The effects of global change upon United States air quality

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Abstract. To understand more fully the effects of global changes on ambient concentrations of ozone and particulate matter with aerodynamic diameter smaller than 2.5 µm (PM2.5) in the United States (US), we conducted a comprehensive modeling effort to evaluate explicitly the effects of changes in climate, biogenic emissions, land use and global/regional anthropogenic emissions on ozone and PM2.5 concentrations and composition. Results from the ECHAM5 global climate model driven with the A1B emission scenario from the Intergovernmental Panel on Climate Change (IPCC) were downscaled using the Weather Research and Forecasting (WRF) model to provide regional meteorological fields. We developed air quality simulations using the Community Multiscale Air Quality Model (CMAQ) chemical transport model for two nested domains with 220 and 36 km horizontal grid cell resolution for a semi-hemispheric domain and a continental United States domain, respectively. The semi-hemispheric domain was used to evaluate the impact of projected global emissions changes on US air quality. WRF meteorological fields were used to calculate current (2000s) and future (2050s) biogenic emissions using the Model of Emissions of Gases and Aerosols from Nature (MEGAN). For the semi-hemispheric domain CMAQ simulations, present-day global emissions inventories were used and projected to the 2050s based on the IPCC A1B scenario. Regional anthropogenic emissions were obtained from the US Environmental Protection Agency National Emission Inventory 2002 (EPA NEI2002) and projected to the future using the MARket ALlocation (MARKAL) energy system model assuming a business as usual scenario that extends current decade emission regulations through 2050. Our results suggest that daily maximum 8 h average ozone (DM8O) concentrations will increase in a range between 2 to 12 parts per billion (ppb) across most of the continental US. The highest increase occurs in the South, Central and Midwest regions of the US due to increases in temperature, enhanced biogenic emissions and changes in land use. The model predicts an average increase of 1–6 ppb in DM8O due to projected increase in global emissions of ozone precursors. The effects of these factors are only partially offset by reductions in DM8O associated with decreasing US anthropogenic emissions. Increases in PM2.5 levels between 4 and 10 µg m–3 in the Northeast, Southeast, Midwest and South regions are mostly a result of increase in primary anthropogenic particulate matter (PM), enhanced biogenic emissions and land use changes. Changes in boundary conditions shift the composition but do not alter overall simulated PM2.5 mass concentrations.
1 Introduction

Despite extensive efforts to reduce anthropogenic emissions, air pollution continues to be a public health issue in the United States (US EPA, 2010). Elevated concentrations of pollutants in the troposphere, such as ozone (O$_3$) and particulate matter, degrade air quality and have been associated with, among other things, increasing human respiratory diseases in urban areas (WHO, 2005), and in the case of particulate matter (PM), with low birth weights across the world (Dadavand et al., 2012).

High concentrations of tropospheric O$_3$ and PM matter with aerodynamic diameter smaller than 2.5 µm (PM$_{2.5}$) are caused by a combination of adverse meteorological conditions and the atmospheric emissions of their primary precursors. While regulatory controls are expected to reduce emissions of many emitted pollutants in the United States (US) in the future, the negative effects of global climate change may offset the positive effects of such reductions. Furthermore, global emissions of greenhouse gases and other pollutant precursors are projected to increase (IPCC, 2007). Moreover, recent research has provided evidence of increasing long-range transport of O$_3$ and PM$_{2.5}$ precursors from Asia and their influence over the western US (Leelieveld and Dentener, 2009; Wuebbles et al., 2007; Cooper et al., 2010, 2012; Zhang et al., 2010; Ambrose et al., 2011; WMO, 2012; Lin et al., 2012).

In the United States, regulations and technological changes in the transportation and energy sectors are projected to reduce regional atmospheric pollutants in the future (Loughlin et al., 2011). However, the interplay between climate change, increasing global emissions, and intercontinental transport pose challenges that air quality managers will have to address in order to maintain regional air quality standards (Ravishankara et al., 2012). To provide a foundation for building effective management strategies and public policies in a changing global environment, modeling approaches that link global changes with regional air quality are required. The general approach has been to use output from general circulation models (GCMs) to drive regional climate models (RCMs) and regional or global chemical transport models (CTMs/GTMs; Giorgi and Meleux, 2007; Jacob and Winner, 2009).

This downscaling approach has been used in a variety of studies in Europe, Canada and Asia at different timescales of climate change – e.g., Liao et al., 2006 (2000 to 2100); Langner et al., 2005 (2000 to 2060); Forkel and Knoche, 2006 (2990 to 2030); Meleux et al., 2007 (1975 to 1985); Kunkel et al., 2007 (1990 to 2090); Lin et al., 2008 (2000 to 2100); Spracklen et al., 2009 (2000 to 2050); Kelly et al., 2012 (2000s to 2050s). These investigations based the global emissions on future anthropogenic emissions scenarios developed from the Intergovernmental Panel on Climate Change (IPCC) assessment reports. Despite the differences in emission scenarios, timescales, modeling frameworks and future climate realizations, increases in ozone concentrations on the order of 2 to 10 parts per billion (ppb) in polluted regions were consistently predicted from these studies as a result of climate change alone. By contrast, there is little consistency among the model predictions of climate change effects on PM (Jacob et al., 2009; Dawson et al., 2013).

In the US, a combined effort between the Environmental Protection Agency and the academic community resulted in a set of modeling studies that adopted a variety of modeling methods (Hogrefe et al., 2004; Leung and Gustafson, 2005; Liang et al., 2006; Steiner et al., 2006; Tagaris et al., 2007; Liao et al., 2006; Racherla and Adams, 2006, 2008; Tao et al., 2007; Huang et al., 2007, 2008; Nolte et al., 2008; Wu et al., 2008a, b; Chen et al., 2009b; Avise et al., 2009; Dawson et al., 2009). These US investigations based their current and future climate realizations on the results of GCMs using the various IPCC emissions scenarios (IPCC, 2007) projected to the 2050s. In some of the studies, the global climate realizations were subsequently downscaled to a higher resolution using the PSU (Pennsylvania State University)/NCAR (National Center for Atmospheric Research) Mesoscale Model version 5 (MM5; Grell et al., 1994) to horizontal resolutions that ranged from 90 to 36 km. Many of these studies based their analysis on the effects of climate change on summer air quality in the continental US (CONUS). In summary, despite the differences in modeling elements, all studies found increases in the summer average of the daily maximum 8 h average ozone concentrations over large regions of CONUS on the order of 2 to 8 ppb (Weaver et al., 2009). In contrast, PM concentrations showed changes between ±0.1 and ±1 µg m$^{-3}$, with little consistency between studies, including the sign of the differences (Day and Pandis, 2015; Trail et al., 2015; Jacob and Winner, 2009).

It is important to note that variations between modeling frameworks did result in very diverse regional patterns of key weather drivers for ozone and PM formation. Thus, while most of the studies mentioned above projected an average increase in ozone concentrations for the simulated domains, reductions or insignificant changes in certain regions of the domain were also simulated. Generally, temperature and solar radiation reaching the surface were the major meteorological drivers for regional ozone concentrations. For PM concentrations, most of the studies found a direct link between changes in precipitation and relative humidity and changes in PM concentrations (Liao et al., 2006; Unger et al., 2006; Racherla and Adams 2006; Tagaris et al., 2007; Avise et al., 2009; Chen et al., 2009b). Nevertheless, the direct impacts of changes in meteorological conditions are not the only factors of change for ozone and PM concentrations. Changes in emissions of biogenic volatile organic compounds (BVOCs), due to climate and land cover change, and the treatment of isoprene nitrates in the chemical mechanism were found to be a key factor in the regional variability of ozone and PM, particularly in areas of the southeastern US (Jacob and Winner, 2009; Weaver et al., 2009).
In this work, we present a continuation of the work described by Avise et al. (2009) and Chen et al. (2009a, b), who downscaled the Parallel Climate Model (PCM; Washington et al., 2000) and MOZART (Model for OZone And Related chemical Tracers; Horowitz, 2006) global model output for the A2 IPCC scenario using MM5 and the Community Multiscale Air Quality Model (CMAQ; Byun and Schere, 2006) to simulate current and future air quality in the US. For this update, we implemented a semi-hemispheric domain for the Weather Research and Forecasting (WRF) mesoscale meteorological model (http://www.wrf-model.org) and CMAQ simulations in lieu of using MOZART output for chemical boundary conditions for our CONUS CMAQ simulations. We used the ECHAM5 global climate model (Roecker et al., 1999, 2003) output for the A1B scenario to drive these simulations for 2 decadal periods: the current decade 1995–2004 and the future decade 2045–2054. In presenting our results, we follow the attribution approach described in Avise et al. (2009), where the separate and combined effects of changes in climate, US anthropogenic emissions, global anthropogenic emissions and biogenic emissions due to changes in regional meteorology and land use are investigated. Ideally, this framework should include feedback from changes in atmospheric chemistry to the climate system (Raes et al., 2010). However, due to the computational requirements of an online approach, we did not incorporate feedback between the atmospheric chemistry and transport simulations from the CTM to the RCM. Furthermore, despite the observed sensitivity of tropospheric ozone to regional emissions and global burden of methane (Zhang et al., 2011; Fiore et al., 2008; Wu et al., 2008a; Nolte et al., 2008), in this work, we do not address the direct effect of emissions of methane on the air quality simulations.

In Sect. 2, we provide an overview of the modeling framework and emissions scenarios. Evaluation of the model performance for the climate simulations and results of the changes in meteorological fields are also presented in Sect. 2. Assessment of air quality changes and the individual and combined effects from changes in model components are presented in Sect. 3. Finally, we present a summary of the results and conclusions in Sect. 4.

2 Methodology

2.1 General framework

Results from the global climate model ECHAM5 under the IPCC Special Report on Emissions Scenarios (SRES) A1B scenario (Nakicenovic et al., 2000) were downscaled using the WRF model separately to a semi-hemispheric (S-HEM) 220 km domain and nested CONUS domains of 108 km (not shown) and 36 km (Fig. 1). Although, it has been suggested that periods of 10 to 30 years are required to fully determine climatological conditions (Andersson and Engardt, 2010), the fact that emission inventories can substantially change from one decade to the next suggests that using 5 to 10 year periods for air quality assessment is more appropriate. Thus, five representative summers (June–July–August; with May as a spin-up period) for the present (1995 to 2004) and the future (2045 to 2054) decades were selected. Ranked in terms of their CONUS-mean maximum temperature of the year, the summers of the warmest and coldest years, as well as the second, fifth and seventh warmest years in each decade were selected for CMAQ simulations. Comparison of the meteorological conditions of these five selected summers to those of the full decades is presented in Sect. 3.1. These five representative summers for the present and future periods were processed with the meteorology–chemistry interface processor v3.4.1 (MCIP; Otte and Pleim, 2010) for the S-HEM and 36 km CONUS domains. Meteorological fields generated from MCIP for both domains were used to estimate biogenic emissions using the Model of Emissions of Gases and Aerosols from Nature v2.04 (MEGANv2.04; Guenther et al., 2006) and to calculate the temporal profiles within the Sparse Matrix Operator Kernel Emissions (SMOKE) v2.7 (http://www.smoke-model.org). With the elements described above, a framework to perform air quality simulations using the CMAQ4.7.1 (Foley et al., 2010) was created. The overall schematic for the modeling system is shown in Fig. 2.

2.2 Climate and meteorology

The regional weather model WRF includes advanced representations of land-surface dynamics and cloud microphysics to simulate complex interactions between atmospheric processes and the land surface characteristics. Detailed descriptions of WRF can be found at http://wrf-model.org and a discussion of its range of regional climate modeling applica-
Figure 2. Schematic of the modeling framework.

The MEGANv2.04 biogenic emission model (Guenther et al., 2006; Sakulyanontvittaya et al., 2008) was used to estimate current and future BVOC and soil NO\textsubscript{x} emissions based on the WRF meteorology with current and future estimates of land use and land cover. For the current decade, the default MEGANv2.04 land cover and emission factor data (Guenther et al., 2006) were used. For the future decade, cropland distributions were estimated by combining three data sets: the IMAGE 2100 global cropland extent data set, (Zuidema et al., 1994), the SAGE maximum cultivable land data set (Ramankutty et al., 2002) and the MODIS-derived current cropland data (as used in MEGANv2.04 and described in Guenther et al., 2006). The IMAGE 2100 data set was created from the output of a land cover model, which forms part of a sub-system of the IMAGE 2.0 model of global climate change (Alcamo, 1994). The SAGE cultivable data set was created using a 1992 global cropland data set (Ramankutty and Foley, 1998) modified by characterizing limitations to crop growth based on both climatic and soil properties. The future global cropland extent distribution was generated by analyzing predicted changes in agriculture on a continent-by-continent basis (using the IMAGE data). These changes were then applied to the MODIS based cropland map (used for present-day MEGAN simulations) using the SAGE maximum cultivable data set as an upper limit to cropland extent. The resulting land cover data have considerably lower cropland fraction than the original IMAGE data, which likely overestimates future cropland area by not considering whether a location is cultivable.

In addition to generating a future crop cover data set to simulate potential biogenic VOC emissions using MEGAN, future data sets representing several other MEGAN driving variables were developed. These included geo-gridded potential future plant functional type (PFT)-specific emission factor (EF) maps for isoprene and terpene compounds, as well as future-extent maps of four non-crop PFTs: broadleaf trees, needle-leaf trees, shrubs and grasses. For regions outside of the US, the non-crop PFT distributions were generated by reducing the current extent of each non-crop PFT map by an amount that would appropriately offset the predicted cropland expansion for a given continent. For the US, future non-crop PFT maps were generated using the Mapped Atmosphere–Plant–Soil System (MAPSS) model output (http://www.fs.fed.us/pnw/mdr/mapss/; Neilson, 1995), based on three GCM future scenarios. Present-day MAPSS physiognomic vegetation classes were associated with current PFT fractional coverage estimates by dividing the US into sub-regions and by averaging existing (MODIS-derived) geospatially explicit PFT data within each sub-region as a function of MAPSS class. Sub-regions were created based on Ecological Regions of North America (http://www.epa.gov/wed/pages/ecoregions.htm). After every current MAPSS class had been assigned PFT-specific fractional coverage estimates, future PFT cover was determined by reclassifying future distribution maps for the three MAPSS data sets using the fractional PFT cover estimates for each MAPSS class (within each ecological region), and averaging the three resultant future data sets into a single estimate of future cover for each PFT.

For the eastern US, future isoprene and monoterpene PFT-specific EF maps were constructed using changes in tree species composition predicted by the US Department of Agriculture “Climate Change Tree Atlas” (CCTA; http://nrs.fs.fed.us/atlas/tree/). The CCTA data are based on ecosystem changes driven by the average of three GCMs that represented the most conservative emissions scenarios available.

Using existing speciated EF data (Guenther, 2013), we applied anticipated changes in the average species composition of each PFT to generate species-weighted PFT-specific EF maps on a state-by-state basis (the CCTA data are organized by state). As data were lacking on predicted species-level changes for areas outside the eastern US, we did not attempt to alter EF maps outside the eastern US.
2.4 Anthropogenic emissions

For S-HEM domain CMAQ simulations, global emissions of ozone precursors from anthropogenic, natural and biomass burning sources were estimated for the period 1990–2000 applied to the POET emission inventory (Granier et al., 2005). Non-US anthropogenic emissions (containing 15 sectors) were projected based on national activity data and emission factors. Gridded maps (e.g., population maps) were applied to spatially distribute the emissions within a country. The global emission inventory includes emissions from fossil fuels, biofuels, open burning of biomass, and urban waste. Emissions are varied by combustion practices, which consider combinations of fuel, combustion type and emission controls, as well as their prevalence on a regional basis.

Global emissions for the year 2000 from the POET, MEGAN and Bond et al. (2004) inventories were combined, and the 16 gas-phase POET and MEGAN species, along with the OC and BC species were adapted to the Statewide Air Pollution Research Center (SAPRC)-99 (Carter, 1990, 2009) chemical mechanism. Diurnal patterns were developed and applied to the gridded emission inventories and processed using SMOKE. For the future decade hemispheric domain simulations, current decade emissions were projected to the year 2050 based on the IPCC A1B emission scenario. The percent change in emissions was summarized according to the regions and countries in the S-HEM domain that surround the CONUS domain (Fig. 3).

US anthropogenic emissions for the 36 km CONUS current decade CMAQ simulations were developed using the 2002 National Emission Inventory. The Emission Scenario Projection (ESP) methodology, version 1.0 (Loughlin et al., 2011), was applied to project future decade US anthropogenic emissions. A primary component of ESP v1.0 is the MARKet Allocation (MARKAL) energy system model (Loulou et al., 2004). MARKAL is an energy system optimization model that characterizes scenarios of the evolution of an energy system over time. In this context, the energy system extends from obtaining primary energy sources, through their transformation to useful forms, to the variety of technologies (e.g., classes of light-duty personal vehicles, heat pumps, or gas furnaces) that meet “end-use” energy demands (e.g., projected vehicle miles traveled, space heating). Within ESP 1.0, MARKAL is used to develop multiplicative factors that grow energy-related emissions from a base year to a future year. Surrogates, such as projected population growth or industrial growth, are used to develop non-energy-related growth factors. The resulting factors were applied within SMOKE to develop a future decade inventory from the 2002 National Emissions Inventory (NEI).

For the work presented here, the EPAUS9r06v1.3 database (Shay et al., 2006) was used with MARKAL to develop growth factors for CO2, NOx, SO2 and PM with aerodynamic diameter smaller than 10 µm (PM10). The PM10 growth factors were also applied to PM2.5, and the CO2 growth factors were used as a surrogate for energy system CO, NH3, VOC, HCl and chlorine. For mobile sources, NOx growth factors were used for CO, VOC and NH3. Non-combustion industrial emission growth factors were developed from projections of economic growth. Growth factors for non-combustion emissions from the residential and commercial sectors are linked to population growth. The resulting energy and non-energy factors were then used within SMOKE to multiply emissions from the 2002 NEI to 2050.

EPAUS9r06v1.3 originally was calibrated to mimic the fuel use projections of the US Energy Information Administration’s 2006 Annual Energy Outlook (AEO06; US DOE, 2008). Energy demands were adjusted to account for population growth consistent with the A1B storyline. The results reflect business as usual assumptions about future environmental and energy regulations as of 2006. Thus, while electric sector emissions are capped to capture the effects of the Clean Air Interstate Rule (CAIR; US EPA, 2005), the impacts of increases in natural gas availability, the 2007 economic downturn, and the relatively new 54.5 Corporate Average Vehicle Efficiency (CAFE) standard (US CFR, 2011) are not reflected. More recent versions of the MARKAL database reflect these factors with expanded pollutant growth coverage and refined emission factors (US EPA, 2013). The ESP v1.0, including the MARKAL database EPAUS9r06v1.3, was selected here to maintain compatibility with previous and ongoing activities.

The differences between the base year and future-year US inventories were summarized at the pollutant and regional level (Fig. 4). Using the ESP v1.0 methodology, emissions of...
NO\textsubscript{x} and SO\textsubscript{2} are projected to decrease between 16\% in the South and Southwest to 35\% in the Northeast and Northwest. On the other hand, emissions of pollutants that were not captured endogenously by MARKAL, such as carbon monoxide (CO), non-methane volatile organic compounds (NMVOCs) and ammonia (NH\textsubscript{3}) are projected to increase in nearly all regions across the CONUS domain. The use of surrogates for growth factors as described above means the projected changes in CO and VOC emissions are likely too high. The largest increase (70\%) in emissions of CO is projected in the Midwest; this is co-located with an increase of about 20\% of NMVOC. The smallest increase of CO is projected for the South; however, the same region was projected to increase NMVOC by about 12\%. The largest increase in PM\textsubscript{2.5} emissions is projected in the Northwest (< 20\%) and the smallest increase (3\%) of PM\textsubscript{2.5} is projected in the Central region, which also has a projected 34\% increase in NMVOC.

2.5 Air quality simulations

The CMAQ model version 4.7.1 was employed to simulate the potential impact of climate change on surface ozone and PM\textsubscript{2.5} over the CONUS at 36 km horizontal grid spacing and covering 18 vertical layers from the surface up to 100 mb. The model configuration included the use of the SAPRC-99 chemical mechanism and version 5 of the aerosol module, with secondary organic aerosol (SOA) parameters of Carlton et al. (2010). Methane concentration is fixed at 1.85 parts per million (ppm) for all CMAQ simulations. Stratospheric intrusion (STE) was not included in the CMAQ simulations; high STE events are mostly relevant during the spring season; thus, lack of STE in our summer simulations is not expected to have a significant effect in our results.

Using the framework components described above, a matrix of CMAQ simulations that included changes in predicted meteorological conditions and potential emission scenarios was constructed (Table 1). For each set of simulations shown in Table 1, five representative summers were modeled. Simulation 0 represents the base case simulation, where all model inputs are set to current decade conditions. Simulation 1 is used to investigate the impact of climate change alone: all model inputs are set to current decade conditions except for meteorology (biogenic emissions are not allowed to change with the future climate for this case). Simulation 2 is the same as Simulation 1, except that biogenic emissions are allowed to change with the future climate, and in Simulation 3, future land use is also incorporated into the biogenic emission estimates. Simulation 4 is used to investigate the impact of future decade US anthropogenic emissions, where all inputs are set to current decade levels except for US anthropogenic emissions. The impact of future global emissions is investigated in Simulation 5. Finally, Simulation 6 represents the combined impacts of Simulations 1–5.

2.6 Evaluation of model performance

To aid in summarizing model results, the 36 km domain was divided geographically into seven regions (Fig. 4, lower right). Since the WRF simulations used to drive CMAQ are based on a climate realization rather than reanalysis data, a direct comparison between the modeled output and observations cannot be made. Instead, the frequency distributions of simulated and observed values are compared. For the simulated meteorological fields, daily maximum temperature and
Table 1. List of simulations to assess the effect of global climate changes upon air quality in the United States.

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<th>Climate</th>
<th>Biogenic emissions</th>
<th>Anthropogenic emissions</th>
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<td>climate</td>
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<td>Current</td>
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<td>1</td>
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<td>6</td>
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Figure 6. Shown are the 2nd, 25th, 50th, 75th, 98th percentiles of observed vs. modeled values of DM8O for each region.

Figure 7. Shown are the 2nd, 25th, 50th, 75th, 98th percentiles of observed vs. modeled values of 24 h average PM$_{2.5}$ for each region.

daily precipitation are compared against a decade of summer observations (1995–2004) from the United States Historical Climatological Network (US-HCN; http://cdiac.ornl.gov/ftp/ushcn_daily/; Karl et al., 1990) in Fig. 5. The model distributions of temperature and precipitation agree reasonably well with the observations, and provide a good representation of the regional variability of precipitation and temperature. Except for the Northwest and Southwest regions, the observed mean and maximum temperatures are over predicted, with the largest overprediction in the Midwest. For all analyzed regions the model successfully simulates the seasonal trend of summer temperatures, showing the observed increase in mean temperature from June to July and subsequent decrease in mean temperature from July to August (not shown).

The modeled daily maximum 8 h ozone concentrations (DM8O) from the five representative summers (Fig. 6) from the current decade CMAQ simulations (Simulation 0) were compared to the range of observations from the AIRNow network (http://airnow.gov/). As seen in Fig. 6, DM8O is overestimated in regions where temperature maxima are also over predicted, most noticeably in the Midwest, the South and the Southeast but also in the Northeast. Except for the less populated Central region, DM8O shows a bias that ranges between +10 ppb (+15 %) and +25 ppb (+37 %) across the domain. This is consistent with previous climate downscaled results by Tagaris et al. (2007), who found a bias of +15 %, and with Avise et al. (2009), who found regional biases as high as +39 %. Despite the bias, results from the modeling framework presented here have been shown to accurately represent the correlation between ozone and temperature at rural Clean Air Status and Trends Network (CASTNET) sites throughout the US (Avise et al., 2012), suggesting that the bias in temperature is the main cause of the bias in DM8O. This implies that the CTM is responding to the meteorological driver of ozone production and thus can predict the impact of climate change on DM8O.

Simulations for the current decade show a domain-average DM8O of 66 ± 20 ppb (standard deviation between simulated DM8O for the five summers), while the observed average at the AIRNow sites was 56 ± 18 ppb. The simulations successfully captured the enhanced DM8O concentrations over the major urban areas and regions with high biogenic sources (Fig. 12a). Interannual variability of the simulated summer DM8O concentrations is on the order of 10 % (not shown) in highly populated areas and as little as 1 % in less populated areas, with the greatest variability found in the Northeast region.
Simulated concentrations of current decade PM$_{2.5}$ (PM$_{2.5}$ with no water content, unless otherwise specified) show a five summer average of $12.05 \pm 10.8 \, \mu g \, m^{-3}$, compared to $14.3 \pm 9.2 \, \mu g \, m^{-3}$ observed at the Speciation Trends Network (STN; US EPA, 2000). In general, the model slightly overestimates PM$_{2.5}$ in the Midwest, the Southeast and the Northeast and significantly underestimates PM$_{2.5}$ in the western half of the US (Fig. 7). Several factors contribute to the underestimation of PM$_{2.5}$ in the western US, including a lack of windblown dust and fire smoke emissions, and an underestimation of SOA formation (Carlton et al., 2010; Foley et al., 2010; Appel et al., 2012; Luo et al., 2011). Another important factor that influenced the underestimation of PM$_{2.5}$ is the overprediction of precipitation as shown in Fig. 5.

Figure 8. (Top panel) mean regional temperature for the five chosen summers (red) and ten summers (blue) of the current (C) and future (F) decades. (Bottom panel) total regional precipitation per day for the five chosen summers (red) and ten summers (blue) of the current (C) and future (F) decades. Each box-and-whisker indicates median, 5, 25, 75 and 95 % quartiles within each region.

3 Results and discussion

3.1 Projected changes in meteorology

For these types of climate change simulations, it is important to consider whether the five selected summers represent the climatological conditions for the 1995–2004 and 2045–2054 periods. To address this, we compared the regional mean temperature and total precipitation (Fig. 8) as well as maximum daily insolation and mean relative humidity (not shown) for all ten summers versus the five selected summers. Based on the two sample $t$ test, except for the Northwest region, we found no statistical difference in the overall regional average conditions between the five and ten summer samples ($p > 0.01$). For the purposes of this air quality assessment, this comparison of the meteorological conditions for the five selected summers to the full ten summer set of data suggest that the five summers provide a reasonable representation of decadal summer meteorological conditions. While no statistical difference was found between the five and ten summer samples, some distinct features should be highlighted: (1) for the current decade, except for the Southeast, the chosen set of five summers on average is slightly warmer than the average of the ten summers; (2) the five summers chosen for current and future decades led to a projection of cooling in the Northwest. The effects of the higher average temperature as result of the five summer sample, and the projected decrease in future temperature in the Northwest are discussed below.

Similar to the 30 year meteorological variability assessment carried out by Andersson and Engardt (2010), the differences between current and future summer meteorological conditions, based on the five representative periods, were found to be significant at the 99 % confidence level for all regions except for the Northwest. This further supports the use of five representative summers as the basis for the air quality assessment of current and future conditions.

Projected changes in selected meteorological parameters are shown in Fig. 9. Except for some minor cooling along the Pacific coast, mean summer temperature across the continental US is projected to increase between 0.5 and 4 °C (Fig. 9a). This increase falls within the lower bound of the warming predicted by the ensemble of 20 GCMs under the A1B emission scenario described by Christensen et al. (2007), but differs in the regional variability due to the higher resolution of our simulations. When compared to similar studies of equal resolution using a GCM (e.g., Goddard Institute for Space
Figure 9. Projected changes in summertime meteorological fields (future decade–current decade): (a) changes in 2 m temperature (°C); (b) percent change in solar radiation reaching the ground; (c) percent change in precipitation; (d) change in relative humidity.

Studies (GISS GCM II) driven by the A1B IPCC emission scenario and downscaled with MM5 to 36 km resolution, our simulated temperatures show higher temperature differences between future and current decades (Leung and Gustafson, 2005; Tagaris et al., 2007). Tagaris et al. (2007) and Leung and Gustafson (2005) predicted an average increase of between 1 and 3 °C for most of CONUS, and temperature reductions in the border states of the Central and South regions. Nevertheless, despite the differences in physical parameterizations contained in the GCMs and the driving IPCC emission scenarios that were used, similar temperature differences (2 to 4 °C) between our study and previous investigations were simulated for the Northeast and Southeast regions (Leung and Gustafson, 2005; Tagaris et al., 2007; Avise et al., 2009).

Projected increases in solar radiation reaching the ground vary by region. A decrease in solar radiation in the Northwest that extends to the northern boundaries of the Central regions is simulated. Small changes in the Southwest, South and Midwest are also predicted, with the largest increase experienced in the Northeast and Southeast regions (Fig. 9b). Similar results for the Northeast regions are reported by previous investigations (Leung and Gustafson, 2005; Tagaris et al., 2007; Avise et al., 2009); however, these same investigations had higher reductions in solar radiation at the border states between the Central and South regions.

Projected changes in precipitation across the US also vary depending on the region. With the exception of the Northwest and the northern boundary of the Central region, summertime precipitation is projected to decrease between −10 and −80 %. The largest decrease is projected in the Southwest region. Our results show greater precipitation reductions than those presented in Christensen et al. (2007), who projected between a 5 to 15 % decreases in the South and Southwest regions. Also, previous investigations agreed with our projected mean precipitation reductions across the domain (Fig. 9c). In the Northwest, the modeled increase in precipitation is consistent with Leung and Gustafson (2005), who projected an increase in precipitation throughout the Northwest region. In contrast, the Southeast and Northeast regions show disparities in the magnitude and the sign of the change in precipitation. While our simulations show a reduction in precipitation between 10 to 20 %, the ensemble of 20 GCMs in Christensen et al. (2007) resulted in an increase between 5 to 10 % across these regions. The disparities may be a result of the differences in resolution and parameterization schemes between our study and those used for the 20 GCMs.

Changes in relative humidity are shown in Fig. 9d. Relative humidity is shown to decrease in most of the domain except for the regions where decreases in solar radiation were projected. The greater decrease in relative humidity occurs in the Southwest and Central regions of the domain, and the largest increase is observed in the Northwest region.

3.2 Changes in biogenic emissions

Average summertime isoprene emissions over five summers of simulation for each decade are shown in Fig. 10a. Isoprene emissions occur at relatively high rates (> 50 metric tons day⁻¹) in the eastern US and at much lower rates in the western US (< 10 metric tons day⁻¹). Under future climate conditions and current land use, isoprene and monoterpene emissions are projected to increase in all regions except for the Northwest (Figs. 4 and 10b); this follows the spatial pattern of projected temperature changes (Fig. 9a). The most noticeable increases occur in the Northeast and Southeast
regions. The model projects a larger percentage increase in monoterpenes than isoprene across the domain; however, total isoprene emission is an order of magnitude higher and thus dominates the changes in total BVOC. The increase in total BVOC ranges between 17 and 45%. The only region that is projected to have reduced total BVOC emissions is the Northwest, where the model simulates a 7% reduction in isoprene emissions (Fig. 4) that in absolute amounts is greater than the 20% increase in simulated monoterpene emissions. The reduction in isoprene emissions in the Northwest is a result of the decrease in temperatures in the coastal area where the higher isoprene emissions are encountered (Fig. 9a). Previous investigations (Liao et al., 2006; Nolte et al., 2008) show the greatest increase in BVOC emissions in the Southeast region (10–50%). Similarly, Nolte et al. (2008) predicted the greatest increase in BVOC in the Southeast, but did not show any significant changes in the Northwest region.

When future climate is combined with future land use to project biogenic emissions, the spatial extent of isoprene emission increase is reduced, reflecting the expansion of low isoprene-emitting croplands into regions of high isoprene-emitting deciduous forests. In this case, the domain-average increase was approximately 12% of current decade emissions, compared with a 25% increase when changes in land use are not included (Fig. 11a). Thus, future expansion of cropland and subsequent reduction of broadleaf forested lands are projected to lessen the overall increase in US isoprene emissions that result from a warmer climate. When the future decade meteorology is combined with future land use, an increase of over 100% of current decade monoterpene emissions is predicted (Fig. 11b). The growth is most noticeable in the Central, South and Midwest regions. Also, an overall increase between 25 and 50% for the western and eastern regions is simulated. This limited increase is primarily driven by the projected changes in land use predicted for those regions.

Since the version of MEGAN used in this work does not include the suppression of isoprene emissions due to elevated concentrations of CO$_2$ (Rosenstiel et al., 2003; Heald et al., 2009), the future estimates in this study are likely to be an upper bound on isoprene emissions, and it is likely that future isoprene emissions will be lower than predicted by this work. Monoterpene emissions from US landscapes are not expected to be suppressed by increasing CO$_2$ and so are not impacted by omitting this process.

3.3 Effects of global changes upon ozone concentrations

Results for how the various global changes affect summertime DM8O are summarized in Table 2 and Fig. 12. Simulations for the future decade (Simulation 6) show a domain average of 48 ± 11 ppb with higher DM8O in the Northwest, Central and South regions than the current decade simulation (Simulation 0). In general, increases in DM8O are due to growing global anthropogenic emissions and climate change, while decreasing US emissions reduce DM8O. Changes in biogenic emissions as a result of a changing climate and land use have less of an influence on DM8O than an increase of
Table 2. Percent change in DM8O between each future scenario and the current decade base case.

<table>
<thead>
<tr>
<th>Region</th>
<th>Climate (1)</th>
<th>Climate &amp; BVOC (2)</th>
<th>Climate, BVOC, land use (3)</th>
<th>US anthropogenic emissions (4)</th>
<th>Boundary conditions (5)</th>
<th>Combined (6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Northwest</td>
<td>0.4</td>
<td>-1.0</td>
<td>-0.8</td>
<td>-0.6</td>
<td>8.1</td>
<td>4.5</td>
</tr>
<tr>
<td>Southwest</td>
<td>2.0</td>
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<td>0.0</td>
<td>-3.5</td>
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<td>4.2</td>
</tr>
<tr>
<td>Central</td>
<td>5.6</td>
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<td>-0.1</td>
<td>8.9</td>
<td>12.3</td>
</tr>
<tr>
<td>South</td>
<td>6.2</td>
<td>4.3</td>
<td>6.1</td>
<td>-0.9</td>
<td>9.6</td>
<td>13.0</td>
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<tr>
<td>Midwest</td>
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<td>8.5</td>
<td>0.0</td>
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<td>10.0</td>
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<td>Northeast</td>
<td>8.2</td>
<td>6.6</td>
<td>7.6</td>
<td>-2.3</td>
<td>1.4</td>
<td>5.3</td>
</tr>
<tr>
<td>Southeast</td>
<td>8.6</td>
<td>6.1</td>
<td>7.7</td>
<td>-3.0</td>
<td>3.3</td>
<td>6.1</td>
</tr>
</tbody>
</table>

global anthropogenic emissions. Factors that influence future DM8O are discussed in the following sections.

3.3.1 Contributions from changes in global and US anthropogenic emissions

The effects of increased long-range transport of global emissions are shown in Fig. 12f. The changes in chemical boundary conditions (the difference between Simulations 0 and 5) increase DM8O between 2 to 6 ppb across the CONUS domain. The general west-to-east and south-to-north gradients of the change in DM8O reflects intercontinental and regional transport of ozone and its precursors from the west and from Mexico at the south. The greatest impact occurs in the South (6 ppb) and Southwest (5 ppb) regions. These results show a smaller influence in DM8O from the intercontinental transport than the simulations presented in Avise et al. (2009), who report increases between 3 and 6 ppb of DM8O across the domain, with the greatest increase in the Southwest and South regions. The higher effect from intercontinental transport presented in Avise et al. (2009) is due to larger increases NOx emissions from global anthropogenic sources under the SRES A2 emission scenario. The effects of future global emissions and intercontinental transport of ozone precursors in the continental US have also been investigated by Hogrefe et al. (2004), who predicted an increase of 5 ppb in the Northeast region under the SRES A2 emission scenario.

Changes in regional US emissions of ozone precursors (difference between Simulations 0 and 4) reduce DM8O concentrations between 2 and 15 ppb in most of eastern US, most of western US and Texas. Projected increases in ozone in urban areas near the coasts are mainly due to the limited representation of the heavy-duty, shipping and rail sectors on the ESPv1.0 (Loughlin et al., 2011) by which local steady or increase in emissions of NOx and VOCs in ports are the main cause of increase in ozone in those urban areas. Regionally, larger reductions are observed in the Southeast (−3 %) and Southwest (−3.5 %) regions with a reduction of 5 ppb and the Northeast (−2.3 %) and South (−0.9 %) re-
regions with a reduction of 3 ppb (Fig. 12c, Table 2). Similar results are shown in Nolte et al. (2008) and Tagaris et al. (2007) despite a difference in the magnitude of projected emissions reductions. Tagaris et al. (2007) simulated similar ozone reductions (about 9%), with a higher nationwide reduction of 51% in NO\textsubscript{x} emissions and a slight increase (about 2%) in VOC emissions from projections based on the CAIR emission inventory. Nolte et al. (2008) showed a decrease in ozone across the domain (−12 to −16 ppb) as a result of projected reductions of 45% for NO\textsubscript{x} and 21% for VOC emissions from the NEI 2002, following the SRES A1B emission scenario. In contrast, our future simulations included a 38% reduction in NO\textsubscript{x} emissions and a slight increase (about 2%) in VOC emissions. Avise et al. (2009) predicted an average contribution of +3 ppb across the domain as a result of projecting the NEI 1999 (NEI-1999) with the Economic Growth Analysis System (EGAS) and the SRES A2 emission scenario, increasing emissions by 5% for NO\textsubscript{x} and 50% for VOCs in the future. The smaller reduction in ozone concentrations between the future and the current decade in comparison to Nolte et al. (2008) is likely to be a consequence of the increase in VOC and CO emissions from the business-as-usual scenario of MARCHAL, which, as explained in Sect. 2.4, uses diverse surrogates for growth factors for CO and VOC (Loughlin et al. 2011).

3.3.2 Contributions from changes in meteorological fields

Figure 12b shows the difference between simulations that include changes in meteorological conditions (without the effect of biogenic emissions or land use) and the current decade base case (Simulations 0 and 1). The local reductions in DM8O concentrations in the Northwest resulted from an increase in cloud cover and lower solar radiation reaching the ground, which cause a reduction in photochemistry (Fig. 9b). For other regions, increases in DM8O concentrations were projected (+5 ppb) because of increases in temperature and solar insolation; this is particularly evident in the eastern half of the US.

3.3.3 Contributions from changes in biogenic emissions and future land use

When biogenic emissions are allowed to change with the future meteorology, an average increase of DM8O with respect to the current decade base case simulations is predicted (Simulations 0 and 2). Increases of as much as 7 ppb in DM8O concentrations are mainly predicted in areas with substantial biogenic sources (Fig. 12c). Similar results are shown by Nolte et al. (2008) and Tagaris et al. (2007); both predicted an increase of DM8O above 5 ppb in the east coast. Simulated reductions between 2 and 4 ppb of DM8O in the coastal areas of the western regions are probably due to cooler temperatures and reduced solar insolation (Fig. 9a, b). Minor changes in DM8O concentrations are shown over the Southwest and Northwest regions. This is in agreement with Avise et al. (2009) and Nolte et al. (2008) who predicted reductions in DM8O concentrations from 1 to 4 ppb in the western regions, while Tagaris et al. (2007) also predicted similar reductions in ozone in the Central and Midwest regions. The disparities between this investigation and Avise et al. (2009) are reasonable due to the different climate realizations used (A2 vs. A1B; storyline scenario A2 considers higher emissions of CO\textsubscript{2} by 2050 than the scenario A1B). However, the difference in geographical features of DM8O changes with Nolte et al. (2008) and Tagaris et al. (2007) suggests that the source of disparities reside in the simulated regional meteorological fields resulting from different global climate models, RCMs and the methods used to estimate emissions from biogenic sources. We used the ECHAM5 global climate model results while both Nolte et al. (2008) and Tagaris et al. (2007) used results from the GISS global climate model. For regional climate simulations, we used WRF; both Nolte et al. (2008) and Tagaris et al. (2007) used MM5. In contrast with Nolte et al. (2008) and Tagaris et al. (2007) who use the BEIS/BELD3 (Hanna et al., 2005; http://www3.epa.gov/ttn/chief/emch/biogenic/) tool to compute biogenic emissions, this investigation estimates the biogenic emissions with MEGAN v2.04, which generally predicts higher isoprene emissions than BEIS (Hogrefe et al., 2011; Sakulyanontvittaya et al., 2012). Hogrefe et al. (2011) showed that MEGAN leads to higher DM8O in the Northeast by upwards of 7 ppb under the scenario of 2005 anthropogenic emissions; however, for a scenario by which anthropogenic NO\textsubscript{x} emissions were reduced by ~ 60%, the difference in DM8O was generally 3 ppb due to greater sensitivity to NO\textsubscript{x} emissions when MEGAN was used.

When the results from Simulation 2 (Fig. 12c) are compared to the climate-only simulations (Simulation 1, Fig. 12b), our results suggest that changes in the meteorological fields are the main driver of DM8O enhancement in Simulations 2 and 3 (Fig. 12c and d) across the domain. Even though BVOC emissions are higher in Simulation 2 relative to Simulation 1, Simulation 2 resulted in 2–4 ppb lower DM8O in the Southeast. This decrease is associated with a reduction in NO\textsubscript{x} concentrations (Fig. 14a). This decrease in NO\textsubscript{x} suggests that the effect of sequestration of NO\textsubscript{x} by the biogenic VOCs as organic nitrates (RNO\textsubscript{3}) is predominant over the effect of recycling of NO\textsubscript{x} considered in SAPRC-99, which lumps all non-peroxyacetyl (non-PAN) organic nitrates as one compound that has a NO\textsubscript{x} recycling efficiency of about 30%. The simulated reduction in ozone is consistent with the results of Xie et al. (2013), who reported increases in NO\textsubscript{x} and ozone in the Southeast when sequestration by isoprene nitrates was reduced relative to the base SAPRC-07T mechanism that has the same RNO\textsubscript{3} treatment as SAPRC-99. Evidence of the predominant effect of sequestration over the recycling of NO\textsubscript{x} in the eastern US is seen in Fig. 14, which shows an increase in RNO\textsubscript{3} and reduction in the NO\textsubscript{x} con-
concentrations in most of the eastern US for Simulation 2 relative to Simulation 1. When land use changes are included along with biogenic emissions, the increase in BVOC emissions is projected to be less while NO emission is projected to increase in areas where natural vegetation is converted to cropland. This combination leads to higher DM8O in Simulation 3 than Simulation 2 (Table 2, Fig. 12d).

### 3.3.4 Contributions from combined global change to future changes in DM8O concentrations

When the combined global changes are considered (Simulation 6), DM8O is projected to increase in all regions, except along the western coastlines. Increases of DM8O between 1 to 3 ppb in the Northwest, Southwest and Northeast regions are shown along with a local increase of 1 to 6 ppb in parts of the South, Midwest and Central regions (Fig. 12g). The increase in DM8O is mostly due to an increase in global emissions of ozone precursors from the semi-hemispheric domain, which contributes to an increase of 2–6 ppb under current climate conditions (Fig. 12f). The other contributing factors to increasing DM8O are a combination of meteorological changes (Fig. 12b) and higher BVOC emissions (with current and future land use; Fig. 12c, d). Reductions in DM8O in the urban areas resulted generally from reductions in ozone precursors from regional anthropogenic sources (Fig. 12e). However, in the western regions, lower DM8O are the result of a combination of favorable meteorological conditions (e.g., reduction in temperature and solar radiation reaching the ground) and reductions in regional ozone precursors.

### 3.4 Effects of global changes upon PM$_{2.5}$ concentrations

Results for how the various global changes affect PM$_{2.5}$ concentrations and composition are summarized in Tables 3–5 and Fig. 13. Overall, projected increase in US anthropogenic emissions have the largest impact on PM$_{2.5}$, leading to an increase in concentrations in all regions. Changes in global emissions do not have a significant impact on PM$_{2.5}$ concentrations, while changes in the climate and biogenic emissions can lead to both increases and decreases in PM$_{2.5}$ depending on the region.

#### 3.4.1 Contribution to PM$_{2.5}$ concentrations from changes in global and regional anthropogenic emissions

The results from this study are similar to those reported by Avise et al. (2009), who predicted a change in PM$_{2.5}$ of less than 1 µg m$^{-3}$ as a result of changes in future chemical boundary conditions. In our simulation, the highest increase in PM$_{2.5}$ concentrations is found in the South region (< 1 %). This increase in the South region is indicative of the effects of increased emissions from Mexico (Fig. 13f).

When the chemical composition is analyzed, Table 3 shows an increase in aerosol nitrate (NO$_3^-$) in nearly all regions except for the South; these increases are less than 0.1 µg m$^{-3}$, similar to the results of Avise et al. (2009). In our simulation, increases between 3 and 8 % in SO$_4^{2-}$ and NH$_4^+$ in the Southwest, Central and South regions are mostly a result of increase in emissions of SO$_2$ and NH$_3$ from Mexico. Similarly, Avise et al. (2009) showed higher future concentrations (by 7 to 25 %) of SO$_4^{2-}$ for the same regions resulting from higher global SO$_2$ emissions. In our simulations, changes in global anthropogenic emissions cause reductions in SOA in the Southwest, Central and South regions and increases in the Northwest, Midwest, Southeast and Northeast regions (Table 4); the simulated changes in SOA are very small (< 1.3 % and < 0.05 µg m$^{-3}$) and the variation may be due to small differences in modeled OH radical concentrations.

![Figure 13.](image-url)
Table 3. Percent change in the aerosol $\text{NH}_4^+$, $\text{SO}_4^{2-}$ and $\text{NO}_3^-$ between each future scenario and the current decade base case. The corresponding simulation number for each sensitivity simulation is shown in parenthesis.

<table>
<thead>
<tr>
<th>Region</th>
<th>Climate &amp; BVOC (1)</th>
<th>Climate, BVOC, land use (2)</th>
<th>Climate, BVOC, anthropogenic emissions (3)</th>
<th>US anthropogenic emissions (4)</th>
<th>Boundary conditions (5)</th>
<th>Combined (6)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\text{NH}_4^+$</td>
<td>$\text{SO}_4^{2-}$</td>
<td>$\text{NO}_3^-$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Northwest</td>
<td>15.7</td>
<td>-0.6</td>
<td>-0.9</td>
<td>12.8</td>
<td>-0.2</td>
<td>12.2</td>
</tr>
<tr>
<td>South</td>
<td>3.4</td>
<td>-8.8</td>
<td>-7.9</td>
<td>4.2</td>
<td>8.2</td>
<td>4.8</td>
</tr>
<tr>
<td>Central</td>
<td>12.5</td>
<td>2.1</td>
<td>2.7</td>
<td>6.9</td>
<td>3.3</td>
<td>14.8</td>
</tr>
<tr>
<td>South</td>
<td>9.1</td>
<td>4.3</td>
<td>5.8</td>
<td>7.5</td>
<td>4.8</td>
<td>22.9</td>
</tr>
<tr>
<td>Midwest</td>
<td>5.1</td>
<td>0.6</td>
<td>3.3</td>
<td>12.2</td>
<td>0.4</td>
<td>18.1</td>
</tr>
<tr>
<td>Northeast</td>
<td>1.8</td>
<td>-5.0</td>
<td>-4.2</td>
<td>17.5</td>
<td>-0.3</td>
<td>12.7</td>
</tr>
<tr>
<td>Southeast</td>
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<td>5.0</td>
<td>4.8</td>
<td>12.4</td>
<td>0.5</td>
<td>21.3</td>
</tr>
</tbody>
</table>

In the US, reductions in regional $\text{SO}_2$ and $\text{NO}_x$ emissions from regulatory curtailment on electricity generation are offset by the projected increase in emissions of PM$_{2.5}$ and NH$_3$ from other sources, thus resulting in an overall increase in PM$_{2.5}$ concentrations between 1 and 4 $\mu$g m$^{-3}$ across CONUS. Similarly, Avise et al. (2009) predicted an average increase of 3 $\mu$g m$^{-3}$ across the domain but as a result of increasing NO$_x$ and SO$_2$ from anthropogenic sources. The greatest increase, between 2 and 4 $\mu$g m$^{-3}$, is found in the urban areas across the Northwest, Northeast, Midwest and Southeast region (Fig. 13e) as a result of increase in primary emissions of PM$_{2.5}$. Similarly, Trail et al. (2015) find an increase in PM$_{2.5}$ concentrations between 1 and 2 $\mu$g m$^{-3}$ as a result of a scenario that consider changes in fuel use. In contrast, Tagaris et al. (2007) predicted a decrease of 23 % as a result of decreasing emissions. Increase in SOA concentrations resulted from higher emissions of NMVOC and an increase in primary organic aerosol from anthropogenic sources in the US (Table 4).

In terms of the inorganic PM$_{2.5}$ components, reductions in SO$_2$ and NO$_x$ emissions in the US are offset by higher emissions of primary sulfate and nitrate and ammonia, leading to an increase in both sulfate and ammonium. Compared to Tagaris et al. (2007), our investigation shows no reduction in sulfate concentrations as a result of smaller reduction in SO$_2$ emissions from anthropogenic sources. Furthermore, similar to Shimadera et al. (2014) the increase in nitrate concentrations in the form of ammonium nitrate is highly dependent on the increase in NH$_3$ emissions and less sensitive to changes in emissions of NO$_x$.

3.4.2 Contribution to PM$_{2.5}$ concentrations from global climate change alone

Despite the effect of precipitation on PM loading, as it washes out the precursors and the existing PM from the atmosphere (Seinfeld and Pandis, 2006), the effect of climate change alone (with no change to biogenic emissions) on total PM$_{2.5}$ concentrations over land is a change of less than...
1 \mu g m^{-3} (Fig. 13b). However, the change in PM$_{2.5}$ composition due to climate change is highly variable and depends on changes in temperature, relative humidity and precipitation. Increases in reaction rate constants of SO$_2$ and higher oxidant concentrations from increased temperature and solar insolation lead to an increase in aerosol sulfate formed (Dawson et al., 2007). Relative humidity and temperature affect the thermodynamic equilibrium of SO$_4^{2-}$–NH$_4^+$–NO$_3^-$, especially the partitioning of HNO$_3$ between the gas and particulate phases.

For all regions, sulfate concentrations are predicted to increase by 3–10%. Except for the Northwest regions, this change in concentrations is consistent with decreased precipitation, which reduces wet deposition, and increases in temperature and solar insolation, which increase radical production rates and increase the oxidation of SO$_2$ to produce aerosol sulfate. The same increase in temperature leads to nitrate being more volatile and thus decreases aerosol nitrate concentrations in regions where sulfate concentrations are predicted to increase. For the same regions where sulfate is projected to increase, higher concentrations of radicals also lead to higher oxidation of VOC, thus increasing SOA concentrations in the same regions.

While increasing precipitation is generally associated with decreasing PM$_{2.5}$, results here for the Northwest region showed an increase in PM$_{2.5}$ despite an increase in precipitation (Fig. 13b). This suggests the effects of slightly colder temperature and higher relative humidity in this region, leading to an enhanced formation of ammonium nitrate (Table 3). Furthermore, the increase in relative humidity in the Northwest and the coastal areas of the Southwest regions leads to the increase in production of sulfate aerosol via aqueous reaction (Luo et al., 2011). Higher concentrations of ammonium nitrate and higher concentrations of SOA (Table 4) indicate increased aerosol formation dominate over the effect of precipitation.

**3.4.3 Contribution to PM$_{2.5}$ concentrations from changes in biogenic emissions and future land use**

Simulations that consider projected climate change as well as the associated change in biogenic emissions (Simulation 2) show an increase in PM$_{2.5}$ between 0.5 and 3 \mu g m^{-3}; these changes are mainly reflected in areas with high biogenic sources (Fig. 13c). When the effects of future land use are considered (Simulation 3), an increase in the geographical extent of PM$_{2.5}$ is observed in comparison to the climate and biogenic emissions case, and higher increases (up to 6 \mu g m^{-3}) of PM$_{2.5}$ are predicted in parts of the South, Southeast, Midwest and Northeast regions (Fig. 13d). This is primarily due to the increase in emissions of sesquiterpenes (not shown) and monoterpenes (Fig. 11b), leading to more SOA being formed.

In terms of the inorganic components of PM$_{2.5}$, the effect of climate change is still the predominant factor for the change in sulfate concentrations for the Central, South, Midwest and Southeast regions (Table 3). The smaller increase or absolute reduction in sulfate in comparison to the climate-only case is due to the competition between BVOC and SO$_2$ for the availability of OH, which is an oxidant for both. Additionally, a smaller decrease in NO$_x$ in most of the domain and increase in the Northwest is predicted due to changes in the availability of OH as a result of the changes in emissions of BVOC and soil NO. The increase in availability of OH and increase in soil NO emissions lead to a higher formation of ammonium nitrate in Simulations 2 and 3 than in Simulation 1.

SOA concentrations are predicted to increase as a result of higher emissions of BVOC across the domain (Table 4). Furthermore, when climate change and biogenic emissions are combined with future land use, concentrations of SOA are predicted to increase up to 121% in the Central region and up to 188% in the Southeast due to increased biogenic monoterpenes and sesquiterpene emissions (not shown).

**3.4.4 Changes in precursors and PM$_{2.5}$ concentrations from the combined global changes**

Table 5 shows the summary of changes to PM$_{2.5}$ for all simulations. The differences in PM$_{2.5}$ between the future decade and current decade base case are greater in the eastern half of the US compared to the western half. In the eastern half of the US, the largest increases in PM$_{2.5}$ occur in the Southeast. Our results show that the 2 to 10 \mu g m^{-3} increase in PM$_{2.5}$ in the Southeast region is dominated by higher concentrations of SOA due to increased biogenic emissions as a result of increased biogenic emissions.
Table 4. Percent change of SOA between each future scenario and the current decade base case. The corresponding simulation number for each sensitivity simulation is shown in parenthesis.

<table>
<thead>
<tr>
<th>Region</th>
<th>Climate &amp; BVOC (1)</th>
<th>Climate BVOC &amp; land use (2)</th>
<th>Climate BVOC anthropogenic conditions (3)</th>
<th>US anthropogenic emissions (4)</th>
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<td>SOA</td>
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<td>0.1</td>
<td>131.0</td>
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<tr>
<td>Northeast</td>
<td>17.9</td>
<td>59.5</td>
<td>108.8</td>
<td>9.8</td>
<td>0.2</td>
<td>119.1</td>
</tr>
<tr>
<td>Southeast</td>
<td>14.2</td>
<td>73.2</td>
<td>135.1</td>
<td>8.1</td>
<td>0.3</td>
<td>143.5</td>
</tr>
</tbody>
</table>

Table 5. Percent change of PM$_{2.5}$ between each future scenario and the current decade base case. The corresponding simulation number is shown in parenthesis.

<table>
<thead>
<tr>
<th>Region</th>
<th>Climate &amp; BVOC (1)</th>
<th>Climate BVOC, land use (2)</th>
<th>Climate, BVOC, anthropogenic emissions (3)</th>
<th>US anthropogenic emissions (4)</th>
<th>Boundary conditions (5)</th>
<th>Combined (6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PM$_{2.5}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Northwest</td>
<td>7.0</td>
<td>2.1</td>
<td>7.3</td>
<td>43.2</td>
<td>−0.8</td>
<td>51.7</td>
</tr>
<tr>
<td>Southwest</td>
<td>3.3</td>
<td>3.3</td>
<td>7.1</td>
<td>20.7</td>
<td>0.7</td>
<td>27.8</td>
</tr>
<tr>
<td>Central</td>
<td>10.5</td>
<td>12.6</td>
<td>31.0</td>
<td>14.5</td>
<td>0.0</td>
<td>46.5</td>
</tr>
<tr>
<td>South</td>
<td>5.4</td>
<td>21.3</td>
<td>40.5</td>
<td>17.6</td>
<td>1.0</td>
<td>60.8</td>
</tr>
<tr>
<td>Midwest</td>
<td>7.8</td>
<td>15.2</td>
<td>37.6</td>
<td>22.4</td>
<td>0.1</td>
<td>61.2</td>
</tr>
<tr>
<td>Northeast</td>
<td>7.8</td>
<td>16.0</td>
<td>30.4</td>
<td>28.5</td>
<td>0.0</td>
<td>58.3</td>
</tr>
<tr>
<td>Southeast</td>
<td>10.6</td>
<td>29.8</td>
<td>52.4</td>
<td>24.3</td>
<td>0.4</td>
<td>78.5</td>
</tr>
</tbody>
</table>

of climate change (Fig. 13c), changes in land use (Fig. 13d; Table 4) and increase in anthropogenic emissions (Fig. 13e). Table 5 indicates that the combination of climate change, biogenic emissions and land use, and increase in anthropogenic emissions increases the concentrations of PM$_{2.5}$ between 27 and 78 % depending on the region.

4 Conclusions

We have investigated the individual and combined contributions of factors that impact US air quality by dynamically downscaling future climate projections using the WRF model and using the regional chemical transport model CMAQ version 4.7.1. Decreases in future US anthropogenic ozone precursor emissions are the only consistently beneficial influence that improves the air quality in the US and updated assumptions to generate scenarios of future US anthropogenic emissions may show even more positive influence. However that positive influence is offset by (1) increasing global emissions and changes in long-range transport, which have a negative impact on air quality across the domain; (2) climate changes (namely, increased temperatures and solar radiation), which increase ozone concentrations in the Central, South, Midwest, Northeast and Southeast regions of the domain; and (3) increases in US BVOC emissions, which also increase ozone concentrations in regions with high biogenic emissions such as the South, Midwest, Northeast and Southeast.

In the case of the overall concentrations of PM$_{2.5}$, our results indicate that the effects of increasing biogenic emissions in addition to increased primary PM from anthropogenic sources have an overall negative impact on air quality by increasing PM$_{2.5}$ concentrations between 27 and 78 %. In terms of the PM$_{2.5}$ composition, we show a regionally dependent mixture of inorganic aerosols and SOA. For the case of the Southeast, our findings indicate that increases in BVOC may result in higher concentrations of SOA. This effect extends to the Midwest and Northeast regions due to changes in land use. Furthermore, meteorological changes or regulatory curtailment, as incorporated in these simulations do not offset the increasing concentrations of primary PM.
and BVOC. In addition, synergistic effects of changes in meteorological parameters and changes in emission may shift the composition of the inorganic fraction of PM$_{2.5}$ in the US. The synergistic effects of increase of sulfate and SOA in the urban areas of the coastal regions of the Northwest and Southwest lead to an increase in PM$_{2.5}$ in those regions, offsetting decreases due to increased precipitation and temperature, and reduced primary anthropogenic emissions of SO$_2$ and NO$_x$.

In conclusion, the results of this study suggest that the efforts to improve air quality through low emission technologies and public policy directed to the electricity generation sector may not have a major effect, if future emissions from other sectors are allowed to increase. In addition, higher global anthropogenic emissions, a warmer future world and the effects of these changes on emissions from biogenic sources may increasingly undermine all regulatory efforts. Consequently, additional measures may be necessary to improve air quality in the US.

Much of the modeling components used for this research carry different levels of complexity and have reached diverse stages of development; thus, subsequent research intended to assess the effect of climate change and future regional emissions upon air quality would benefit from newer versions of the emission inventories (e.g., 2011), updated assumptions on the US emission projections (e.g., New versions of MARKAL with the use of the ESP 2.0 methodology), newer versions of MEGAN that take into account the isoprene emission suppression due to CO$_2$ concentrations and more realistic estimates of land use change, and the inclusion of emissions from wildfires and the consequent effect upon air quality.

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References


IPCC: Intergovernmental Panel on Climate Change and Intergovernmental Panel on Climate Change: Climate change 2007: the


