Climatic effects of 1950–2050 changes in US anthropogenic aerosols – Part 2: Climate response

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Abstract. We investigate the climate response to changing US anthropogenic aerosol sources over the 1950–2050 period by using the NASA GISS general circulation model (GCM) and comparing to observed US temperature trends. Time-dependent aerosol distributions are generated from the GEOS-Chem chemical transport model applied to historical emission inventories and future projections. Radiative forcing from US anthropogenic aerosols peaked in 1970–1990 and has strongly declined since due to air quality regulations. We find that the regional radiative forcing from US anthropogenic aerosols elicits a strong regional climate response, cooling the central and eastern US by 0.5–1.0 °C on average during 1970–1990, with the strongest effects on maximum daytime temperatures in summer and autumn. Aerosol cooling reflects comparable contributions from direct and indirect (cloud-mediated) radiative effects. Absorbing aerosol (mainly black carbon) has negligible warming effect. Aerosol cooling reduces surface evaporation and thus decreases precipitation along the US east coast, but also increases the southerly flow of moisture from the Gulf of Mexico resulting in increased cloud cover and precipitation in the central US. Observations over the eastern US show a lack of warming in 1960–1980 followed by very rapid warming since, which we reproduce in the GCM and attribute to trends in US anthropogenic aerosol sources. Present US aerosol concentrations are sufficiently low that future air quality improvements are projected to cause little further warming in the US (0.1 °C over 2010–2050). We find that most of the warming from aerosol source controls in the US has already been realized over the 1980–2010 period.

1 Introduction

Global mean surface temperatures increased by 0.74 ± 0.18 °C between 1906 and 2005 due to increasing greenhouse gases (Trenberth et al., 2007). However, temperature trends on regional scales are more complicated. For example, the eastern US experienced a cooling between 1930 and 1990 (Fig. 1). The net cooling effect of anthropogenic aerosols is known to have mitigated some of the global warming from greenhouse gases (Hegerl et al., 2007), but the importance of aerosol cooling on temperature trends in the US has received little attention. As US aerosol sources are increasingly controlled to improve air quality, the associated cooling is undone resulting in accelerated warming (Andreae et al., 2005; Brasseur and Roeckner, 2005; Kloster et al., 2009; Mickley et al., 2012). Air quality
improvement thus comes with climate consequences (Raes and Seinfeld, 2009). In Leibensperger et al. (2012), we reconstructed and projected the aerosol trends and associated radiative forcing from US anthropogenic sources over the 1950–2050 period. US aerosol concentrations peaked in 1970–1990 and have decreased rapidly since. We use here a general circulation model (GCM) to study the associated climate response.

Anthropogenic aerosols directly affect the climate system by scattering and absorbing solar radiation, and indirectly by altering cloud microphysical properties. Aerosol scattering cools and absorption warms the atmosphere, but both cause a reduction in surface solar radiation. Observations of surface solar radiation over the US show a widespread decrease over the 1950–1990 period followed by a more recent increase (Liepert and Tegen, 2002; Long et al., 2009). These trends have been identified in clear and all-sky scenes, suggesting a role for both direct and indirect aerosol effects. These trends in surface solar radiation are qualitatively consistent with changes in aerosol sources (Streets et al., 2009), but cannot be entirely explained by anthropogenic aerosols (Liepert and Tegen, 2002; Long et al., 2009; Wild, 2009b; Koch et al., 2011).

Aerosols are scavenged from the atmosphere by precipitation on a time scale of days, so that their radiative forcing is strongly localized over source regions (Schulz et al., 2006). A critical issue is whether the regional radiative forcing of aerosols elicits a correspondingly regional climate response. Observation-based studies have related changes in surface solar radiation with other climate variables as a method of deducing the local effects of aerosol forcing. They show evidence that aerosols have lowered surface air temperature and temporarily offset greenhouse warming (Qian and Giorgi, 2000; Wild et al., 2007; Ruckstuhl et al., 2008; Philipona et al., 2009), reduced the diurnal temperature range by dampening daily maximum temperatures (Liu et al., 2004b; Wild et al., 2007; Makowski et al., 2009), lowered evaporation rates (Peterson et al., 1995; Liu et al., 2004a; Roderick et al., 2007), and increased soil moisture (Robock et al., 2005).

Some GCM studies have found strong regional climate sensitivity to aerosols including in India (Menon et al., 2002; Wang et al., 2009a), southeast Asia (Chang et al., 2009; Zhang et al., 2011; Lee and Kim, 2010), the North Atlantic (Fischer-Bruns et al., 2009), and the western US (Jacobson, 2008). However, other studies have found that the climate response to aerosol radiative forcing is more hemispheric or global in scale with patterns similar (but opposite in sign) to greenhouse gas forcing (Mitchell et al., 1995; Shindell et al., 2007, 2008; Levy et al., 2008; Kloster et al., 2009). Shindell et al. (2010) estimated the spatial extent of the climate response to aerosol radiative forcing in four GCMs and found it to extend ~3500 km in the meridional direction and ~12,000 km in the zonal direction. Fischer-Bruns et al. (2010) investigated the climate impacts of removing North American aerosols and found a 1.0–1.5°C summer warming in surface air over the US and North Atlantic Ocean. They also found a 1.5–2.0°C warming of the Arctic in winter. However, Mickley et al. (2012) simulated the climate response of completely removing aerosols over the US and found a 0.4–0.6°C regional warming in the US with little effect elsewhere.

There is strong motivation for air quality agencies to decrease aerosol concentrations to improve public health. Better understanding of the associated climate response is necessary. The US is of particular interest in this regard because aerosol concentrations rose in the 20th century, peaked in the 1980s, and have been decreasing rapidly since due in large part to a 56% reduction of SO2 emissions between 1980 and 2008 (US EPA, 2010). Here we use the 1950–2050 time series of US aerosol concentrations from Leibensperger et al. (2012) to conduct 1950–2050 transient climate simulations with the NASA Goddard Institute for Space Studies (GISS) GCM 3 (Rind et al., 2007). Our objective is to investigate the regional climate effects of historical and projected changes in US anthropogenic aerosol sources. An important advance compared to previous work is the use of a realistic evolution of aerosol sources.

2 Methods

We conduct sensitivity simulations with the GISS GCM 3 to study the evolving 1950–2050 climate response to changing US anthropogenic aerosol sources. The GCM uses archived global 3-D monthly mean concentrations of different aerosol components from the GEOS-Chem chemical transport model (CTM) with time-dependent emissions based on historical data and future projections. The CTM

Fig. 1. Observed change in surface air temperature between 1930 and 1990. Temperature change is based on the linear trend as in Hansen et al. (2001). Observations are from the NASA GISS Surface Temperature Analysis (GISTEMP; http://data.giss.nasa.gov/gistemp/).
2.1 Aerosol simulations

We conduct a 2-yr GEOS-Chem simulation of coupled ozone-NO\textsubscript{x}-VOC-aerosol chemistry (http://geos-chem.org; Bey et al., 2001; Park et al., 2004) for each decade between 1950 and 2050. The first year is used as model initialization. Monthly mean fine aerosol concentrations are archived from the second year for use in the GCM including sulfate-nitrate-ammonium (SNA), primary organic aerosol (POA), secondary organic aerosol (SOA), and black carbon (BC) (Park et al., 2006; Liao et al., 2007). Simulations for all years use the same assimilated meteorological data from 2000–2001 so that changes in concentrations over the 1950–2050 period are due to emissions only. The meteorological data are from the NASA Goddard Earth Observing System (GEOS)-4 with $1^\circ \times 1.25^\circ$ horizontal resolution, 55 vertical levels, and 6-h temporal resolution (3-h for surface variables and mixing depths). The horizontal resolution is degraded to $2^\circ \times 2^\circ$ for input to GEOS-Chem. The effect of US anthropogenic sources is determined by parallel sensitivity simulations for the 1950–2050 period with zero US anthropogenic emissions of SO\textsubscript{2}, NO\textsubscript{x}, POA, and BC. “Anthropogenic” here includes fuel and industrial sources but not open fires.

We use 1950–1990 global anthropogenic emissions of SO\textsubscript{2} and NO\textsubscript{x} from EDGAR Hydro 1.3 (van Aardenne et al., 2001) and 2000 emissions from EDGAR 3.2 FT (Olivier and Berdowski, 2001). Historical emissions of BC and POA are from Bond et al. (2007). Emissions past 2000 are calculated using growth factors derived from the Integrated Model to Assess the Greenhouse Effect (IMAGE; Streets et al., 2004) following the IPCC A1B scenario (Nakićenović and Swart, 2000). As in Fiore et al. (2002) and Wu et al. (2008), growth factors are calculated for different countries and fuel types (fossil fuel and biofuel). Ammonia emissions are from Bouwman et al. (1997) as modified by Park et al. (2004), except for East Asia (Streets et al., 2003). Additional sources include climatological biomass burning (Duncan et al., 2003), fertilizer (Wang et al., 1998), aircraft (Chin et al., 2000), the biosphere, volcanoes, and lightning. See Leibensperger et al. (2012) for more detail.

2.2 Climate simulations

We conduct transient climate simulations with the GISS GCM 3 using a horizontal resolution of $4^\circ \times 5^\circ$ and 23 vertical levels that extend from the surface to 0.002 hPa. GISS GCM 3 shares a common history with another NASA GISS GCM, Model E (Schmidt et al., 2006), but the two differ in their parameterizations of gravity wave drag, convection, and the boundary layer (Rind et al., 2007). GISS GCM 3 has been previously used in studies investigating air quality-climate interactions (Leibensperger et al., 2008; Wu et al., 2008; Pye et al., 2009; Chen et al., 2010b; Mickley et al., 2012), tracer transport (Rind et al., 2007), climate response to solar forcing (Rind et al., 2008), and stratospheric ozone-climate interactions (Rind et al., 2009a, b). The model contains a Q-flux ocean, in which monthly oceanic heat transports are held constant but sea surface temperature and sea ice coverage are allowed to respond to energy exchange with the atmosphere (Hansen et al., 1988). We calculate ocean heat transport using equilibrium climate simulations forced with observed sea surface temperature (SST) and sea ice distributions for 1946–1955 (Rayner et al., 2003), following the method outlined by Hansen et al. (1988).

To compute the aerosol direct effect, we import 3-D monthly mean aerosol concentrations archived from GEOS-Chem. Concentrations are interpolated between decadal time slices. Aerosol optical properties are calculated assuming an internal mixture. Following Chung and Seinfeld (2002) and Liao et al. (2004), we assume a standard gamma aerosol size distribution with an effective dry radius of 0.3 µm and area-weighted variance of 0.2. Optical properties of the internal mixture are calculated using the volume-weighted mean of the refractive indices of the individual components. The resulting direct forcing from US anthropogenic aerosol sources in 1980 (peak aerosol loading) is only $-0.07 \text{ W m}^{-2}$ globally but $-2.0 \text{ W m}^{-2}$ over the eastern US (Leibensperger et al., 2012). Dust and sea salt aerosols are considered externally mixed and held constant for 1950–2050 at the climatological radiative forcing values of Hansen et al. (2002).

Aerosol indirect radiative effects are computed following the approach previously implemented by Chen et al. (2010b) in the GISS GCM 3. This includes the first indirect effect (cloud albedo) and the second indirect effect (cloud lifetime), both on liquid stratiform clouds only. Aerosol mass concentrations affect the cloud droplet number concentration $N_c$ (m$^{-3}$), which in turn determines the effective radius of cloud droplets and the rate of autoconversion to precipitation. We calculate $N_c$ from the archived GEOS-Chem aerosol mass distributions using the method of Chen et al. (2010b):

$$\log N_c = A + B \log m_i$$

where $m_i$ is the molar concentration of dissolved aerosol ions (mol m$^{-3}$) and $A$ and $B$ are gridded 3-D monthly mean coefficients derived from detailed simulations of aerosol microphysics and activation within the GCM (Adams and Seinfeld, 2002; Nenes and Seinfeld, 2003; Fountoukis and Nenes, 2005; Pierce and Adams, 2006). Cloud optical depth scales as the inverse of the area-weighted mean effective radius $r_e$ of the cloud droplet size distribution (Del Genio et al., 1996). We obtain $r_e$ from:

$$r_e = \kappa^{-\frac{1}{3}} \left[ \frac{3L}{4\pi N_c} \right]^\frac{1}{2}$$

where $L$ is the liquid water content of the cloud (cm$^3$ water per cm$^{-3}$ air), and $\kappa$ is a constant (0.67 over land, 0.80 over...
ocean; Martin et al., 1994) that relates the volume-weighted and area-weighted mean radii. Autoconversion rates are calculated using the parameterization of Khairoutdinov and Kogan (2000), which fits results from an explicit microphysical model:

$$\frac{dg}{dt} = -1350\gamma q_{1}^{2.47} N_{c}^{-1.79}$$  \hspace{1cm} (3)

where $q_{1}$ is the cloudwater mass content (kg kg$^{-1}$). Hoose et al. (2008) and Chen et al. (2010b) added the tuning parameter $\gamma$ in order to retain GCM climate equilibrium. We find that a $\gamma$ value of 12 retains top-of-atmosphere (TOA) radiative balance in a climate equilibrium simulation for 1950 conditions including fixed SST and sea ice (Leibensperger et al., 2012). $\gamma = 12$ is consistent with the recent results of Morales and Nenes (2010), who found that autoconversion rates can be underestimated by a factor of 2 to 10 when models use gridbox-scale values of $N_{c}$.

Two sets of control simulations were conducted. The first includes only the direct radiative forcing from aerosols, using the imported aerosol concentrations from GOM-Chem. The second additionally imports $N_{c}$ as calculated in Eq. (1) to account for the aerosol indirect effects. Sensitivity simulations were conducted relative to each of these controls using the aerosol and $N_{c}$ fields from the GOM-Chem simulations with US anthropogenic sources of $SO_{2}$, $NO_{x}$, POA, and BC shut off. Differences between the control and sensitivity simulations then measure the climate response to US anthropogenic sources through the direct and indirect effects of aerosols.

Each set of simulations consists of a five-member ensemble conducted from 1950 to 2050 using the trends in natural and greenhouse gas forcing described by Hansen et al. (2002). Future greenhouse gas concentrations follow the IPCC SRES A1B scenario with $CO_{2}$, $N_{2}O$, and $CH_{4}$ reaching 522, 0.350, and 2.40 ppm, respectively, by 2050. The A1B scenario provides a radiative forcing for the 21st century comparable to the more recent RCP6 scenario from the IPCC (Moss et al., 2010). Initiating climate simulations from 1950 equilibrium conditions neglects the small radiative imbalance that occurred at that time. A previous study with the GISS GCM using the same climate forcing as here (except for tropospheric aerosols) found the Earth to be out of radiative balance by +0.18 W m$^{-2}$ for 1951 conditions (Sun and Hansen, 2003). This suggests that our simulations underestimate post-1950 global warming by approximately 0.1 °C, a small effect that does not impact our assessment of the climate response to US anthropogenic aerosols since it is common to both the control and sensitivity simulations.

We conducted an additional sensitivity simulation to isolate the climate effects of US anthropogenic sources of BC. This simulation uses the sulfate, nitrate, POA, and SOA distributions from the control simulation, but no BC emission from US anthropogenic sources. Results indicate that the climate effects of US anthropogenic BC are small and indistinguishable from interannual variability. This is consistent with the weak radiative forcing from US anthropogenic BC (+0.02 W m$^{-2}$ globally and +0.4 W m$^{-2}$ over the eastern US for 1970–1990; Leibensperger et al., 2012). More generally, we find that a regional radiative forcing of about 1.0 W m$^{-2}$ is necessary to produce a climate response greater than natural variability in the GISS GCM 3 with a 5-member ensemble. The climate effects of BC may differ when considering co-emitted species and microphysical effects (Bauer et al., 2010; Chen et al., 2010a; Jacobson, 2010). The climate effects of BC are additionally sensitive to the vertical profile of BC and modeled cloud cover (Koch and Del Genio, 2010). Koch et al. (2011) found surface air temperature to be less sensitive to BC than a similar amount of sulfate radiative forcing due to compensating changes in stability and cloud cover. Previous studies have shown a significant climate response in Asia where BC radiative forcing is greater (Menon et al., 2002; Wang et al., 2009a), but this region has larger BC emissions. We do not discuss this simulation further.

The use of archived monthly mean aerosol distributions as input to our climate simulations does not allow for feedbacks of changing climate on aerosol concentrations. These feedbacks are likely very small relative to the source driven aerosol perturbations implemented here, considering that both models and observations indicate little direct sensitivity of aerosol air quality to climate change (Jacob and Winner, 2009; Tai et al., 2011). The use of monthly mean aerosol concentrations does not introduce significant bias in the calculation of the direct radiative effect (Koch et al., 1999), but it may affect the aerosol indirect effect due to the nonlinear relationship between aerosol amount and cloud droplet number (Jones et al., 2001) as seen in Eq. (1).

We test the statistical significance of all our results using a modified version of Student’s t-test to account for serial correlation (Zwiers and von Storch, 1995). Results presented here are significant at the 95th percentile and represent the mean of ensemble members unless specified.

### 3 Climate response to US anthropogenic aerosols

#### 3.1 Radiation

Figure 2 shows the annual mean change in net TOA and surface solar radiation due to the direct and indirect radiative effects of US anthropogenic aerosols in 1970–1990, the period when US aerosol loadings were at their peak. These deficits from the radiative forcings reported by Leibensperger et al. (2012) in that they include the effects of climate response, such as changes in cloud cover. We find that the radiative effect is strongly concentrated over the eastern US, and define a mid-Atlantic US region boxed in Fig. 2 where the effect is maximum. The annual mean TOA radiative effect averages $-6$ W m$^{-2}$ over that region, whereas the global TOA radiative effect is only $-0.08$ W m$^{-2}$. 

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Fig. 2. Changes in net top-of-atmosphere (TOA) radiation (left) and surface solar radiation (right) due to US anthropogenic aerosol sources. The top panels show the annual mean aerosol effect (direct and indirect) averaged over 1970–1990. The bottom panels show the 1950–2050 evolution of the direct effect only (dashed) and the sum of direct and indirect effects (solid) averaged over the mid-Atlantic US (boxed region). Values are differences between the control simulation and a simulation with US anthropogenic aerosol sources shut off. In the top panels, dots indicate differences significant at the 95th percentile, downward triangles indicate observation sites used by Liepert and Tegen (2002), and upward triangles indicate observation sites used by Long et al. (2009). In the lower panels, black squares indicate the annual mean aerosol direct radiative forcings over the mid-Atlantic US region from US anthropogenic sources calculated by Leibensperger et al. (2012). Time series have been filtered by a 15-yr moving average.

The bottom panels of Fig. 2 show the 1950–2050 time series of the TOA and surface radiative effects from US anthropogenic aerosol sources over the mid-Atlantic region. Results are shown for the direct effect only and for the sum of direct and indirect effects. The magnitudes and trends of the direct effect match closely the corresponding instantaneous radiative forcings computed in Leibensperger et al. (2012) (squares in Fig. 2). The indirect effect is comparable in magnitude to the direct effect. Sulfate is the largest contributor to US anthropogenic aerosol forcing (Leibensperger et al., 2012). We see from Fig. 2 that the radiative perturbations largely follow the evolution of SO$_2$ sources: increase until 1980 and rapid decrease afterward. Changes level off after 2020, by which time SO$_2$ emissions are 80% lower than their 1980 peak.

The surface radiative effects in Fig. 2 are about 50% larger than the TOA effects due to aerosol absorption. Liepert and Tegen (2002) used six observation sites east of 95°W (downward triangles in Fig. 1) from the National Solar Radiation Database (NSRDB; http://rredc.nrel.gov/solar/old_data/nsrdb/) to estimate a clear-sky trend in surface solar radiation of −7 W m$^{-2}$ between 1961 and 1990 that they attributed to the aerosol direct effect. Long et al. (2009) calculated an observed 1996–2007 increase in surface solar radiation of +5 W m$^{-2}$ (clear sky) and +8 W m$^{-2}$ (all sky) averaged over 11 Department of Energy Atmospheric Radiation Measurement (ARM) and National Oceanic and Atmospheric Administration (NOAA) US Surface Radiation Budget (SURFRAD) sites (upward triangles in Fig. 2). However, and as previously noted (Liepert and Tegen, 2002; Long et al., 2009; Wild, 2009a, b; Koch et al., 2011), we find that anthropogenic aerosols cannot explain the magnitude of the observed surface radiation trends. Sampling our control simulation at the six sites analyzed by Liepert and Tegen (2002), we find a simulated all-sky decrease of surface solar radiation of 1.1 W m$^{-2}$ between 1961 and 1990 due to the aerosol direct effect, much smaller than the observed value. Conversion of our all-sky trend to the clear-sky value would not reconcile the difference. Similarly, our control simulation underestimates the 1996–2007 all-sky increase of Long et al. (2009) (+2.4 W m$^{-2}$ vs. +8 W m$^{-2}$).
We previously showed that our simulated trend of sulfate over the US is in good agreement with observations, at least for 1990–present, while the simulated decreasing trend of black carbon is too low by a factor of 2 (Leibensperger et al., 2012). Correcting for this discrepancy does not reconcile the simulated and observed 1996–2007 trends in solar radiation. The observed surface solar radiation trends remain larger than our simulated trends and thus seem much larger than can be explained from aerosol trends. The discrepancy seems unlikely to arise from underestimated aerosol radiative effects since models consistently underestimate the observed trend (Liepert and Tegen, 2002; Long et al., 2009; Wild, 2009a, b; Koch et al., 2011). The 1961–1990 NSRDB surface solar radiation data suffer from inconsistent data quality, but the more recent measurements presented by Long et al. (2009) have consistent annual calibration and daily performance monitoring. The cause of the model-observation discrepancy is unclear but suggests that observed trends in surface radiation are not driven solely by aerosols.

3.2 Temperature

Figure 3 shows the annual mean temperature changes in surface air and at 500 hPa due to the direct and indirect effects of US anthropogenic aerosol sources over the 1970–1990 period when US anthropogenic aerosol forcing was at its peak. Changes in surface air temperature are largest in the eastern US (0.5–1.0 °C cooling), collocated with the maximum radiative effect (Fig. 2). The cooling influence of US anthropogenic aerosols extends over much of the Northern Hemisphere, but beyond the US and North Atlantic Ocean it is only marginally significant against modeled interannual variability. The annual mean cooling averages 0.1 °C for the Northern Hemisphere and 0.05 °C for the Southern Hemisphere.

We find that the cooling effect of US anthropogenic aerosols is largely confined to the US and North Atlantic. As pointed out in the Introduction, Shindell et al. (2010) estimated the spatial extent of the climate response to aerosol radiative forcing to span ~3500 km in the meridional direction and ~12 000 km in the zonal direction. We find in our simulation that some cooling from US anthropogenic aerosols extends to these distances but is only marginally significant or insignificant. The cooling within the US caused by US anthropogenic aerosols shown in Fig. 3 is not apparent in the simulations of Shindell et al. (2010), which simulated the climate response of all anthropogenic aerosols. However, cooling within the US can be found in other simulations of the climate effects of anthropogenic aerosols (Mitchell et al., 1995; Hansen et al., 2005; Wilcox et al., 2009; Mickley et al., 2012).

Cooling is more diffuse at higher altitudes, reflecting the faster transport of heat (Fig. 3, top). At 500 hPa, the largest cooling is 0.3 °C over the eastern US and North Atlantic Ocean. Statistically significant cooling covers more of the Northern Hemisphere at 500 hPa than at the surface. Annual mean cooling at 500 hPa averages 0.1 °C for the Northern Hemisphere and 0.02 °C for the Southern Hemisphere, similar to the hemispheric changes in surface air temperature. Cooling is even more diffuse at 300 hPa (not shown), but Northern Hemisphere cooling is similarly 0.1 °C.

Figure 4 shows the surface air cooling over the US due to US anthropogenic aerosols for the 1970–1990 period for the simulations including only the aerosol direct effect (top) and the combination of direct and indirect effects (bottom). The bottom panel is a zoomed version of the bottom panel of Fig. 3. Significant cooling extends over much of the US even when the aerosol direct effect alone is considered. The magnitude of the cooling doubles when the indirect effects are included but the spatial patterns are similar. Thus the significance and localization of the US cooling due to US anthropogenic aerosol sources is not contingent on the indirect effects, which are far more uncertain than the direct effect.
Cooling is strongest in the Midwest, shifted westward relative to the surface and TOA radiative effects shown in Fig. 2. This is due to hydrological factors and is discussed further in Sect. 3.3.

Figure 5 shows seasonal statistics of the effects of US anthropogenic aerosols on surface air temperatures in the mid-Atlantic US (boxed region in Fig. 2). The change in surface air temperature is largest in summer and autumn, with regionally averaged cooling of more than 1.0 °C in autumn. Aerosol radiative forcing is largest in summer when solar radiation is strongest. The larger surface air temperature changes in autumn reflect the drier conditions typical of that time of year, so that less of the radiative effect is buffered by changes in surface evaporation (latent heat flux).

Figure 5 also presents the changes in extreme temperatures. We find that aerosol cooling has a larger effect on daily maximum than daily minimum temperatures, in all seasons, as expected since the forcing is due to scattering of solar radiation. The cooling is largest during heat waves of summer and autumn (95th percentile daily maximum temperatures, corresponding to the 5 hottest days of each season). These heat waves occur under cloud free conditions when the aerosol direct effect is especially effective. We find that the warmest days are cooler by 1.0 °C in summer and 1.3 °C in autumn. The coldest nights of each season (5th percentile daily minimum temperatures) are least sensitive to aerosols except in winter when they are typically associated with synoptic-scale clear-sky conditions.

The model pattern of aerosol-driven cooling over the US in Fig. 4 is remarkably similar to the observed pattern in the 1930–1990 trend in surface air temperature shown in Fig. 1. The largest area of cooling in the central US has previously been referred to as a “warming hole” (Pan et al., 2004; Kunkel et al., 2006). Previous GCM studies have associated this warming hole with variations in SSTs in the tropical Pacific (Robinson et al., 2002; Kunkel et al., 2006; Wang et al., 2009b). Kunkel et al. (2006) additionally point out a strong association between the observed variability of North Atlantic SSTs and central US surface temperatures. Our results indicate that the warming hole could be due to US anthropogenic aerosols, as SO$_2$ anthropogenic emissions in 1930 were only 60 % of those in 1980. We find that US anthropogenic aerosols lower SSTs in the North Atlantic region outlined by Kunkel et al. (2006) by up to −0.3 °C. Lower SSTs over the North Atlantic enhance the anticyclonic transport of marine air over the Gulf Coast of the US, magnifying the cooling as discussed below.
Aerosols affect the hydrological cycle by reducing evaporation (due to reduced surface solar radiation) and by altering cloud cover and precipitation. Figure 6 shows the annual mean response of evaporation, precipitation, soil moisture availability, and cloud cover to US anthropogenic aerosols for the 1970–1990 period. The changes shown in Fig. 6 are for the total aerosol effect (direct + indirect), but similar responses with less statistical significance are found when only the aerosol direct effect is considered.

Reduced solar radiation at the surface decreases annual mean evaporation rates in the eastern US and the North Atlantic (Fig. 6a). The change is greatest in summer, when evaporation along the eastern seaboard decreases by up to 0.4 mm day\(^{-1}\). As mentioned in Sect. 3.2, the decrease in latent heat flux associated with lower evaporation rates acts as a buffer to surface temperature changes, reducing the magnitude of aerosol cooling. We find that US anthropogenic aerosols decrease latent heat fluxes much more in summer (6.6 W m\(^{-2}\) for 1970–1990 averaged over mid-Atlantic US) than in autumn (0.6 W m\(^{-2}\)), reflecting the greater availability of soil moisture in summer. In contrast to the general decrease in evaporation rates over the US, we find an increase in the south-central region. This is due to changes in soil moisture as discussed below.

The reduction in evaporation in the eastern US results in a decrease of downwind precipitation along the eastern seaboard (Fig. 6b). The decrease in precipitation is additionally promoted by the aerosol cloud lifetime effect (second aerosol indirect effect), which reduces the precipitation efficiency of clouds. The cloud lifetime effect is applied in the model to liquid stratiform clouds only, and this appears to be the dominant cause for the increase in eastern US cloud cover in Fig. 6c. We find little net change in moist convective cloud cover.

In contrast to the general slowdown of the hydrological cycle over the US, we find that evaporation and precipitation increase in the south central US. This is mostly driven by a summertime aerosol-induced change in circulation. Figure 7 shows the effect of US anthropogenic aerosols on 850 hPa geopotential heights in summer. Aerosols cool the North Atlantic (Fig. 2), strengthening the Bermuda High and thus the onshore flow of marine air from the Gulf of Mexico that is the principal source of moisture for the central and eastern US in summer. This enhances cloud cover, precipitation, and soil moisture over the south-central US (Fig. 6). A similar increase in central US precipitation due to anthropogenic sulfate was presented but not discussed by Jones et al. (2007). Increased cloud cover in the central US due to anthropogenic aerosols explains the particularly strong radiative and surface cooling effects in that region (Figs. 2 and 3).
This is consistent with previous observational studies of the US warming hole which found it to be associated with additional moisture from the Gulf of Mexico causing enhanced evapotranspiration (Pan et al., 2004) and cloud cover (Robinson et al., 2002). Our work suggests that US anthropogenic aerosols may be the drivers of changes in circulation causing central US cooling.

4 Aerosol effects on 1950–2050 trends in US surface air temperature

Our analysis of the climate effects of US anthropogenic aerosols has focused so far on the 1970–1990 period when the US aerosol loading was the largest. Leibensperger et al. (2012) presented a detailed analysis of US anthropogenic aerosol trends for the 1950–2050 period. US aerosol loading (and corresponding radiative forcing) increased from 1950 to 1980, decreased since then, and is projected to continue decreasing in the future (Fig. 2). Most of the anthropogenic aerosol radiative forcing is due to sulfate produced by oxidation of SO\textsubscript{2}. Emissions of SO\textsubscript{2} in the US decreased by 56% from their peak in 1980 to 2008 according to EPA (2010) and this is verified by observed trends in sulfate wet deposition (Leibensperger et al., 2012). US sources of SO\textsubscript{2} and other aerosol precursors in the IPCC A1B scenario are projected to continue to decrease until 2020 and then level off.

Figure 8 shows the 1950–2050 simulated cooling trends over the mid-Atlantic US due to US anthropogenic aerosol sources. Cooling increased from −0.3°C in the 1950s to −0.65°C in the 1970s, remained flat until 1995, and then decreased back to −0.3°C by 2010. Further decrease in aerosol cooling is projected over the coming decades but at a much slower rate, to −0.2°C by 2030. This trend in cooling is well correlated with the model trend in US aerosol sources (Leibensperger et al., 2012).

The observations show no significant warming between 1960 and 1979 (+0.01 ± 0.20°C decade\textsuperscript{−1}). Our control simulation, which incorporates US anthropogenic aerosols, reproduces this result (−0.02 ± 0.20°C decade\textsuperscript{−1}). The simulation without US anthropogenic aerosols, however, produces a warming trend over this period (+0.30 ± 0.19°C decade\textsuperscript{−1}). We conclude that the increasing abundance of US anthropogenic aerosols effectively offset greenhouse warming before 1980 (difference in trends significant at the 90th percentile). Figure 9 shows significant observed warming over the eastern US for the 1980–2009 period (+0.21 ± 0.20°C decade\textsuperscript{−1}). Anthropogenic aerosol sources in the US peaked in 1980 and decreased afterward. Our control simulation reproduces the post-1980 warming with a rate of +0.41 ± 0.08°C decade\textsuperscript{−1} for the 1980–2009 period. The sensitivity simulation without US anthropogenic
Aerosols show slower warming (+0.30 ± 0.10 °C decade⁻¹, difference from control simulation significant at the 90th percentile), which represents a continuation of the 1960–1979 trend due to greenhouse warming. The larger trend in the control simulation reflects the acceleration of positive radiative forcing due to loss of the aerosol radiation shield. Beyond 2010 the rate of warming in the control simulation eases and eventually approaches that of the sensitivity simulation with no US anthropogenic aerosol sources.

### 5 Conclusions

Aerosol concentrations over the US peaked in the 1970–1990 period, have decreased rapidly since, and are projected to continue decreasing in the future as a result of air quality regulations to protect public health. We used a general circulation model (GISS GCM 3) to study the regional climate response to this time-dependent regional aerosol radiative forcing. The climate response to US anthropogenic aerosol sources was diagnosed through 1950–2050 control climate simulations including our best estimates of time-dependent greenhouse gas concentrations and aerosol distributions (Leibensperger et al., 2012) and through sensitivity simulations with US anthropogenic aerosol sources shut off. Our goal was to determine how aerosol trends have contributed to recent climate trends over the US and to examine the climate consequences of further US aerosol reductions in the future.

We find that peak US anthropogenic aerosol loadings (1970–1990) cooled the central and eastern US by 0.5–1.0 °C on an annual mean basis. This cooling is strongest in summer and autumn, and especially strong during heat waves. Surface solar radiation was reduced by up to 8 W m⁻² (5 W m⁻² from the direct effect alone) during the same period of peak aerosol loading. The reduction of solar radiation and prolonged cloud lifetimes slowed the hydrological cycle in the eastern US by reducing evaporation and precipitation (0.2 mm day⁻¹ annual mean, 0.4 mm day⁻¹ in summer).

We find the cooling effect of US anthropogenic aerosols to be largest in the central US. This spatial pattern is found in observations (Fig. 1) and has been reported previously as a “warming hole” (Pan et al., 2004; Kunkel et al., 2006). This is due in the model to aerosol-driven cooling of the western North Atlantic that enhances the southerly flow of moist air from the Gulf of Mexico. This hydrological contribution to the central US “warming hole” has been previously identified from observations by Robinson et al. (2002) and Pan et al. (2004), and attributed to sea surface temperature (SST) anomalies. We show that this mechanism is consistent with the expected effect of US anthropogenic aerosols on North Atlantic SSTs.

Our model results show that US anthropogenic aerosols can explain the observed lack of warming over the eastern US from 1930 to 1980 followed by very rapid post-1980 warming. Without US anthropogenic aerosol sources, we find in the model a relatively constant rate of warming over the 1950–2050 period, driven by increasing greenhouse gases. Increasing aerosols until 1980 offset the warming. Decreasing aerosol after 1980 accelerated the warming due to the loss of the aerosol cooling shield. We find that the observed warming from 1990 to 2010 is significantly greater than would have been expected from greenhouse gases alone.
We project that future reductions in US aerosol sources will increase warming over the eastern US by 0.1 °C. However, we find that most of the warming due to reducing US aerosol sources for air quality objectives has in fact already been realized (0.35 °C over the eastern US in 1980–2010).

Our results have several implications for US air quality policy. We find that reductions in aerosol sources to improve air quality have elicited a strong regional warming response over the past 20 yr. We also find that future aerosol reductions should have relatively little additional climate impact because aerosol sources are already low by now. It has been suggested that future black carbon (BC) emission controls could provide relief from future warming (Bond, 2007; Grieshop et al., 2009; Penner et al., 2010), but we find that BC sources in the US are too small for their climatic impact to be significant.

Relating aerosol radiative forcing to regional climate change is challenging. There are many model uncertainties involved in the mechanisms of aerosol-cloud interactions, the response of the hydrological cycle, the lateral transport of heat in the ocean (the Q-flux parameterization used here does not allow for change in that transport), and other aspects of the climate model. Multi-model analyses are needed to address the robustness of results (National Research Council, 2005). Our ability to reproduce observed 1950–2010 temperature trends lends some confidence to our conclusions.

Although our results are specific to the US, they also warn of possibly strong regional warming over East Asia in the coming decades as China embarks on vigorous emission controls to address its pressing aerosol pollution problem. The climate response to anthropogenic aerosols in East Asia may be very different from the US because of the greater contribution of BC to the aerosol mix and because of specific meteorological features such as the monsoon. Application of our approach to that region would be of considerable interest.

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