systems Technology, University of Maryland Baltimore County, Baltimore, MD 21250

Edited by Mark H. Thiemens, University of California, San Diego, La Jolla, CA, and approved December 22, 2014 (received for review September 28, 2014)

We report fluxes and dry deposition velocities for 16 atmospheric compounds above a southeastern United States forest, including: hydrogen peroxide (H2O2), nitric acid (HNO3), hydrogen cyanide (HCN), hydroxymethyl hydroperoxide, peroxycetic acid, organic hydroxy nitrates, and other multifunctional species derived from the oxidation of isoprene and monoterpenes. The data suggest that dry deposition is the dominant daytime sink for small, saturated oxygenates. Greater than 6 wt % C emitted as isoprene by the forest was returned by dry deposition of its oxidized products. Peroxides account for a large fraction of the oxidant flux, possibly eclipsing ozone in more pristine regions. The measured organic nitrates comprise a sizable portion (15%) of the oxidized nitrogen input into the canopy, with HNO3 making up the balance. We observe that water-soluble compounds (e.g., strong acids and hydroperoxides) deposit with low surface resistance whereas compounds with moderate solubility (e.g., organic nitrates and hydroxycarbonyls) or poor solubility (e.g., HCN) exhibited reduced uptake at the surface of plants. To first order, the relative deposition velocities of water-soluble compounds are constrained by their molecular diffusivity. From resistance modeling, we infer a substantial emission flux of formic acid at the canopy level (~1 mmol m2 s-1). GEOS–Chem, a widely used atmospheric chemical transport model, currently underestimates dry deposition for most molecules studied in this work. Reconciling GEOS–Chem deposition velocities with observations resulted in up to a 45% decrease in the simulated surface concentration of trace gases.

Significance

Dry deposition is an important removal mechanism for oxidized atmospheric compounds. This process remains, however, poorly understood due to the scarcity of direct flux observations for all but small, inorganic molecules in the atmosphere. The chemically speciated fluxes presented here comprise a unique and novel dataset that quantifies the dry deposition velocities for a variety of trace gases in a typical forested ecosystem. The data illustrate the key role of molecular diffusion in the atmosphere—biosphere exchange of water-soluble species. Furthermore, this work enabled evaluation of the dry deposition parameterization in a global chemical transport model. The results aid in resolving key discrepancies within the global model, resulting in more-accurate predictions of trace gas lifetimes and surface concentrations.

Author contributions: P.O.W. designed research; T.B.N., J.D.C., A.P.T., J.M.S.C., F.P., and G.M.W. performed research; J.D.C. contributed new reagents/analytic tools; T.B.N. analyzed data; and T.B.N. and P.O.W. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission. Freely available online through the PNAS open access option.

1B.N. and J.D.C. contributed equally to this work.

2To whom correspondence may be addressed. Email: wennberg@caltech.edu or tbn@caltech.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1418702112/-/DCSupplemental.
exchange behavior of OVOCs rely on parameterizations that have been optimized for O₃ and SO₂ (14). The higher water solubility of oxygenates compared with hydrocarbons increases their propensity to deposit to the surface by interacting with surface water layers; thus, it is unclear if the parameterization developed for less soluble species is sufficient to describe the flux behavior of water-soluble compounds. Direct measurements of dry deposition for key VOC oxidation products are clearly needed to complete the understanding of their atmospheric fates.

As the biosphere exchange of chemical species within the boundary layer is driven by turbulence, surface fluxes can be estimated with micrometeorological methods such as eddy covariance (EC). Calculations using the EC method require mixing ratio measurements performed fast enough to capture the high-frequency timescale of turbulent eddies; e.g., measurements should exceed 1 Hz sampling frequency at this site (SI Appendix, Fig. S7).

The recent development of fast mass spectrometry-based detectors facilitates EC flux measurements for a wide variety of species, including complex OVOCs (15). The EC flux of the desired species x \((F_x)\) above a surface, such as a forest canopy, can be expressed as the correlation of the mixing ratio of x with the vertical wind component (w),

\[
F_x = \overline{w'x'}
\]

where primes denote deviation from the mean value, and overbars denote an average over a typical flux period (~30 min). Important underlying assumptions in Eq. I are that (i) the measurement is stationary (i.e., no significant variations in the turbulent statistics in time or in the point of measurement relative to meteorological events), (ii) the surface is homogeneous so that the horizontal wind components are unimportant, and (iii) the scale of chemical change is slow in relation to flux variation (16). In this work, we focus on the downward exchange of trace gases, where the deposition velocity \((V_d, \text{cm s}^{-1})\) is defined in relation to the flux and the mean mixing ratio of x,

\[
V_d = \frac{F_x}{x}
\]

This velocity determines the deposition lifetime of compound x in the tropospheric boundary layer \((\tau_{dep} = hV_d, \text{where} \ h \text{ is the boundary layer height in centimeters})\). We report EC flux observations for 16 biogenic trace gases and energy balance closure above a southeast US temperate mixed forest in the summer. The organic and inorganic species studied here were chemically resolved with time-of-flight chemical ionization mass spectrometry using CF₃O⁻ reagent anions (CIMS), a soft-ionization method specific for the detection of hydroperoxides, acids, multifunctional nitrates, and multifunctional oxygenated compounds (17). The mixing ratios of compounds, meteorology, and three-dimensional wind \((u, v, \text{and} \ w)\) were measured atop a 20-m walk-up tower in the Talladega National Forest in June 2013 as part of the Southern Oxidant and Aerosol Study (SOAS) campaign (SI Appendix, section 1).

Results and Discussion

Fluxes are calculated from the fast measurements of chemical mixing ratios (10 Hz), virtual temperature (8 Hz), and vertical winds (8 Hz). Details of the site, measurement methods, calibration (SI Appendix, Table S1), and quality assessment/quality control of flux data can be found in Materials and Methods and SI Appendix. The site location and tower sampling platform presented challenges to the flux measurements due to inconsistencies in fetch to the south and perturbations of wind trajectory in this direction due to shadowing by the instrument. Accurate flux measurements were possible on several days, however, when the wind direction was ideal to sample air from the forest to the north. The measurement of solar radiation, externally calibrated and converted to net radiation (18), was used as one verification of flux quality (16). The closure condition requires that the observed net radiation downward \((R_{n})\), less the storage heat flux \((S)\) (19), is equivalent to the observed sensible heat flux upward \((SH, \text{heat that can be measured})\) plus latent heat flux upward \((LE, \text{heat that is used to evaporate water})\). The quality constraints (SI Appendix, section 4) were met for only a subset of measurements (~16%). We restrict our analysis to these periods.

Our observations demonstrate that dry deposition is a significant atmospheric sink for many OVOCs formed in the dark chemical and photochemical reactions of isoprene and monoterpenes. Gas-phase inorganic compounds are also observed to deposit, often quite rapidly. Fig. 2 and SI Appendix, Fig. S17, show downward exchanges of relevant OVOCs and inorganic species and upward exchanges of heat and water for the days in...
Fig. 2. The balance of latent heat (LE) and sensible heat (SH) fluxes compared with net radiation (Rn) minus heat storage (S), virtual temperature and absolute water vapor, and fluxes and mean mixing ratios for select chemical species. SI Appendix, Fig. S17, shows the remaining fluxes and mean mixing ratios measured in this work.

Fig. 3 shows the observed mean deposition velocities ($V_d$) for each compound binned into hourly periods. Daytime $V_d$ values (Fig. 3 legend) were averaged for local hours 1000–1500 across all measurement days. In most cases, the deposition velocities are more rapid than that of $O_3$ (21). Diurnal patterns of deposition velocities largely mirrored those for flux. However, the diurnal patterns for some compounds are asymmetric (peaking before noon), which may indicate that their biosphere exchange is controlled in part by stomatal conductance and/or surface water availability. Although daytime $V_d$ may depend to some extent on environmental conditions and canopy characteristics at each site, it is instructive to compare the values measured here with available data on dry deposition over forests. For example, in this work, the observed $V_d$ for H$_2$O$_2$ ($5.2 \pm 1.1$ cm s$^{-1}$) is in excellent agreement with other direct measurement accounts ($V_d$R$_{H_2O_2} = 5 \pm 2$ cm s$^{-1}$ (22–24)). Measured fluxes of formic acid have been reported to be bidirectional, with an effective range of $V_{d,\text{formic}} = 0.17$–1.0 cm s$^{-1}$ (25, 26) that is in reasonable agreement with our determination ($1.0 \pm 0.4$ cm s$^{-1}$). Measurements of nitric acid (HNO$_3$) dry deposition encompass the largest range in the literature [$V_d\text{HNO}_3 = -1.8$ cm s$^{-1}$ (27–30)]. HNO$_3$ is a challenging compound to measure because of its interactions with instrument surfaces. The CIMS measurement of HNO$_3$ was affected by an instrumental delay and was corrected with a time response perturbation function (SI Appendix, section 5). Our determination for nitric acid deposition velocity ($V_d\text{HNO}_3 = 3.8 \pm 1.3$ cm s$^{-1}$) is within range of other reports and is consistent with results from resistance modeling performed in this work. Other speciated $V_d$ we observe compare well, although indirectly, to previously reported data available as averaged measurements within the class of organic hydroperoxides [$V_d\text{ROOH} = 1.1–2.5$ cm s$^{-1}$ (22, 23)] or organic nitrates [$V_d\text{RONO}_3$ (gas + particles) $\approx 2.0$ cm s$^{-1}$ (29, 30)].

Resistance Model of Deposition. Here we seek to evaluate the physical factors that control dry deposition within the framework of resistance modeling. Deposition velocity is usually parameterized as a series of resistances in the transport from the atmosphere to a surface—in this case, a canopy that is represented as a theoretical large leaf (31). The downward trajectory of a chemical species can be hindered by the step-wise resistances to aerodynamic transfer ($R_a$), molecular diffusion through the quasi-laminar boundary layer of the leaf ($R_b$), and uptake at the surface of the leaf ($R_c$).

$$V_d = \frac{1}{R_a + R_b + R_c} \quad [3]$$

SI Appendix, section 6, describes the resistance model in detail. We used $R_a$ and $R_b$ parameterization as suggested (32–34). $R_c$ is dependent only on the atmospheric stability (i.e., turbulence). $R_a$ is inversely dependent on the turbulence, the molecular diffusivity of compound $x$ in air [$D_x$ (m$^2$·s$^{-1}$)], and canopy characteristics such as the observed leaf area index (LAI) of the canopy at the measurement site (LAI = 4.7; see SI Appendix). $R_b$ is parameterized according to a revised Wescley scheme (35), which, to first order, considers the combined resistances from stomata ($r_s$), mesophyll ($r_m$), and cuticles ($r_w$) of plants. The surface uptake of
a particular compound is represented as functions of \( \text{D}_w \), water solubility (\( H \)), Henry’s Law coefficient, and a reactivity coefficient (\( f_0 \), range 0–1 where 1 is as reactive as ozone),

\[
R_c \sim \left( \frac{1}{r_c(D_w) + r_m(H, f_0)} + \frac{1}{r_m(H, f_0)} \right)^{-1}.
\]

The original Wexley parameterization overestimates \( R_c \) for compounds like \( \text{H}_2\text{O}_2 \) (20), which should be close to zero (22), despite the fact that water solubility of \( \text{H}_2\text{O}_2 \) is not significantly variable below \( \text{pH} \approx 8 \) and \( f_0 \) is set to maximum. With empirical results at hand, we optimized the parameters in the Wexley \( R_c \) scheme to minimize discrepancies between calculated and observed fluxes. We adjusted coefficients only in the \( r_m \) and \( r_c \) terms (SI Appendix, Eqs. 14–16) to be more sensitive to water solubility. Furthermore, we use the simple Henry’s Law coefficients (\( H \)) that describe physical solvation in pure water, instead of effective Henry’s Law coefficients (\( H^* \)). We find that \( H \) satisfactorily reproduces empirical \( R_c \). Furthermore, \( H \) is a more practical parameter as the knowledge of ionic activity (36, 37) and \( \text{pH} \) at surface aqueous layers is not readily available to estimate \( H^* \) for the species where solubility enhancements are important (e.g., aldehydes, acids, and bases). We neglect \( \text{pH} \) dependence for solubility because \( \text{pH} \) inside plant mesophyll (38) and leaf surface (39) tend to be buffered near neutral. \( \text{H}_2\text{O} \) was estimated based on chemical proxies for the larger multifunctional compounds (SI Appendix, Table S2) where measured values were unavailable (40).

Fig. 4 shows that results from the revised resistance model are in excellent agreement with ambient observations for the majority of the compounds studied in this work. The poorer model agreement for the mean of ISOPoOH + IEPOX is likely due to error in flux measurement and not modeling. Deposition velocity for IEPOX may be slightly underrepresented in our work because the specified time response delay in stomata \( r_s \) and \( r_c \) terms was uncorrected (SI Appendix, section 5). Surface resistances are small for the water-soluble and more reactive species (e.g., strong acids, hydroperoxides, and peroxyacids) but become important for other compounds. The parameterization captures observed \( V_d \) successfully on both ends of the water-solubility spectrum, e.g., for poorly water-soluble compounds like HCN (\( H \approx 10 \text{ M} \cdot \text{atm}^{-1} \) (40)) and for highly water-soluble compounds like HMHP (\( H \sim 10^6 \text{ M} \cdot \text{atm}^{-1} \)).

The exception is formic acid, where \( V_d \) predicted by the resistance model is approximately twice the observed value.

The behavior of formic acid cannot be explained by the physical characteristics of this molecule. With the assumption that the resistance model results are robust, poor model agreement implies that error may exist in the formic acid measurement, solubility estimations, or the assumption that formic acid has no emission source. A time response test suggests that >98% of the formic acid flux was captured by the measurement (SI Appendix, Fig. S11) and \( H \) is not expected to be overestimated for formic acid. More likely, we believe that a significant portion of the observed formic acid in a forested setting is produced by secondary, possibly heterogeneous, reactions (6, 41). These secondary emission pathways are additional to the proposed primary emission of formic acid from biogenic, anthropogenic, and pyrogenic sources. The discrepancy between modeled and observed \( V_d \) of formic acid requires a large emission flux—on the order of 1 nmol m\(^{-2}\) s\(^{-1}\) in the daytime.

One possible route to secondary formic acid formation may include water-mediated degradation of larger OVOCs that are deposited on canopy surfaces or on aerosol surfaces, e.g., the oxidation of IEPOX (42) and decomposition of HMHP (43). The deposition of HMHP and IEPOX is likely independent of surface chemistry at this site because surface uptake is not a limiting process for these compounds (Fig. 4, Lower). Thus, their in-canopy chemistry is not expected to lead to model/measurement disagreement, within the error of the measurement, by significantly enhancing deposition. HMHP decomposition (downward flux of \( \sim 0.2 \text{ nmol m}^{-2}\text{s}^{-1} \)) may produce formic acid via thermal, photochemical, or dark decomposition pathways. The lifetime of HMHP with respect to hydrolysis at neutral \( \text{pH} \) has been measured to be roughly 17 min (44), on the timescale of mixing within the canopy. The formation yield of formic acid from IEPOX oxidation is significant in the gas phase (10–30%) (42), although the lifetime of IEPOX in the gas phase is long (>10 h). We speculate the yields are higher in the aqueous phase; however, more work is needed before the contribution of IEPOX to the formic acid flux can be estimated.

Another possible formic acid source is the rapid in-canopy ozonolysis of monoterpenes and other biogenic VOCs (45, 46), which may occur primarily within stomata (47) and is enhanced under humid conditions (43). For example, the in-canopy ozonolysis of conifer terpenes (\( r_s \approx 1–2 \text{ min} \)) has been estimated to consume 0.6–1.5 nmol m\(^{-2}\) s\(^{-1}\) (7–28%) of the downward ozone flux and 46–61% of the sesquiterpenes mass in Amazonia (48). Approximately 6% of the ozone flux at this site is required to account for the entire formic acid flux. Although the kinetics and yields of the myriad formic acid formation routes are not well understood, a 1 nmol m\(^{-2}\) s\(^{-1}\) secondary source of formic acid from these combined pathways appears plausible.

The calculated resistances shed light on a remarkable empirical relationship of highly depositing compounds—their observed daytime deposition scale with inverse molecular mass (\( \sqrt{\text{m}} \) \(^{-1}\)) (Fig. 5). These trends suggest that molecular diffusivity through the leaf boundary layer and, for larger molecules, from the leaf surface itself acts as a constraint on dry deposition at a specific turbulent condition. According to the resistance model, diffusivity restrictions arise from the terms describing transport to the quasi-laminar layer of the leaf surface (\( R_s(R_s) \)) or through stomata (\( r_s \) component of \( R_s \)). We find two “diffusion-limited” upper bounds: when there is no resistance at the surface of the leaf (\( R_{\text{total}} = R_s + R_c \), Fig. 5, dashed line) and when there is a stomata-controlled resistance at the surface of the leaf (\( R_{\text{total}} = R_s + R_c + R_s \), \( r_s \), solid line). The two limits converge for smaller compounds (<80 Da) with high diffusivity such that \( r_s \) is minimal. As expected, the observed daytime \( V_d \) for H\( \text{NO}_3 \) (the prototypical molecule for rapid deposition) is at maximum for its size. It appears that bulkier compounds such as MTNP cannot achieve surface resistance-free deposition because they are hindered by subsequent transport through the stomata.

![Fig. 4.](image-url)
Efficiently-depositing compounds approaching the stomata-controlled diffusion limit generally belong to the chemical classes of simple or multifunctional hydroperoxides and strong acids with high water solubility (threshold $H$ in this model $\approx 3 \times 10^3$ M·atm$^{-1}$). The model sensitivity to various parameters is discussed in SI Appendix, section 6, and shown in SI Appendix, Fig. S16, for $H$. Organic nitrates and peroxyacids are close to the upper bound. Most of the major outliers in Fig. 5, e.g., HCN and ketones, have poor solubility in water or low reactivity at the leaf surface, which produces a substantial nonstomatal $R_d$ term. In particular, uptake of HCN from the surface comprise >90% of its total resistance. Formic acid is also an outlier on this plot, likely due to the aforementioned emission. In addition to diffusion restrictions, the absolute deposition velocities for a specific compound remain highly dependent on atmospheric stability. The aerodynamic resistance ($R_d$) can be significant when the atmospheric stability is high; however, this term is identical for all compounds, so it will materialize as a vertical offset in Fig. 5. Under exceptionally high turbulent mixing conditions ($R_d \rightarrow 0$), we estimate that $V_d$ for HNO$_3$ and H$_2$O$_2$ can reach $\sim$6 cm·s$^{-1}$ and 8 cm·s$^{-1}$, respectively, in this forest setting.

The empirical relationship between mass and deposition velocity for soluble species serves as a useful paradigm for understanding previous and future measurements performed under differing conditions. That a single physical characteristic of atmospheric trace gases constrains their deposition velocities in relation to each other has not yet been fully realized. Current sensitivity simulations of atmospheric loss often exceed these “diffusion-limited maxima” when modeling dry deposition of OVOCs. For example, HNO$_3$ is typically modeled to have the highest relative deposition velocity, and sensitivity studies tend to equate $V_d$ of other compounds that may also deposit quickly (e.g., ISOPN) to $V_d$ of HNO$_3$ (7, 49). These data suggest that it may be more appropriate to scale $V_d$ of highly depositing species in sensitivity simulations with respect to H$_2$O$_2$ or HNO$_3$ based on molecular mass. Furthermore, it appears that H$_2$O$_2$ will always deposit faster than HNO$_3$ at any given turbulent condition.

**Impact on Regional and Global Budgets of Trace Gases.** Global atmospheric transport models used to assess chemistry and climate feedbacks often overpredict surface volume mixing ratios for trace gases such as H$_2$O$_2$ (50, 51), HCN (52), and ISOPN (5). Depending on the molecule, such model discrepancies can affect the entire troposphere or only the boundary layer. These overestimations have motivated hypotheses about unknown atmospheric sinks for these compounds or their precursors, e.g., enhanced aerosol uptake of HO$_2$ due to Fenton chemistry that does not produce H$_2$O$_2$ in the gas phase (51) or an ocean sink for HCN (52).

We performed regional and global simulations using GEOS–Chem v9.2 (53), a global 3D chemical transport model, to assess the importance of dry deposition in model simulations of surface mixing ratios. GEOS–Chem simulations were driven by GEOS-FP (forward processing) meteorology with hourly resolution (SI Appendix, section 7). Compound-specific surface mixing ratio and vertical profile impacts in response to modeled changes in dry deposition are a nonlinear result of chemistry, land use, and meteorology; thus, they are not possible to predict a priori. In one case study, we performed a 1-mo simulation at a fine-grid horizontal resolution of 0.25° × 0.33° to facilitate comparisons with hourly observations of deposition velocities and meteorology at the measurement site. Two year-long global simulations at a coarser resolution (2° × 2.5°) were also performed for a “base” case and “sensitivity” case. In the base simulation, we use the default GEOS–Chem dry deposition velocities, while in the sensitivity simulation, we adjust the average daytime $V_d$ of several compounds to match measurements. We use the most recent version of GEOS–Chem chemical mechanism with a few updates (SI Appendix, Table S3) to reflect revised knowledge of OVOC reactions published very recently, such as updated ISOPN + OH and ISOPN + O$_3$ rate coefficients (54). Model results for HCN and several other species of interest are not available because these compounds are either not included in GEOS–Chem (HMHP, HCN, HPALDs, HDC$_3$, DHC$_3$, INP, and MTNP) or because their dry deposition was neglected in the model (PAA). Among the compounds not included in the simulations, HMHP (55) and HPALDs (56) may have the more significant impact on model predictions because of their high deposition velocities and large abundance in forested settings and in biomass burning plumes.

Fig. 6 shows the meteorological conditions and energy fluxes generated by GEOS-FP and deposition velocities calculated by the fine-grid GEOS–Chem simulation for June 2013 at the coordinates of measurement site (CTR). Excellent agreement in both the mean magnitudes and diurnal pattern was obtained for the comparison of friction velocity (turbulence, Fig. 6A), sensible heat flux (Fig. 6B), and latent heat flux (Fig. 6C) between the model (averaged for 30 d) and measurements (averaged for 5 d). This suggests that the meteorological and surface exchange conditions during the select days used in this work are representative of the site during June. Furthermore, these simulations suggest that flux observations during the SOAS campaign are not atypical for the eastern US region or for the year 2013 (SI Appendix, Figs. S18 and S19, respectively).

The deposition velocities for compounds of interest showed poorer agreement in the fine-grid simulations (Fig. 6 D–F, black lines). As meteorology and land surface processes are reasonably well represented in the atmospheric model, the discrepancy likely resides in the parameterization of surface uptake resistance ($R_d$) by the canopy. The GEOS–Chem model uses the Wexler scheme (Eq. 4) to calculate surface resistance for chemical species. As noted earlier (20), $R_d$ for H$_2$O$_2$ and certain OVOCs are overestimated compared with observations, leading to low values of $V_d$ simulated by the model. For the sensitivity study, we arbitrarily increased the effective solubility inputs for compounds of interest (SI Appendix, Table S4) until the calculated $V_d$ matched observed values (Fig. 6 D–F, blue lines, when an applicable change is made). In some cases, the required solubility adjustments are clearly too large compared with uncertainties and suggest that a change to the parameterization, such as was done in Resistance Model of Deposition, should be implemented to improve the model description of dry deposition in future work. This method, however, is a reasonable temporary solution that can be
immediately implemented in atmospheric models while an updated parameterization is under development.

Table 1 shows simulation results extracted for June 2013 from the base and sensitivity studies in the grid box that encompasses the coordinates of the measurement site. Changes in $V_d$ significantly impact predictions of surface concentration for the depositing species, but secondary chemistry, e.g., ozone production or SO$_2$ oxidation, are not considerably affected. The surface mixing ratios for compounds with adjusted $V_d$ decreased $\sim$10–50%. Although the observed changes in modeled mixing ratios were not found to affect O$_3$ and HO$_3$ (OH + HO$_2$) at this site, season, and altitude (<1 km), effects on O$_3$ and HO$_3$ due to the photochemistry of isoprene nitrates and H$_2$O$_2$, respectively, may be more significant for other regions of the globe and times of the year (51, 57). The adjustment in $V_d$ for H$_2$O$_2$ was the largest (a factor of 7 increase), and its surface mixing ratio was correspondingly reduced by the greatest amount (45%). Higher H$_2$O$_2$ dry deposition significantly affected simulation results of its surface concentration for much of the forested areas of the globe (Fig. 7), e.g., Canadian and European boreal forests, the Amazon basin, and the Congo basin. Measurements at these sites are not available for model comparison; however, the results suggest that underrepresentation of dry deposition for H$_2$O$_2$ is likely responsible for much of past overestimations of its mixing ratios in GEOS–Chem simulations (51).

The sensitivity of H$_2$O$_2$ boundary layer mixing ratio to dry deposition may be expected because this compound has a long lifetime with respect to oxidation and photolysis in the lower atmosphere. To estimate dry, midday sinks, we use measured daytime deposition velocities (Fig. 3), simulated [OH] $\approx$ 2.6 $\times$ 10$^6$ cm$^{-3}$ (twice the 24-h value in Table 1), simulated photolysis under tropical noon with 300 Dobson units ozone (58), and a boundary layer height of 1.5 km. The deposition lifetime of H$_2$O$_2$ ($\tau_{dep}$ $\approx$ 8 h) accounts for $>70\%$ of its total atmospheric loss under dry conditions [$\tau_{hv}$ $\approx$ 1.4 d; $\tau_{OH}$ $\approx$ 2.6 d (59)]. Reacton with ozone, usually several orders of magnitude slower for unsaturated compounds and unimportant for saturated compounds, was neglected as a loss process. Dry deposition of other saturated compounds can be expected to dominate their atmospheric removal in a manner similar to H$_2$O$_2$ because their reactive lifetimes in the atmosphere are long and their water solubilities are high. For example, HMHP ($\sim$4 cm s$^{-1}$) has a deposition lifetime of 10 h, accounting for $\sim$55% of its atmospheric removal [$\tau_{hv}$ $\approx$ 4 d, $\tau_{OH}$ $\approx$ 14 h (estimated based on methylhydroperoxide+OH rate coefficient) (60)]. Similarly, dry deposition may comprise $\sim$60% of the atmospheric loss for PAA (59, 61) and $\sim$45% of the atmospheric loss for IEPOX (42). IEPOX is also removed by reactive uptake to aerosols. Using data from a recent

![Table 1. GEOS–Chem base and sensitivity (sens.) simulation for the grid box representing the Brent, AL, field site, 2° × 2.5° resolution](image)

<table>
<thead>
<tr>
<th>Tracer name</th>
<th>Base $V_d$* cm s$^{-1}$</th>
<th>Sens. $V_d$* cm s$^{-1}$</th>
<th>Base conc. (70 m)</th>
<th>Sens. conc. (70 m)</th>
<th>$\Delta$ %</th>
</tr>
</thead>
<tbody>
<tr>
<td>OH</td>
<td>–</td>
<td>–</td>
<td>1.32 $\times$ 10$^6$</td>
<td>1.30 $\times$ 10$^6$</td>
<td>–2</td>
</tr>
<tr>
<td>HO$_2$</td>
<td>–</td>
<td>–</td>
<td>3.84 $\times$ 10$^8$</td>
<td>3.77 $\times$ 10$^8$</td>
<td>–2</td>
</tr>
<tr>
<td>NO</td>
<td>–</td>
<td>–</td>
<td>0.12</td>
<td>0.12</td>
<td>0</td>
</tr>
<tr>
<td>NO$_2$</td>
<td>–</td>
<td>–</td>
<td>0.98</td>
<td>0.98</td>
<td>0</td>
</tr>
<tr>
<td>O$_3$</td>
<td>–</td>
<td>–</td>
<td>50.97</td>
<td>50.63</td>
<td>–1</td>
</tr>
<tr>
<td>PAN</td>
<td>–</td>
<td>–</td>
<td>0.38</td>
<td>0.37</td>
<td>–2</td>
</tr>
<tr>
<td>HNO$_3$</td>
<td>3.02</td>
<td>3.02</td>
<td>0.46</td>
<td>0.45</td>
<td>–1</td>
</tr>
<tr>
<td>H$_2$O$_2$</td>
<td>0.37</td>
<td>2.66</td>
<td>2.83</td>
<td>1.55</td>
<td>–45</td>
</tr>
<tr>
<td>PPN</td>
<td>–</td>
<td>–</td>
<td>0.019</td>
<td>0.018</td>
<td>–2</td>
</tr>
<tr>
<td>HCHO</td>
<td>–</td>
<td>–</td>
<td>3.64</td>
<td>3.60</td>
<td>–1</td>
</tr>
<tr>
<td>MHP</td>
<td>–</td>
<td>–</td>
<td>1.18</td>
<td>1.15</td>
<td>–2</td>
</tr>
<tr>
<td>SO$_2$</td>
<td>–</td>
<td>–</td>
<td>1.20</td>
<td>1.21</td>
<td>1</td>
</tr>
<tr>
<td>SO$_4$</td>
<td>–</td>
<td>–</td>
<td>0.53</td>
<td>0.52</td>
<td>–2</td>
</tr>
<tr>
<td>ISOPN</td>
<td>0.30</td>
<td>0.82</td>
<td>0.081</td>
<td>0.073</td>
<td>–10</td>
</tr>
<tr>
<td>MACN+MVKN</td>
<td>0.30</td>
<td>0.81</td>
<td>0.075</td>
<td>0.069</td>
<td>–9</td>
</tr>
<tr>
<td>PROPN</td>
<td>0.30</td>
<td>0.48</td>
<td>0.055</td>
<td>0.047</td>
<td>–14</td>
</tr>
<tr>
<td>HAC</td>
<td>0.30</td>
<td>0.71</td>
<td>1.48</td>
<td>1.25</td>
<td>–15</td>
</tr>
<tr>
<td>ISOPOOH</td>
<td>0.76</td>
<td>0.76</td>
<td>0.38</td>
<td>0.38</td>
<td>0</td>
</tr>
<tr>
<td>IEPOX</td>
<td>2.86</td>
<td>2.73</td>
<td>0.58</td>
<td>0.59</td>
<td>1</td>
</tr>
</tbody>
</table>

Concentrations (conc.) are reported in parts per billion by volume, except OH and HO$_3$, where units are in molecules per cubic centimeter. Percent change is defined with respect to the base simulation.

*The 24-h average (approximately half of daytime value).
laboratory study of IEPOX particle uptake probabilities (62), we estimate that the dry deposition fate of IEPOX is significantly affected only in the presence of very acidic inorganic sulfate particles at pH < 1 (reduced to ~20%) and is relatively unaffected if the particles have more moderate acidities (pH > 3).

For unsaturated compounds like ISOPN, deposition may comprise a smaller fraction (<10%) of the daytime sink because its reaction with OH is fast (54). Thus, errors in representation of ISOPN dry deposition will have a much smaller effect on simulated ISOPN surface concentrations. In general, while dry deposition is less important for organic nitrates than peroxides, changes in their mixing ratios obtained from the sensitivity study were significant for both classes of compounds. Mixing ratios for organic nitrates from isoprene were reduced by 9–14% in the sensitivity study. The carbonyl nitrates (in particular MACN) may be reasonably photolabile (63), further diminishing the importance of dry deposition. Smaller multifunctional carbonyls like HAC are longer-lived in the atmosphere. Correspondingly, their surface concentration reductions (15%) in the sensitivity study are larger than for ISOPN.

The differences in H$_2$O$_2$ and other trace gas mixing ratios in the troposphere between the base and sensitivity studies were most significant within the boundary layer (<3 km). Importantly, the concentration gradient is larger near the surface when $V_d$ is higher (SI Appendix, Fig. S20), leading to more pronounced differences between the base and sensitivity studies at the measurement height of ~20 m. As the lowest vertical layer in GEOS–Chem corresponds to an average height of 70 m, we linearly extrapolated the surface gradient to the tower height for compounds of interest to compare with measurements. Compared with mixing ratios measured at SOAS [24-h mean ~ 0.5 (±30%) parts per billion by volume (ppbv) on campaign days without significant precipitation], H$_2$O$_2$ mixing ratios are overpredicted by more than 300% in the base simulation and are much closer in the sensitivity simulations, but still overpredicted by ~85%. Daytime values for other trace gases are still overestimated by 30–300% in the model for the CTR measurement site; however, all mixing ratios in the sensitivity study are substantially closer to measured values at SOAS when extrapolated to the measurement height. The errors in overestimations are reduced by factors of 1.5–4.

The most dramatic overestimation occurred with IEPOX, where 24-h mean values at SOAS are measured to be ~0.09 ppb with tandem mass spectrometry and simulated to be ~0.35 ppb (extrapolated to 20 m). Unlike H$_2$O$_2$, this result is not attributed to underestimation of dry deposition. In fact, the simulated $V_d$ in GEOS–Chem for IEPOX is ~70% higher than results from the adjusted resistance model we used in this work. Interestingly, $V_d$ simulated in GEOS–Chem for ISOPOOH is ~50% lower than our estimates. IEPOX deposition velocity should be limited by molecular diffusion due to its high water solubility (Fig. 5), which suggests a lower daytime $V_d$ than that simulated by GEOS–Chem (~5.5 cm·s$^{-1}$). The average $V_d$ for these isobaric species was not adjusted in our sensitivity simulations because their mean simulated $V_d$ is similar to the observed value due to the neglecting errors. More likely, the low observed mixing ratios of IEPOX at SOAS may be due to error in its production rate or because heterogeneous loss reactions, such as conversion to organic aerosols in anthropogenically impacted particle liquid water (64), were not accounted for in the model. IEPOX uptake clearly involves hydronium-catalyzed ring opening in aerosol liquid water (62, 65, 66). However, field results suggest that more factors than acid catalysis govern the chemistry (67). Laboratory results indicate that hydrated ammonium sulfate particles (but not sodium or magnesium sulfate) produce a similar magnitude of organic aerosol mass due to IEPOX uptake regardless of whether acid was added (68, 69). More work is needed to elucidate field observations and update atmospheric model mechanisms.

Impact on Ecosystems. The field observations suggest that a significant portion of VOC emitted by the biosphere is returned to the ecosystem via dry deposition of OVOCs. Fig. 8, Left, shows the net mean deposition of C, N, and O atomic mass, obtained from the speciated contributions of 16 molecular formulas observed, averaged across multiple days. As our measurements encompass only a subset of organic species present in the troposphere, all estimations of combined fluxes are lower limits. We observed that at least 6% of the reactive carbon mass that was emitted as isoprene in an AL forest (=4 mgC·m$^{-2}$·h$^{-1}$ daytime (72), integrated over 24 h) were removed by the canopy via dry deposition of isoprene’s oxidation products. Our analysis does

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|}
\hline
Component & Mass Flux (X m$^{-2}$ h$^{-1}$) & % Deposited & N Dioxide & Organic Nitrates HNO$_x$
\hline
Carbon & C & 25 & 25 & 50
Oxygen & O & 30 & 30 & 30
Nitrogen & N & 20 & 20 & 20
\hline
\end{tabular}
\end{table}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure8}
\caption{(Left) Net carbon, oxygen, and nitrogen downward flux by mass, calculated from molecular formulas for compounds shown in SI Appendix, Table S1, and averaged over the dates in the study. (Right) Percent contribution of organic nitrates and nitric acid to the measured oxidized nitrogen flux.}
\end{figure}
not include HCN and MTNP. This estimate of the contribution of dry deposition removal of isoprene carbon may be larger if the many isoprene-derived OVOCs that are expected to deposit are included in the analysis, e.g., ethanol nitrates, various acyloxy-nitrates (APNs), acetic acid, glycolaldehyde, formaldehyde, glyoxal, methylglyoxal, acetaldehyde, and other OVOCs.

Oxidation products contribute most of the mass flux due to the combined effect of their larger molecular weight from the incorporation of oxygen during oxidation (73), and the propensity for less-volatile and more-water-soluble compounds to deposit. The oxidant deposition observed by our instrument is attributed to hydroperoxides (−OOH), although we are missing the likely minor contribution of methyl hydroperoxide (MHP). Organic hydroperoxides can be formed primarily from the RO2 reaction with HO2, more prevalent in pristine environments, or from humid ozonolysis, more prevalent in urban-influenced environments (Fig. 1). Thus, oxidant deposition is highly dependent on geographical location and time of day. However, ozone has been considered as the predominant oxidant in modeling dry deposition or in estimating oxidant-induced plant damage. The apparent effects of ozone on forest ecosystems are well documented, but they may not be directly attributable to ozone. Ozone concentration rapidly declines after stomatal uptake, and damaging effects in plants are often linked to the production of secondary stressors that are dependent on ozone dose (74). These secondary stressors were suggested to be, in part, H2O2, HMHP, and other hydroperoxides that are produced as part of alkene ozonolysis (46). It should be noted that both H2O2 and O3 may elicit beneficial defense reactions in plants below their relative phytotoxic limits (75, 76), and the physiological effects are tightly coupled. Thus, it may be useful to consider the total activated oxygen flux as the sum of ozone and peroxides.

We estimate that the daytime molar peroxide flux at this southeast US site (2.4 nmol OOH·m−2·s−1) is 15% of the ozone flux on average (71). The OM of peroxides was measured using [O3] as 50 ppbv and Vh,02 = 0.8 cm·s−1 (77). However, in more pristine areas like the Amazon rainforest, the concentration of total peroxides is higher and ozone is lower than the southeastern United States. The estimated daytime peroxide flux of 5.7 nmol·m−2·s−1 in Amazonia (using [H2O2] and [ROOH] observed by Lelieveld et al. (4) and Vh,H2O2 = 2.3 cm·s−1) can be split into the contribution of observed daytime O3 fluxes during the dry season (78). These results suggest that, across varying degrees of urban influence, peroxides can comprise a significant fraction of the daytime oxidant flux in forest ecosystems.

Exogenous H2O2 (millimolar equivalent) has been shown to inhibit stomatal opening by signaling the production of NO in guard cells (79, 80). Furthermore, as peroxides are potent oxidizers and generators of reactive oxygen species (ROS) like the hydroxyl radical (OH) and the superoxide (O2−) in water, their large deposition flux motivates inquiries into their fates at the surface of plants. Deposited peroxides likely participate in aqueous or heterogeneous chemistry, e.g., the oxidation of deposited SO2 to sulfuric acid (81) or the production of ROS like OH, HO2, and O2− through direct photochemistry and catalytic reaction with trace metals (Fe) (82). During the SOAS campaign, we measured plumes of SO2 in the range of 1–9 ppb (SI Appendix, Fig. S21). The peroxide-mediated production of strong acids and oxidants may play a role in plant damage during these episodes. Additionally, this chemistry may contribute to the aforementioned inferred emission flux of formic acid through heterogeneous oxidation of OVOCs. Whether the observed deposition flux of peroxides can impact atmospheric chemistry by heterogeneous pathways or affect plant function under typical atmospheric conditions remains to be further explored.

The mean daytime dry deposition flux from HNO3, HCN, and speciated organic nitrates was measured to be 27 ± 15 μgN·m−2·h−1. To estimate the ecosystem input of nitrogen, the dry flux of N-containing species that were not measured in this work (NH3, NH4+, etc.) were simulated with GEOS-Chem. The wet-deposited N data for 2013 (measured as NH4+ and NO3−) was provided by the National Atmospheric Deposition Program at the AL03 site corresponding to CTR (nadp.sws.uiuc.edu). We estimate that dry deposition supplied ~52% of the total wet and dry input of nitrogen to the canopy by mass. This result is consistent with average estimates (~46%) at 10 North American forests (83).

HNO3 is commonly assumed to dominate the oxidized nitrogen dry deposition flux; however, our data indicate that organic nitrates (RONO2) measured by our instrument can constitute a sizable portion (15%) of the dry flux of oxidized nitrogen observed at this site (Fig. 8, Right), with nitric acid making up the balance. This is a lower limit of the organic nitrogen input into the canopy. Zhang et al. (49) assumed a deposition velocity of isoprene nitrates to be ~5.5 cm·s−1 in the daytime (greater than a factor of 3 larger than our observations) and arrived at a similar conclusion. If we include an estimation for acyloxy nitrates from data obtained in Northern California (84), the resulting organic nitrogen contribution would be a bit higher (~25%). HCN was, at most, 3% of the N flux observed in this work. However, the main source of HCN is biomass burning and, thus, its flux will vary significantly depending on location and time of year. The minor impact from NO2 deposition (21) was neglected. The data demonstrate that organic nitrogen is an important fraction of the oxidized nitrogen dry deposition flux, similar to wet deposition (85). The organic nitrogen may be taken up by foliage (86, 87) and incorporated into the synthesis of leaf nutrients. As the effects of organic nitrogen on carbon sequestration are inadequately represented in most ecological models, these data motivate further inquiry into the ecosystem ramifications of OVOC deposition.

Materials and Methods

Site and Sampling. This work was performed as part of the SOAS Campaign (soas2013.rutgers.edu) at the CTR Southeastern Aerosol Research and Characterization Study (SEARCH) site operated by the Electric Power Research Institute (Brent, AL, Lat 32.90289 Lon −87.24968) for the US Environmental Protection Agency. The metal walk-up sampling tower was ~20 m in height (measurement height = ~22 m) and the mean canopy height was ~10 m. The CIMS was situated on the topmost platform, facing north toward the forest. The temperatures inside the instrument enclosure and CIMS flow region were controlled, and the instrument was shielded from the elements by an insulated enclosure. An ultrasonic anemometer extending off the north side of the tower separates the airflow from the instrument and measured the 3D wind components and the converted virtual temperature. A weather station comprised of several sensors monitored the relative humidity (RH, percent), air temperature (T, degrees Centigrade), barometric pressure (P, millibars), solar radiation (watts per square meter), wind speed (meters per second), and wind direction (°-360°).

CIMS Mixing Ratios. The ionization mechanism and details about calibration for each species are presented in SI Appendix. Briefly, compounds were measured through a high-flow fluoropolymer-coated glass inlet (~40 cm long, 3.1 cm ID, ~2,000 L·min−1). The analytical method, using reagent ion CF3O−, is specific toward the detection of acids, hydroperoxides, multifunctional nitrates, and other multifunctional species. The ionization mechanism is fluoride transfer for acidic analytes and cluster formation for other analytes. The absolute sensitivity and water vapor dependence of the ionization were calibrated for each species in the laboratory by quantitative techniques (spectroscopy, gravimetry, liquid chromatography, etc.), when calibration standards are available commercially or can be synthesized and purified (11 out of 16 compounds). The sensitivities were calculated by ab initio methods otherwise. High-frequency CIMS measurement of water vapor was calibrated by 1-Hz weather station measurements of RH, T, and P after accounting for the temperature dependence of the ionization efficiencies were performed every 2 h with commercially available standards, and in-field zero backgrounds were measured every half hour, under dry and ambient RH conditions.
Flux Calculations. Data processing protocols, flux quality assessment and criteria, standard corrections, and CIMS-specific corrections are documented in detail in SI Appendix. We limit the fluxes reported here to a few ideas days from the campaign period, characterized by the following conditions: The turbulence (measured by the momentum flux) is well established, the data reproducibility is high, the spectral analysis shows expected behavior for sampling in the inertial subrange, and the energy balance closure at the surface (Fig. 2) is adequately met. Acceptable data generally correspond to measurement periods for which winds were northerly, with exclusively forest fetch of the measurement period and the vertical wind were measured at 8 h or faster. A time response correction was applied for HNO$_3$ (factor of 1.62) as described in SI Appendix, section 5, due the measured flux dampening caused by the time delay of this compound from interactions with instrument surfaces. Measurement uncertainties (±1σ) are largest for species with lower mean concentrations during the daytime, such as MTNP, which has a dominant nighttime source. A short period of atypical nighttime correction was removed from the data for Julian day 165 due to the flux criteria discussed in SI Appendix, section 4. EC fluxes of other species commonly detected by CIMS (notably, SO$_2$, glycolaldehyde, and methyl hydroperoxide) were not calculated in this work because mixing ratios at this site were either too rapidly changing (SO$_2$ plumes) or contain interfering species when measured with the time-of-flight mass detector.

ACKNOWLEDGMENTS. We thank the organizers and committee members of the SOAS campaign: A. G. Carlson, A. H. Goldstein, J. L. Jimenez, R. W. Pinder, J. de Gouw, B. J. Turpin, and A. B. Guenther. We acknowledge C. J. Groff at Purdue University for his assistance with leaf area index measurements and tree surveys. We thank D. J. Jacob and the Atmospheric Chemistry Modeling Group at Harvard University for making GEOS-Chem available for this work. Meteorological data used in the GEOS-Chem simulations were provided by the Global Modeling and Assimilation Office at NASA Goddard Space Flight Center. We acknowledge funding from the National Science Foundation (NSF) under Grant AGS-1240604 and NSF Postdoctoral Research Fellowship program Award AGS-1331360. Financial and logistical support for SOAS was provided by the NSF, the Earth Observing Laboratory at the National Center for Atmospheric Research (operated by NSF), the person at Atmospheric Research and Analysis, and the Electric Power Research Institute.

References