Offsetting Effects of Aerosols on Arctic and Global Climate in the Late 20th Century

Qiong Yang\textsuperscript{1,2,*}, Cecilia M. Bitz\textsuperscript{1} and Sarah J. Doherty\textsuperscript{2}

\textsuperscript{1}Department of Atmospheric Sciences, University of Washington, Seattle
\textsuperscript{2}Joint Institute for the Study of Atmosphere and Ocean, University of Washington,
Seattle

*Correspondence to Qiong Yang (qyang@atmos.washington.edu)
Abstract

We examine the impacts of atmospheric aerosols on Arctic and global climate using a series of 20th century transient simulations from Community Climate System Model version 4 (CCSM4). We focus on the response of surface air temperature to the direct radiative forcing driven by changes in sulfate and black carbon (BC) concentrations from 1975 to 2005 and we also examine the response to changes in sulfate, BC, and organic carbon (OC) aerosols collectively. The direct forcing from sulfate dominates the aerosol climate effect. Globally averaged, changes in all three aerosols produce a cooling trend of 0.015 K/decade during the period 1975-2005. In the Arctic, surface air temperature has large spatial variations in response to changes in aerosol concentrations. Over the European Arctic, aerosols induce about 0.6 K/decade warming which is about 1.8 K warming over the 30 years period. This warming is triggered mainly by the reduction in sulfate and BC emissions over Europe since the 1970s and is reinforced by sea ice loss and a strengthening in atmospheric northward heat transport. Changes in sulfate concentrations account for about two thirds of the warming and BC for the remaining one-third. Over the Siberian and North American Arctic, surface air temperature is likely influenced by changes in aerosol concentrations over Asia. An increase in sulfate optical depth over Asia induces a large cooling while an increase in BC over Asia causes a significant warming.
1. Introduction

The Arctic (the region poleward of 60°N) has warmed dramatically since the 1970s, by ~1.5°C. The warming in the Arctic is at least two times larger than the global mean temperature increase [e.g., Serreze et al., 2009]. This phenomenon is known as Arctic amplification [e.g., Manabe and Stouffer, 1980]. The detailed mechanisms causing the warming are not fully understood [Serreze and Barry, 2011]. Climate model simulations have shown that ice-albedo feedback is likely to account for much of the Arctic warming [e.g., Holland and Bitz, 2003; Screen and Simmonds, 2010], whereby warmer temperatures cause highly reflective snow and sea ice to melt, decreasing the Earth’s planetary albedo and thus inducing further warming. The lapse rate feedback may be equally important [Armour et al., 2013; Pitham and Mauritzen, 2013].

While the feedbacks triggered by greenhouse gas warming may dominate Arctic warming, short-lived aerosols in the atmosphere also are an important forcing agent in this region [e.g., Quinn et al., 2008; Koch et al., 2009; Shindell and Faluvegi, 2009; Serreze and Barry, 2011]. Further, climate changes triggered by aerosol trends will also be enhanced by local feedbacks and modified by circulation changes. Shindell and Faluvegi [2009] conducted sensitivity experiments using the GISS-ER climate model and suggested that decreasing concentrations of sulfate aerosols and increasing concentrations of BC have substantially contributed to Arctic warming over the last three decades. They also found Arctic temperature changes depend on the location of BC in the atmosphere. Increasing concentrations of BC in low/mid-latitude causes warming in the Arctic, while increasing BC in the Arctic itself causes cooling in the Arctic. In another climate model study, Sand et al. [2013] produced similar results to Shindell and Faluvegi [2009] and
attributed the response in Arctic surface air temperature mainly to the changes in atmospheric northward heat transport (NHT). Increasing atmospheric BC in the Arctic leads to a weakening of NHT and thus surface cooling; when BC is located in mid-latitudes, NHT is strengthened leading to Arctic surface warming. Arctic surface temperature is also found to be sensitive to the vertical distribution of BC in the CCSM4 [Flanner, 2013]: a layer of BC centered in the upper troposphere produces surface cooling, while a layer of BC in the lower troposphere causes weak surface warming. While there has been much focus on the role of BC in Arctic climate, we are interested in how Arctic climate has been affected by changes in both sulfate and BC aerosol distributions over the past three decades. As shown below trends in the two are not homogenous in either space or time.

Here, the response of Arctic and global surface air temperature to the trends of atmospheric sulfate and BC aerosols are examined using 20th century simulations from CCSM4. We performed single forcing experiments, in which only direct radiative forcing from sulfate aerosols or BC was included. This enables us to isolate the effects of sulfate and BC and their contributions to the effect of all aerosols on climate. We also examine the response to changes in sulfate, BC, and OC aerosols collectively. In addition to surface air temperature, we investigate the response of sea level pressure, sea ice coverage, cloud radiative forcing and atmospheric NHT to determine the mechanisms that cause the surface air temperature change in the Arctic during the period 1975-2005.

2. Model and Experiments
We use CCSM4 with fully-coupled atmosphere, ocean, land and sea ice components [Gent et al., 2011]. The atmosphere component is the Community Atmosphere Model 4 (CAM4) [Neale et al., 2011] with a horizontal resolution of 0.9°×1.25°. The atmospheric aerosol concentrations were derived from an off-line CAM-Chem [Lamarque et al., 2012] driven by observationally based estimates of aerosol emissions [Lamarque et al., 2010; 2011]. Anthropogenic emissions of sulfur species are an update of Smith et al. [2001; 2004]. The oceanic DMS emission was estimated from Kettle et al. [1999]. Volcanic and biomass sources of sulfur are excluded [Neale et al., 2011]. Emissions of BC and OC represent an update of Bond et al. (2007) and Junker and Liousse (2008). The Mie calculations for sulfate assume that it is comprised of ammonium sulfate with a log-normal size distribution [Neale et al., 2011]. BC and OC are assumed to age from hydrophobic to hydrophilic at an e-folding time of 1.2 days. The optics for BC and OC are identical to the optics for soot and water-soluble aerosols in the Optical Properties of Aerosols and Clouds (OPAC) data set [Hess et al., 1998; Neale et al., 2011]. Total aerosol optical depth comparisons with AERONET observations indicate a reasonably good simulation [Lamarque et al. 2010]. Shindell et al. [2013] used this atmospheric aerosol concentration dataset and demonstrated that it captures total aerosol optical depth trends of 1980-2000 well over the areas of high aerosol emissions (e.g., Europe, eastern North America and south and east Asia), compared with Advanced Very High Resolution Radiometer (AVHRR) observations. Compared with high-latitude aircraft campaigns, simulated BC concentrations, using the same emission sources as Lamarque et al. [2010] but with a previous version of CAM, are within the observed standard deviation [Koch et al., 2009]. CAM4 includes the direct and semi-direct effects
of aerosols, but the aerosol first indirect effect [Twomey et al. 1984] is not included. The same model and 20th century forcing datasets were used for the Coupled Model Intercomparison Project phase 5 (CMIP5) [Taylor et al., 2012] contributions from CCSM4.

Table 1 lists the details of our individual model experiments. The all-aerosol simulations were realized by varying the time- and space-evolving mass concentrations of sulfate, BC and OC aerosols simultaneously. All other forcings were kept fixed at 1850 levels, including surface depositions of BC on snow and sea ice. Three ensemble members of all-aerosol simulations were obtained from CMIP5 [Meehl et al., 2012]. The sulfate-only and BC-only single forcing experiments are analogous to the all-aerosol experiment except for only varying the mass concentrations of sulfate or BC, respectively. For each experiment, six branch runs were carried out from year 1920 and run to year 2005, making six ensemble members in each experiment. The restart files of year 1920 were obtained from 20th century total aerosol forcing only integrations with CCSM4 for CMIP5 [see Meehl et al., 2012]. In addition to six simulations of sulfate-only forcing, we also include surface air temperature fields from three runs of CMIP5 sulfate-only [Meehl et al., 2012]. The 1850 control run was also obtained from CMIP5.

Figure 1 shows linear trends in optical depths of sulfate and BC from 1975 to 2005. The results are ensemble means of six integrations. Since the 1970s, sulfate aerosol optical depth decreased significantly in Europe and North America. However, aerosol optical depth increased in Southeast of Asia, India and the Pacific Ocean region (Fig. 1a). Globally, there is a decrease in sulfate aerosol optical depth (not shown). Due to the emissions reductions in Europe and North America, sulfate optical depth decreased over
the Arctic, especially over the Eurasian Arctic (Fig. 1b). Sulfate aerosol is almost entirely scattering, with a single scattering albedo equal to one in the solar spectrum and a small fraction of absorption in the near-infrared spectrum. Therefore, it causes a net radiative cooling at the surface by scattering solar radiation back to space and letting less solar radiation reach the surface [IPCC, 2007, p160]. The mean surface shortwave radiative flux change in year 2000 due to sulfate is estimated at -0.84 W/m\(^2\) globally and -0.22 W/m\(^2\) for the Arctic in CCSM4. These estimates were performed by running CAM4 shortwave radiative transfer code twice with adjusted temperatures at all levels.

From 1975 to 2005 there were significant reductions in BC optical depth in Europe, but over the same period there were significant increases in optical depth in India and China (Fig. 1c). At the same time, emissions of fossil fuel BC declined in the U.S. [IPCC, 2007, p163]. Thus, the slight increase in BC optical depth over North America seen in Fig. 1c is possibly due to the downstream transport from Asia. In contrast, there was a decline in sulfate optical depth over North America and over all of the Arctic (Fig. 1a and 1b). While global mean sulfate emissions declined, BC emissions increased [Lamarque et al., 2010]. In the Arctic, BC optical depth shows a negative trend over the European Arctic but a positive trend over the rest of the Arctic (Fig. 1d). BC is a light-absorbing aerosol, so it absorbs solar radiation and heats the surrounding air [IPCC, 2007, p163]. The annual mean instantaneous flux change at the surface due to the direct effect of atmospheric BC is -0.46 W/m\(^2\) over the whole globe and -0.14 W/m\(^2\) over the Arctic in model year 2000 in CCSM4.

3. Results
3.1. Surface air temperature trends

Figure 2 depicts the time-evolving surface air temperature response to the change in all aerosols, sulfate-only and BC-only from 1920 to 2005. The all-aerosol, sulfate-only and BC-only runs have ensemble members of three, nine and six, respectively (Table 1). Globally, surface air temperature from all-aerosol forcing shows a significant negative trend of about 0.02 K/decade 1920-2005 and ~ 0.015 K/decade 1975-2005 in CCSM4. This result agrees with the study by Fyfe et al. [2013] that other forcing (primarily aerosol forcing) causes a modest Arctic cooling among CMIP5 models from 1970-2005.

Surface air temperature from sulfate-only forcing resembles that of the all-aerosol forcing. Therefore, the response of global surface air temperature to all aerosols is dominated by the direct forcing by sulfate. BC has a warming effect on global temperature (~ 0.1 K), but this is almost completely offset by the cooling influence from organic carbon, which is co-emitted with BC [e.g., Bond et al., 2013]. In the Arctic, surface air temperature is more variable, with no clear trends in Arctic-averaged surface air temperature from 1975 to 2005 in any of the three cases. The BC-only case indicates some warming from 1980 to 2000, followed by a cooling 2000-2010, but this is not statistically significant. However, there are statistically significant positive and negative temperature trends in different regions of the Arctic, which in this Arctic-wide average offset each other.

Geographic distributions of surface air temperature trends 1975-2000 are shown in Figure 3. We focus primarily on the changes in the Arctic, but show the global maps to aid interpretation of what is driving the Arctic changes. Direct radiative forcing by all aerosols produces a pronounced warming of 0.6 K/decade over the European Arctic, a
cooling of 0.6 K/decade over the Russian Arctic and a slight warming over the North American Arctic (Fig. 3a and b). The pattern of temperature trends in the all-aerosol case has elements in common with both sulfate-only and BC-only cases. In the sulfate-only experiment, there is a strong warming of 0.4 K/decade over the European and western Eurasian Arctic (Fig. 3c and d) where sulfate optical depth has declined (Fig. 1a). In and downstream of Siberia and in the western U.S. there is a significant cooling, which is contrary to what might be expected, given that sulfate concentrations and sulfate optical depth decreased across this region (Fig. 1). Such mismatch in the sign of temperature response and aerosol forcing is not unique to SO$_4$. In BC-only simulations (Fig. 3e and 3f), surface air temperatures warm 0.2 K/decade over the European Arctic, presumably in response to a reduction in BC concentrations aloft. There is a pronounced warming of roughly 0.4 K/decade over the Siberian and Alaskan Arctic and a strong cooling over the far north Atlantic despite small increases in BC optical depth in these regions. While there is some correspondence between the change in aerosol optical depths (Fig.1) and surface air temperatures (Fig. 3), the two are not perfectly correlated. This is because, in addition to the direct impact of aerosols on radiative fluxes, there may be other climate responses to the forcings which themselves affect surface air temperatures. In some cases, quite long-range connections are possible. Previous studies [Shindell and Faluvegi, 2009; Sand et al., 2013; Flanner, 2013] regarding the role of remote aerosols on Arctic temperatures show that an increase in BC concentrations at low latitudes causes a warming in the Arctic. Such remote influence is also shown by Teng et al. [2012] who found surface warming over the Siberian Arctic in response to increasing BC concentrations in Asia in CCSM4 (see their Figure 2). We emphasize that these results
are specific to CCSM4. As shown in Koch et al. [2009], general circulation models have
great variability in simulating BC aerosols. Even when using a fixed set of emissions,
different models will simulate different horizontal and vertical distributions of aerosols,
as well as differences in total atmospheric burden. Therefore it is of great interest for
future studies to examine the climate response of BC using different models.

3.2. Interpreting the climate responses to forcing

To further understand the temperature trends we analyzed sea level pressure, sea
ice coverage, radiative flux changes at top of the atmosphere (TOA) due to changes in
clouds and changes in NHT using the transient sulfate-only and BC-only runs as
described in section 2. NHT is calculated following equation (1) in Sand et al. [2012]. In
the sulfate-only experiment (Fig. 4a-d), there is a dipole in sea level pressure trends, i.e.
in the eastern North Atlantic versus in the European and west Eurasian Arctic. This draws
warmer air northward from lower latitudes, consistent with the strong warming trend
found in the European Arctic. The significant sea ice loss over the Barents Sea amplifies
the warming there. Surface cooling over most of the rest of the Arctic is consistent with
cold-air advection from Siberia, amplified by sea ice gain on the Siberian shelf and into
the Chukchi and Beaufort Sea. Net changes in cloud radiative fluxes at TOA, which are
the summation of shortwave and longwave fluxes, have a similar pattern to the changes
in sea ice coverage. These show a radiative cooling effect over the European Arctic and
warming over Siberia. NHT enhances the warming over the Eurasian Arctic and the
cooling over the Siberian and North American Arctic. These findings suggest that direct
surface radiative cooling from sulfate aerosols is the possible trigger for the surface
cooling while the dynamical response of atmospheric circulation, sea ice, and clouds work together to reinforce such temperature trends. Cloud changes have a weaker influence than sea ice and NHT changes in terms of magnitudes of trends. Again, we emphasize that the cloud changes produced here are only due to a thermodynamic response to the aerosol direct radiative forcing. If cloud microphysical effects were included in the model runs cloud changes might have a much more significant impact on Arctic climate.

The dynamical responses of the atmosphere and sea ice are similarly important in the BC-only experiment (Fig. 4e-h). Sea ice coverage decreases near the Barents Sea and the eastern Siberia shelf, where surface air temperature increases. NHT has strong positive trends over the Eurasian and North American Arctic. Therefore, the responses in both sea ice and NHT to aerosol direct radiative forcing reinforce the surface air temperature changes. Trends in net cloud radiative fluxes are weak and do not show a clear pattern.

4. Summary and discussion

We use fully coupled CCSM4 with CAM4 physics to investigate the Arctic and global climate response to the change in concentrations of all aerosols, sulfate aerosols only and BC only during the three decades from 1975 to 2005. Single-forcing transient simulations were performed in order to isolate the impacts of all aerosols, sulfate only and BC only. The surface air temperature response to all aerosols is dominated by changes in sulfate, while the effects of BC are apparently mostly offset by coincident trends in OC. Globally averaged, trends in all aerosols produce a cooling trend of 0.015
K/decade during the period of 1975-2005, with 0.02 K/decade cooling driven by changes in sulfate aerosols. Averaged across the whole Arctic, surface air temperature shows no significant trend. However, there are pronounced geographical variations in temperature trends. Over the European Arctic, aerosols induce about 0.6 K/decade warming, or about 1.8 K warming over the 30-year period from 1975 to 2005. This warming is triggered by a reduction in sulfate and BC concentrations over that region and is maintained by sea ice loss and a strengthening in NHT. Changes in sulfate concentrations account for about two thirds of the warming and BC for the remaining one-third. A recent study by Cowtan and Way [2014] shows that global temperature rise of the past 15 years has been largely underestimated due to data gaps especially in the Arctic. Based on the simulations presented here, we believe that sulfate aerosol trends have played an important role in the Arctic warming and potentially have prevented the warming “hiatus” seen in global temperature trends [Trenberth and Fasullo, 2013] from being seen in the Arctic temperature trends. Over the Siberian and North American Arctic, surface air temperature is likely influenced by changes of aerosol optical depth over Asia. An increase in sulfate optical depth over Asia induces a large cooling while an increase in BC optical depth over Asia causes a significant warming, consistent with Shindell and Faluvegi [2009]. Thus, full understanding of drivers of Arctic climate change require accounting for changes in all aerosol species – not just BC – and of the climate responses to both local and remote forcings.
Acknowledgements

This research used computing resources at the National Center for Atmospheric Research (NCAR). We acknowledge Julie Arblaster and Adrianne Middleton for providing the restart files and CMIP5 sulfate-only runs. We thank Loretta Mickley, Thomas Breider, Daniel Jacob, Mark Flanner and two anonymous reviewers for helpful discussions. This study was supported by the National Science Foundation grant ARC-1049002.
References


Koch, D., Schulz, M., Kinne, S., McNaughton, C., Spackman, J. R., Balkanski, Y., Bauer, S., Berntsen, T., Bond, T. C., Boucher, O., Chin, M., Clarke, A., De Luca, N.,
Dentener, F., Diehl, T., Dubovik, O., Easter, R., Fahey, D. W., Feichter, J.,

Fillmore, D., Freitag, S., Ghan, S., Ginoux, P., Gong, S., Horowitz, L., Iversen, T.,

Kirkevåg, A., Klimont, Z., Kondo, Y., Krol, M., Liu, X., Miller, R., Montanaro, V.,

Moteki, N., Myhre, G., Penner, J. E., Perlwitz, J., Pitari, G., Reddy, S., Sahu, L.,


Takemura, T., Textor, C., van Aardenne, J. A., and Zhao, Y.: Evaluation of black
carbon estimations in global aerosol models, *Atmos. Chem. Phys.*, 9, 9001-9026,

Lamarque, J.-F., Bond, T. C., Eyring, V., Granier, C., Heil, A., Klimont, Z., Lee, D.,

Liouesse, C., Mieville, A., Owen, B., Schultz, M. G., Shindell, D., Smith, S. J.,

Stehfest, E., Van Aardenne, J., Cooper, O. R., Kainuma, M., Mahowald, N.,

gridded anthropogenic and biomass burning emissions of reactive gases and aerosols:
methodology and application, *Atmos. Chem. Phys.*, 10, 7017-7039, doi:10.5194/acp-

Lamarque, J.-F., Kyle, G. P., Meinshausen, M., Riahi, K., Smith, S. J., van Vuuren, D. P.,

Conley, A., and Vitt, F.: Global and regional evolution of short-lived radiatively-
active gases and aerosols in the representative concentration pathways, *Climatic

Lamarque, J.-F., Emmons, L. K., Hess, P. G., Kinnison, D. E., Tilmes, S., Vitt, F.,

Heald, C. L., Holland, E. A., Lauritzen, P. H., Neu, J., Orlando, J. J., Rasch, P. J., and

Tyndall, G. K.: CAM-chem: description and evaluation of interactive atmospheric
chemistry in the Community Earth System Model, *Geosci. Model Dev.*, 5, 369-411,


**Table 1.** List of experiments, number of ensemble members, whether the run was obtained from CMIP5 or it is a new run conducted in this study, run period, trend analysis period and aerosols that vary in the run.

<table>
<thead>
<tr>
<th>Model experiment</th>
<th># of ensemble members</th>
<th>CMIP5 or new run</th>
<th>Run period</th>
<th>Trend analysis Period</th>
<th>Aerosols that vary</th>
</tr>
</thead>
<tbody>
<tr>
<td>All aerosols</td>
<td>3</td>
<td>CMIP5</td>
<td>1850 -- 2005</td>
<td>1975 -- 2005</td>
<td>SO$_4$, BC and OC</td>
</tr>
<tr>
<td>SO$_4$-only</td>
<td>3*</td>
<td>CMIP5</td>
<td>1850 -- 2005</td>
<td></td>
<td>SO$_4$</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>New run</td>
<td>1920 -- 2005</td>
<td>1975 -- 2005</td>
<td></td>
</tr>
<tr>
<td>BC-only</td>
<td>6</td>
<td>New run</td>
<td>1920 -- 2005</td>
<td>1975 -- 2005</td>
<td>BC</td>
</tr>
<tr>
<td>Control</td>
<td>1</td>
<td>CMIP5</td>
<td>1850 -- 2005</td>
<td>N/A</td>
<td>None</td>
</tr>
</tbody>
</table>

* only surface air temperature field was available.
Figure 1. Linear trends in optical depths per decade for sulfate (a, b) and black carbon (c, d) for the period 1975-2005, both globally and for the Arctic.
Figure 2. Time series of area-weighted, annual-mean surface air temperature (SAT) over the globe (a) and Arctic (b) for all aerosol forcing (black), sulfate-only forcing (blue) and BC-only forcing (red). Shading indicates one standard deviation of ensemble members. A 9-year running mean was applied. A year 1850 control run is shown in black dashed line.
Figure 3. Linear trends in surface air temperature for the period 1975-2005 over the globe and Arctic due to changes in all aerosols (a and b), sulfate only (c and d) and black carbon only (e and f), respectively. Gray dots indicate trends that are statistically significant at the 95% level (p < 0.05) based on an F-test.
Figure 4. Linear trends in sea level pressure (PSL), sea ice coverage, cloud net radiative fluxes at top of the atmosphere, and atmospheric northward heat transport (NHT) over the period 1975-2005 in the Arctic due to direct radiative forcing by sulfate only (a, b, c and d) and black carbon only (e, f, g and h), respectively.