Land cover change impacts on atmospheric chemistry: simulating projected large-scale tree mortality in the United States

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Abstract. Land use and land cover changes impact climate and air quality by altering the exchange of trace gases between the Earth’s surface and atmosphere. Large-scale tree mortality that is projected to occur across the United States as a result of insect and disease may therefore have unexplored consequences for tropospheric chemistry. We develop a land use module for the GEOS-Chem global chemical transport model to facilitate simulations involving changes to the land surface, and to improve consistency across land–atmosphere exchange processes. The model is used to test the impact of projected national-scale tree mortality risk through 2027 estimated by the 2012 USDA Forest Service National Insect and Disease Risk Assessment. Changes in biogenic emissions alone decrease monthly mean O3 by up to 0.4 ppb, but reductions in deposition velocity compensate or exceed the effects of emissions yielding a net increase in O3 of more than 1 ppb in some areas. The O3 response to the projected change in emissions is affected by the ratio of baseline NOx : VOC concentrations, suggesting that in addition to the degree of land cover change, tree mortality impacts depend on whether a region is NOx-limited or NOx-saturated. Consequently, air quality (as diagnosed by the number of days that 8 h average O3 exceeds 70 ppb) improves in polluted environments where changes in emissions are more important than changes to dry deposition, but worsens in clean environments where changes to dry deposition are the more important term. The influence of changes in dry deposition demonstrated here underscores the need to evaluate treatments of this physical process in models. Biogenic secondary organic aerosol loadings are significantly affected across the US, decreasing by 5–10% across many regions, and by more than 25% locally. Tree mortality could therefore impact background aerosol loadings by between 0.5 and 2 µg m$^{-3}$. Changes to reactive nitrogen oxide abundance and partitioning are also locally important. The regional effects simulated here are similar in magnitude to other scenarios that consider future biofuel cropping or natural succession, further demonstrating that biosphere–atmosphere exchange should be considered when predicting future air quality and climate. We point to important uncertainties and further development that should be addressed for a more robust understanding of land cover change feedbacks.

1 Introduction

Land use and land cover changes impact climate by altering energy exchange at the surface of the Earth, and by altering the composition of the atmosphere through changes in biogeochemical cycling (Feddema et al., 2005; Pielke et al., 2011). Though recognized as a crucial factor in future climate scenarios (van Vuuren et al., 2011), projections of land use and land cover change remain highly uncertain (Smith et al., 2010). The terrestrial biosphere also plays an important role in surface–atmosphere exchange of reactive trace species that control the oxidative chemistry of the troposphere (Arneth et al., 2010; Laathawornkitkul et al., 2009; Mellouki et al., 2015; Monson and Holland, 2001), so that...
changes in vegetation can further impact air quality and climate (Heald and Spracklen, 2015; Unger, 2014). These changes can be human-driven (e.g., urbanization, forestry management, and agricultural practices) or natural (e.g., wildfires, insect infestations, and biome shifts). Over the 21st century, variations in biogenic volatile organic compound (BVOC) emissions due to climate change and crop management will likely impact surface ozone (O₃) and secondary organic aerosol (SOA) concentrations (Ashworth et al., 2012; Smith et al., 2001; Ganzeveld et al., 2010; Hardacre et al., 2013; Heald et al., 2008; Wu et al., 2012). Here we consider the air quality and atmospheric chemistry implications of another form of land cover change on relatively shorter timescales: large-scale insect- and disease-driven tree mortality.

Modifications to vegetation distribution, plant type, canopy characteristics, and soil properties alter the regional emission and deposition of reactive trace gases from the terrestrial biosphere. For example, large-scale deforestation of the Amazonian rainforest, the expansion of oil palm plantations in Asia, and cultivation of biofuel feedstocks can significantly alter BVOC emissions, with various implications for secondary pollutants (Ashworth et al., 2012; Beltman et al., 2013; Ganzeveld, 2004; MacKenzie et al., 2011). In the eastern US, harvest practices and forest management have likely resulted in a net increase in BVOC emissions since the 1980s, counteracting successful anthropogenic emission reductions (Purves et al., 2004). Ecological succession, either from anthropogenic land management or natural processes, could also impact regional chemistry (Drewniak et al., 2014). Some changes in land cover have compensating impacts. For example, higher vegetation density could lead to increased O₃ precursor emissions but also faster depositional losses (Ganzeveld et al., 2010; Wu et al., 2012). Consequently, we require models that account for the combination of these mechanisms in a consistent manner to understand the relevant net impacts on air quality and climate.

Almost a third of the Earth’s land surface is covered by forests, providing a variety of economic, recreational, and ecosystem services including regulating climate through complex biogeophysical and hydrological feedbacks and by taking up CO₂ from the atmosphere (Bonan, 2008; MEA, 2005). A prominent risk to forests in the near future (< decades) is tree mortality resulting from insect attack and disease (Krist et al., 2014). Biotic disturbances resulting in tree mortality occur naturally at low and predictable rates (Smith et al., 2001), but in the coming decades many forests across the US are predicted to experience tree mortality well above background. Between 2013 and 2027, over 80 million acres of treed land in the United States are projected to experience basal area mortality rates exceeding 25%, with some tree species at risk of losing more than 50% of their volume (Krist et al., 2014). The dominant contributing hazards are expected to be root diseases, bark beetles, and oak decline, with highest risks occurring in Idaho, Montana, and Oregon in the western US and in Rhode Island, Connecticut, and Massachusetts in the eastern US (Krist et al., 2014). The wood volume lost from insects and pathogens can cost the US several times more than losses by wildfire (Dale et al., 2001), and can have a major impact on carbon cycling (Hicke et al., 2012), but the atmospheric chemistry impacts have not been fully explored. Berg et al. (2013) simulated the impact of past bark beetle infestations in the western US using a decade of tree mortality data. They found large changes to monoterpenes emissions, and subsequently SOA concentrations, that could potentially affect background aerosol concentrations and visibility in pristine regions.

Given the important role of natural emissions in the chemistry of the atmosphere (Zare et al., 2014), large-scale future tree mortality may influence ozone production and organic aerosol concentrations. Nonattainment of O₃ air quality standards in the US is more sensitive to BVOC emissions than anthropogenic VOC emissions (Hakami et al., 2006), and secondary organic aerosol mass can be dominated by biogenic sources (Pye et al., 2010). The main anticipated effect of tree mortality is a reduction of the BVOC emissions from the species that die, but a change in local vegetation density would also be expected to impact dry deposition, since this is directly related to the surface area available for deposition. Vegetation changes can also affect the local microclimate, further impacting depositional processes. Changes in dry deposition may be significant for species (such as O₃) whose depositional losses are competitive with chemical sinks near the surface of the earth. Finally, since soil NOₓ emissions to the atmosphere depend not only on available nitrogen and soil conditions but also on the extent of uptake to vegetation canopies, changes to forests driven by tree mortality could impact these emissions as well.

Here we use the GEOS-Chem chemical transport model to investigate the impact of projected tree mortality on atmospheric composition. We harmonize the description of land cover across the relevant surface–atmosphere exchange processes, and use this adapted model to simulate the impacts of predicted tree losses as a result of insect and disease in the United States from 2013–2027. We explore how changes in dry deposition might compensate for changes in chemical production by separating these impacts in individual simulations. We highlight that even modest tree mortality could impact regional atmospheric chemistry across the US, and identify specific regions for further investigation. We also discuss directions for future development to better understand the influence of vegetation changes on atmospheric reactivity and composition.
2 Model description

2.1 General description of GEOS-Chem

We use the GEOS-Chem model (Bey et al., 2001; www.geos-chem.org) v9-02 to simulate the impact of changes in vegetation on atmospheric chemistry. GEOS-Chem is a global 3-D atmospheric chemical transport model driven by assimilated meteorology from the NASA Global Modeling and Assimilation Office. Our simulations are driven by GEOS-5 meteorological data for the year 2010 and performed over North America at the nested horizontal resolution of 0.5° × 0.667°, with dynamic boundary conditions supplied from a global simulation at 2° × 2.5°.

The model includes a detailed HOx-NOx-VOC-O3 chemical scheme originally presented by Bey et al. (2001). The chemical mechanism includes over 90 species (including the following lumped categories: > C3 alkanes, > C2 alkenes, > C4 alkanes, > C3 alkenes, > C4 aldehydes, > C1 alcohols, and > C1 organic acids), over 200 chemical reactions, and over 50 photolysis reactions, incorporating the latest JPL and IUPAC recommendations. Detailed isoprene oxidation chemistry is included, following Paulot et al. (2009a, b) as implemented for GEOS-Chem by Mao et al. (2013). Explicit oxidation pathways are not yet included for terpenes. Given that isoprene dominates biogenic OH reactivity over the continental US, we assume terpenes play a minor role outside of SOA formation (see below) in our land cover change simulations. Gas-aerosol partitioning in the sulfate-nitrate-ammonium system is described according to the thermodynamic ISORROPIA II equilibrium model (Fountoukis and Nenes, 2007).

Carbonaceous aerosol sources include primary emissions from fossil fuel, biofuel, and biomass burning (Park et al., 2003) and reversible SOA formation following Pye et al. (2010). Secondary organic aerosol is lumped into five species based on the parent hydrocarbons (terpenes, isoprene, light aromatics, and intermediate volatile organic compounds, semivolatile organic compounds (SVOCs), and oxidized SVOCs). Aerosol yields are parameterized using a volatility basis set (Donahue et al., 2006) for aerosol systems with multiple parent hydrocarbons or aerosol formation pathways, or an Odum 2-product approach (Odum et al., 1996) for systems with one parent hydrocarbon. Emitted biogenic parent hydrocarbons are lumped in the following manner: (1) α-pinene + β-pinene + sabine + carene; (2) limonene; (3) t-β-cymene + myrcene + other monoterpenes; (4) farnesene + carphyllene + other sesquiterpenes; and (5) isoprene. SOA yields from ozonolysis (at high and low NOx) and nitrate radical oxidation are represented in the model for groups (1)–(4), while yields from photooxidation (initiated by OH) and nitrate radical oxidation are represented for isoprene. Further gas-aerosol phase coupling occurs for example through N2O5 uptake (Evans and Jacob, 2005) and HO2 uptake (Mao et al., 2013).

We use anthropogenic emission inventories according to the NEI-2005 inventory for the United States (http://www.epa.gov/ttnchie1/trends/), CAC for Canada (http://www.ec.gc.ca/pdb/cac/), and BRAVO (Kuhlms et al., 2005) for Mexico, and scale these to the year 2010 following van Donkelaar et al. (2008). The model also includes biomass burning emissions (GFED3, Mu et al., 2011), lightning NOx (Murray et al., 2012), and volcanic SO2 emissions (Fisher et al., 2011). Soil NOx and BVOC emission coefficients are described below.

2.2 Default land–atmosphere exchange in GEOS-Chem

Here we briefly describe the main mechanisms in the model by which vegetated land cover impacts atmospheric chemistry.

GEOS-Chem v9-02 includes the Berkeley-Dalhousie Soil NOx Parameterization (Hudman et al., 2012). In this parameterization, the flux of NOx from soils is a function of temperature, soil moisture, and emission coefficients which depend on available nitrogen and biome type. Biomes (and basal emission coefficients) are defined according to Steinkamp and Lawrence (2011), with 24 different land cover types. Dry spell length is also included to account for pulsing. A canopy reduction factor is calculated according to leaf area index (LAI), wind speed, and surface resistance, and is designed to simulate the uptake of NOx by vegetation following soil emission (Wang et al., 1998).

Biogenic VOC emissions from vegetation are calculated using the Model of Emissions of Gases and Aerosols from Nature (MEGAN v2.02: Guenther et al., 2006, with updates from Sakulyanontvittaya et al., 2008). In GEOS-Chem v9-02, mapped basal BVOC emission factors are provided as an input to the model and are modulated online by activity factors that are a function of temperature, LAI, photosynthetically active radiation (PAR), and average leaf age.

Dry deposition is calculated by the resistance-in-series scheme of Wesely (1989), using a “big-leaf” approximation where the deposition surface is treated as a single uniform surface (or leaf). Dry deposition velocities are calculated as a combination of aerodynamic resistance ($R_A$), laminar layer resistance ($R_L$), and surface resistance ($R_s$). $R_s$ is calculated separately for unstable, moderately stable, and very stable atmospheric conditions, and is a function of roughness heights (which would be a function of land cover type) that are provided by the meteorological input data. $R_L$ depends on meteorological data and the identity of the gas-phase species being deposited. The $R_c$ parameterization depends on the solubility and reactivity of individual chemical compounds and on land type according to Wesely (1989), and is scaled by LAI. Land types are derived by the USGS global land characteristics database (http://edc2.usgs.gov/glc2/globdoc2_0.php), known also as the Olson Land Map. Over 70 land types are represented and mapped to the 11 deposition surface types given by Wesely (1989). Aerosol deposition is also parameterized by the
resistance-in-series scheme according to Zhang et al. (2001), with deposition to snow and/or ice as presented by Fisher et al. (2011). Gravitational settling of dust and sea salt is described according to Fairlie et al. (2007) and Alexander et al. (2005) respectively.

As described above, the parameterizations of soil NO₃ emissions, BVOC emissions, and dry deposition all depend on LAI in some way. By default, GEOS-Chem uses a MODIS-derived monthly LAI product (Myneni et al., 2007) that is mapped to the GEOS-Chem grid (year-specific or a climatology), and linearly interpolated to daily values.

2.3 Modifications to land–atmosphere exchange in GEOS-Chem

Here we document the development of a land use module to describe land–atmosphere exchange in GEOS-Chem and to facilitate simulations involving changes in land cover and land use, such as the tree mortality being explored here.

To increase the flexibility in the BVOC emissions, basal emission factors are now mapped at simulation initialization using input land cover data. As a base input, we use present-day (year 2000) land cover from the Community Land Model (CLM) v. 4 (http://www.cgd.ucar.edu/tss/clm/ and Lawrence et al., 2011). Vegetation is divided into 16 plant functional types (PFTs, Table A1 in the Appendix) and their fractional coverage is mapped globally at a native resolution of 0.23° × 0.3125°. We incorporate updated emission factors following MEGAN v2.1 (Guenther et al., 2012).

We also eliminate the dependence of the dry deposition velocities on the Olson Land Map. Instead, the same PFTs that drive BVOC emissions are mapped directly to the 11 deposition types from Wesely (1989). We replace the roughness heights provided by the assimilated meteorological product with values that are specific to the land cover or plant functional type (Table A1). Furthermore, rather than basing dry deposition on the dominant land type at a certain native resolution, the complete sub-grid fractional coverage of all PFT/land types are accounted for. In this way, deposition in the model should be less dependent of the horizontal resolution of the simulation or land cover data set. For soil NO₃ emissions, we map the same set of PFTs to the 24 biomes of Steinkamp and Lawrence (2011) based on plant type and latitude (Fig. A1).

To achieve consistency between our land type description and the LAI used in the model, we replace the monthly MODIS-derived gridded LAI with the sub-grid PFT-specific monthly LAI from the CLM4 land cover description, also based on MODIS observations as well as additional cropping data (Lawrence et al., 2011).

In this way, BVOC emissions, soil NO₃ emissions, dry deposition, and surface roughness are all newly harmonized to the same land cover input and vegetation density. These changes make it possible to alter the specified PFT distributions and/or fractional coverages, and self-consistently investigate the impact on biosphere–atmosphere exchange.

2.4 Impact of updates and land use harmonization on GEOS-Chem simulation

Our modifications to GEOS-Chem impact the emissions, deposition, and simulated concentrations compared to the default model, demonstrating the important role of land cover on atmospheric chemistry. GEOS-Chem and other chemical transport models have previously shown a large sensitivity to land cover data sets (Li et al., 2013) and biogenic emission models (Fiore, 2005; Kim et al., 2014; Zare et al., 2012). Globally, we find annual emissions of isoprene decrease by 14 % from 531 to 459 Tg yr⁻¹ with land use harmonization and updated emission factors. The emissions of some monoterpenes decrease (e.g., β-pinene, limonene, sabineone, and carene by 10 % or less; ocimene by 36 %), while others increase significantly (α-pinene by 64 %, myrcene by 145 %). Sesquiterpene emissions increase between 20 and 60 % depending on the species. These changes result from the new maps of PFTs, the updated emission factors from MEGAN v2.1 (Guenther et al., 2012), and the new LAI values used. Our modified global emissions are generally consistent with those for MEGAN v2.1 as formulated by Guenther et al. (2012). For example, our α-pinene emissions increase from 40 to 66 Tg yr⁻¹, compared to 66 Tg yr⁻¹ estimated by Guenther et al. (2012). Global soil NO₃ emissions, which depend on biome mapping from the PFT data set and LAI, decrease by 2 % compared to the default model (from 9.8 to 9.6 Tg yr⁻¹).

Figure 1 shows how all of the modifications impact predicted global monthly mean O₃ concentrations for August 2010. The spatial agreement between the simulations is very high (r = 0.99), suggesting that our modifications have not resulted in significant changes to predicted O₃. While the changes that we made to the model were not in principle intended to improve the accuracy of the GEOS-Chem O₃ simulation (rather the priority was to more easily enable land-cover change experiments), the updated land cover data and the new consistency in the descriptions modestly improve the spatial correlation (from r = 0.54 in the default model to r = 0.56 in the modified model) between the simulated and gridded monthly mean O₃ observed over North America, Europe, and other locations worldwide (Evans and Sofen, 2015) for the whole year. The modifications tend to decrease the high O₃ concentrations at midlatitudes of the Northern and Southern hemispheres. In particular, the high summer bias in monthly mean O₃ drops by 0.5–0.9 ppb (e.g., from RMSE =15.6–14.8 in August) while making little difference to winter month O₃ (RMSE changed by < 0.3 ppb).
Figure 1. Simulated global surface O$_3$ concentrations for August 2010 in the (top) default, and (middle) modified GEOS-Chem configuration. (Bottom) Difference between the modified and default simulations.

3 Predicted tree mortality in the United States

To simulate national-scale tree mortality across the US, we use projected tree mortality rates from the 2012 National Insect and Disease Risk Forest Risk (NIDR) Assessment for 2013–2027, assembled by the Forest Health Technology Enterprise Team of the United States Department of Agriculture Forest Service (Krist et al., 2014). This assessment includes results from 186 individual insect and disease hazard models. We gridded the 240 m spatially resolved total tree mortality data (http://www.fs.fed.us/foresthealth/technology/nidrm.shtml) to the native resolution of the new GEOS-Chem land input file ($0.23^\circ \times 0.31^\circ$) and focused on the conterminous United States. We use these data to contrast atmospheric chemistry before vs. after the change in tree cover.

Figure 2 shows the default fractional area covered by the sum of all tree PFT categories, and the resulting loss in tree-covered fractions due to projected mortality after applying the fractional loss from the NIDR. We applied mortality losses predicted by the NIDR to all tree species in a particular input grid box, instead of accounting for losses specific to one plant functional type only. The magnitude and spatial distribution of tree loss is qualitatively consistent with the agent- and species-specific summaries in the NIDR assessment (Krist et al., 2014), since certain PFT categories usually dominate in specific regions or grid boxes. We briefly summarize the major agents driving projected mortality in the NIDR assessment here. In the western US, insects causing evergreen mortality include the mountain, western, and Jeffrey pine beetles, spruce and Douglass fir beetles, the Douglass fir tussock moth, and the Western spruce budworm. In the east, insect-driven evergreen mortality is driven by the Eastern spruce and Jack pine budworm and hemlock woolly adelgid in the north, and the southern pine beetle in the south. Engraver beetles and the balsam woolly adelgid affect evergreens in both the west and east. Deciduous tree mortality is large in the northeast and eastern US, where oak and maple decline is high. Deciduous tree mortality by diseases such as beech bark, oak wilt, and Dutch elm is also large. Aspen and cottonwood declines are significant in the western US and Great Plains. While root diseases, which impact both needleleaf and broadleaf tree categories, represent the largest single agent-level hazard, the impact of all bark beetles together are projected to cause the highest basal area losses (Krist et al., 2014).

4 Impact of tree mortality on atmospheric chemistry in the US

We perform four simulations (Table 1) to investigate the role of insect- and disease-driven tree mortality on atmospheric chemistry: (1) a base scenario in which the vegetation is not altered; (2) a scenario where the BVOC emissions respond to the scaled tree cover, but where soil NO$_x$ and dry deposition are calculated using the land cover in the base scenario; (3) a scenario where the BVOC and soil NO$_x$ emissions respond to the scaled tree cover, but where dry deposition is calculated using the land cover in the base scenario; and (4) a full tree mortality scenario where the BVOC emissions, soil NO$_x$ emissions, and dry deposition are all calculated using the scaled tree cover. The combination of these simulations allows us to decouple the effects of changing BVOC and soil NO$_x$ emissions from the effects of changing deposition. We focus our analysis on June–August since this is the seasonal peak in impacts of changes in biogenic emissions on O$_3$ and SOA formation across the United States.

4.1 Impacts on biogenic emissions and on deposition velocity

Figure 3 shows the simulated emissions of isoprene, total monoterpenes, and total sesquiterpenes, and the change in emissions due to tree mortality. The impact to total emissions across the US is a 6–7% decrease for isoprene, monoterpenes, and sesquiterpenes, with much larger impacts locally. Over the continental US, isoprene emissions are projected to decrease by more than 5% for more than 25% of the model grid boxes (762 out of a total of 2693). The highest relative impact to isoprene emissions occurs in the Rocky Mountain forests of the northwestern US, where mortality is pro-
Figure 2. Fraction of grid box covered by trees in present day (left), and the loss in tree cover due to predicted mortality from 2013 to 2027 based on the National Insect and Disease Risk Map (right). (a, b) Total tree cover; (c, d) needleleaf tree cover only; (e, f) broadleaf tree cover only.

Table 1. Descriptions of the individual simulations used here to test the impacts of large-scale tree mortality.

<table>
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<tr>
<th>Simulation</th>
<th>Description</th>
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<tr>
<td>1</td>
<td>Base land cover simulation (no tree mortality)</td>
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<tr>
<td>2</td>
<td>Tree mortality-driven BVOC emissions (soil NO\textsubscript{x} and dry deposition using base land cover)</td>
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<tr>
<td>3</td>
<td>Tree mortality-driven BVOC and soil NO\textsubscript{x} emissions (dry deposition using base land cover)</td>
</tr>
<tr>
<td>4</td>
<td>Tree mortality-driven emissions and dry deposition.</td>
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jected to be high. For example, the largest relative decrease occurs in Idaho [46.0° N, 115.3° W] where isoprene emissions decrease by 47% (1.8 µmol m\textsuperscript{-2} h\textsuperscript{-1}), compared to the base simulation. These pine-, spruce-, and fir-dominated forests of the northwest are relatively low isoprene emitters compared to the deciduous forests of the eastern US. The reduction in mean OH reactivity due to tree mortality-induced isoprene changes in the northwest is \sim 0.2–0.5 s\textsuperscript{-1} at most. In the oak-dominated Ozarks of Arkansas and Missouri [\sim 36° N, 92° W], and the central Appalachian region [\sim 38° N, 81° W], baseline isoprene emissions are an order of magnitude higher; the corresponding reduction in mean OH reactivity due to tree mortality-induced isoprene changes exceeds 3 s\textsuperscript{-1}. The highest absolute impact of mortality on isoprene emissions occurs at the border of West Virginia and Virginia [38.0° N, 80.0° W], where emissions decrease by 8.6 µmol m\textsuperscript{-2} h\textsuperscript{-1} (relative decrease of 26%).

Likewise, the highest relative impacts to total monoter-
more and total sesquiterpene emissions occur in the Rocky
Mountain forests of the western and northwestern US
(the largest relative decrease occurs in Colorado [38° N, 
106.7° W] where the monoterpene and sesquiterpene emis-
sions decrease by 48–50%). Significant relative impacts
are also predicted in the pine forests of the Sierra Nevada
(10–20%). In terms of absolute magnitude, the difference
in monoterpene and sesquiterpene emissions is largest in
pine-dominated forests of the southern US. The highest
absolute impacts in the country occur in eastern Texas
[31.0° N, 94.7° W] where the monoterpene emissions de-
crease by 1.4 µmol m\textsuperscript{-2} h\textsuperscript{-1} (or 22%), and in Arkansas.
Figure 3. Mean JJA (June-July-August) biogenic VOC emissions in the base scenario (left), and the change in emissions resulting from predicted tree mortality (right). (a, b) Isoprene emissions; (c, d) total monoterpane emissions; (e, f) total sesquiterpene emissions.

Figure 4. Mean JJA soil NO\textsubscript{x} emissions in the base scenario for (left), and the change in emissions resulting from predicted tree mortality (right).

In some of these areas (the largest relative difference occurs in western Montana [46° N, 115.3° W] where soil NO\textsubscript{x} emissions increase by 15%). In projecting changes to soil NO\textsubscript{x} emissions, we allow the canopy reduction factor to respond to changes in LAI, but we assume that the tree mortality does not impact the basal soil NO\textsubscript{x} emission factors (nor soil temperature or moisture). The increase in net emission therefore arises from a decrease in canopy reduction factor only, representing the impact of less NO\textsubscript{2} uptake by the canopy before export into the lower mixed layer. A better understanding of the canopy reduction factor, and accounting for canopy chemistry, would facilitate a more thorough assessment of these projected increases in soil NO\textsubscript{x} emissions.

Figure 5 shows the dry deposition velocity of O\textsubscript{3} in the baseline scenario, and the simulated change resulting from tree mortality. In the northeast, where deciduous forests dominate and vegetation is dense, O\textsubscript{3} deposition velocities are highest (0.6–0.7 cm s\textsuperscript{-1}), whereas the deposition velocity over needleleaf forests is lower (0.3–0.4 cm s\textsuperscript{-1}). Lowest deposition velocities occur over the arid and sparsely vegetated regions of the country. Where projected tree mortality is high, O\textsubscript{3} deposition velocity decreases by up to 0.08 cm s\textsuperscript{-1} due to reduced stomatal uptake and change in roughness height. The highest absolute impact occurs in the eastern US, along the border of Virginia and West Virginia [38.0° N, 80.0° W]. On a relative basis the impact is largest in the northwest (deposition velocity in northern Idaho [47.5° N, 116.0° W] decreases...
by 16 %, or 0.06 cm s⁻¹). Spatially, the impact on the deposition velocity for other constituents is similar. For example, the deposition velocity of HNO₃ (which is largely limited by aerodynamic resistance instead of surface resistance as in the case for O₃) also decreases in the same regions due to the change in roughness heights in the tree mortality scenario. In this case, decreases in HNO₃ deposition velocity exceeding 20 % are predicted in the northwest and eastern US.

4.2 Impacts on surface ozone concentrations

Figure 6a shows the June-July-August mean surface O₃ concentrations simulated in the base scenario (Simulation 1). The high concentrations in the western US are consistent with previous work and are a consequence of the elevation and the dry climate resulting in a deep boundary layer and slow deposition velocities (Fiore et al., 2002; Wu et al., 2008). High concentrations are also simulated in the eastern US. Figure 6b shows the change in simulated O₃ concentrations as a result of changes in BVOC and soil NOₓ emissions in a tree mortality scenario where deposition is calculated using baseline land cover (Simulation 3 – Simulation 1). Changes in soil NOₓ emissions alone increase O₃ slightly (Simulation 3 – Simulation 2), but this response is an order of magnitude smaller (or less) than the response to decreased BVOC emissions. The result is a net decrease in O₃ on the order of 0.2–0.4 ppb across a large area of the eastern US and in parts of the northwest and California. The largest change occurs in eastern Texas [32.5° N, 94.7° W] where mean O₃ decreases by 0.44 ppb. Concentrations increase slightly over the Ozarks of Arkansas and Missouri [≈ 36° N, 92° W] and the Appalachian region in West Virginia [≈ 38° N, 81° W].

Figure 6c shows the simulated change in surface O₃ due to tree mortality including the impact of changes to dry deposition (Simulation 4 – Simulation 1). The increase in concentrations due to slower deposition velocities counteracts the decrease in O₃ concentrations that result from changes in BVOC emissions alone. In some regions these influences are predicted to be roughly equal so that the net change in O₃ is close to zero. However, in many parts of the country including the northeast (e.g., Vermont, New Hampshire, and Maine), and the northwest (northern Idaho and western Montana), the predicted change in deposition is large compared to the change from emissions alone, resulting in net increases to O₃ approaching 1 ppb or greater. Over the central Appalachian region (most notably West Virginia) and Ozarks the predicted change including dry deposition is also very large compared to the small increase from emissions alone. The highest increase in O₃ occurs at the tristate intersection of Kentucky, West Virginia and Virginia [37.5° N, 82.0° W], where mean O₃ is 1.4 ppb higher than in the base simulation. The substantial effect of slower dry deposition underscores the importance of understanding canopy deposition and the potential impact of canopy processes on chemical losses in predictions of land cover change impacts. Given the influence of changes in dry deposition in our simulations, exploring the uncertainties in dry deposition calculations should be a priority for model development.

Since regions where the impact on tree cover is largest are heavily forested and removed from pollution sources, they tend to have relatively low NOₓ concentrations. In such situations, O₃ production is expected to be NOₓ-limited so that decreases in VOC emissions weakly impact O₃ formation. This is the case over the central Appalachian and Ozarks regions, where NOₓ concentrations are below 1 ppb and BVOC emissions decreased by 10–20 %, but where O₃ is minimally impacted in the scenario with altered emissions only (Fig. 6b). In these forest environments, the change to dry deposition velocity will be the dominant mechanism impacting O₃ concentrations, and indeed we find that O₃ increases when all mechanisms are considered (Fig. 6c). On the other hand, in high-NOₓ (or polluted) regions, O₃ production can be expected to be more sensitive to changes in VOC emissions, and since these areas tend to be more developed, deposition plays a smaller role. As a result, in the scenario considering only changes in emissions we find that the predicted impact to O₃ concentrations is relatively large in the heavily populated regions along the coast of the mid-Atlantic (Fig. 6b, ≈ 40° N, 74° W).

In general, we find that the ratio of NOₓ to VOC concentrations (ppb NOₓ/ ppb C) in a grid box can explain some of the O₃ response to changes in tree cover across the US, despite varying degrees of predicted land cover change. Figure 7 shows histograms of the change in surface O₃ concentrations for two populations of grid boxes that had changes in
isoprene emissions of at least 0.1 µmol m$^{-2}$ h$^{-1}$ ($N = 1115 \text{ grid boxes from a total of } 2693 \text{ grid boxes in the continental US}$). These two distributions ($N = 111$ in both) are grid boxes with the lowest 10% NO$_x$ : VOC concentrations in the base scenario, and grid boxes with the highest 10% NO$_x$ : VOC concentrations in the base scenario. These distributions are statistically different ($p < 0.001$, Wilcoxon rank-sum test), and represent the general pattern of impact on “clean” and “polluted” regions respectively. The top panel displays results based on the scenario where only biogenic emissions change (Simulation 3 – Simulation 1). Grid boxes with the highest NO$_x$ to VOC ratios tend towards stronger changes in O$_3$ concentrations than the grid boxes with lowest NO$_x$ to VOC ratios. This suggests more generally that in addition to the extent of land cover change, the impacts of tree mortality on O$_3$ can depend on whether the conditions are NO$_x$-limited (low NO$_x$ : VOC) or VOC-limited (high NO$_x$ : VOC). This NO$_x$-dependence of the regional chemistry impacts resulting from land system changes has also been identified by Wiedinmyer et al. (2006) and Hardacre et al. (2013) for example. The bottom panel displays the results based on the scenario where changes to biogenic emissions and dry deposition are accounted for (Simulation 4 – Simulation 1). Here we find that the change in O$_3$ is more frequently positive (increasing O$_3$ compared to the base scenario) in the low NO$_x$ : VOC grid boxes, since the deposition response tends to be large compared to the impact of emissions. In contrast, while slower deposition counteracts some of the decrease in O$_3$ concentrations in the more polluted grid boxes, the net impact remains largely negative (decreasing O$_3$ compared to the base scenario).

The changes in monthly mean ozone mask even larger impacts on shorter timescales (hours) that may be of importance to air quality standards. The magnitude of the impact on surface O$_3$ in the scenario that considered changes to both emissions and deposition is highest during the day and less significant at night due to the diurnal pattern of chemical O$_3$ production and biogenic emissions, and to the strong dependence of modeled deposition velocities on time of day. As a result, the number of days with O$_3$ above a specific threshold changes in many locations depending on the land cover scenario. We consider for example daily maximum 8 h averages. The EPA has recently revised the O$_3$ air quality standard to be based on 8 h averages exceeding a threshold of 70 ppb instead of the previous 75 ppb (http://www3.epa.gov/ttn/naaqs/standards/ozone/s_o3_index.html), so we investigate the number of days during June-July-August in each grid box of the US where the 8 h average O$_3$ exceeds 70 ppb. In the scenario considering only a change in emissions (Simulation 3 – Simulation 1), the number of days exceeding an 8 h O$_3$ concentration of 70 ppb decreases in 16% of the grid boxes in the lowest NO$_x$ : VOC
decile (“clean” regions of the US), and in 45 % of the grid boxes in the highest NO$_x$: VOC decile (“polluted” regions of the US). Across the US, the number of exceedances decreases by 4 or more days in several regions such as central South Carolina (34.0° N, 81.3° W), central Kentucky (37.5° N, −86.0° W), central Indiana (38.5° N, −90.7° W), northern Ohio (41.5° N, 83.3° W), and southwest Michigan (42.0° N, 71.3° W). In the scenario considering both the change in biogenic emissions and also the change to deposition rates (Simulation 4 – Simulation 1), many grid boxes experience a net increase in the number of days exceeding an 8 h O$_3$ concentration of 70 ppb. The increase impacts clean regions disproportionately (30 % of lowest NO$_x$: VOC grid boxes) compared to polluted regions (5 % of high NO$_x$: VOC grid boxes). The largest increase is 4 days, which occurs north of Richmond, VA (38.0° N, 77.3° W). In the same scenario, less than 1 % of the low NO$_x$: VOC grid boxes experiences a decrease in the number of days exceeding an 8 h O$_3$ concentration of 70 ppb, compared to 26 % of the high NO$_x$: VOC grid boxes.

4.3 Impacts on reactive nitrogen oxide compounds

Figure 8 shows the mixing ratios of reactive nitrogen oxides in the base scenario (Simulation 1), and the simulated changes resulting from tree mortality (Simulation 4 – Simulation 1) on a relative scale (% change). We plot the results for the sum of all reactive nitrogen oxides (NO$_y$, Fig. 8a), in addition to the individual contributions from NO$_x$ (Fig. 8c), HNO$_3$ (Fig. 8e) and the sum of all alkyl-, peroxy-, and acylperoxy-nitrates (or “organic nitrates”, Fig. 8g). We find that the relative impacts on NO$_y$ and its partitioning as a result of the tree mortality could be locally significant, and are a complex result of all three mechanisms (changes in BVOC emissions, changes in soil NO$_x$, and changes to the deposition velocities), depending on the chemical species. Total NO$_y$ increases by up to 8 % in the northwest (the largest relative increase of 120 ppt is along the Idaho-Montana border [47.5° N, 115.3° W]). The increases here consist of roughly equal increases in NO$_x$ (79 ppt) and HNO$_3$ (66 ppt) mixing ratios with a smaller decrease in organic nitrates (29 ppt). Over the rest of the country, changes in total NO$_y$ are small, in part because the increases in NO$_x$ and HNO$_3$ are counteracted by decreases in organic nitrate species. Significant changes in NO$_x$ abundance and NO$_y$ partitioning could alter the transport and removal of O$_3$ precursors, and alter the peroxy radical chemistry involved in O$_3$ production.

We find that the increases in NO$_x$ are largely a result of elevated soil NO$_x$ emissions (Simulation 3 – Simulation 1). On the other hand, the increases in HNO$_3$, which are up to 18 %
on a relative scale, are due to both slower deposition and increasing soil NO$_x$ emissions (Simulation 4 – Simulation 1). Small increases in HNO$_3$ (locally up to 3–4%) are also observed in the BVOC emissions only scenario (Simulation 2 – Simulation 1). Broad decreases in the organic nitrate concentrations (approaching 10%) are found across large parts of the country. This result is nearly entirely due to the reduction in BVOC emissions alone, with only a small counteracting effect of lower deposition velocities. For example, where the relative impact was largest (a 10% decrease near Missoula MT [47° N, 114.7° W]), the decrease from the BVOC emissions alone is 36 ppt, while the decrease after accounting for dry deposition and soil NO$_x$ emissions is 29 ppt.

4.4 Impacts on organic aerosol

Figure 9a shows the predicted biogenic SOA (BSOA) surface mass concentrations in the base simulation (Simulation 1). The dominant contributors to BSOA over the United States in these simulations are terpenes, consistent with the results of Pye et al. (2010). This results from nitrate radical oxidation included in the SOA mechanism, since terpenes are emitted at night (in addition to during the day) and model aerosol yields from nitrate oxidation are relatively high. The baseline simulation predicts BSOA greater than 3 µg m$^{-3}$ throughout most of the southeast US, approaching 10 µg m$^{-3}$ near the Mississippi–Alabama and Missouri–Arkansas borders. Biogenic SOA contributes 80% or more of the modeled total OA mass concentration in this region and season. In parts of the northeast and on the west coast, BSOA can also exceed 3 µg m$^{-3}$ and the model predicts the biogenic contribution to total organic aerosol to exceed 50% there. In the northwest, BSOA approaches 1–2 µg m$^{-3}$.

Figure 9b shows the change in BSOA predicted due to tree mortality (Simulation 4 – Simulation 1). In contrast to O$_3$ and NO$_x$ species (where the relative importance of deposition and chemical production could vary), the simulation predicts consistent decreases in BSOA from the tree mortality scenario as a result of decreasing BVOC emissions. The change in atmospheric lifetime as a result of slower dry deposition is negligible (Simulation 4 – Simulation 3). Across the eastern US, BSOA decreases by 5–10%. The relative impacts are highest where terpene emissions are significant and projected tree mortality is high due to the dominance of terpenes as precursors to BSOA in these simulations. The impact on BSOA due to tree mortality generally exceeds 10% where the contributions of terpene emissions represent 50% or more of total BVOC emissions (in mass carbon). The spatial pattern in ΔBSOA corresponds most to the relative contribution of the lumped MTTP category of terpenes (α-pinene + β-pinene + sabinene + carene). In some parts of the southeast the change exceeds 25% (1–2 µg m$^{-3}$ in terms of absolute mass). The largest absolute impact occurs in southern Arkansas [33.5° N, 92.7° W], where BSOA decreases by 2.0 µg m$^{-3}$ (or 20%). The relative impact is also high in the northwest, where BSOA decreases by 0.5–1 µg m$^{-3}$ (the highest relative difference of 39% occurs in northern Idaho [46.0° N, 115.3° W]).

Given the dominance of BSOA in much of the US, these changes appreciably impact total OA (and consequently total aerosol mass). Relative impacts to the sum of all organic aerosol are on the order of 20% or greater in many parts of the south, northwest, and northern California. These simulations suggest that tree mortality and the concomitant change in biogenic emissions has the potential to impact background aerosol levels by up to 2 µg m$^{-3}$ in some regions. This may be of particular relevance to the EPA Regional Haze Program, aimed at improving visibility in national parks and wilderness areas (http://www3.epa.gov/visibility/program.html).

5 Discussion

In this study we develop and apply a land use module for GEOS-Chem to demonstrate that projected tree mortality in the coming decades could impact air quality across the US. We find that the changes in BVOC emissions, soil NO$_x$ emissions, and dry deposition can impact O$_3$ mixing ratios, reactive nitrogen oxide speciation and abundance, and biogenic secondary organic aerosol formation. The magnitude of change in mean O$_3$ (−0.4 to +1.4 ppb depending on the simulation) and SOA (up to −2.0 µg m$^{-3}$) in some grid boxes is similar to regional changes predicted by examples of biofuel cropping or natural succession scenarios (Ashworth et
al., 2012; Porter et al., 2012; Drewniak et al., 2014), and comparable with the tree mortality effect from past bark beetle infestations simulated in western North America by Berg et al. (2013).

In the case of O$_3$, we find that lower deposition velocities resulting from the change in tree cover could reverse the impact of decreased chemical production. This produces regional variability in the sign of the O$_3$ response depending on which effect dominates locally. Generally, our simulations predict that high levels of O$_3$ could be exacerbated in the low-NO$_x$, densely forested areas where mortality is projected to be high. This increase in O$_3$ could have further feedbacks given the documented negative effect of O$_3$ on forest health (Ashmore, 2005; Taylor et al., 1994). Using the number of days when 8 h O$_3$ exceeds 70 ppb, we find that tree mortality generally reduces the number of exceedances for high-NO$_x$ environments.

Our simulations also predict large impacts on organic aerosol. While the exact yields and SOA composition are uncertain (Hallquist et al., 2009) and depend on the SOA model used, the post-disturbance impact is a robust and direct response to a reduction in biogenic emissions (and is not sensitive to changes in deposition). Similar to the reduction in O$_3$ that favors polluted regions, the projected tree mortality could decrease background aerosol levels by up to 1–2 µg m$^{-3}$ locally, inadvertently making progress in other air quality objectives (e.g., long-term visibility at national parks and wilderness areas where mortality is in some cases projected to be high).

These results do not account for changes in anthropogenic emissions that may occur over the same period of time as the changes to vegetation. We therefore performed a subsequent test where the same land cover change was applied, using anthropogenic emissions from 2005 (instead scaling the emissions to 2010 as was performed for Simulations 1–4). Between 2005 and 2010, modeled anthropogenic emissions of NO$_x$ and SO$_2$ over the continental US decreased by 30 and 44% respectively. Despite this large perturbation in anthropogenic emissions, the predicted impacts due to the land cover change were fundamentally the same. The range of impact on simulated mean O$_3$ over the US due to both emissions and dry deposition combined (Simulation 4 - Simulation 1) went from $\Delta$O$_3 = [-0.24, +1.45]$ ppb for the 2010 emissions, to $\Delta$O$_3 = [-0.34, +1.35]$ ppb for the 2005 emissions. Likewise, the maximum impact on SOA changed very little, from $\Delta$BSOA = -2.05 µg m$^{-3}$ in the 2010 simulation, to $\Delta$BSOA = -1.94 µg m$^{-3}$ in the 2005 simulation. Nevertheless, simultaneous changes in both anthropogenic and biogenic emissions increase the uncertainty in the exact magnitude of projected changes in secondary pollutants.

Many opportunities exist for development and incorporating further complexity. For example, these simulations have not accounted for the temporal dynamics of forests undergoing disturbances from insect attack and disease. In the case of insect infestation, VOC emissions can be enhanced during the attack (Amin et al., 2012), and Berg et al. (2013) found that the spatiotemporal patterns in tree mortality can greatly affect the relative impacts of the attack effect vs. the mortality effect on BVOC emissions. Numerous compounds have been observed to be emitted by trees when under stress (Faiola et al., 2015; Joutsensaari et al., 2015) that GEOS-Chem does not yet represent. Not only have we compared simple “pre-” and “post-” disturbance scenarios ignoring attack effects, but we have not considered forest succession. Extensive mortality caused by insects and disease may be compared to forest fires (Hicke et al., 2012), with growth of surviving trees and understory potentially accelerating (Brown et al., 2010). In such cases, BVOC emissions may not necessarily decrease universally, but the composition of those emissions could change over time. Forest recovery after an outbreak may be possible within decades, as has been predicted in the case of bark beetle outbreak in the western US using a forest vegetation simulator (Pfeiffer et al., 2011). Successional dynamics could for example be simulated by an individual-based model (e.g., Shuman et al., 2014), and used as inputs at various time points in the chemical transport model. We have also assumed that basal BVOC emission factors for the surviving vegetation are the same as pre-disturbance, but experiments have shown for example that monoterpane basal emission can increase significantly after forest thinning (Schade and Goldstein, 2003), which may or may not be a temporary effect.

Improvements in the parameterization of O$_3$ deposition should also be explored. While we find changes in dry deposition velocity to be an important (and in the majority of cases overriding) factor in our simulation of O$_3$ change, other hypothetical simulations where European crop- and grasslands were converted to poplar plantations for biomass production found that changes from altered dry deposition velocity were an order of magnitude lower than the change in biogenic emissions (Beltman et al., 2013). Dry deposition rates can depend strongly on the choice of model (Hardacre et al., 2015; Park et al., 2014; Wu et al., 2011), making predictions that depend on this uncertain. Improvements can be expected by more accurate representations of land cover (and subsequent changes) (Hardacre et al., 2015), or by including a more process-based model of deposition that depends on soil moisture and vapor deficit (Büker et al., 2012; Pleim et al., 2001). There is also evidence that a significant fraction of the O$_3$ uptake observed over forest canopies is actually an unaccounted for chemical sink (Kurupius and Goldstein, 2003; Rannik et al., 2012; Schade and Goldstein, 2003; Wolfe et al., 2011), but changes in this above-canopy chemistry are not captured in our current set of simulations.

Likewise, canopy chemistry and stand development post-disturbance will affect the predicted impacts on soil NO$_x$ emissions. The impacts of canopy uptake and canopy chemistry resulting from changes in vegetation density and composition could be explored in more detail with future work using a 1-D forest canopy-chemistry model (e.g., Wolfe,
2011; Ashworth et al., 2015) for the regions where we project large impacts. We have assumed that the basal emissions from the soil after the disturbance will be the same as those prior to the disturbance, but large-scale tree mortality and forest succession have the potential to alter soil biogeochemistry (Gao et al., 2015; Norton et al., 2015; Trahan et al., 2015).

We anticipate the impacts of tree mortality that are simulated here to be conservative. Future climate change is not included in the NIDR assessment, but is expected to increase the risk of mortality from several pests (Krist et al., 2014). Likewise, insect attack could make some tree species more sensitive to climate stresses, resulting in mortality despite what might have been an otherwise non-lethal insect attack (Anderegg et al., 2015). Predictions over the timescale of years and decades will depend on how the insect and/or disease disturbances interact with other abiotic environmental disturbances (e.g., drought, extreme heat), but these interactions are rarely fully coupled (Anderegg et al., 2015). Furthermore, tree mortality from many other factors outside of pests and pathogens are not considered (e.g., competition from invasive exotic plants, drought, or other disturbances). As a result, the actual tree loss in the coming decades, and the concomitant impacts on atmospheric chemistry, may be higher than simulated here. We have also ignored any potential feedback between tree mortality and fire incidence or severity, which is not well understood (Bond et al., 2009). Increases in wildfire (and associated emissions) due to climate change have been predicted to have important consequences for ozone air quality (Yue et al., 2015). Finally, our simulations only explored tree mortality across the United States. No similar large-scale projection of mortality risk exists for Canada, despite insect outbreak already being the dominant cause of tree mortality in boreal forests of eastern Canada (Zhang et al., 2014), and severe (although decreasing) mountain pine beetle infestations in western Canada (Buston and Maclachlan, 2014). Increasing insect outbreaks are also a potential threat to forests elsewhere in the world (Lindner et al., 2010). We note that our simulations neglect any potential human intervention in response to these risks.

6 Conclusions

Land use and land cover change is expected to be a major driver of global change that remains difficult to constrain. The change in vegetation that we have explored in these simulations represents one of a myriad of changes that are occurring (and are projected to occur) to the Earth’s land surface. We anticipate that these GEOS-Chem model developments will enable investigation of a wide range of land cover and land use change impacts (e.g., vegetation succession, deforestation or afforestation, and crop conversions). Properly representing changes in land cover by including accurate and timely updates to chemical transport models will be an important part of simulating global change. By linking all terrestrial biosphere exchange to plant functional type, our GEOS-Chem developments bring the model a step closer to eventual coupling with dynamic vegetation and/or Earth system models.

Our results add to the literature demonstrating that changes to vegetation can have significant impacts on local chemistry due to changes in biosphere-atmosphere fluxes of reactive trace species, with consequences for controlling regional air quality. Given the general tightening of air quality standards to improve the health of global populations, understanding how changes in land cover will aid or abet these achievements could become increasingly important.
Table A1. Mapping of CLM-input land types used in the modified version of GEOS-Chem to the Wesely deposition surfaces for deposition, and the associated roughness ($Zo$) heights for each.

<table>
<thead>
<tr>
<th>Land type</th>
<th>Wesely surface</th>
<th>$Zo$ (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lake/Ocean</td>
<td>Water</td>
<td>0.001</td>
</tr>
<tr>
<td>Bare Ground</td>
<td>Desert</td>
<td>0.001</td>
</tr>
<tr>
<td>NET Temp</td>
<td>Coniferous forest</td>
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</tr>
<tr>
<td>NET Boreal</td>
<td>Coniferous forest</td>
<td>1</td>
</tr>
<tr>
<td>NDT Boreal</td>
<td>Coniferous forest</td>
<td>1</td>
</tr>
<tr>
<td>BET Temp</td>
<td>Amazon Rainforest</td>
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</tr>
<tr>
<td>BET Temp</td>
<td>Deciduous forest</td>
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<tr>
<td>BDT Temp</td>
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<tr>
<td>BDT Temp</td>
<td>Deciduous forest</td>
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<tr>
<td>BDT Boreal</td>
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<td>1</td>
</tr>
<tr>
<td>BES Temp</td>
<td>Shrub/grassland</td>
<td>0.01</td>
</tr>
<tr>
<td>BDS Temp</td>
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<td>BDS Boreal</td>
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<tr>
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<td>0.002</td>
</tr>
<tr>
<td>C3 Other GR</td>
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<tr>
<td>Wetland</td>
<td>Wetland</td>
<td>0.05</td>
</tr>
</tbody>
</table>

Appendix A: Land cover classification system

Table A1 lists the land and plant functional types in the CLM4 land cover description which we use as a base land cover input for our simulations. The table also shows how we have mapped these land cover types to the original Wesely deposition surfaces and to roughness heights for the dry deposition parameterization.

Figure A1 schematically lays out how we have defined biomes in accordance with the nomenclature used for soil NO$_x$ emissions based on the CLM4 land and plant functional type coverage.

Figure A1. Mapping of native CLM land input classes to soil-NO$_x$ biomes (according to Steinkamp and Lawrence, 2011) for land cover harmonization in GEOS-Chem.
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References


