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In-canopy gas-phase chemistry during CABINEX 2009: sensitivity of a 1-D canopy model to vertical mixing and isoprene chemistry

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Abstract. Vegetation emits large quantities of biogenic volatile organic compounds (BVOC). At remote sites, these compounds are the dominant precursors to ozone and secondary organic aerosol (SOA) production, yet current field studies show that atmospheric models have difficulty in capturing the observed HO_x cycle and concentrations of BVOC oxidation products. In this manuscript, we simulate BVOC chemistry within a forest canopy using a one-dimensional canopy-chemistry model (Canopy Atmospheric CHemistry Emission model; CACHE) for a mixed deciduous forest in northern Michigan during the CABINEX 2009 campaign. We find that the base-case model, using fully-parameterized mixing and the simplified biogenic chemistry of the Regional Atmospheric Chemistry Model (RACM), underestimates daytime in-canopy vertical mixing by 50-70% and by an order of magnitude at night, leading to discrepancies in the diurnal evolution of HO_x, BVOC, and BVOC oxidation

products. Implementing observed micrometeorological data from above and within the canopy substantially improves the diurnal cycle of modeled BVOC, particularly at the end of the day, and also improves the observation-model agreement for some BVOC oxidation products and OH reactivity. We compare the RACM mechanism to a version that includes the Mainz isoprene mechanism (RACM-MIM) to test the model sensitivity to enhanced isoprene degradation. RACM-MIM simulates higher concentrations of both primary BVOC (isoprene and monoterpenes) and oxidation products (HCHO, MACR+MVK) compared with RACM simulations. Additionally, the revised mechanism alters the OH concentrations and increases HO₂. These changes generally improve agreement with HO_x observations yet overestimate BVOC oxidation products, indicating that this isoprene mechanism does not improve the representation of local chemistry at the site. Overall, the revised mechanism yields smaller changes in BVOC and BVOC oxidation product concentrations and gradients than improving the parameterization of vertical mixing with observations, suggesting that uncertainties in vertical mixing parameterizations are an important component in understanding observed BVOC chemistry.

1 Introduction

There is increasing evidence of the important role of forest canopies and biogenic volatile organic compound (VOC) emissions on tropospheric composition and atmospheric chemistry (Goldstein and Galbally, 2007; Lelieveld et al., 2008). VOC oxidation, in the presence of reactive nitrogen oxides $(NO_x = NO + NO_2)$ and sunlight, is critical for ozone formation (Logan, 1985) and condensation of their oxidation products can yield secondary organic aerosols (SOA) (Claeys et al., 2004; Carlton et al., 2009; Hallquist et al., 2009). Additionally, VOC can control the oxidation capacity of the troposphere through the regulation of hydrogen radicals $(HO_x = OH + HO_2)$ (Poisson et al., 2000; Tan et al., 2001). Forest canopies are an important VOC source both globally and regionally, contributing to nearly half the global VOC budget (Guenther et al., 1995). To affect the troposphere, biogenic VOC (BVOC) emissions and their oxidation products must be mixed effectively out of the forest canopy. This forest-atmosphere exchange is highly sensitive to turbulent mixing and chemistry because BVOC oxidation and transport occur on similar timescales (Molemaker and Vilà-Guerau de Arellano, 1998; Krol et al., 2000; Pugh et al., 2010).

To investigate the role of BVOC on tropospheric chemistry, several recent field campaigns have involved chemical measurements at multiple heights throughout the forest canopy (e.g., Carroll et al., 2001; Hewitt et al., 2010; Martin et al., 2010). Results from these field campaigns highlight gaps in our understanding of BVOC oxidation. For example, the hydroxyl radical (OH) is underestimated in most forest ecosystem types (Carslaw and Carslaw, 2001; Tan et al., 2001; Butler et al., 2008; Karl et al., 2009). Modeling studies of remote forest sites that add a HO_x recycling mechanism through BVOC oxidation reactions (Lelieveld et al., 2008; Hofzumahaus et al., 2009) or OH regeneration from epoxides (Paulot et al., 2009) display some improvement in measuredmodeled agreement in some locations, yet these do not show consistent improvement in all studies (Karl et al., 2009; Barkley et al., 2011). Additionally, BVOC oxidation products are poorly simulated in a number of forest regimes. Pugh et al. (2010) show that first-generation oxidation products of isoprene (C₅H₈, 2-methyl-1,3-butadiene)—the dominant BVOC emission in many broadleaf ecosystems-are overestimated by box-model simulations of a Malaysian tropical rainforest. Karl et al. (2009) compare several mechanisms with enhanced isoprene oxidation and find that some isoprene oxidation products such as hydroxyacetone are underpredicted compared to observations. While revised chemical mechanisms can explicitly account for more detailed isoprene chemistry (Pöschl et al., 2000; Paulot et al., 2009; Peeters et al., 2009; Stavrakou et al., 2010), difficulties remain in simulating isoprene degradation and oxidation products under low-NO_x conditions (Karl et al., 2009).

In addition to uncertainties in the pathways of BVOC oxidation, vertical transport within and above the canopy sub-layer is an additional source of uncertainty in forestatmosphere exchange (Finnigan, 2000; Hurst et al., 2001). Here, we define the canopy sub-layer as the thin atmospheric layer nearest the surface containing forest roughness elements. Turbulence occurs over a range of scales in the planetary boundary layer (PBL) from the mesoscale to the sub-grid scale, and thus, must be either represented with higher-order turbulence schemes or parameterizations. Among the most common parameterizations is the first-order flux-gradient relationship, known as K-theory, in which turbulent exchange is a function of the eddy diffusivity parameter, K (Blackadar, 1979). Because mixing strength in the mid-PBL peaks at two orders of magnitude higher than in the canopy roughness layer (Gao et al., 1993), turbulent transport tends to be much smaller within plant canopies than above the roughness elements. In fact, K-theory has been found to break down completely within forest canopies due to the existence of intermittent coherent structures that encompass the entire depth of the canopy (Raupach et al., 1996), yet the parameterization continues to be used for its computational efficiency in many models (Forkel et al., 2006). Though high-resolution canopy models may have the vertical resolution to capture fine-scale turbulence within the canopy, many models do not have the detailed description of higher-order turbulence to simulate the effects of coherent structures and other canopy-scale turbulence. Therefore, most models have large uncertainties in the role of vertical mixing on BVOC gradients and forestatmosphere exchange (e.g., Ganzeveld et al., 2006). Nearfield effects, often represented by a scaling factor (Makar et al., 1999; Stroud et al., 2005; Wolfe and Thornton, 2011), have been shown to improve modeled in-canopy and abovecanopy turbulence (Raupach, 1989). Large-eddy simulation models (e.g., Heus et al., 2010; Patton et al., 2001) can capture these dynamical changes, though development with sufficient detail in chemical mechanisms is still underway (e.g., Kim et al., 2012). The use of a 1-D model in this study represents a computationally efficient preference towards a detailed chemical mechanism over that of a detailed turbulence parameterization.

Despite the uncertainties in vertical mixing, onedimensional (1-D) models are still useful tools for studying the vertical transport in the context of atmospheric composition because they focus on the implications of in-canopy chemistry on vertical concentrations and gradients (e.g., Wolfe et al., 2011; Boy et al., 2011). Wolfe et al. (2011) examine daytime biogenic chemistry at a ponderosa pine plantation, concluding that chemistry may outweigh the effect of turbulent mixing on forest-atmosphere exchange, even for compounds with long chemical lifetimes relative to their transport timescales. Boy et al. (2011) applies a more detailed turbulence scheme to understand BVOC oxidation and tracer transport and their effect on particle formation in a Scots Pine forest in Finland and find that this improved mixing parameterization can reproduce observed vertical profiles of BVOC.

Here, we focus on chemistry and turbulence within and above a deciduous hardwood forest in Northern Michigan. A number of field campaigns as part of the Program for Research on Oxidants: PHotochemistry, Emissions, and Transport (PROPHET, Carroll et al., 2001) have been conducted since 1997 at the University of Michigan Biological Station (UMBS). Results from the summer 1998 PROPHET intensive show that models underestimate OH (Faloona et al., 2001; Sillman et al., 2002), while measured HO₂ concentrations compare well with model results (Tan et al., 2001). Measurements of OH reactivity suggest an unknown BVOC source, which may contribute to these OH discrepancies (Di Carlo et al., 2004). In 2009, the Community Atmosphere-Biosphere INteractions Experiment (CABINEX) was conducted to provide new insights into the role of BVOC chemistry and its relationship to HO_x chemistry observed at the PROPHET site. Branch enclosure measurements from CAB-INEX 2009 show that identified primary emissions reasonably reflect current emission estimates (Ortega et al., 2007), suggesting that the missing ambient OH reactivity could be explained by secondary BVOC oxidation products (Kim et al., 2011).

In this manuscript, we compare measurements from the CABINEX 2009 campaign with a 1-D Eulerian Canopy Atmospheric CHemistry Emission model (CACHE, Forkel et al., 2006) to investigate the role of in-canopy chemistry and turbulence on HO_x and BVOC concentrations and vertical gradients. To examine the relative sensitivities of the model to mixing and chemistry, we compare a base-case model scenario with the original turbulence and chemistry description within CACHE against a revised mixing scheme and isoprene-focused chemical mechanism. We explore the relative impacts of HO_x pathways in an atmospheric chemical mechanism versus the effects of vertical mixing on ambient concentrations to isolate and highlight the key processes of biosphere-atmosphere interactions at this deciduous forest ecosystem.

2 Methods

2.1 CABINEX 2009 campaign

CABINEX 2009 was conducted at the PROPHET site at UMBS near Pellston, MI $(45^{\circ} 33' 31.66'' \text{ N}, 84^{\circ} 43' 52.40'' \text{ W})$ at the transition between mixed hardwood

to boreal forest (Schmid et al., 2003). Depending on the wind direction, the site can be controlled by local emissions and chemistry or regional transport from urban areas (Milwaukee, WI (\sim 378 km SW); Detroit, MI (\sim 385 km SSE); and Chicago, IL ($\sim 475 \text{ km SW}$), as shown in Fig. S2). The local vegetation is diverse (Pressley et al., 2005), containing varieties of aspen, oak, beech, birch, maple, and pine with an average canopy height (h) of approximately 22.5 m. Climate conditions at the PROPHET site are generally cold but with warm summers (FLUXNET database, Baldocchi et al., 2001). The average daily maximum temperatures for Pellston, MI, in July and August are 26 °C and 25 °C with average precipitation of 63.2 mm and 81.8 mm, respectively. The summer of 2009 was unseasonably cool and cloudy with an average high temperature of 22 °C and with rain or fog occurring on 62% of the days within the 1 July-8 August 2009 observational period, which may reduce BVOC emissions and photochemical activity.

O₃, NO₂, NO, isoprene, monoterpenes, formaldehyde (HCHO), methacrolein and methyl vinyl ketone (MACR+MVK), OH, HO₂, and glyoxal (GLY) were measured at multiple heights within and above the canopy. Due to limited instrumentation during the campaign, simultaneous measurements at multiple heights could not be obtained for all compounds. Therefore, data availability at a given height widely varies over the campaign. Primary BVOC species and BVOC oxidation products were measured, alternating at 10-min intervals between 6 m, 20.4 m, and 34 m, using a proton transfer reaction mass spectrometer (PTR-MS) with dehumidified sampling to allow more sensitive measurement of HCHO (de Gouw and Warneke, 2007; Jobson and McCoskey, 2010). NO and NO₂ were measured, alternating between the same three heights, using a 2-channel chemiluminescence instrument with a blue light photolytic converter for NO₂ (Air Quality Design). HO_x species were measured only at 32 m during the two-day simulation presented here, with OH measured using laser-induced fluorescence with the Fluorescence Assay by Gas Expansion (FAGE) technique and HO₂ measured by chemical conversion to OH with added NO (Stevens et al., 1994; Dusanter et al., 2009). Total OH reactivity was measured at 30.9 m using a turbulent flow tube technique similar to that described in Kovacs and Brune (2001). Glyoxal (GLY) was measured at 35.4 m using laser-induced phosphorescence (Huisman et al., 2011). Wind speed and direction were measured via propeller anemometer at 36.4 m. Additionally, two sonic anemometers measuring the three-component wind field (u, u)v, and w) and temperature were mounted at 20.6 m and 34 m (Steiner et al., 2011). Temperature was also measured at 6 m, 20.4 m, and 31.2 m using R. M. Young relative humidity and temperature probes. Photosynthetically active radiation (PAR) was measured at 32.6 m using a BF-3 Sunshine Sensor.

2.2 Model description and simulations

CACHE is a 1-D multilayer model that simulates the vertical distribution of concentrations and vertical fluxes of heat, moisture, and gas-phase chemical species over time using the prognostic equations for potential temperature (θ) and volume mixing ratio for compound *i* (c_i):

$$\frac{\partial \theta}{\partial t} = \frac{\partial}{\partial z} \left(K_{\rm H} \frac{\partial \theta}{\partial z} \right) + S_{\rm H} \tag{1}$$

$$\frac{\partial c_i}{\partial t} = \frac{\partial}{\partial z} \left(K_{\rm H} \frac{\partial c_i}{\partial z} \right) + S_{\rm c} + C \tag{2}$$

Equations (1) and (2) are solved for each model layer, where $K_{\rm H}$ denotes the turbulent exchange coefficient for heat, $S_{\rm H}$ and S_c denote sources and sinks for heat and mass, respectively, and *C* denotes chemical transformation. Turbulent exchange, chemistry, emissions, deposition, and advection are described in subsequent sections.

For our simulations of the PROPHET site, we use a model domain consisting of 40 vertical layers spanning 4.4 km in the vertical (z) direction with eight layers in the 6-m trunk space and ten layers in the 16.5-m crown space. The grid resolution decreases exponentially with height with a spacing of 0.81 m at ground level and 1061.1 m at the top. Simulations are run for 48 h at a time step of 60 s. Initial conditions are provided to the model for vertical profiles of chemical concentrations based on observed near-canopy concentrations at the start of the simulation; the initial vertical temperature profile is interpolated using radiosonde data from Gaylord, MI (~59.5 km S). Model input includes (Fig. S1 in the Supplement): (1) observed PAR to drive the prognostic temperature profile and photochemistry and account for cloud cover, and (2) observed wind speed and direction at 36.4 m to drive the vertical wind profile and NOx and VOC advection. Additional input required for the revised turbulence scheme (see Sect. 2.2.1) include u_* and σ_w at 20.6 and 34 m. In addition, we prescribe a total leaf area index (LAI) of $3.8 \text{ m}^2 \text{ m}^{-2}$ (as observed at PROPHET, Ortega et al., 2007) and typical values for leaf reflectance and transmittance observed in deciduous broadleaf ecosystems (see Table 1, Asner, 1998). Model output is interpolated at the instrument heights and data collection times for precise comparison with measurements.

2.2.1 Turbulent exchange

Vertical transport is parameterized in CACHE using a firstorder flux-gradient relationship, or K-theory, in which heat and mass are transported by eddy diffusion at a rate proportional to the turbulent exchange coefficient for heat, $K_{\rm H}$. Vertical fluxes of heat and mass are computed at each model time step as follows:

$$\overline{w'\theta'} = -K_{\rm H} \frac{\partial\bar{\theta}}{\partial z} \tag{3}$$

Table 1. Leaf and soil reflectance and transmittance by waveband (visible/near infrared/thermal) used in CACHE. Absorptivities are 1 - (reflectance + transmittance). Values are derived from Asner (1998).

	Soil	Leaf
Reflectance	0.15/0.20/0.10	0.20/0.45/0.10
Transmittance	0.00/0.00/0.00	0.10/0.30/0.10

$$\overline{w'c'_i} = -K_{\rm H} \frac{\partial \bar{c_i}}{\partial z}.$$
(4)

Modeled $K_{\rm H}$ (hereinafter denoted $K_{\rm H,mod}$) is derived empirically according to Forkel et al. (1990) given a length scale l, the vertical wind shear $|\partial v/\partial z|$, and a stability parameter f:

$$K_{\rm H,mod} = l^2 \left| \frac{\partial v}{\partial z} \right| \cdot f.$$
 (5)

l varies within and above the canopy according to the following parameterization:

$$l = \frac{\kappa(z-d)}{1 + (\kappa(z-d)/\lambda)} \tag{6}$$

where κ is the von Kármán constant (0.4), d is the zero-plane displacement height (0.85 h for $z \ge h$ and zero for z < h), and λ is the maximum mixing length (given by Forkel et al. (1990, Eq. (10)) for $z \ge h$ and set constant at 2 m for z < h). The vertical wind profile for computing $|\partial v/\partial z|$ derives from the common logarithmic expression for the PBL (Stull, 1988); within the canopy, winds dissipate as a function of u_* and canopy structure according to a modified logarithmicwind equation following Baldocchi (1988, Eq. 6). f is a function of the Richardson number Ri:

$$f(\text{Ri}) = \begin{cases} 1.35\sqrt{1 - 11\text{Ri}}\frac{1 - 5.5\text{Ri}}{1 - 3\text{Ri}} & : \text{Ri} < 0\\ 1.35[(1 + 6\text{Ri})\sqrt{1 + 6\text{Ri}}]^{-1} & : \text{Ri} \ge 0 \end{cases}.$$
 (7)

In the original model configuration—hereafter referred to as the "BASE" model scenario – the vertical turbulence profile is driven solely by Eqs. (5–7).

To evaluate the sensitivity of BVOC gradients to in-canopy vertical mixing, we apply a modified K-theory parameterization in a modeling scheme hereafter referred to as the "MIX" simulation. We first define "observed" $K_{\rm H}$ (hereafter denoted by $K_{\rm H,obs}$) following Makar et al. (1999):

$$K_{\rm H,obs} = \sigma_w^2 T_{\rm L} \tag{8}$$

where $T_{\rm L}$ is the Lagrangian timescale ($T_{\rm L} = 0.3h/u_*$), u_* is the friction velocity, and σ_w is the vertical velocity standard deviation. u_* and σ_w are computed using half-hour Reynolds

averages of raw (10 Hz) sonic anemometer measurements of u, v, and w above the canopy (34 m; 1.5 h) and in the top of the canopy (20.6 m; 0.9 h):

$$u_* = \left(\overline{u'w'}^2 + \overline{v'w'}^2\right)^{1/4}$$
(9)

$$\sigma_w = \sqrt{\frac{1}{n-1} \sum_{i=1}^n (w_i - \bar{w})^2}$$
(10)

In the MIX simulations, we adjust the modeled turbulence profile (i.e. the BASE case) to match the near-surface observations estimated in Eq. (8) as follows. At each time step, we first compute a full vertical profile of $K_{\rm H,mod}$ using Eq. (5). Then, we linearly interpolate between $K_{\rm H,mod}$ at the canopy base (6 m) and $K_{\rm H,obs}$ at the first measurement level (20.6 m). At the model grid points between the two measurement heights (20.6–34 m), we linearly interpolate u_* and σ_w and compute $K_{\rm H,obs}$ using Eq. (8). Above 34 m, modeled values are adjusted to remove any discontinuity between the measured and modeled values. Near-canopy $K_{\rm H}$ ($z \le 1.64 h$) is scaled by an *R* factor to account for nearfield effects of the canopy, where *R* is:

$$R = \frac{\left[1 - \exp(-\tau/T_{\rm L})\right](\tau - T_{\rm L})^{3/2}}{\left[\tau - T_{\rm L} + T_{\rm L}\exp(-\tau/T_{\rm L})\right]^{3/2}}$$
(11)

and the transport timescale τ is derived using a $\tau/T_{\rm L}$ ratio of 4 (Stroud et al., 2005; Wolfe and Thornton, 2011). We evaluate the revised mixing scheme in Sect. 3.1.

2.2.2 Chemistry

Gas-phase chemical transformation in the original CACHE model implements the Regional Atmospheric Chemistry Mechanism (RACM, Stockwell et al., 1997), which includes a suite of 77 chemical species and 237 reactions. The mechanism includes explicit treatment of three BVOC categories: isoprene (denoted by the RACM surrogate, ISO), monoterpenes with one double bond (i.e. α -pinene; denoted API), and monoterpenes with two double bonds (i.e. *d*-limonene; denoted LIM). The remaining VOC are lumped into four alkane categories, four alkene categories, and three aromatic categories based on reactivity with OH. In the RACM mechanism, MACR+MVK are contained in the surrogate for all unsaturated C4 carbonyls (denoted MACR in Stockwell et al. (1997)), which derive from both anthropogenic and biogenic diene oxidation; however, we note that measurements from the campaign only account for MACR+MVK alone.

In this study, we perform a sensitivity test (hereafter referred to as the "MIM" simulation) comparing RACM against the Mainz isoprene mechanism (MIM) adaptation of RACM (RACM-MIM, Geiger et al., 2003). RACM-MIM treats an additional seven species explicitly that are split from surrogate species in RACM. By using this greater speciation, RACM-MIM includes an additional twelve reactions. Ultimately, this mechanism provides more specific pathways of isoprene oxidation, the production of second-generation isoprene oxidation products, and further detail on the MACR chemistry under low-NO_x conditions.

2.2.3 Emissions

BVOC emissions are controlled by site-specific emission rates that depend on ecosystem-specific emission factors, temperature, and PAR (Steinbrecher et al., 1999). Isoprene emission fluxes were not routinely measured during the campaign; therefore, we use the mean isoprene surface emission flux observed at PROPHET during 2003 and 2005 (Ortega et al., 2007). Bigtooth aspen (Populus grandidentata) and northern red oak (Quercus rubra) account for 99% of the isoprene budget, emitting at mean basal emission rates of 46.3 and $53.5 \,\mu g C g^{-1} h^{-1}$ at standard conditions (PAR = 1000 μ mol m⁻² s⁻¹; temperature = 30 °C). Since 2009 was colder than average summers, we subtract one standard deviation from the mean isoprene emission fluxes, corresponding to $24.8 \,\mu g C g^{-1} h^{-1}$ for aspen and 23.8 μ gCg⁻¹h⁻¹ for oak. The net emission of isoprene per model level is determined by scaling the given emission factors by PAR and temperature, according to the parameterization described in Forkel et al. (2006) following Guenther et al. (1995), and a prescribed vertical LAI distribution. Monoterpene emission rates derive from tree branch enclosure measurements in 2003, 2005 and 2009 (Ortega et al., 2007; Ortega and Helmig, 2008). Normalized (20 °C) foliage emission rates for the sum of monoterpenes for northern red oak (Quercus rubra), eastern white pine (Pinus strobus), red pine (Pinus resinosa), and American beech (Fagus grandifolia) measured during CABINEX 2009 are 0.16, 0.38, 0.56, and 7.46 μ g C g⁻¹ h⁻¹, respectively. For beech, the predominant emission is *d*-limonene (27%), followed by sabinene (17%), α -pinene (12%), and cymene (12%). Ortega et al. (2007) measure a mean basal emission rate for paper birch (*Betula papyrifera*) of $0.5 \,\mu g C g^{-1} h^{-1}$. For white and red pine, we scale the emission factors by 2.56 (Perterer and Körner, 1990) to account for the conversion from projected to total leaf area. We sum these species contributions and split this total monoterpene emission factor into the RACM species API (56.5%, or $0.086 \,\mathrm{nmol}\,\mathrm{m}^{-2}\,\mathrm{leaf}\,\mathrm{area}\,\mathrm{s}^{-1}$) and LIM (43.5%, or $0.066 \text{ nmol m}^{-2}$ leaf area s⁻¹) according to the measured fractional contribution of similar species.

We note that these emission estimates are based on available data from the site collected by several investigators over several seasons. Due to the high variability seen in these data, it is difficult to define representative values particularly given the cool conditions during the summer of 2009. Previous studies have noted that emission factors can vary based on prior temperatures on the span of weeks (Pétron et al., 2001) and can vary based on the plants acclimation, particularly for isoprene (Hanson and Sharkey, 2001). Consequently, these estimates may have uncertainties of a minimum of a factor of two. Measurements indicate a dependence on temperature for monoterpene emissions; therefore, we presume monoterpene emissions to be from pools within the foliage, and are, thus, scaled according to the temperaturedependent parameterization described in Forkel et al. (2006). Soil NO emissions are parameterized according to Forkel et al. (2006), following Simpson et al. (1995), based on an emission rate of 180 nmolm⁻² h⁻¹ observed previously at PROPHET (Alaghmand et al., 2011). Foliage NO₂ emission from nitrate photolysis at the leaf surface (Hari et al., 2003) is not considered in this study due to uncertainties in emission factors.

2.2.4 Deposition

Dry deposition of chemical species to canopy foliage $(S_{c,dep,i}; nmol m^{-2} s^{-1})$ is parameterized following Meyers and Baldocchi (1988):

$$S_{c,dep,i}(z) = -LAI(z)c_i(z)v_{d,i}(z)$$
(12)

where LAI and c_i are the leaf area index per unit height and concentration for gas *i*, respectively. Deposition velocity ($v_{d,i}$; m s⁻¹) is represented by four resistances: the quasi-laminar boundary layer ($R_{b,i}$), stomatal (R_s), mesophyll ($R_{m,i}$), and cuticular ($R_{c,i}$):

$$v_{d,i}(z) = \frac{1}{R_{b,i}(z) + R_s(z)D_{H_2O}/D_i + R_{m,i}} + \frac{2}{R_{b,i}(z) + R_{c,i}}.$$
 (13)

where $D_{\rm H_2O}/D_i$ is the ratio of the molecular diffusivities of water to gas *i* (Gao et al., 1993). $R_{\rm b,i}$ is a function of $D_{\rm H_2O}/D_i$ and the layer-mean horizontal wind \bar{u} (Meyers, 1987):

$$R_{\rm b,i}(z) = \frac{r_{\rm b}(z)}{D_{\rm H_2O}/D_i}.$$
(14)

where

$$r_{\rm b}(z) = \begin{cases} 500 \\ 180\sqrt{l/\bar{u}(z)} : \bar{u}(z) \le 0.01 \text{ m s}^{-1} \\ : \bar{u}(z) > 0.01 \text{ m s}^{-1} \end{cases}$$
(15)

given an estimated typical leaf length l of 8 cm. R_s is dependent on light, leaf temperature, and water potentials during the day, following Jarvis (1976), and is equivalent to the cuticular resistance for O₃ ($R_{c,O_3} \approx 3000 \text{ m s}^{-1}$, Wesely, 1989) at night. $R_{m,i}$ and $R_{c,i}$ are parameterized according to Wesely (1989) as a function of the Henry's law constant H_i^* and a reactivity factor $f_{0,i}$ describing the extent that gas *i* decomposes in the plant mesophyll relative to O₃, ranging from non-reactive ($f_{0,i} = 0$) to full decomposition ($f_{0,i} = 1$, i.e. O₃):

$$R_{\mathrm{m},i} = (H_i^*/3000 + 100 f_{0,i})^{-1}$$
(16)



Fig. 1. Simulated canopy-integrated deposition velocity (v_d) for methacrolein (C₄H₆O), formaldehyde (CH₂O), ozone (O₃), and hydrogen peroxide (H₂O₂).

$$R_{c,i} = R_{c,O_3} (10^{-5} H_i^* + f_{0,i})^{-1}.$$
(17)

 $f_{0,i}$ follows Wesely (1989) with the exception of MACR and HCHO, which are set to that of O₃ ($f_{0,i} = 1$) in response to recent suggestions that oxidized VOC can decompose in the leaf more rapidly than previously believed (Karl et al., 2004, 2010). Following this change, CACHE simulates canopy-integrated deposition velocities for MACR within the expectations of Karl et al. (2010, $v_{d,MACR} < 2.4 \text{ cm s}^{-1}$) and comparable to that of O₃ (Fig. 1). H₂O₂ deposition rates are consistent with previous modeling studies, yet may be underestimated according to recent observational studies (Ganzeveld et al., 2006). Deposition to the ground follows Gao et al. (1993).

2.2.5 Advection

Horizontal advection of anthropogenic NO_x and long-lived VOC is represented in CACHE as a function of wind direction. Cooper et al. (2001) attribute elevated concentrations of O_3 , CO, NO_x, and other oxidized nitrogen species (NO_z) observed at PROPHET to southerly flow from Chicago or Detroit and lower mixing ratios to flow from clean Canadian air masses over the Great Lakes. We estimate the directiondependent advection rate of eight RACM species (NO2, HCHO, MACR, KET, HC3, HC5, OLT, and OLI; see Stockwell et al. (1997) for full definitions) according to the geographical location of PROPHET relative to nearby major urban centers. Chicago (pop. ~ 2.9 million) and Detroit (pop. \sim 950000) are the major contributors of anthropogenic emissions affecting northern Michigan, with emission inventories for NO_x and VOC totaling over $20\,000\,\text{kg}\,\text{day}^{-1}$, whereas peak NO_x and VOC emissions from Milwaukee (pop. ~ 950000) total in the range of 12500–15000 kg day⁻¹

(LADCO, 2010). Therefore, we presume signatures of anthropogenic advection observed at PROPHET to be more pronounced with air originating in Chicago or Detroit over Milwaukee by assuming the strongest advection rates when winds are directly from the south. Westerly advection from Lake Michigan has been associated with lower isoprene concentrations than advection from forests to the south (Sillman et al., 2002); therefore, we add advection of biogenic HCHO and MACR from isoprene oxidation under southerly winds. We incorporate advection of the above species between the heights 45-106 m (22-26 model levels) at the hourly, winddirection-dependent rates shown in Table 2. Measured NO₂, HCHO, and MACR concentrations are used to tune advection rates for the model scenario that incorporates both the revised turbulence scheme and RACM-MIM chemistry (hereafter, the MIX+MIM case). Due to the lack of ambient observations, anthropogenic hydrocarbon RACM categories, OLI, OLT, KET, HC3, and HC5, are added to reflect average concentrations of regional pollution events.

2.2.6 Case study: 4–5 August

The case study simulation period includes 48 h starting at 00:00 Eastern Standard Time (EST) on 4 August 2009 and ending at 23:59 EST on 5 August 2009. This two-day period offers the clearest daytime skies within the period when the most chemistry observations are available. We select these clear-sky days because they represent the time period most conducive to BVOC emissions and photochemistry, as well as providing the best measured-modeled agreement for the turbulence parameterization. NCEP surface reanalysis indicates a weak frontal passage occurring at $\sim 07:00$ EST on 4 August (Fig. S2 in the Supplement), visible in the observed wind direction (Fig. S1 in the Supplement) through a shift from southerly to northwesterly winds. Back-trajectory data (Fig. S2 in the Supplement) show that air originated in northern Illinois (southwest of the PROPHET site) prior to the frontal passage. Following the frontal passage, the source region ranges from northern Minnesota and southwestern Ontario, Canada (west and northwest of the site, respectively). This meteorological scenario allows us to evaluate our advection scheme presented in Sect. 2.2.5 and the ability of the model to capture the shift from polluted- to clean-air advection, while also providing a good comparison between chemistry at PROPHET under the influence of regional transport versus predominantly local chemistry.

3 Results and discussion

We compare micrometeorological and chemistry observations from CABINEX 2009 against CACHE simulations during the 4–5 August 2009 case period using four model scenarios:

Table 2. Advection rates for NO_x , VOC, and other hydrocarbons as a function of wind direction. Rates (in $ppbvh^{-1}$) are scaled by the geostrophic wind speed.

RACM species name	90–135°, 225–270°	135–225°
NO ₂	0.05	0.25
MACR	0.00	1.00
HCHO (anthropogenic)	0.00	0.03
HCHO (biogenic)	0.00	1.00
KET	0.00	0.25
HC3	0.00	$2.50 imes 10^{-4}$
HC5	0.00	0.25
OLT	0.00	0.13
OLI	0.00	0.01

- 1. BASE: a control run in which turbulence follows $K_{\rm H,mod}$ (Eq. 5) alone and chemistry is according to RACM;
- 2. MIX: sensitivity of the model to turbulence is tested by modifying $K_{\text{H,mod}}$ with $K_{\text{H,obs}}$ (Eq. 8) and R (Eq. 11) to account for near-field effects (Sect. 2.2.1) while using BASE chemistry;
- MIM: sensitivity of the model to varying treatments of isoprene degradation is tested by replacing RACM with RACM-MIM while using BASE turbulence; and
- MIX+MIM: the combined effects of enhanced turbulence and biogenic representation are examined by applying MIX turbulence and MIM chemistry.

We first evaluate the modifications made to the turbulent exchange parameterization implemented in the MIX model scenario (Sect. 3.1), followed by an analysis of model-measurement comparisons throughout the canopy and the surface layer (to approximately 3 h) for O₃ and NO_x (Sect. 3.2), BVOC and their oxidation products (Sect. 3.3), and HO_x concentrations and OH reactivity (Sect. 3.4).

3.1 Evaluation of turbulent exchange

We estimate observed eddy diffusivity ($K_{\rm H,obs}$, Fig. 2) from sonic anemometer measurements (Eq. 8) at two heights on the PROPHET tower (0.9 h and 1.5 h) for the 4–5 August case study. At 34 m, $K_{\rm H,obs}$ has a strong diurnal cycle ranging from 3 m² s⁻¹ at night to 10 m² s⁻¹ at midday with nighttime and daytime standard deviations of around 1 and 2 m² s⁻¹, respectively. $K_{\rm H,obs}$ in the upper canopy is only 1–3 m² s⁻¹ lower than above the canopy, with a similar magnitude of standard deviations. The majority of the canopy foliage resides below the 0.9 h measurement, and the absence of lower canopy micrometeorological measurements is a limiting factor in our estimates of in-canopy mixing.

The original CACHE parameterization of $K_{H,mod}$ (Eq. 5) is below one standard deviation of $K_{H,obs}$ at both



Fig. 2. Observed ($K_{\text{H,obs}}$, given by Eq. (8, diamonds)) and modeled ($K_{\text{H,mod}}$, given by Eq. (5), solid lines) time series of the turbulent exchange coefficient in the upper-canopy (20.6 m, 0.9 *h*, red) and above the canopy (34 m, 1.5 *h*, black) for 4–5 August 2009. Standard deviations of the mean diurnal cycle of $K_{\text{H,obs}}$ for the sunny and partly sunny days (21, 29 July, 2, 4, 5, and 7 August) are shaded. $K_{\text{H,obs}}$ is calculated using sonic anemometer estimations of u_* and σ_w (see Sect. 2.2.1).

measurement heights 95 % of the simulation time (Fig. 2). At 1.5 *h*, daytime $K_{\rm H,mod}$ is underestimated by a factor of two, whereas nighttime values are two orders of magnitude below $K_{\rm H,obs}$. $K_{\rm H,obs}$ at 0.9 *h* are an order of magnitude greater than model estimates over the full simulation period. In addition, the modeled onset of the stable nocturnal boundary layer, represented by the shift from higher, buoyancy-driven mixing to lower, mechanically-driven mixing at the end of the day, occurs two hours earlier and more abruptly than observed. This suggests that the empirical piecewise stability function (Eq. 7) may inadequately capture the formation of the stable layer and/or its effect on turbulence.

We evaluate the MIX turbulence scheme by comparing the BASE and MIX model simulations against observed time series of near-canopy temperatures (Fig. 3) and midday vertical profiles of $K_{\rm H}$, temperature, primary BVOC (isoprene and monoterpenes), and BVOC oxidation products (formaldehyde, MACR+MVK, and acetaldehyde) (Fig. 4). Following the frontal passage, which CACHE cannot simulate, nearcanopy temperatures are reproduced by the model, especially in the MIX simulation (Fig. 3). This indicates that the PARdriven heating source and vertical exchange of heat at the surface are captured fairly well by the model. $K_{\rm H,obs}$, which drives in- and near-canopy turbulence for the MIX case, is twice as large as the BASE-case turbulence (i.e. $K_{H,mod}$) at 1.5 h and an order of magnitude larger at 0.9 h (Fig. 4a), indicating missing turbulence such as coherent structures (as observed during CABINEX 2009 Steiner et al., 2011), countergradient terms, or other processes that cannot be captured by



Fig. 3. Measured and modeled temperatures at 34, 20.4, and 6 m.

the original K-theory parameterization. The standard deviations of the $K_{H,obs}$ (denoted by the error bars) are based on daily averages for the sunny and partly sunny days during the period of available u_* and σ_w measurements (21, 29 July, 2, 4, 5, and 7 August). Of these "clear-sky" days, observed turbulence was stronger than average on 5 August, leading to nearly uniform temperatures with height that is well captured by the MIX model case (Fig. 4b). In the BASE scenario, $K_{\rm H,mod}$ decreases to 0.1 m² s⁻¹ at the displacement height, creating an unrealistic artificial boundary. This is an artifact of the use of two different equations to construct the in- and above-canopy wind profiles and turbulence schemes, which creates a discontinuity at the forest-atmosphere interface and prevents BVOC transport out of the canopy sub-layer in the model. Temperature (Fig. 4b) decreases with height within and above the canopy according to the observations, yet the BASE model case imposes a stabilizing inversion induced by



Fig. 4. Measured and modeled vertical profiles of (a) eddy diffusivity, (b) temperature, (c) isoprene, (d) monoterpenes, (e) formaldehyde, (f) MACR+MVK, and (g) acetaldehyde at 14 EST 5 August 2009. Whiskers denote the standard deviations for the sunny and partly sunny days (21, 29 July, 2, 4, 5, and 7 August).

heating of the upper canopy. Consequently, in-canopy mixing is weak in the BASE simulation.

Observed midday vertical gradients of BVOC and oxidation products are also compared against the BASE and MIX simulations in Fig. 4c-g. Enhanced turbulence effectively improves the agreement of modeled and measured concentrations for the more reactive species (isoprene and monoterpenes) by reducing in-canopy concentrations and weakening the vertical gradient. For the longer-lived species (formaldehyde, MACR+MVK, and acetaldehyde), concentrations are decreased and gradients are weakened by the enhanced mixing, leading to a nearly uniform modeled vertical profile in the lowest 60 m (3 h). Oxidized VOC (e.g., MACR+MVK) can deposit as efficiently as O₃ (Karl et al., 2010), yet simulated gradients of these species in the MIX case are weak. This suggests that either a compensation point has been reached or there are some primary sources of oxidized VOC from the forest canopy.

Above the canopy, CACHE accurately captures the characteristic features of the PBL, as indicated by the diurnal cycles of stability and turbulence. During the day, surface heating from canopy absorption of solar radiation induces instability and buoyancy-driven mixing, generating the convective mixed layer. The maximum height of the mixed layer varies with surface heating, but generally peaks mid-day around 1 km above the surface with a maximum strength occurring at approximately 500 m as modeled in Gao et al. (1993). At the end of the day, buoyant turbulence ceases in the absence of surface heating, and the stable nocturnal boundary forms beneath a well-mixed residual layer. CACHE does not explicitly model conditions in the free atmosphere nor the existence of an entrainment zone along a capping inversion. Therefore, our study focusses on exchange across the forest-atmosphere interface where we expect little influence from these upper PBL features.

3.2 O₃ and NO_x

We assess the performance of the model in simulating nearcanopy O₃, NO₂, and NO during the 4–5 August case period (Fig. 5). At all three heights (6, 20.4, and 34 m), observed O₃ mixing ratios peak at 40–50 ppbv shortly after the start of the simulation period and subsequently drop by 20 ppbv in 5 h. At that time, backward trajectories produced by the Hybrid Single Particle Lagrangian Integrated (HYSPLIT) model (Fig. S2 in the Supplement) indicate that air parcels observed at PROPHET prior to 08:00 EST on 4 August originated near Milwaukee and Chicago, and parcels arriving after 08:00 EST originated in northern Minnesota and Canada. Additionally, the local wind direction observed at PROPHET (Fig. S1) shifted from southerly to northwesterly over a 3- to 5-h period just after midnight. This transport, concurrent with a peak in observed NO₂ concentrations (Fig. 5), highlights the influence of large-scale advection on local O₃ and NO_x concentrations. Despite the inclusion of the same advection scheme in the BASE and MIX cases, the BASE-simulated O₃ is reduced after initialization while the concentrations in the MIX case increase to near-observed values. This indicates the influence of turbulent mixing on advection from aloft.

Under clean-air advection (i.e. following the frontal passage discussed in Sect. 2.2.6), above- (Fig. 5a) and uppercanopy (Fig. 5d) O_3 measurements show only small variability (e.g., approximately 10 ppbv) over the remainder of the simulation. BASE simulates a strong diurnal cycle, whereas MIX shows no clear diurnal cycle, indicating that locally



Fig. 5. Measured versus modeled time series for $(a, d, g) O_3$, $(b, e, h) NO_2$, and (c, f, i) NO at 34 m, 20.4 m, and 6 m for the 4–5 August 2009 simulation period, showing observed concentrations (OBS, black) and modeled concentrations as follows: the original model configuration with modeled turbulence and RACM chemistry (BASE; red); the corrected mixing simulation with RACM chemistry (MIX; green); the original mixing scheme with RACM-MIM chemistry (blue; MIM); and the revised mixing scheme with RACM-MIM chemistry (MIX+MIM; orange).

produced ozone is also highly sensitive to mixing. In the lower canopy (Fig. 5g), O₃ measured after the frontal passage is more variable, which is not captured by any of the model simulations. Wind directions are weakly variable from the west to northwest, showing no relationship with changes in O₃ concentration, suggesting that advection does not explain the observed variability. At the minima of the variability, lower-canopy concentrations are lower than the upper two measurements, indicating the potential relevance of an important O₃ canopy sink. In the canopy layer, deposition is the primary sink of O₃, exceeding modeled chemical loss by a factor of 10. Modeled O3 deposition velocity peaks mid-day at $1.6 \,\mathrm{cm \, s^{-1}}$ (Fig. 1), which is higher than other modeling studies (0.4 cm s^{-1}) , Finkelstein et al., 2000; Stroud et al., 2005), yet yields deposition fluxes that compare quite well with previous observations at PROPHET $(15-20 \text{ nmol m}^{-2} \text{ s}^{-1})$, Hogg et al., 2007). This suggests O₃ deposition is fairly well represented in the model. However, midday ozone deposition velocity increases by less than 0.02 % with enhanced mixing, indicating that deposition is not the main driver in the large concentration differences in the BASE and MIX ozone simulations. Lastly, we note that micrometeorological observations were not available in the lower canopy and may be underestimated by the MIX simulation, leading to an uncertainty in lower-canopy turbulence. Consequently, we are unable to evaluate whether these uncertainties in mixing can explain the lower canopy variability. In contrast with turbulent mixing, the enhancement in isoprene degradation between BASE and MIM show little impact on O₃ concentrations, indicating that O₃ is more sensitive to mixing than chemistry.

Like O3, observed NO2 displays a signature of anthropogenic advection on the early morning of 4 August. In the model, we have tuned the NO₂ advection rate (Sect. 2.2.5) to capture the higher concentrations observed at the beginning of the simulation and subsequent lower concentrations as the winds shift following the frontal passage. Observed NO_2 ranges from 1.2 ppbv at the beginning of the simulation to approximately 0.5 ppbv on the second night, with midday concentrations of less than 0.1 ppbv after photolysis. MIX+MIM underpredicts NO₂ slightly in the early part of the second night, likely due to an oversimplification in our NO2 advection scheme, missing downward transport from the residual layer, or an additional NO₂ source (e.g., emission from foliage). NO mixing ratios show a distinct diurnal pattern in the observations, peaking at nearly 180 pptv around mid-morning 5 August at 34 m, consistent with measurements taken over multiple years at PROPHET (Alaghmand et al., 2011). At night, mixing ratios reach as low as 1-5 pptv. The model reproduces the NO diurnal cycle well at the top of the canopy in all model scenarios, with concentrations overestimated on the first day in the BASE and MIM cases and good agreement for the MIX cases, and underestimated on the second day in the MIX and MIX+MIM cases. NO concentrations decrease with increasing canopy depth as a result of light attenuation lowering the NO yields from NO₂ photolysis. All four model scenarios are able to capture this effect of attenuation reasonably well.

3.3 Biogenic VOC and oxidation products

Figure 6 evaluates the four model scenarios against observations for primary BVOC, isoprene and monoterpenes.



Fig. 6. Same as Fig. 5, but for (a, c, e) isoprene and (b, d, f) monoterpenes.

Observations of isoprene show a strong diurnal cycle ranging from 0.2 ppbv at night to up to 2–3 ppbv at midday at all height levels. MIM simulations show slightly increased concentrations of isoprene over the BASE simulation, owing to lower OH availability as enhanced MACR+MVK concentrations increase the competition for OH. However, the BASE and MIM model scenarios have difficulty reproducing this diurnal pattern in two respects: (1) modeled concentrations increase later in the morning than observed, and (2) a rapid increase in concentrations occurs at sunset, causing a large discrepancy between the model and observations. Near the ground, this modeled pattern is dampened with a less pronounced evening peak and better drawdown of concentrations at night. Such a pattern has been frequently observed (cf. Martin et al., 1991), and is prevalent in many modeling studies (e.g., Sillman et al., 2002; Forkel et al., 2006; Barkley et al., 2011), which attribute the end-of-the-day increase to decreased mixing associated with the development of the stable nocturnal boundary layer. As noted in Sect. 3.1, CACHE captures the formation of the stable layer through the development of positive potential temperature gradients and highly reduced turbulent mixing below a well-mixed residual layer. In the BASE simulation, however, rapidly reduced mixing associated with an early and abrupt onset of the stable layer (Sect. 3.1, Fig. 2) leads to accumulation of isoprene (and other BVOC) near the canopy at the end of the day, further indicating the inability of the stability parameter (Eq. 7) to realistically capture the stable layer formation and its effect on turbulent exchange. As demonstrated in the MIX simulation, amplified turbulence and the gradual dissipation of daytime buoyancy-driven turbulence dampens these accumulations. We note, however, that underestimations of modeled in-canopy turbulence due to a lack of micrometeorological measurements in the lower canopy may result in insufficient exchange across the forest-atmosphere boundary. Our evaluation of the BASE and MIX turbulence schemes presented in Sect. 3.1 discusses several observed model-measurement discrepancies in the BASE $K_{\rm H}$ that are corrected in the MIX scenario, including the end-of-day decrease in mixing at sundown. This indicates the sensitivity of the top-of-canopy BVOC flux to turbulence and emphasizes the importance of an accurate representation of in-canopy mixing in models. Overall, these results highlight the importance of an accurate representation of in-canopy mixing in models and the need for improved simulation the transition from the convective to the nocturnal PBL.

While enhanced mixing improves the diurnal evolution of isoprene, modeled mixing ratios exceed observations on the second day by 1-3 ppbv, likely due to an underestimate of isoprene oxidation. Oxidation of isoprene in the model is primarily controlled by reaction with OH, with loss rates in the model peaking at midday around 11 pptv min⁻¹ (Fig. 7). Oxidation by ozone follows at much smaller rates (up to $2 \text{ pptv} \text{min}^{-1}$ around sunset) and small contributions by loss with the nitrate radical at night (approximately $0.5 \text{ pptv min}^{-1}$). Simulated NO₃ mixing ratios at 34 m (not shown) peak at 4 pptv the first night under elevated NO₂ concentrations from regional transport and decrease to 0.3 pptv on the second night when chemistry is dominated by local emissions. Past estimates of NO₃ at PROPHET range from 0.4 pptv (Pratt et al., 2012) to 2-3 pptv (Faloona et al., 2001), although we note that NO₃ has not been measured at the site before, making it difficult to evaluate the model. The primary model NO₃ source is the nighttime oxidation of NO₂ by O₃, yet observed NO₂ concentrations at the site remain relatively low; therefore, we should expect relatively low concentrations of NO₃ and low isoprene-NO₃ oxidation rates. A potential explanation for the lack of nighttime oxidation may be OH concentrations, as discussed in Sect. 3.4. In addition to the effect of nighttime chemistry, Ganzeveld et al. (2008) suggest that downward transport of oxygenated VOC from the residual layer may lower OH availability, further reducing isoprene oxidation, particularly in the morning. This downward transport is captured in our model for oxygenated VOC during the second morning of the simulation, yet model overestimations of isoprene begin as early as sundown on the prior day, suggesting that the effect of entrained oxygenated VOC from the residual layer on OH concentrations is likely small relative to the observed discrepancies in nighttime chemistry.

Monoterpenes ($C_{10}H_{16}$) are grouped together as a total monoterpene concentration by the PTR-MS. As noted at other forest sites (Bouvier-Brown et al., 2009), observations show a different diurnal cycle than isoprene. The early morning and late evening peak (Fig. 6) and higher concentrations at night than during the day have been attributed to high photooxidation during the daytime and an accumulation at night as these temperature-dependent emissions continue in the absence of sunlight. Above the canopy, higher concentrations



Fig. 7. Modeled isoprene loss rates with respect to reaction with OH, NO_3 , and O_3 at 34 m from the MIX case.

of monoterpenes are observed at the beginning of the simulation, suggesting a potential advective source of terpenes to the site in the first six hours of the simulation, which is consistent with air traveling over the forested state. RACM-MIM simulates terpene concentrations to be very similar to the RACM case due to no changes in the MIM terpene oxidation scheme. As with isoprene, however, enhanced mixing greatly improves the model-measurement agreement in terms of magnitude of concentrations, though the diurnal cycle is only weakly captured.

Formaldehyde (HCHO) is an important VOC oxidation product and is typically produced in relatively large quantities from the oxidation of isoprene. While anthropogenic VOC can also provide a substantial source of HCHO (Pang et al., 2009), their effect on local HCHO concentrations observed at the PROPHET site is minor (Sumner et al., 2001) unless under advective conditions from the south. Observed mixing ratios at the site are 0.5-1 ppbv (Fig. 8), reflecting values that are slightly lower than the 1999 field campaign observations (0.5-12 ppbv, Sumner et al., 2001), consistent with the expected reduced photochemistry in the summer of 2009. An advection source is apparent at the beginning of the simulation with higher HCHO concentrations both above and below the canopy (Fig. 8a, d, g). When an advective HCHO source is added at 45-106 m, measured-modeled comparisons improve above the canopy but not below the canopy, suggesting either that there is in-canopy production that the model does not capture, in-canopy deposition is overestimated, or mixing within the canopy is stronger than simulated. HCHO midday deposition velocity is higher than other studies $(2.3 \text{ cm s}^{-1} \text{ in our model as compared to})$ $1.5 \,\mathrm{cm}\,\mathrm{s}^{-1}$ in Sumner et al. (2001)), yet the model still overestimates HCHO in all simulations at all heights, even after enhancing the in-leaf reactivity to that of O_3 , as suggested for other oxidized VOC (Karl et al., 2010). This indicates the possible existence of a compensation point, similar to that observed of other compounds (e.g., NO₂, acetone, and methanol, Ganzeveld et al., 2002, 2008), at which the concentration in the plant mesophyll matches or exceeds that of the ambient air, restricting further deposition and potentially inducing emissions.HCHO deposition velocity decreases by less than $0.1 \,\mathrm{cm}\,\mathrm{s}^{-1}$ with enhanced mixing, as with that of O₃. When advection is not playing a role (the second day of the simulation), modeled HCHO exhibits a diurnal cycle with higher concentrations during the day especially in the BASE and MIM cases. Both above and below the canopy, the change to MIM increases midday HCHO concentrations by about 15 % due to larger HCHO yields from BVOC oxidation (Geiger et al., 2003). MIM makes additional HCHO from the isoprene peroxy self-reaction (ISOP+ISOP), plus the new methacrolein peroxy radicals (MACP). Enhanced mixing (MIX and MIX+MIM) weakens the diurnal pattern of HCHO, better reflecting observations.

Other key BVOC oxidation products are the lumped species methacrolein and methyl vinyl ketone (MACR+MVK or C_4H_6O). These compounds are detected at the same nominal mass on the PTR-MS and are also lumped in the RACM mechanism. Observed concentrations peak in the early portion of the simulation both above and below the canopy (Fig. 8) due to advection of oxidation products from the south. Adding an advective source of MACR aloft improves measured-modeled agreement at the beginning of the simulation at all measurement heights. The MIX case improves concentrations as compared to observations, yet removes the observed diurnal pattern. Changing to the MIM mechanism doubles the BASE-case concentrations of MACR+MVK due to the increased yield in MACR+MVK by the reaction of first-generation oxidation products of isoprene with NO. Consequently, modeled concentrations of the MACR RACM-MIM surrogate overestimate measured MACR+MVK by a factor of three throughout the profile. This finding is consistent with chamber study comparisons of RACM and RACM-MIM by Geiger et al. (2003), who attribute the result to measurements only accounting for MACR+MVK while the RACM species also includes all other unsaturated C₄ carbonyls. Past studies suggest that dry deposition rates for MACR+MVK may be underestimated (Karl et al., 2004, 2010; Pugh et al., 2010), yet the modeled MACR+MVK deposition velocity mid-day peak of $1.6 \,\mathrm{cm}\,\mathrm{s}^{-1}$ (Fig. 1) compares well with observations by Misztal et al. (2011, $1-2 \text{ cm s}^{-1}$) and Karl et al. (2010, < $2.4 \,\mathrm{cm\,s^{-1}}$) after modifying the reactivity factor for oxidized VOC following Karl et al. (2010). Enhanced mixing reduces the MACR deposition velocity by about 5%.

We also evaluate the simulation of the biogenic oxidation product hydroxyacetone ($C_3H_6O_2$) added to the RACM-MIM mechanism (denoted as HACE) representing a major product of MACR+MVK oxidation. While calibrated observations of HACE are not available, the model simulates a diurnal cycle with mixing ratios ranging up to 100 pptv that



Fig. 8. Same as Fig. 5, but for (**a**, **d**, **g**) formaldehyde, (**b**, **e**, **h**) MACR+MVK, and (**c**, **f**, **i**) hydroxyacetone. Observations of hydroxyacetone are not available.



Fig. 9. Measured and modeled correlations between MACR+MVK and isoprene (left), and hydroxyacetone and MACR+MVK (right) at 34 m for 5 August 2009 between 11:00-17:00 EST. The squared correlation coefficients (R^2) and slopes of the regression lines are given in the upper right-hand corners in colors corresponding to the appropriate model scenario.

decrease slightly with increased mixing (Fig. 8), which is slightly lower than the range of the uncalibrated measurements (200–500 pptv). Glyoxal (GLY) was also measured at the site, with mixing ratios reaching up to 25 pptv during midday, and with a clear advective signal on the first day of the simulations (Fig. S4). However, both RACM versions only form GLY from anthropogenic precursors and do not include the production from any biogenic species; therefore, modeled mixing ratios are on the order of 0.01–1 pptv. Observations of GLY suggest local biogenic production of GLY, a source that could be included in future models.

We compare the daytime (11:00–17:00 EST) ratios of (MACR+MVK)/isoprene and HACE/(MACR+MVK) to evaluate the ability of the mechanisms to reproduce observed BVOC oxidation. Over the full field campaign (not shown), the observed (MACR+MVK)/isoprene ratio is 0.18, substantially lower than observed in the Amazon (0.44,Karl et al., 2009), yet comparable to observations from previous PROPHET studies (0.12, Apel et al., 2002). Apel et al. (2002) observe lower MACR+MVK concentrations under westerly flow associated with clean-air advection from Lake Michigan, leading to lower MACR+MVK/isoprene ratios. Additionally, as shown by comparing the ratios of MACR+MVK/isoprene with wind direction over the full two-day simulation (Fig. S4 in the Supplement), we observe elevated ratios (0.71) under southerly flow associated with polluted advection. Due to this strong variability in MACR+MVK with respect to wind direction at the PROPHET site, the correlation between MACR+MVK and isoprene is substantially weaker ($R^2 = 0.25$) than observed by Karl et al. (2009, $R^2 = 0.86$). Figure 9 compares the observed daytime ratios against the four model scenarios for the second day (5 August) to examine local chemistry in the absence of pollution transport. The observed (MACR+MVK)/isoprene ratio of 0.03 is much lower than the mean daytime ratio for the full campaign (0.18), likely due to enhanced clean-air advection from the northwest. The BASE and MIM scenarios yield negative ratios, indicating inefficient oxidation of isoprene. This is consistent with the overestimation of isoprene by the BASE and MIM cases, particularly on the second day, as a result of inefficient mixing out of the canopy. In these scenarios, OH is depleted before isoprene is completely oxidized, leading to insufficient production of MACR+MVK given the amount of isoprene available. This is consistent with past studies, who propose the need for a OH recycling mechanism (Lelieveld et al., 2008). With enhanced mixing (MIX and MIX+MIM scenarios), isoprene is oxidized more effectively leading to ratios that correlate well with observations. Additionally, the HACE/(MACR+MVK) relationship can highlight the added oxidation capacity when including the new MACR oxidation pathways in MIM. For local conditions on 5 Aug, the modeled ratios are 0.08 and 0.09 for the MIM and MIX+MIM cases, respectively, both of which are substantially lower than observed in the Amazon (0.3, Karl et al., 2009) due to lower oxidant concentrations. While the MIX+MIM case ratios are lower than the MIM, we note that the correlation is weaker in the MIX+MIM case ($R^2 = 0.67$) than in the MIM case ($R^2 = 0.82$), indicating a large uncertainty with these ratios.

To summarize, the original CACHE BASE simulations strongly overestimate isoprene concentrations, particularly in the early evening and at nighttime. This is a known problem in models that occurs at all scales (e.g., 1-D models, 3-D models, etc.) and we attribute this increase at the end of the day to improper mixing in the model. The revised mixing scheme, which is based on observed friction velocities and vertical velocity standard deviations, greatly improves the simulation of primary BVOC at most model levels. Therefore, a realistic representation of boundary layer turbulence is critical for modeling forest-canopy exchange and its effect on BVOC chemistry accurately. Oxidation products such as HCHO and MACR+MVK are overestimated by the BASE model simulations, with the greatest measured-modeled improvement resulting from the change in mixing parameterization versus the chemical mechanism. In general, the more detailed biogenic oxidation scheme (RACM-MIM) increases the oxidation products to three times more than observed, although the mechanism does improve modeled HO_x as will be discussed in the next section.

3.4 HO_x and OH reactivity

Modeled OH concentrations reproduce the diurnal cycle and magnitude of observed OH $(1-2.5 \times 10^6 \text{ molecules cm}^{-3} \text{ at})$ midday) (Fig. 10). Difficulties associated with transmission of laser power to the top of the tower led to few measurements of OH greater than the limit of the detection of the instrument (approximately 1×10^6 molecules cm⁻³) during this time period; therefore, Fig. 10 displays an average diurnal cycle of OH of the two simulation days with a peak value of 2×10^6 molecules cm⁻³. The model produces higher OH concentrations on the first day of simulation due to the higher oxidation from incoming advection, while modeled concentrations on the second day are approximately half the observed values. The MIX and MIM cases decrease modeled OH by about 10 and 20%, respectively, on the first day, and on day two, MIX increases modeled OH by about 10%. From the vertical profiles (Fig. 11), in-canopy OH concentrations are low, suggesting small OH production rates. OH concentrations are highest above the canopy where substantial production from O_3 photolysis and subsequent reaction with H₂O occurs. In general, enhanced mixing increases the modeled OH concentrations at all heights, whereas the change from RACM to RACM-MIM decreases OH from the surface to 3 *h*. An exception is during the anthropogenic advection event on the morning of 4 August, when enhanced mixing increases above canopy OH and decreases OH within the canopy; implementing RACM-MIM increases OH throughout the column, due to increased production from the HO_2 + NO reaction.

For HO₂, the measurements show a strong diurnal cycle that is reproduced by the model (Fig. 10). In general, the model underestimates HO₂ in the BASE case simulation, with a slight increase in HO₂ from the MIM simulation. However, recent studies suggest that the detection of HO₂ radicals using chemical conversion to OH by reaction with added NO may be sensitive to the detection of a fraction of hydroxyalkyl peroxy radicals produced from the OHinitiated oxidation of alkenes (Fuchs et al., 2011). Calibrations of the Indiana University FAGE instrument indicate that approximately 90% of isoprene-based hydroxyalkyl peroxy radicals are detected in addition to HO₂, while only 5 % of propane-based alkyl peroxy radicals are detected. Given that isoprene dominates the HO₂ radical chemistry at this site, the measured HO₂ concentrations (HO₂^{*}) likely reflect the sum of both HO₂ and isoprene peroxy radicals (ISOP). In Fig. 10, we compare measured HO₂^{*} with a similar metric from the model ($HO_2^* = HO_2 + ISOP$) and this greatly improves measured-modeled agreement. The increase in late evening modeled HO_2^* is due to an accumulation of the ISOP radicals, an artifact from the end-of-day increase in isoprene concentrations (see Sect. 3.3). When changing to RACM-MIM, the reaction rate for HO₂ + ISOP increases and the self-reaction ISOP + ISOP is explicitly added, leading to greater HO_2^* destruction and improving the model-measurement agreement. In the vertical profiles (Fig. 11), the model produces a strong source of HO₂ above the canopy with some in-canopy production. RACM-MIM increases HO2 throughout the vertical profile, particularly in the daytime. The enhanced mixing (MIX case) increases both in- and above-canopy HO₂ concentrations in the morning and above the canopy during the night. Otherwise, slight decreases in HO₂ occur. During the advection event on 4 August, increased NO_x leads to decreased HO₂ in both the MIX and MIM cases due to loss with NO.

OH reactivity (R_{OH}) represents the total first order loss rate of OH (inverse of the OH lifetime). Measured R_{OH} values during CABINEX 2009 range from 0–2 s⁻¹ at night to up to 10 s⁻¹ during the day (Fig. 12, left panel), which compare well with previous measurements at PROPHET (Di Carlo et al., 2004). Modeled R_{OH} is calculated by summing the product of the rate constant and reactant concentrations for all species that consume OH. Modeled R_{OH} compares best with observations for the MIX scenario, due to the poor reproduction of the observed diurnal cycle of isoprene simulated by the BASE and MIM simulations. This again suggests the dependence of modeled R_{OH} on the vertical mixing



Fig. 10. Same as Fig. 5, but for OH (left), HO₂ (middle), and HO₂^{*} (right) at 32 m.



Fig. 11. Vertical profiles of modeled OH (top) and HO₂ (bottom) for the BASE case (left) and the absolute difference between the BASE and MIX cases (middle) and BASE and MIM cases (right). Blue values indicate higher concentrations in the BASE case, and red values indicate higher concentrations in the MIX or MIM cases.

in the model. For modeled R_{OH} , we speciate contributions from BVOC (isoprene, API, LIM) and oxidation productions (HCHO and MACR+MVK). During the afternoon, BVOC account for approximately 85% of the reactivity in the model, followed by the CO at 15%, whereas the oxidation products and CO dominate at night and in the early morning (Fig. 12, right panel). Contributions from methane and NO₂ are relatively small. Kim et al. (2011) note that the oxidation products can account for about 8% of the reactivity if NO concentrations are low. However, as noted by Kim et al. (2011) and Karl et al. (2009), the R_{OH} tends to increase when photochemically aged air masses arrive at the observation site, which is evident on the first day of the simulation. Because we are including the advection of some primary anthropogenic and secondary oxidation products (Table 2), we correctly model this increase in $R_{\rm OH}$ on the first day of simulation. The second day of the simulation reflects the local conditions, with slightly lower $R_{\rm OH}$ that is overestimated by the model.

4 Summary and conclusions

This manuscript presents results from a 1-D canopychemistry model, CACHE, applied to a northern Michigan



Fig. 12. Total OH reactivity measured (OBS) and modeled (BASE, MIX, MIM, and MIX+MIM) at 30.9 m for 4–5 August 2009 (left); modeled total OH reactivity and from speciated contributions of NO₂, CO, CH₄, BVOC (ISO+API+LIM), HCHO, and MACR for the MIX case (right).

mixed hardwood forest to elucidate in-canopy atmospheric chemistry during the CABINEX 2009 field campaign. CACHE calculates vertical mixing within and above the forest canopy using K-theory, a parameterization used by many 1-D and 3-D models despite its limitations in the canopy roughness layer. Chemical transformation is modeled using RACM, a condensed mechanism that can cover a broad range of chemical situations but with limited BVOC chemistry particularly under low-NO_x conditions. In this study, we test the model sensitivity of vertical gradients of BVOC and their oxidation products to (1) turbulent exchange and (2) chemistry. First, we account for turbulence in the canopy roughness layer by applying the modified K-theory parameterization of Makar et al. (1999) and adjusting the model with hightime-resolution sonic anemometer measurements of friction velocity and vertical velocity standard deviation. Second, we implement an expanded version of RACM with more explicit BVOC chemistry, RACM-MIM.

Traditional K-theory (i.e. BASE) underestimates forest canopy exchange by 0.5-2 orders of magnitude, leading to an overly-strong diurnal cycle of ozone, and an overestimate of NO_x, BVOC and their oxidation products that accumulate within and above the canopy to 2-3 times higher than observed. This highlights the issue that models with differing turbulence parameterizations in and above the canopy may inhibit exchange across the top of the canopy due to a discontinuity in mixing between the two equations. In addition, traditional K-theory, in which turbulence is driven by a prognostic temperature profile, does not capture the observed gradual onset and termination of convective mixing due to uncertainties with the empirical stability parameter at the transition between stable and unstable conditions. This leads to anomalous spikes in primary BVOC near the canopy, particularly around sunset, that are not present in the observations. Driving near-canopy vertical mixing with micrometeorological observations (e.g., MIX) improves the representation of vertical mixing as evidenced in the improved vertical profiles and diurnal cycles of BVOC and their oxidation products. While this parameterization cannot account for asymmetric transport associated with coherent structures and other non-Fickian diffusion processes, this method provides substantial improvement in the model simulations. Adding additional BVOC oxidation pathways with the RACM-MIM mechanism slightly increases isoprene and HCHO (15%) with greater changes in MACR+MVK (80%), although these concentrations were about five times higher than observed at all heights. Past research suggests that MACR+MVK may constitute only a fraction of the MACR RACM surrogate, and that models may underestimate MACR+MVK surface deposition; however, deposition velocity for MACR+MVK in the model compares well with observations. Changes in O_3 and NO_x concentrations with enhanced isoprene chemistry were negligible. Overall, the parameterizations tested in this study suggest BVOC and their oxidation products can be very sensitive to the mixing parameterization.

The impact of vertical mixing on HO_x chemistry is dependent on the advection conditions. Advection from polluted regions (e.g., the first day of our simulation) increases OH in the region of advection (45-106 m) and decreases OH below the level of advection. When local chemistry dominates, an increase in mixing increases OH concentrations suggesting that the canopy can be a HO_x source. For HO₂, an increase in mixing tends to decrease concentrations regardless of advection conditions. With changes to the RACM-MIM chemistry, OH decreases due to increased secondary oxidation of biogenic oxidation products and HO₂ decreases throughout the profile. While the additional BVOC oxidation pathways of RACM-MIM improve HO₂, the overestimation of MACR+MVK suggests that the mechanism pathway may not be properly capturing the oxidation of the biogenic oxidation products. Additionally, we find that GLY is underestimated in the model by an order of magnitude (Fig. S3 in the Supplement), suggesting a missing primary biogenic oxidation source.

Typically, 1-D models are subject to several aspects of uncertainty, including (1) the emissions from the canopy and soil, (2) the reactions described by the chemical mechanism, (3) the exchange driven by the turbulence parameterization, and (4) the sink to surface deposition. We have evaluated each of these aspects in this paper, while focussing our sensitivity study on mixing and chemistry. The BVOC emissions have been fairly well-constrained by multiple measurements at the site; however, we find that observed emission factors for isoprene are likely on the lower end of the spectrum due to the unusually cool summer at UMBS. If emission rates were higher than modeled in this study, BVOC concentrations accumulate in the model to unrealistic concentrations. In theory, this could be matched by higher reactivity in the forest, with increased oxidation by OH through the implementation of an OH recycling mechanism (Lelieveld et al., 2008) or enhanced deposition of compounds competing for OH (e.g., oxygenated VOC, Karl et al., 2004, 2010). With enhanced oxygenated VOC deposition according to Karl et al. (2010)

applied in this study, BVOC still accumulate. OH concentrations were well-matched with the observations; therefore, we did not implement a recycling mechanism in this study. Additionally, underestimations in peroxide (e.g. H₂O₂) deposition may inhibit in-canopy photochemistry and exchange (Ganzeveld et al., 2006); however, peroxide measurements were not available to constrain deposition rates. We tested the sensitivity of the model to higher NO_x , potentially from a local or advective source by increasing NO₂ advection rates until the NO_x concentrations match urban levels (not shown). With higher isoprene emissions (e.g., the mean value of Ortega et al., 2007), higher NO_x can increase the oxidation and reduce BVOC concentrations to observed values, however modeled NO_x concentrations then exceed observed values by an order of magnitude. It is also possible that OH concentrations are too low in the model (e.g., Fig. 10, left panel); however, our modeled BVOC oxidation products are already higher than observed and we have good measured-modeled agreement in OH reactivity. Therefore, we have evidence to show that the modeled emissions and chemistry balance in the model represents the observed conditions fairly accurately.

Overall, we find that an improved representation of incanopy turbulent transport based on micrometeorological observations and a consideration for near-field effects improves the simulation of concentrations and vertical gradients of BVOC and their oxidation products observed during the CABINEX 2009 campaign. The change to a mechanism with more specific BVOC pathways slightly improves agreement with observations for HO₂, but produces more BVOC oxidation products than observed (e.g., HCHO, MACR+MVK). While the yields of BVOC oxidation products in MIM may be too high (e.g., Geiger et al., 2003) or surface deposition rates may be too low (Pugh et al., 2010), observed concentrations of these primary oxidation products suggest that our in-canopy oxidation is within the observational constraints. However, we note that implementation of other isoprene oxidation mechanisms may yield different results. Our results show that mixing in the canopy may be more important than changes to BVOC chemistry mechanisms for accurate modeling of BVOC chemistry and forest-atmosphere exchange, and point to the need for a revised in-canopy turbulence parameterization in existing 1-D and 3D atmospheric models. A thorough intercomparison of turbulence and BVOC chemistry data from a variety of forest ecosystems is required to assess the applicability of our results on the global scale. Other aspects of the forest canopy, including the turbulence structure of the lower canopy and the effect of vertical heterogeneity of vegetation (i.e. an understory and overstory of differing plant type) on the oxidation capacity of the canopy and forest-atmosphere exchange of BVOC may provide further information for understanding the vertical profiles of BVOC, their oxidation products, and their contribution to tropospheric chemistry.

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