Regional radiative impact of volcanic aerosol from the 2009 eruption of Redoubt volcano

C. L. Young, I. N. Sokolik, and J. Dufek

School of Earth and Atmospheric Sciences, Georgia Institute of Technology, Atlanta, Georgia, USA

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Correspondence to: I. N. Sokolik (isokolik@eas.gatech.edu)

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Abstract

High northern latitude eruptions have the potential to release volcanic aerosol into the Arctic environment, perturbing the Arctic's climate system. In this study, we present assessments of shortwave (SW), longwave (LW) and net direct aerosol radiative forcings (DARFs) and atmospheric heating/cooling rates caused by volcanic aerosol from the 2009 eruption of Redoubt Volcano by performing radiative transfer modeling constrained by NASA A-Train satellite data. The Ozone Monitoring Instrument (OMI), the Moderate Resolution Imaging Spectroradiometer (MODIS), and the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model for volcanic ash were used to characterize aerosol across the region. A representative range of aerosol optical depths (AODs) at 550 nm were obtained from MODIS, and the Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) was used to determine the altitude and thickness of the plumes. The optical properties of volcanic aerosol were calculated using a compositionally resolved microphysical model developed for both ash and sulfates. Two compositions of volcanic aerosol were considered in order to examine a fresh, ash rich plume and an older, ash poor plume. Optical models were incorporated into a modified version of the Santa Barbara Disort Atmospheric Radiative Transfer (SBDART) model. Radiative transfer calculations were made for a range of surface albedos and solar zenith angles (SZA) representative of the region. We find that the total DARF caused by a fresh, thin plume (~2.5–7 km) at an AOD (550 nm) range of 0.16–0.58 and SZA = 55° is –46 Wm
\(^{-2}\)AOD
\(^{-1}\) at the top of the atmosphere (TOA), 110 Wm
\(^{-2}\)AOD
\(^{-1}\) in the aerosol layer, and –150 Wm
\(^{-2}\)AOD
\(^{-1}\) at the surface over seawater. However, the total DARF for the same plume over snow and at the same SZA at TOA, in the layer, and at the surface is 170, 170, and –2 Wm
\(^{-2}\)AOD
\(^{-1}\), respectively. We also see that the total DARF when SZA = 75° for the same layer over snow is 35 Wm
\(^{-2}\)AOD
\(^{-1}\) at TOA, 64 Wm
\(^{-2}\)AOD
\(^{-1}\) in the layer, and 11 Wm
\(^{-2}\)AOD
\(^{-1}\) at the surface. These results indicate that environmental conditions, such as surface albedo and SZA, control the sign of the radiative forcing at TOA and at the surface.
and the magnitude of the forcing in the aerosol layer. An older plume over snow at SZA = 55° would have total DARFs of 25, 31, and −5 Wm⁻²AOD⁻¹ at TOA, in the layer, and at the surface, respectively. Our results demonstrate that plume aging can alter the magnitude of the radiative forcing. We also compare results for the thin plume to those for a thick plume (~3–20 km) with an AOD (550 nm) range of 1 to 3. The fresh, thin plume with AOD = 0.58, over seawater, and SZA = 55° will heat the atmosphere in the SW by ~2.5 Kday⁻¹ and cool the atmosphere in the LW by ~0.3 Kday⁻¹. The fresh, thick plume with AOD = 3 under the same environmental conditions will produce SW heating in the atmosphere by ~31 Kday⁻¹ and atmospheric LW cooling of ~6.7 Kday⁻¹. These calculations convey the importance of vertical plume structure in determining the magnitudes of the radiative effects. We compare our assessments with those reported for other aerosols typical to the Arctic environment (smoke from wildfires, Arctic haze, and dust) to demonstrate the importance of volcanic aerosols.

1 Introduction

Mt. Redoubt is a stratovolcano located in Alaska, USA. Rising to an elevation of 3108 m (10 197 ft), Redoubt is the second tallest volcano in the Aleutian Range. Redoubt has been active five times since 1900, with the most recent eruptions occurring in 1989 and 2009. The last period of explosive eruptions began on 23 March 2009 UTC and lasted through 4 April 2009 UTC. Eruptive plumes reached into the stratosphere on several occasions, reduced air quality and posed a threat to aviation (Carlile and Nelson, 2009; Coombs and Schaefer, personal communication). However, little attention has been given to the regional radiative impact of volcanic aerosols. Here, we use the 2009 eruption of Redoubt as an opportunity to assess a range of the radiative impacts that may be expected from high northern latitude eruptions.

The primary mechanism by which volcanic eruptions alter the Earth’s climate is through the formation of sulfate aerosols. Sulfate aerosol may persist in the stratosphere for up to three years (Clarisse et al., 2008), causing the stratosphere to warm
and the surface to cool (Robock, 2000). Ash expelled by volcanic eruptions can also interact with electromagnetic radiation, but the lifetime of fine ash in the stratosphere is only on the order of a few weeks (Niemeier et al., 2009). It is for this reason sulfates are the principal aerosol used to predict the global climate perturbation expected from a given eruption (Stenchikov et al., 1998; Ramachandran et al., 2000). However on regional scales, volcanic aerosols in the troposphere become important. In this case, sulfates and ash have similar residence times, on the order of days to weeks. Therefore, ash particles must be considered along with sulfate aerosol in estimating the regional radiative impact, as well as specific environmental conditions including surface albedo, solar zenith angle (SZA), regional weather, and other aerosol types present in the region.

Volcanic aerosols may be expected to have a particularly strong influence on the regional Arctic climate because of the sensitivity of the Arctic environment to radiative perturbations, as indicated by numerous studies focusing on other aerosol types, both natural (Myhre et al., 2007; Stone et al., 2007, 2008) and anthropogenic (Ritter et al., 2005; Quinn et al., 2007, 2008). The resulting forcing an aerosol will have on the Arctic environment is strongly controlled by the seasonality of many related factors, such as the amount of incoming solar radiation, surface albedo, precipitation, and transport of pollutants (Quinn et al., 2008; Sokolik et al., 2011). There is a large increase in tropospheric aerosols in the Arctic in late winter and early spring each year, which leads to the creation of Arctic haze (Quinn et al., 2008; Shaw, 1995; Sirois and Barrie, 1999). The haze-producing emissions are largely from Europe and Asia and consist of sulfate and nitrate aerosols, organic carbon, and soot (Klonecki et al., 2003; Stohl, 2006; Quinn et al., 2008). The importance of surface albedo has been investigated, and in general, an absorbing aerosol (i.e., smoke) over a highly reflective surface tends to reduce the amount of solar energy reaching the surface, while leading to warming in the aerosol layer (Stone et al., 2008; Shaw and Stamnes, 1980; Quinn et al., 2008). Given the high surface albedos common during Arctic springtime, even an aerosol with a moderate absorbing capability can have this effect (Quinn et al., 2008). However, the
effects of natural aerosols, such as volcanic aerosol, were not considered by Quinn et al. (2008) in making recommendations to mitigate Arctic warming due to anthropogenic aerosols.

The impact of volcanic aerosol on the Arctic climate is difficult to assess because of seasonal variations that dramatically alter the amount of incoming solar radiation and surface reflectivity. Additionally, volcanic aerosol may have an indirect effect on clouds (Lohmann et al., 2003; Sokolik et al., 2011). It is also possible that ash deposits may lower the albedo of ice and snow covered surfaces, which would be expected to perturb the Arctic's radiation budget just as soot deposits (Flanner et al., 2007) and dust deposits (Painter et al., 2010) have been shown to do.

Performing optical and radiative transfer calculations requires information on aerosol composition, size distributions, and types of aerosol present. Composition must be known in order to determine the spectral refractive index. Satellite data has been employed to determine size distributions of sulfates after eruptions (Stenchikov et al., 1998; Ramachandran et al., 2000). Particle counter measurements on balloons and aircraft have aided in calculating size distributions of stratospheric sulfuric acid aerosol (Jager and Deshler, 2002; Deshler et al., 1992, 1993; Deshler, 2008; Pueschel et al., 1994); however, these measurements have typically been made several weeks to months after the eruption, which does not help to determine the size distributions of “fresh” volcanic sulfate or ash. Therefore, in efforts to study the formation and aging process of volcanic sulfate, models have been developed to calculate how the size distribution evolves (Stier et al., 2005) and have been incorporated into general circulation models (Niemeier et al., 2009). For volcanic ash, there have been few airborne measurements of size distribution, mainly due to the dangers involved in making this measurement. Although size distributions have been measured on ashfall samples (Munoz et al., 2004; Martin et al., 2009; Riley et al., 2003; Scott and McGimsey, 1994), these values will not be representative of the actual size distribution during an eruption due to sorting which occurs during transport in the atmosphere (Rose and Durant, 2009). As the volcanic plume ages, the radiative properties of the plume are signif-
icantly altered by the changing size and composition of volcanic aerosol. Thus, it is important to examine how the aging of volcanic aerosol affects the radiative impact.

The goal of this study is to assess the range of regional radiative impact of volcanic aerosol from the 2009 eruption of Redoubt volcano using satellite data and a radiative transfer model. We examine both the longwave (LW) and shortwave (SW) components of the radiative forcing at the top of the atmosphere (TOA) and at the surface, and compute SW and LW atmospheric heating/cooling rates to determine how radiation is distributed in the atmosphere. This work employs a satellite multiple sensor approach to constrain the transport, areal extent, and characteristics of volcanic plumes. The NASA A-Train satellite constellation provides the unique capability of passive and active colocated satellite sensors to study volcanic eruptions (Carn et al., 2009; Thomas et al., 2009). The sensors MODIS and OMI facilitate in tracking volcanic aerosol plumes, and the CALIPSO lidar provides vertical profiles of aerosol. The organization of the paper is as follows. Section 2 presents the data used in this study and introduces a methodology to create a compositionally resolved microphysical model for volcanic aerosol. The satellite sensors chosen for this study and retrieval methods are described in further detail. The size distributions and refractive indices selected for optical calculations are discussed, as well as the satellite data products utilized in the radiative transfer model. Section 3 discusses results of direct aerosol radiative forcings (DARFs) and atmospheric heating/cooling rate profiles in terms of sensitivity to plume-specific and environmental input parameters for the Arctic region. Section 4 summarizes the important findings of this study and makes recommendations for future research.

2 Data and methodology

2.1 Satellite data

The CALIOP lidar flies aboard the CALIPSO (Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations) platform and measures backscatter at 532 and...
1064 nm. Past studies have successfully employed CALIPSO to investigate the vertical structure of volcanic plumes (Thomason and Pitts, 2008; Yuan et al., 2011). Vertical profiles are useful in determining the vertical placement of the plume in the atmosphere, as well as plume thicknesses and heights. Vertical profiles of aerosol were obtained from CALIPSO version 3.01 data. The sensors OMI and MODIS have greater spatial coverage and were used both to identify and track the plumes and to validate some of the products retrieved from CALIPSO.

The OMI instrument on the Aura spacecraft has an ultraviolet – visible spectral range of 0.27–0.50 µm and provides the UV aerosol index (AI) (Torres et al., 2007). The AI is a semi-quantitative indicator of the presence of UV-absorbing aerosols, such as volcanic ash, smoke, and dust. It is important to note that sulfate aerosol does not absorb UV radiation and will have negative values of AI, in contrast to UV-absorbing aerosols which have positive AI values.

The MODIS sensor flies on both the Terra and Aqua satellites. In this study, we employed true color MODIS images, aerosol optical depth (AOD) at 550 nm, and aerosol fine mode fraction products. The MODIS level 3 Collection 5 AODs were used to constrain the areal extent of volcanic plumes and determine a range of AODs for calculation of the radiative forcing. Fine mode fraction is the ratio of small mode optical depth to the total AOD (Remer et al., 2005). We used MODIS level 2 Collection 5 fine mode fraction to determine approximate proportions of fine and coarse particles present in a plume.

We performed analyses of satellite data for the entire period of explosive volcano activity between 22 March and 4 April to identify the cases with the best data availability and sensors’ synergy, which were utilized for integrative analysis. For each day in this time frame, CALIPSO overpasses were analyzed for a boxed region located 43° N to 69° N and 130° W to 160° W. A large collection region was needed to obtain a sufficient amount of data, because of CALIPSO’s narrow swath. Therefore, it was expected if CALIPSO were to detect a plume, it would be after transport had carried it some distance from the source. Only data that showed clear detection of an aerosol plume
with well-developed vertical structure in CALIPSO’s vertical cloud feature mask was selected. Aerosol index and \( \text{SO}_2 \) measurements from OMI and AOD from MODIS were then used to determine the likely source region of the detected aerosol plume and to provide evidence that it is volcanic in origin.

### 2.2 HYSPLIT model for volcanic ash transport

The HYSPLIT model for volcanic ash was used to aid in the interpretation of satellite data (Draxler et al., 2011; Rolph, 2011). The model computes the dispersion and transport of an evenly distributed vertical column of ash with a given size distribution placed directly over the volcano summit. The following parameters are needed to run the model: volcano latitude and longitude, volcano summit height, eruption date and time, eruption duration, ash column height, and ash reduction level.

### 2.3 Radiative transfer model

The 1D radiative transfer model used in this study is a modified version of the Santa Barbara Disort Atmospheric Radiative Transfer (SBDART) model, which solves the radiative transfer equation for a plane parallel atmosphere within Earth’s atmosphere and at the underlying surface (Ricchiazzi et al., 1998). We considered conditions representative of the Arctic environment in late winter/early spring. A subarctic winter atmospheric profile was used in radiative transfer calculations over seawater and snow surfaces with SZA ranging from 55° to 75°. In addition, the SBDART model was augmented by the optical properties of volcanic aerosols that were computed in this study.

### 2.4 Microphysical model

The optical properties of volcanic aerosol were determined through the development of a compositionally resolved microphysical model. Volcanic aerosol may be composed of ash, sulfate, hydrometeors, and ash particles coated with sulfate, water, and/or ice (Rose et al., 2004; Prata et al., 2007; Textor et al., 2006; Pueschel et al., 1994).
Here, we considered only ash and sulfate, both of which have distinct size distributions and spectral refractive indices. We incorporated representative size distributions and refractive indices for ash and sulfate using data reported in the literature, as well as a range of realistic ratios of fine to coarse modes and ash to sulfate deduced from satellite data (Kearney and Watson, 2009; Niemeier et al., 2009; Pollack et al., 1973; Spinetti et al., 2008; Stenchikov et al., 1998). A summary of the considered cases is presented in Table 1.

We used lognormal distributions for both ash and sulfate in the following form (Kearney and Watson, 2009)

$$n_c(r) = \frac{1}{\sqrt{2\pi\sigma}} e^{-\frac{(\ln(r)-\mu)^2}{2\sigma^2}}$$  \hspace{1cm} (1)

where $n_c$ is the particle number concentration, $r$ is the particle radius, and

$$\mu = (\ln R_{\text{eff}}) - 2.5\sigma^2$$  \hspace{1cm} (2)

where $R_{\text{eff}}$ is the effective radius.

We assume that sulfate occurs only in the fine mode, but ash can exist in both coarse and fine modes. To simulate fresh sulfate aerosol in a young volcanic plume, an effective radius of 0.5 µm and $\sigma$ of 0.5 were used (Kearney and Watson, 2009). The size distribution of sulfate was held constant in order to study the effects of ash fallout. For ash, $\sigma$ was set to 0.59 and held constant for all simulations, as in Niemeier et al. (2009). Effective radius for ash was allowed to vary from 5 µm to 1.5 µm, according to satellite retrievals of effective radii of ash in an aging plume reported by Spinetti et al. (2008). It should be noted that these previous studies do not include a truly “fine” ash portion (aerodynamic diameter < 2.5 µm), and therefore we only considered ash in the coarse mode.

Refractive indices for both SW (0.3–4 µm) and LW (4–20 µm) spectral regions for ash and sulfate were taken from the literature. The OPAC data were used for 70 % sulfate solution (Hess et al., 1998). For ash, refractive indices of andesite from
Pollack et al. (1973) were used. Sulfate solutions around 70% are commonly chosen for radiative transfer modeling of volcanic aerosol and are valid for most situations (e.g., Stenchikov et al., 1998). However, measurements of temperature and relative humidity in the Arctic suggest that the sulfate solution can be as low as 40–50% (Yue et al., 1994; Russell et al., 1996; Lambert et al., 1997). These measurements are based on temperature and relative humidity of ambient Arctic air. Temperatures and relative humidities in a fresh volcanic plume are much higher, as magmatic volatiles in arc volcanoes are water rich (Williams and McNutt, 2005). Therefore, we felt the 70% sulfate solution was more realistic for a fresh volcanic plume than the lower values occurring in Arctic environments.

Ratios of fine and coarse mode aerosols were constrained with MODIS fine mode fraction retrievals. For the purposes of our study, the fine mode fraction represents the proportion of sulfate present in an externally mixed aerosol mixture containing ash and sulfate. In order to realistically bracket the range of radiative forcing associated with ash fallout as a plume ages, an ash rich mixture (containing smaller fractions of sulfate and larger proportions of ash) was used as a proxy for a fresh plume, and a sulfate rich mixture (consisting of higher proportions of sulfate and a smaller ash fraction) was chosen to simulate a more evolved, aged plume.

2.5 Radiative calculations

The radiative transfer model computes the profiles of downward/upward radiative fluxes, including radiative fluxes at the surface and TOA. Net fluxes for both the SW and LW components at TOA and at the surface were computed by subtracting the upward flux from the downward flux:

\[ F_{\text{net}} = F_{\text{down}} - F_{\text{up}} \]  \hspace{1cm} (3)

The total net flux at TOA and at the surface were then calculated by adding the respective SW and LW net fluxes

\[ F_{\text{total net}} = F_{\text{net,SW}} + F_{\text{net,LW}} \]  \hspace{1cm} (4)
Atmospheric heating rates in units of Kday$^{-1}$ were calculated as follows:

\[
\frac{dT}{dt} = -\frac{1}{c_p} \rho \frac{dF_{net}}{dz}
\]  

(5)

where \(c_p\) is the specific heat at constant pressure (\(c_p = 1004.67\text{Jkg}^{-1}\text{K}^{-1}\)), and \(\rho\) is the air density of the aerosol layer. The change in net flux with respect to altitude is represented by \(dF_{net}/dz\). The change in \(F_{net}\) is the flux divergence, or the change in net flux between atmospheric layers of thickness \(z\).

The direct aerosol radiative forcing, DARF, is defined as the change in the net flux with respect to the change in AOD (550 nm)

\[
\text{DARF} = \frac{\Delta F_{net}}{\Delta \text{AOD}(550 \text{ nm})}
\]  

(6)

This calculation was done for net SW, net LW and total net fluxes at TOA and at the surface. To calculate DARF for an aerosol layer, the surface DARF was subtracted from the DARF at TOA

\[
\text{DARF}_{\text{atm}} = \text{DARF}_{\text{TOA}} - \text{DARF}_{\text{surface}}
\]  

(7)

For each modeling scenario, we computed DARF at TOA, at the surface, and within the atmosphere for SW, LW, and net forcing. The units of DARF are Wm$^{-2}$AOD$^{-1}$. The definition of DARF used in our study is similar to that of Stone et al. (2008) to facilitate the comparison with their results for other aerosol types. Many studies report direct radiative forcings in Wm$^{-2}$ at a particular AOD. To produce forcings comparable to these studies, the DARF is multiplied by AOD(500 nm).

3 Results and discussion

3.1 Satellite data analysis to constrain a radiative transfer model

The 17:24 UTC eruption on 26 March and the 13:58 and 14:16 UTC eruptions on 4 April were among the most violent eruptions, spewing ash well into the stratosphere.
MODIS (Fig. 1) and OMI AI (Fig. 2) help confirm the presence of ash in these plumes, as well as provide insight into the overall transport of aerosol. The plume detected on 4 April (Figs. 1b and 2b) appears to be moving southeast of the volcano, over the Gulf of Alaska. However, the plume from 26 March (Figs. 1a and 2a) appears to be transporting ash both north of the volcano over Alaska, and southeast of the volcano. Analysis of this plume with HYSPLIT indicates that atmospheric conditions on March 26 favor transport of ash to the southeast of the volcano at lower levels in the atmosphere, but changes to a northeasterly direction above approximately 11 km (Fig. 3). The transport of plumes on 26 March and 4 April over both seawater and snow surfaces illustrates the need for assessing the radiative forcing for varying surface albedos.

Extensive cloud and land-snow cover during much of March and early April made satellite retrievals challenging. After examining satellite data for the entire span of the explosive eruptions within the domain chosen, it was determined that the best data available was obtained for a plume detected on 2 April. The aerosol profiles from CALIPSO reveal that the plume is completely contained in the troposphere and extends from ~2.5–7 km (Fig. 4a). The plume thickness obtained from CALIPSO (Fig. 4a), together with total column AODs from MODIS, which range from 0.16 to 0.58 (Fig. 4b), suggest that this plume is moderately thin. Out of 22 CALIPSO overpasses in the selected domain, only the aerosol profile detected on 2 April clearly displays a well-defined vertical volcanic aerosol structure (Fig. 4a). In the absence of true color images on this day, the other sensors employed in this study, along with HYSPLIT, can confirm the volcanic origin of aerosol detected by CALIPSO. Both MODIS (Fig. 4b) and OMI (Fig. 4c) were able to detect volcanic aerosol, which appears to be traveling southeast of the volcano. On 2 April, there were intermittent low level emissions throughout the day, according to plume heights compiled by the Alaska Volcano Observatory (AVO) from ground based radar estimates (Carlile and Nelson, 2009). The HYSPLIT model predicts that ash from a low level eruption travels to the southeast of...
the volcano (Fig. 4d). Combined evidence from MODIS, OMI, and HYSPLIT confirms that the aerosol plume detected by CALIPSO is dominated by volcanic aerosols and that it has been carried away from its source.

Clues into the composition of the volcanic aerosol detected on 2 April can be gleaned from CALIPSO, MODIS, and OMI. For Mie calculations to determine optical properties of aerosol, it is essential to know if both ash and sulfate are present and proportions of ash to sulfate. Nonsphericity was detected in the 2 April plume using particle depolarization ratios from CALIPSO. Fragmented ash particles are expected to register nonspherical. The OMI AI image (Fig. 4c) also supports the presence of UV-absorbing aerosol, such as volcanic ash. The presence of ash or sulfate cannot be determined by MODIS AOD (550 nm), because it detects both types of volcanic aerosol. However, proportions of fine and coarse mode aerosol can be constrained by MODIS fine mode fraction (Fig. 5). In order to conservatively bracket the range of radiative forcing possible from this eruption, two volcanic aerosol compositions are considered, based on the highest and lowest values in the MODIS fine mode fraction (Fig. 5). The first composition was built to simulate a young, ash rich plume with an ash to sulfate ratio of 9 to 1, while the other composition represents an older, ash poor plume having an ash to sulfate ratio of 1 to 9.

However, the 2 April case study alone does not allow us to comprehensively assess the full range of radiative effects possible from this eruption. In order to better examine the role of vertical structure, we constructed a profile for a thick, optically opaque plume, forming directly over the volcano based on both the largest value of AOD and the maximum height for plumes detected throughout the course of the eruptions. The highest MODIS daily average AOD occurred on 27 March, and values of MODIS AOD chosen ranged from 1 to 3 (Fig. 6). The tallest plume height was produced by the 17:24 UTC eruption on 26 March, reaching ~20 km (Carlile and Nelson, 2009; M. Coombs and J. Schaefer, personal communication, 2011). The same aerosol compositions introduced for the plume detected on 2 April were used in this vertical structure in order to simulate plume aging.
3.2 Optical properties of volcanic aerosol

Based on satellite data, two externally mixed aerosol mixtures with varying proportions of ash and sulfate were selected to simulate plume aging. The normalized extinction coefficients, $K_e$, for the two aerosol mixtures and their sulfate and andesite constituents are shown in Fig. 7a as a function of wavelength. The normalized extinction coefficient is a measure of scattered and absorbed radiation by particles per unit concentration. Andesite absorbs and scatters both SW and LW radiation. The ash rich mixture is dominated by andesite ash. Sulfate scatters SW radiation and absorbs LW radiation, with negligible scattering in the LW because of small particle size. It is clear from Fig. 7a that the ash rich mixture resembles that of pure andesite, and the sulfate rich mixture more closely resembles that of sulfate. It is also apparent from Fig. 7a that the extinction coefficient per unit concentration is larger for both pure andesite and the ash dominated mixture at all wavelengths.

Figure 7b shows the single scattering albedo, $\omega_0$, for both aerosol compositions. The single scattering albedo is the fraction of photons that would be scattered by aerosols. In the SW, the ash rich mixture absorbs light more effectively than the sulfate rich one until about 2.5 µm. At longer wavelengths, the ash rich mixture scatters radiation better than the sulfate dominated composition.

3.3 The effect of volcanic aerosols on the Arctic radiation balance

The volcanic aerosol plume detected by CALIPSO, OMI, and MODIS on 2 April gives the most complete data set for a case study, because on no other occasions did all three sensors detect the same plume. Therefore, the 2 April plume provides us with a case study for a thin, optically diffuse plume detected after traveling some distance from the source volcano. This case also helps to determine the role environmental factors and plume aging has on the radiative impacts. We first consider the 2 April case and then examine the extent of the radiative effects by introducing a plume with a different vertical structure that was described in Sect. 3.1. Radiative forcings and
heating/cooling rates were computed for the two plumes as they age and for a range of SZAs and surface albedos. Comparisons between the two plume profiles aid in determining the sensitivity of the radiative impact to plume structure and opacity. Model input is summarized in Table 1.

3.3.1 Case study of volcanic plume detected on 2 April

The plume detected on 2 April 2009 was an optically thin volcanic aerosol plume, stretching from 2.5 to 7 km in altitude and having total column AOD at 550 nm ranging from 0.16 to 0.58. We use this case study to investigate the nature of volcanic aerosol and how plume aging and environmental parameters, such as surface albedo and SZA, influence the extent of the radiative impact. This is done for both SW and LW components. For a given set of environmental conditions, the effect of plume aging alone is to control the magnitudes of the radiative impact. Figure 8 displays the relationship between the net SW flux and AOD (550 nm) at the surface and TOA at SZA = 55°. Over both seawater and snow surfaces and for each AOD, the TOA net SW flux is greater for an ash rich plume than for a plume consisting primarily of sulfate (Fig. 8). As shown in Fig. 7, the reason for this is that the sulfate rich mixture scatters SW radiation more efficiently than does the volcanic aerosol enriched in ash. Conversely, over both types of surfaces for each AOD, the net flux at the surface is less for an ash rich plume than for a plume dominated by sulfate. Due to more SW radiation being scattered by a sulfate rich plume, a smaller downward SW flux at the surface is to be expected, producing a lesser net flux at the surface than for an ash rich plume.

Figure 9a illustrates how aerosol composition and AOD affect the distribution of radiation in the atmosphere. Ash rich mixtures lead to more heating in the layer and cooling at the surface, as do larger values of AOD. Compositions rich in ash warm more in the layer and cool more at the surface due to the ability of ash to absorb SW radiation. It is for this reason a volcanic layer containing ash will produce more positive SW DARF in the atmosphere, contributing to warming, and more negative SW DARF at the surface, contributing to cooling (Table 2). However, the extent of warming in the layer and
cooling at the surface in SW not only depends on AOD and the relative age of volcanic aerosol, but also on surface albedo and SZA.

The surface albedo of the Arctic region in late winter or early springtime ranges from optically black seawater surfaces to highly reflective snow and ice surfaces. For both compositions of volcanic aerosol, surface and TOA net SW fluxes over snow are less than surface and TOA net fluxes over seawater (Fig. 8). This is because the higher reflectivity of snow causes the upward flux both at the surface and TOA to be greater, leading to smaller net fluxes. The TOA net fluxes for both compositions of volcanic aerosol increase with increasing AOD over snow and decrease with increasing AOD over seawater. This generally leads to positive TOA DARF over snow and negative TOA DARF over seawater for both volcanic compositions (Table 2). This tendency is due to multiple scattering between the surface and the volcanic layer over snow when AOD is increased. The exception to this rule, however, is the case of a sulfate rich plume over seawater at low SZA. In this case, TOA DARF is negative because the effect of multiple scattering is drowned out by opposing effects of the composition and SZA.

The effects of multiple scattering over highly reflective surfaces on heating rates can be observed in Fig. 9b. A volcanic layer over snow will absorb more SW radiation due to multiple scattering, warming more in the atmosphere than the same layer over seawater. For a more absorbing aerosol, such as one comprised mostly of ash, this effect is even more pronounced.

In the Arctic, the changing of seasons brings dramatic shifts of incoming SW radiation. In the summer, the sun never sets, and in winter, the region is in perpetual darkness. Here, DARF is computed for the highest SZA for a day in mid March (55°) and a lower SZA chosen to represent a middle point between the highest and lowest daily sun angles (75°). A higher SZA leads to a larger downward SW flux at TOA. Figure 9c illustrates well that when more radiation is entering the system, the heating within the atmosphere is enhanced, especially for a volcanic layer containing larger proportions of absorbing ash. The effect of a lower SZA on SW DARF is to decrease
the extent of warming or cooling at TOA, at the surface and in the atmosphere (Table 2). Therefore, an eruption occurring very close to the winter solstice would have a negligible SW radiative effect.

However, both SW and LW radiation must be considered to determine the net radiative impact of a volcanic plume. In some cases, such as during boreal winter, the LW radiation may be more important than SW radiation. The LW forcing is sensitive to opacity of the volcanic aerosol layer and surface emissivity. From Fig. 7, it is apparent that ash absorbs more SW radiation. Both ash and sulfate absorb and scatter LW radiation, and ash is better overall at attenuating it (Fig. 7). This results in net LW TOA fluxes and net surface LW fluxes for sulfate that are more negative than for ash (Fig. 10). The effect of volcanic aerosol on the LW cooling rate profile is shown in Fig. 11. Overall, volcanic aerosol causes cooling in the atmosphere and warming at the surface, the extent to which is dependent upon the aerosol composition and AOD. Compositions rich in ash produce more pronounced cooling in the atmosphere and warming at the surface, because ash is a better absorber of SW radiation and emitter of LW radiation. This effect can also be observed in LW DARF calculations, which range from 19.5 to 6.3, 37.1 to 13.6, and \( -19.6 \) to \( -7.3 \) in units of \( \text{W m}^{-2} \text{AOD}^{-1} \) for ash and sulfate rich cases at the TOA, surface and within the atmosphere, respectively.

The overall radiative forcing associated with this volcanic plume at TOA, at the surface, and in the atmosphere is reported in Table 2. The total DARF is the sum of SW and LW DARFs for each case. In most cases, the SW radiation contributes more to the total DARF. However, there were three instances in which the LW radiation was significant enough to change the sign of the forcing either at the surface or in the atmosphere. These cases were all at a lower zenith angle and either consisted of ash rich plumes over snow or sulfate rich plumes over seawater and snow. The surface SW DARF for an ash rich plume over snow at a lower zenith angle produces less cooling than a similarly composed plume under other environmental conditions because of the multiple scattering effect between the aerosol layer and the snow and the reduction of solar radiation entering the system due to the lower zenith angle. In this case, LW
radiation heats the surface more than the SW radiation cools it, causing a net heating of the surface. A sulfate rich plume under low SZA will tend to warm the atmosphere less if placed over seawater or snow. This is due to both a reduction in incoming solar radiation and the inability of sulfate to absorb SW radiation. For these cases, LW radiation cools the atmosphere more than the SW radiation warms it, which leads to a net cooling of the atmosphere. These calculations demonstrate the importance of both SW and LW components of the forcing when determining the radiative effects of an aging volcanic plume, especially in the Arctic.

3.3.2 Examining the role of vertical structure

The simulated plume for an optically and physically thick eruption column was based on satellite data collected for the entire span of the eruption. A range of AODs from 1 to 3 was selected, and the plume thickness stretched from 3 km (the volcano summit) to 20 km altitude. We compared calculated direct aerosol radiative forcings and heating rates for an optically thick plume to those produced for the optically thin plume. In this section we demonstrate that, in general, the trends for surface type, SZA, and plume composition discussed above for the thin plume still hold for the thicker plume. However, we also show that the magnitude and, in a few cases, the sign of the forcings and the magnitudes of the heating/cooling rates are very much dependent on plume thickness and AOD.

Net SW fluxes at each AOD for the thick plume over both snow and seawater are shown in Fig. 12. Comparing these results to those for the thin plume in Fig. 8, it immediately becomes apparent that net SW fluxes for the thin plume over seawater, both at the surface and TOA, are considerably less than those for the thin plume at the ranges of AOD chosen for each plume. This is because more of the incoming SW radiation is scattered and absorbed in the thick plume case, resulting in less incoming SW radiation reaching the surface, and therefore less SW radiation reflected by the surface. Surface net SW fluxes over snow are greater for the thin plume than for the thick one for the same reasoning. However, net SW fluxes are greater for the thick...
plume at TOA. This is due to less outgoing radiation at TOA because of enhanced multiple scattering occurring between the snow surface and the more opaque plume. Another important comparison in net SW fluxes between plumes is the differences between the two volcanic aerosol compositions at each AOD. For the thicker plume, it seems that plume composition, and therefore plume age, can cause net SW fluxes at the surface and TOA to differ substantially more than for a thinner plume. Calculations of net SW flux must be more sensitive to composition of volcanic aerosol for a thicker plume.

Shortwave DARFs for the thick plume are calculated from SW net fluxes and presented in Table 3. Like SW DARF for the thin plume (Table 2), DARF is negative at the surface, indicating cooling, and positive in the atmosphere, indicating warming. However, DARF values for the thick plume cases are much smaller in magnitude than those for thin plumes. This is a result of the range of AOD over which net flux calculations were performed. The relationship between net flux and AOD can only be approximated as linear when AOD values are small. Because DARF is computed as the slope of this line, it will depend on the ranges of AOD chosen and must always be reported with this AOD range attached in order to be a meaningful quantity. For the thin plume, the range of AOD used to compute DARF is narrower and the AOD values are smaller. The net fluxes for the thin plume are more sensitive to changes in AOD at the range chosen, leading to larger DARF values. However, DARF is reported in units of Wm$^{-2}$AOD$^{-1}$, and therefore the thick plumes may still produce a larger forcing in Wm$^{-2}$ depending on the exact values of AOD present.

Shortwave heating rates for the thick plume also exhibit the same general dependencies on composition, surface albedo, and SZA as described for the thin plume, with warming in the atmosphere and cooling at the surface (Fig. 13). However, SW heating rates for the thick plume exhibit significant perturbations to clear sky conditions (Fig. 13a), which dwarf those due to the thin volcanic plume discussed above. Heating rates for the thick plume are particularly sensitive to aerosol composition (Fig. 13a), but have lesser sensitivities to surface reflectivity (Fig. 13b) and SZA (Fig. 13c). This find-
ing emphasizes how strongly plume age controls the way SW radiation is distributed within the aerosol layer and throughout the atmosphere.

The importance of plume age is magnified in the LW part of the spectrum. Given a fixed temperature profile, the LW radiative impacts are entirely dependent on composition and opacity of the aerosol layer, which change as the plume evolves. Net LW fluxes for the thick plume increase with increasing AOD (Fig. 14), similarly to net LW fluxes for the thin plume. However, the differences at TOA and at the surface between the two compositions of volcanic aerosol at each AOD are greater for the thick plume than for the thin plume, indicating that plume age plays a larger role in determining net LW fluxes for a thicker plume. Net LW fluxes also tend to be less negative, both at the surface and TOA, for the thick plume than for the thin plume of the same aerosol composition. A thick layer will emit more LW radiation, producing a net LW flux at the surface which is less negative than for a thin plume. The thick plume will also scatter and absorb more outgoing LW radiation coming from both the surface and within the lower layers of the plume, producing net LW fluxes at TOA that are typically less negative.

The slopes of the lines in Fig. 14 give the LW DARF for the thick plume cases, which range from 9.8 to 7.4, 8.0 to 5.3, and 1.8 to 2.1 Wm\(^{-2}\)AOD\(^{-1}\) for ash and sulfate rich cases at the TOA, surface, and within the atmosphere, respectively. Longwave DARF for the thick plume is less than those for the thin plume, because net LW fluxes for the thick plume are less sensitive to changes in AOD at the ranges of AOD chosen. Unlike the thin plume cases, DARF at TOA is greater than DARF at the surface for both compositions of volcanic aerosol, leading to positive DARF in the atmosphere for the thick plume. For the thick plume cases, the way in which LW radiation is distributed in the atmosphere is very different. Longwave cooling rates for each aerosol composition and AOD are displayed in Fig. 15. Emission of LW radiation in the upper layers causes cooling in the upper atmosphere, while the surface layers warm due to emission by middle to lower layers and the surface. Longwave emission from the upper and lower layers causes enhanced warming in middle layers.
The overall radiative effect of volcanic aerosol in a thick plume is given by the total DARF, and this is reported for each case in Table 3. For most cases, SW radiation was larger than LW. However, for four of the cases under lower sun conditions, LW radiation was large enough to change the sign of the total forcing. At the surface, a net warming was produced for both ash rich and sulfate rich compositions over snow. In these cases, surface SW cooling is reduced due to a reduction of solar energy entering the system and multiple scattering between the snow surface and the aerosol layer. TOA warming was produced for an ash rich plume over seawater and a sulfate rich plume over snow. In these cases, LW warming at TOA was enough to counteract SW cooling. For all of the thick plume cases, net warming in the atmosphere due to SW radiation is enhanced by positive values of LW DARF.

3.4 Comparing radiative impacts of major aerosol types in Arctic

The radiative effect of aerosol in a given environment depends upon aerosol-specific properties, such as optical characteristics and vertical loading. Volcanic aerosol is unique in that volcanoes are capable of producing huge loadings, forming plumes that can stretch from near surface level to well into the stratosphere. Therefore, it is obvious from the nature of volcanic eruptions that they may produce thicker plumes and may thereby drown out the signal from any other aerosols present. Figures 16 and 17 compares direct radiative forcings and heating rates for two volcanic aerosol compositions for the 2 April case to other aerosols typical to the Arctic region: mineral dust (Stone et al., 2007), wildfire smoke (Stone et al., 2008), and haze (Quinn et al., 2007; Ritter et al., 2005). While there are multiple factors which control the regional radiative impact, the most comparable surface albedo, AOD, SZA, and plume thickness for each study were selected in efforts to only assess the effects of composition (Table 4).

Figure 17a compares the surface SW forcing. The ash rich volcanic mixture attenuates more solar radiation per AOD than any other compositions, producing the most negative surface forcing. Mineral dust runs a close second to the ash rich mixture, indicating dust must have similar SW extinction coefficients. The ash rich mixture also
produces the most warming at TOA and within the atmosphere, which is because the ash rich mixture is the better absorber of SW radiation, as further illustrated by the SW heating rates in Fig. 17b. In many cases, the forcings and heating rates of the sulfate rich mixture and haze are very similar. This may be due to the haze presented in Quinn et al. (2007) consisting of large proportions of sulfate.

LW forcing for the two compositions of volcanic aerosol and haze (Ritter et al., 2005), and LW cooling rates for the two volcanic mixtures and dust (Stone et al., 2007) are compared in Fig. 17. The LW forcing for smoke is very small because of small particle sizes (Myhre et al., 2007) and is not shown here. LW forcings of dust and volcanic aerosols are important, although seldom calculated in radiative studies of Arctic aerosol. The ash rich mixture seems to dominate the LW radiation as well. Both SW and LW comparisons indicate that even a moderately thin layer of ash rich volcanic aerosol can be a main driver of the aerosol induced radiative impact in the Arctic region.

4 Conclusions

The 2009 eruption of Redoubt volcano provided a source of aerosol to the Arctic environment. The extent to which a given volcanic aerosol layer perturbs the radiation balance depends upon environmental factors, such as surface reflectivity and SZA, and plume specific factors, such as aerosol composition and vertical profile of AOD. We used a satellite integrated approach to investigate the role these factors play in determining the radiative impact of volcanic aerosol in the Arctic. In general, environmental factors can govern both the sign and magnitude of the DARF and heating/cooling rates. The ability of environmental conditions to change whether an aerosol layer will warm or cool at TOA and the surface is especially significant when considering the Arctic region, where surfaces vary from ocean to snow, and seasons dramatically shift the amount of incoming solar radiation. The deposition of volcanic ash onto ice and snow surfaces also has strong potential to greatly alter surface reflectivity and thereby cause
a perturbation that may last long after the aerosol plume has gone.

Plume specific factors influence the magnitudes of DARF and heating/cooling rates. The compositions, thicknesses and opacities of volcanic plumes vary greatly and are in many cases difficult factors to constrain. Due to the remote locations of many volcanoes and the dangers involved in making direct measurements, satellites are absolutely necessary to monitor the spatial and temporal development of volcanic plumes. However, eruption specific size distributions and ash to sulfate ratios are challenging to constrain, even with the help of satellite sensors. The development of multiphase models (Dufek et al., 2007; Neri et al., 2007) to study eruption dynamics may assist in creating better microphysical models for volcanic aerosol. Other plume related simplifications that are often made in radiative transfer modeling which need to be addressed include: the partitioning of volcanic aerosol types at different altitudes and the non-sphericity of ash.

Volcanic eruptions are high intensity events, capable of providing a huge, sporadic aerosol signal compared to predictable, lesser sources. However, volcanic eruptions typically have durations shorter than the season long occurrences of Arctic haze or wildfire smoke. Like other Arctic aerosols, volcanic aerosols from high northern latitude eruptions can cover large areas, as in the recent eruption of Eyjafjallajökull in Iceland, which disrupted air travel across Europe. According to the AVO, Alaskan volcanoes have had an average eruption frequency of 2 per year over the past 40 years, but the appearance of Arctic haze, wildfire smoke, and dust occur only yearly. Therefore, Alaskan volcanic eruptions are on average more frequent than events responsible for the emission of Arctic haze, wildfire smoke, and dust into the Arctic environment. Depending on the time of year, volcanic aerosol may be present along with other aerosol types. Following an eruption in Arctic springtime, dust, haze and volcanic aerosol can contribute to the overall aerosol signal. For eruptions in boreal summer, wildfire smoke will be present along with volcanic aerosols. In boreal winter, these other aerosol types will not be significant contributors, and the LW effects of volcanic aerosol will dominate the regional radiative impacts.
Our results demonstrate a potential for volcanic aerosol to provide a sizeable contribution to the radiative effects and even outshine other types of aerosol when significant proportions of volcanic ash are present, as in young volcanic plumes. Therefore, volcanic aerosol, although sporadically present, can have a significant radiative impact in the region. We recommend that volcanic aerosol be included in future assessments of the Arctic radiation budget. This will facilitate efforts to understand the radiative impacts of natural aerosols on the Arctic environment in order to create successful mitigation strategies for warming due to anthropogenic sources.

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Deshler, T., Johnson, B. J., and Rozier, W. R.: Balloonborne measurements of Pinatubo aerosol
Martin, R. S., Mather, T. A., Pyle, D. M., Power, M., Tsanev, V. I., Oppenheimer, C.,


Rolph, G. D.: Real-time environmental applications and display system (ready), online available at: http://ready.arl.noaa.gov, last access: 26 February 2010, 2011. 26698


Table 1. Input parameters for SBDART. Sulfate size distribution was held constant at an effective radius of 0.5 µm and σ of 0.5 (Kearney and Watson, 2009). For ash, σ is set to 0.59 and held constant for all simulations (Niemeier et al., 2009). Refractive indices for sulfate from OPAC were used for 70% sulfate solution. For ash, refractive indices of andesite from Pollack et al. (1973) were used.

<table>
<thead>
<tr>
<th>Model Parameter</th>
<th>Input</th>
<th>Source of input</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vertical placement and physical thickness of the aerosol layer</td>
<td>2.5 to 7 km&lt;sup&gt;a&lt;/sup&gt;</td>
<td>CALIPSO&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>3 to 20 km&lt;sup&gt;b&lt;/sup&gt;</td>
<td>AVO estimate based on radar&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>0.18, 0.38, 0.58&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Aerosol Optical Depth</td>
<td>1, 2, 3&lt;sup&gt;b&lt;/sup&gt;</td>
<td>MODIS&lt;sup&gt;a, b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Surface albedo</td>
<td>Seawater (0.07) and snow (0.8)</td>
<td>SBDART</td>
</tr>
<tr>
<td>Solar zenith angle</td>
<td>55°–75°</td>
<td>Realistic sun angles</td>
</tr>
<tr>
<td>Range of ash to sulfate ratio</td>
<td>9:1 to 1:9</td>
<td>MODIS fine mode fraction</td>
</tr>
<tr>
<td>( R_{\text{eff}} ) of ash</td>
<td>1.5–5 µm</td>
<td>Spinetti et al. (2008)</td>
</tr>
</tbody>
</table>

<sup>a</sup> indicates values used for the 2 April case study.

<sup>b</sup> indicates values used in sensitivity study. For both plumes, aerosol optical depth was distributed uniformly within the layer.
Table 2. Shortwave (SW) and total DARF for a thin (~2.5–7 km) plume in Wm$^{-2}$AOD$^{-1}$. DARF was calculated for a range of AOD between 0.16 and 0.58.

<table>
<thead>
<tr>
<th>Aerosol mixture</th>
<th>Spectrum</th>
<th>Solar zenith angle</th>
<th>Surface type</th>
<th>DARF TOA</th>
<th>DARF surface</th>
<th>DARF atm</th>
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</thead>
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<td>SW</td>
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<td>−194.4</td>
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<td>seawater</td>
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<td>−131.9</td>
<td>67.9</td>
</tr>
<tr>
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<td>snow</td>
<td>151.0</td>
<td>−41.4</td>
<td>192.4</td>
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<td>snow</td>
<td>55.3</td>
<td>−28.2</td>
<td>83.5</td>
</tr>
<tr>
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<td>−123.8</td>
<td>25.7</td>
</tr>
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<td>snow</td>
<td>18.7</td>
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<td>snow</td>
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<td>−18.3</td>
<td>7.0</td>
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<td>seawater</td>
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<td>−155.3</td>
<td>109.1</td>
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<td>−92.8</td>
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<td>snow</td>
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<td>172.8</td>
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<td>ash rich</td>
<td>Total</td>
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<td>snow</td>
<td>74.7</td>
<td>10.9</td>
<td>63.8</td>
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<td>−110.1</td>
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<td>−81.7</td>
<td>−2.6</td>
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<td>snow</td>
<td>25.0</td>
<td>−5.2</td>
<td>30.2</td>
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<tr>
<td>sulfate rich</td>
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<td>snow</td>
<td>−5.0</td>
<td>−4.6</td>
<td>−0.3</td>
</tr>
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Table 3. Shortwave (SW) and total DARF for a thick (~3–20 km) plume in Wm$^{-2}$AOD$^{-1}$. DARF was calculated for a range of AOD between 1 and 3.

<table>
<thead>
<tr>
<th>Aerosol mixture</th>
<th>Spectrum</th>
<th>Solar zenith angle</th>
<th>Surface type</th>
<th>DARF TOA</th>
<th>DARF surface</th>
<th>DARF atm</th>
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<tbody>
<tr>
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<td>snow</td>
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<td>-64.8</td>
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<td>seawater</td>
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<td>-18.1</td>
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<td>snow</td>
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<td>-7.2</td>
<td>25.6</td>
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<td>Total</td>
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<td>snow</td>
<td>5.1</td>
<td>0.7</td>
<td>4.3</td>
</tr>
</tbody>
</table>

* For this case, DARF at the top of the atmosphere was not linear at broader AOD ranges, so it was calculated in an AOD range of 1 to 1.5.
Table 4. Studies chosen for comparisons of direct radiative forcings of Arctic aerosols, along with aerosol types, surface albedo, AOD at 500 or 550 nm, $\omega_0$ (single scattering albedo) at 550 nm, solar zenith angle (SZA), vertical plume thickness and placement within the atmosphere that were used. Surface albedo and SZA are not applicable for Ritter et al. (2005), because this was a LW only study. Boxes filled with an X indicate information was not available.

<table>
<thead>
<tr>
<th>Study</th>
<th>Aerosol type</th>
<th>Albedo</th>
<th>AOD</th>
<th>$\omega_0$ (550 nm)</th>
<th>SZA</th>
<th>Placement (km)</th>
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</thead>
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<td>Quinn et al. (2007)</td>
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<td>0.92</td>
<td>0.12</td>
<td>0.94</td>
<td>62</td>
<td>X</td>
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<tr>
<td>Stone et al. (2007)</td>
<td>dust</td>
<td>0.825</td>
<td>0.16</td>
<td>0.94</td>
<td>62</td>
<td>2–4</td>
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<td>Stone et al. (2008)</td>
<td>smoke</td>
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<td>0.16</td>
<td>0.95</td>
<td>55</td>
<td>0.5–3.5</td>
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<td>Ritter et al. (2005)</td>
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<td>X</td>
<td>X</td>
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<td>This study</td>
<td>sulfate rich</td>
<td>0.8</td>
<td>0.18</td>
<td>0.98</td>
<td>55</td>
<td>2.5–7</td>
</tr>
<tr>
<td>This study</td>
<td>ash rich</td>
<td>0.8</td>
<td>0.18</td>
<td>0.89</td>
<td>55</td>
<td>2.5–7</td>
</tr>
<tr>
<td>This study</td>
<td>volcanic</td>
<td>0.8</td>
<td>0.18</td>
<td>0.89</td>
<td>55</td>
<td>2.5–7</td>
</tr>
</tbody>
</table>
Fig. 1. Aqua MODIS true color images on (a) 26 March at 22:40 UTC and (b) 2 April at 22:35 UTC. The aerosol plume is circled in red. The location of Redoubt Volcano is marked by a star.
Fig. 2. OMI AI for (a) 26 March and (b) 4 April. The location of Redoubt volcano is marked by a star.
Fig. 3. Transport of ash from the 26 March eruption as predicted by the HYSPLIT model for volcanic ash. The volcanic ash cloud is colored in pink. A duration of 1 h and a ash column height to flight level 650 were selected. The left column shows location of the ash plume 6 h after the eruption. The right column shows position of the plume 12 h after the eruption. No ash reduction was selected.
Fig. 4. (a) Vertical feature mask from CALIPSO between 11:31 UTC and 11:45 UTC on 2 April. Aerosols are colored in orange. The plume of interest is circled. (b) MODIS AOD at 550 nm daily average for 2 April. Redoubt Volcano is marked with a star. The path of CALIPSO is shown with a line. (c) OMI Al for 2 April. The location of Redoubt Volcano is marked with a star. The path of CALIPSO is shown with a line. (d) Transport of ash from the 2 April eruption as predicted by the HYSPLIT model for volcanic ash. The volcanic ash cloud is colored in pink. A duration of 12 h and a ash column height to flight level 150 were selected. The left column shows location of the ash plume 6 h after the eruption. The right column shows position of the plume 12 h after the eruption. No ash reduction was selected.
Fig. 5. Terra MODIS fine mode fraction AOD on 2 April at 20:55 UTC. Redoubt Volcano is marked by a star.
Fig. 6. MODIS AOD at 550 nm daily average for 27 March. These AOD values were the highest over the entire course of the explosive eruptions, due to aerosol produced by the very powerful eruption at 17:24 UTC on 26 March and the eruptions that followed on 27 March.
Fig. 7. (a) Concentration normalized extinction coefficients ($K_e$) versus wavelength for an ash rich mixture containing an ash to sulfate ratio of 9 to 1 and an effective radius for ash of 5 µm compared to andesite with an effective radius of 5 µm (right axis) and a sulfate rich mixture containing an ash to sulfate ratio of 1 to 9 and an effective radius for ash of 1.5 µm compared to sulfate with an effective radius of 1.5 µm (left axis). (b) Single scattering albedo ($\omega_0$) versus wavelength for an ash rich mixture containing an ash to sulfate ratio of 9 to 1 and an effective radius of 5 µm and a sulfate rich mixture containing an ash to sulfate ratio of 1 to 9 and an effective radius of 1.5 µm. Values for andesite and sulfate fall indistinguishably close to ash rich and sulfate rich mixture lines and were not included.
Fig. 8. The relationship between net shortwave flux and AOD (550 nm) at an AOD range of 0.18–0.58 for two aerosol mixtures at the top of the atmosphere (TOA) and surface over seawater (a) and snow (b). Solar zenith angle is held constant at 55°. The $R^2$ regressions for all lines are greater than 0.99.
Fig. 9. (a) The effect of aerosol composition and opacity of a thin plume (~2–7 km) on shortwave heating rate. The solar zenith angle $= 55^\circ$ and surface type is seawater for all cases in the plot. (b) The effect of aerosol composition of a thin plume (~2–7 km) and surface type on shortwave heating rate. The solar zenith angle $= 55^\circ$ and AOD = 0.58 for all cases in the plot. (c) The effect of aerosol composition of a thin plume (~2–7 km) and solar zenith angle (SZA) on shortwave atmospheric heating rate. The AOD = 0.58 and the surface type is seawater for all cases in the plot.
Fig. 10. The relationship between net longwave flux and AOD (550 nm) at an AOD range of 0.18–0.58 for two aerosol mixtures at the top of the atmosphere (TOA) and surface. The $R^2$ regressions for all lines are greater than 0.99.
Fig. 11. The effect of aerosol composition and opacity of a thin plume (~2–7 km) on longwave cooling rate.
Fig. 12. The relationship between net shortwave flux and AOD (550 nm) at an AOD range of 1–3 for two aerosol mixtures at the top of the atmosphere (TOA) and surface over seawater (a) and snow (b). Solar zenith angle is held constant at 55°. The $R^2$ regressions for all lines are greater than 0.95.
Fig. 13. (a) The effect of aerosol composition and opacity of a thick plume (~3–20 km) on shortwave heating rate. The solar zenith angle = 55° and surface type is seawater for all cases in the plot. (b) The effect of aerosol composition of a thick plume (~3–20 km) and surface type on shortwave heating rate. The solar zenith angle = 55° and AOD = 3 for all cases in the plot. (c) The effect of aerosol composition of a thick plume (~3–20 km) and solar zenith angle (SZA) on shortwave heating rate. The AOD = 3 and the surface type is seawater for all cases in the plot.
Fig. 14. The relationship between net longwave flux and AOD (550 nm) at an AOD range of 1–3 for two aerosol mixtures at the top of the atmosphere (TOA) and surface. The $R^2$ regressions for all lines are greater than 0.96.
**Fig. 15.** The effect of aerosol composition and opacity of a thick plume (~3–20 km) on longwave cooling rate.
Fig. 16. (a) Shortwave direct radiative forcings and (b) heating rates for haze (Quinn et al., 2007), dust (Stone et al., 2007), smoke (Stone et al., 2008), and ash rich and sulfate rich volcanic aerosols (this study). Values of forcing for dust were only available at the surface.
Fig. 17. (a) Longwave direct radiative forcings for haze (Ritter et al., 2005) and ash rich and sulfate rich volcanic aerosols (this study). (b) Longwave cooling rates for dust (Stone et al., 2007) and ash rich and sulfate rich volcanic aerosols (this study).