

Physical and Optical Properties of Aged Biomass Burning Aerosol from Wildfires in Siberia and the Western US at the Mt. Bachelor Observatory

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10 **Abstract.** The summer of 2015 was an extreme forest fire year in the Pacific Northwest. Our
sample site at the Mt. Bachelor Observatory (MBO, 2.7 km a.s.l.) in central Oregon observed
biomass burning (BB) events more than 50% of the time during August. In this paper we
characterize the aerosol physical and optical properties of 19 aged BB events during August
2015. Six of the nineteen events were influenced by Siberian fires originating near Lake Baikal
15 that were transported to MBO over 4-10 days. The remainder of the events resulted from
wildfires in Northern California and Southwestern Oregon with transport times to MBO ranging
from 3-35 hours. Fine particulate matter (PM₁), carbon monoxide (CO), aerosol light scattering
coefficients (σ_{scat}), aerosol light absorption coefficients (σ_{abs}), and aerosol number size
distributions were measured throughout the campaign. We found that the Siberian events had a
20 significantly higher $\Delta\sigma_{\text{abs}}/\Delta\text{CO}$ enhancement ratio, higher mass absorption efficiency (MAE;
 $\Delta\sigma_{\text{abs}}/\Delta\text{PM}_1$), lower single scattering albedo (ω), and lower Absorption Ångström exponent
(AAE) when compared with the regional events. We suggest that the observed Siberian events
represent that portion of the plume that has hotter flaming fire conditions and thus enabled strong
pyro-convective lofting and long-range transport to MBO. The Siberian events observed at MBO
25 therefore represent a selected portion of the original plume that would then have preferentially
higher black carbon emissions and thus an enhancement in absorption. The lower AAE values in
the Siberian events compared to regional events indicate a lack of brown carbon (BrC)
production by the Siberian fires or a loss of BrC during transport. We found that mass scattering
efficiencies (MSE) for the BB events ranged from 2.50-4.76 m² g⁻¹. We measured aerosol size
30 distributions with a scanning mobility particle sizer (SMPS). Number size distributions ranged
from unimodal to bimodal and had geometric mean diameters (D_{pm}) ranging from 138-229 nm
and geometric standard deviations (σ_g) ranging from 1.53-1.89. We found MSEs for BB events

to be positively correlated with the geometric mean of the aerosol size distributions ($R^2 = 0.73$), which agrees with Mie Theory. We did not find any dependence on event size distribution to transport time or fire source location.

1. Introduction

Biomass burning (BB) is a major source of aerosol in the atmosphere [Andreae and Merlet, 2001; Bond et al., 2004]. BB particles are predominantly organic carbon (OC) and black carbon (BC), with some inorganic material [Reid et al., 2005b; Vakkari et al., 2014]. These particles can significantly impact the Earth's radiative balance and climate through direct and indirect aerosol effects. The direct effects on radiative forcing are due to the light scattering and absorption of the aerosol [Boucher et al., 2013; Haywood and Boucher, 2000], and the indirect effects are caused by particles acting as cloud condensation nuclei (CCN) which affects cloud albedo [Pierce et al., 2007; Spracklen et al., 2011]. According to the IPCC 2013 report the largest uncertainty in determining global radiative forcing comes from quantifying the direct and indirect effects of aerosols [Boucher et al., 2013]. Biomass burning is a major contributor to global aerosol burden and it has been predicted that these emissions are likely to increase due to climate change, particularly in the boreal forests of North America and Russia [Flannigan et al., 2009; Stocks et al., 1998] and in the western US [Y Liu et al., 2014b; Westerling et al., 2006]. This makes the proper characterization of aged BB emissions even more important.

Currently there are few field measurements of well-aged BB emissions. Our knowledge of BB aerosol primarily comes from laboratory experiments and near-field measurements taken within a few hours of a wildfire [May et al., 2015; May et al., 2014; Okoshi et al., 2014; Vakkari et al., 2014; Yokelson et al., 2013b; Yokelson et al., 2009]. Holder et al. [2016] showed that laboratory measurements of aerosol optical properties do not accurately reproduce field measurements. Freshly emitted BB particles are small in diameter (30-100 nm) [Hosseini et al., 2010; Levin et al., 2010]. As the plume ages, the aerosol undergoes rapid chemical and physical changes on the time scale of minutes to hours [Reid et al., 2005a; Reid et al., 2005b; Vakkari et al., 2014]. The change in particle size is due to coagulation and the condensation of organic material onto the existing particles [Reid et al., 2005b; Seinfeld and Pandis, 2006]. The coagulation rate can be very high in fresh BB plumes since this is equivalent to the square of

particle number concentration. This process increases the size of the particles while decreasing the number concentration. Condensation of secondary organic aerosol (SOA) onto particles in BB plumes also increases particle size. The condensation of SOA is counterbalanced by the loss of primary organic aerosol (POA), which can evaporate during plume dilution [May *et al.*, 2015; May *et al.*, 2013]. The net condensation/evaporation effect is highly variable. Some studies have observed an increase in mass with plume age due to SOA production [Briggs *et al.*, 2016; Hobbs, 2003; Vakkari *et al.*, 2014; Yokelson *et al.*, 2009], while others have observed limited SOA formation [Akagi *et al.*, 2012; Jolleys *et al.*, 2015]. All of these uncertainties in the aging process of biomass burning underscores the importance of characterizing the physical and optical properties of well-aged biomass burning aerosol.

In this study we analyze 19 aged BB events observed in the summer of 2015 at Mt. Bachelor in Oregon. The BB events consisted of Regional events (fires in Northern California and Southwestern Oregon; transported 3-35 hours) and Siberian fire events (fires around Lake Baikal; transported 4-10 days). We investigated the aerosol optical and physical properties of these events and explored their variation with source location and transport time. This study addresses the following questions:

- What are the differences in the optical properties of regional and Siberian BB events observed at MBO?
- What is the range of mass scattering efficiencies for BB events and what explains their variability?
- What is the range in aerosol size distributions of BB events at MBO and how does this vary with plume age?

2. Methods

2.1. Mt. Bachelor Observatory

The Mt. Bachelor Observatory (MBO) is a mountaintop site that has been in operation since 2004 [Jaffe *et al.*, 2005]. It is located at the summit of Mt. Bachelor in central Oregon, US (43.98° N, 121.69° W, 2,764 m a.s.l.). A suite of measurements (including carbon monoxide (CO), ozone (O₃), aerosol scattering coefficients, and more) have been made continuously at the summit site. Previous studies have observed BB plumes in the free troposphere from regional

and distant sources in the spring, summer, and fall [Baylon *et al.*, 2015; Briggs *et al.*, 2016; Collier *et al.*, 2016; Timonen *et al.*, 2014; Weiss-Penzias *et al.*, 2007; Wigder *et al.*, 2013], and long-range transport of Asian pollution in the spring [Ambrose *et al.*, 2011; Fischer *et al.*, 2010a; Fischer *et al.*, 2010b; Gratz *et al.*, 2014; Jaffe *et al.*, 2005; Reidmiller *et al.*, 2010; Timonen *et al.*, 2014; Timonen *et al.*, 2013; Weiss-Penzias *et al.*, 2006]. During the summer of 2015 an intensive field campaign was performed at MBO to measure aerosol physical and optical properties of wildfire emissions.

2.2. CO, CO₂, and Meteorological Data

CO and CO₂ measurements were made using a Picarro G2302 Cavity Ring-Down Spectrometer. Calibrations were performed every 8 hours using three different NOAA calibration gas standards, which are referenced to the World Meteorological Organization's (WMO) mole fraction calibration scale [Gratz *et al.*, 2014]. Total uncertainty based on the precision of calibrations over the campaign was 3%. Basic meteorology measurements, such as temperature, humidity and wind speed were also measured continuously [Ambrose *et al.*, 2011].

2.3. Aerosol Instruments

We measured dry (relative humidity (RH) less than 35%) aerosol scattering and absorption coefficients, aerosol number size distribution, and particle mass during the 2015 summer campaign in 5 minute averages. An inline 1 μm impactor was located prior to the aerosol instruments. The aerosol instruments were located in a temperature-controlled room within the summit building, situated approximately 15 m below the inlet. The aerosol sample line was situated such that the last 2.5 m was located within a space that was temperature controlled at $20 \pm 3^\circ\text{C}$, typically 10°C – 20°C warmer than ambient. Relative humidity of the sampled air was less than 35% throughout the campaign. The temperature increase from going outside into the heated building reduced the RH of the sample. RH was measured in the sample airstream by the nephelometer and SMPS. The average RH during the campaign measured by the nephelometer and SMPS was 22.1% and 22.6%, respectively. Ninety-five percent of the 5 minute averaged samples had an RH less than 30%.

We measured multi-wavelength aerosol light scattering coefficients (σ_{scat}) using an integrating nephelometer (model 3563, TSI Inc., Shoreview, MN) at wavelengths 450, 550, and

120 700 nm. During the 2015 campaign the TSI nephelometer was periodically switched to measure both particle free air and CO₂. The measured values were corrected for offset and calibration drift in addition to angular nonidealities [Anderson and Ogren, 1998]. The filtered air and CO₂ were measured approximately every two weeks [Anderson and Ogren, 1998]. The data reduction and uncertainty analysis that we followed for the scattering data are outlined by Anderson and
 125 Ogren [1998]. Sources of uncertainties associated with the nephelometer include photon counting noise, zeroing and calibration, and the correction for angular nonidealities. Combined these uncertainties yielded total uncertainties of ~15% during BB events.

We measured aerosol light absorption coefficients (σ_{abs}) with a 3 λ tricolor absorption photometer (TAP, Brechtel Inc., Hayward, CA) at wavelengths 467, 528, and 660 nm.

130 Throughout the paper σ_{scat} and σ_{abs} values represent measurements taken at 550 nm and 528 nm, respectively. The TAP is a new instrument that uses the same operating principle as the Particle Soot Absorption Photometer (PSAP) and the same filters (47 mm PALL E70-2075W). Unlike the PSAP, the TAP rotates through 8 filter spots per individual filter along with two reference spots. During deployment at MBO, the TAP was set to rotate to the next filter spot when a filter
 135 spot's transmission reached 50%. The absorption coefficients were corrected using the filter loading and aerosol scattering correction factors derived for the 3 λ PSAP by Virkkula [2010]. Uncertainty calculations were based on those used in a previous study at MBO for measurements with a 3 λ PSAP [Fischer et al., 2010a]. Sources of uncertainty include noise, instrument drift, errors in the loading function, the correction for the scattering artifact, and uncertainty in the
 140 flow and spot size corrections [Anderson et al., 1999; Bond et al., 1999; Virkkula et al., 2005]. Combining these uncertainties yielded total uncertainties of ~25-40% during BB events.

Single scattering albedo (ω) for each event was calculated as the Reduced Major Axis (RMA) regression of scattering and total extinction (scattering + absorption) coefficient at 528 nm. To adjust the σ_{scat} value from 550 nm to 528 nm, a power law relationship was assumed
 145 between scattering and wavelength. The 450-550 nm pair was used to adjust the 550 nm σ_{scat} measurement to 528 nm using equation (1):

$$\sigma_{\text{scat}}^{528} = \sigma_{\text{scat}}^{550} * \left(\frac{\lambda_{550}}{\lambda_{528}} \right)^{SAE_{450,550}} \quad (1)$$

where λ is wavelength and SAE is the scattering Ångström exponent calculated with the two wavelengths specified. The SAE values were calculated for each 5-minute interval using the

150 scattering coefficients measured at 450 nm and 550 nm. Mean SAE values for the BB plumes
ranged from 1.61 to 2.15. Uncertainties for ω were calculated the same as the enhancement
ratios, which is discussed in Section 2.4.

Absorption Ångström exponent (AAE) values were calculated for the absorption
coefficient pair of 467 and 660 nm using equation (2):

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$$AAE = -\log(\sigma_{abs}^{467}/\sigma_{abs}^{660})/\log(467/660) \quad (2)$$

Uncertainties for AAE values were calculated by propagating the uncertainties from the
measurements used to calculate AAE using addition in quadrature [Fischer *et al.*, 2010a].

We measured 5-minute averaged dry aerosol number size distribution with a TSI 3938
Scanning Mobility Particle Sizer (SMPS). The SMPS system consisted of a TSI 3082
160 electrostatic classifier with a TSI 3081 Differential Mobility Analyzer (DMA) and a TSI 3787
water-based condensation particle counter. A total of 107 bins were used to measure a diameter
range from 14.1-637.8 nm. A sheath to aerosol flow ratio of 10:1 was used for the DMA. A
multiple charge correction and diffusion loss correction were applied to the SMPS particle
number concentration data using the TSI software. An additional diffusion correction for the
165 inlet tube (15 m, 12 LPM) was applied assuming a laminar flow [Hinds, 1999]. Prior to
deployment we confirmed the sizing accuracy of the SMPS using polystyrene latex spheres
(PSL).

We measured dry particle mass under 1 μm (PM1) with an Optical Particle Counter
(OPC, model 1.109, Grimm Technologies, Douglasville, GA). This is a U.S. EPA equivalent
170 method for measuring PM2.5 mass concentration. The OPC was factory calibrated prior to
deployment.

All particle measurements (σ_{scat} , σ_{abs} , PM1, number size distribution) were corrected to
standard temperature and pressure (STP; T = 273.15, P = 101.325 kPa).

2.4. Enhancement Ratio Calculations

175 Enhancement ratios ($\Delta Y/\Delta X$) were calculated from the slope of the RMA regression of Y plotted
against X. Briggs *et al.* [2016] calculated enhancement ratios (ERs) of BB plumes using three
different methods, one method using the RMA slope of the linear correlation of two species, and
two others calculating absolute enhancement above local background using two different
definitions of background. All three methods produced similar results for $\Delta\sigma_{\text{scat}}/\Delta\text{CO}$,

180 $\Delta\text{NO}_y/\Delta\text{CO}$, and $\text{PAN}/\Delta\text{CO}$, but differing results for $\Delta\text{O}_3/\Delta\text{CO}$. In our study we used the RMA regression method for calculating ERs of $\Delta\sigma_{\text{scat}}/\Delta\text{CO}$ and $\Delta\sigma_{\text{abs}}/\Delta\text{CO}$.

Mass scattering and mass absorption efficiencies (MSE and MAE) were calculated as the enhancement ratios of $\Delta\sigma_{\text{scat}}/\Delta\text{PM1}$ and $\Delta\sigma_{\text{abs}}/\Delta\text{PM1}$, respectively, at 550 nm for σ_{scat} and 528 nm for σ_{abs} . As previously mentioned, ω was calculated as the RMA regression of scattering and
185 total extinction (scattering + absorption). In all cases the enhancements (Δ) are large compared to background, thus avoiding the problems described by *Briggs et al.* [2016] for small enhancements above background.

We determined the uncertainties for the enhancement ratio calculations from the uncertainties in the extensive properties used in calculating the enhancement ratios and the
190 uncertainty of the RMA regression using addition in quadrature. For example, the uncertainty in $\Delta X/\Delta Y$ was calculated by adding in quadrature the uncertainty in the RMA regression, the uncertainty in the X measurement, and the uncertainty in the Y measurement.

We present both precision uncertainty and total uncertainty as described by [*Anderson et al.*, 1999] for all values derived from optical measurements. Precision uncertainty includes
195 uncertainty associated with noise and instrument drift. This is best used when comparing measurements collected using the same instruments and protocols. It is the appropriate uncertainty to consider when comparing individual BB events seen at MBO in this study. Total uncertainty includes precision uncertainty, the uncertainty associated with the corrections we applied to the data, and the uncertainty associated with the calibration method. This is the
200 appropriate uncertainty to consider when comparing the measurements presented in this study with data collected using other measurement methods.

2.5. Biomass Burning Event Identification

We identified BB events as time periods during which 5-min ambient aerosol scattering coefficients $\sigma_{\text{scat}} > 20 \text{ Mm}^{-1}$ for at least one hour, 5-min CO $> 150 \text{ ppbv}$ for at least one hour, and
205 there was a strong correlation ($R^2 > 0.80$) between σ_{scat} and CO. To determine fire locations we calculated back-trajectories using the National Oceanic and Atmospheric Administration Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model, version 4 [*Draxler*, 1999; *Draxler and Hess*, 1997; 1998; *Stein et al.*, 2015]. We used the Global Data Assimilation System (GDAS) $1^\circ \times 1^\circ$ gridded meteorological data from the National Oceanographic and Atmospheric

210 Administration's Air Resources Laboratory (NOAA-ARL). Within GDAS, the grid containing
MBO is located at ~1500 m amgl (above model ground level) so back-trajectory starting heights
of 1300, 1500, and 1700 m amgl were chosen [Ambrose *et al.*, 2011]. We identified fire locations
using Moderate Resolution Imaging Spectroradiometer (MODIS) satellite-derived active fire
counts [Justice *et al.*, 2002], and Fire INventory from NCAR (FINN) data [Wiedinmyer *et al.*,
215 2011]. Similar criteria for identifying BB events has been used by Baylon *et al.* [2015] and
Wigder *et al.* [2013] from data collected at MBO.

3. Results and Discussion

3.1. Identified BB Events and Fire Source Identification

The summer of 2015 was a very active fire season in the Pacific Northwest. During the month of
220 August 2015, 51% of the 5-minute averages met the criteria for a BB event, having $\sigma_{\text{scat}} > 20$
 Mm^{-1} and $\text{CO} > 150$ ppbv, including several multi-day periods (Figure 1). We split these multi-
day events up if discernable plumes within the event could be identified. Altogether we
identified 19 events, ranging from 1.5-45 hours in duration. We use the term *event*, not *plume*,
because of the long duration of some of the events and the fact that most BB events observed in
225 2015 were influenced by emissions from multiple fires.

Two large multi-day events of regional BB smoke from fires in Northern California and
Southwestern Oregon dominated the sampling period (dotted box in Figure 1). Transport time
from these regional fires to MBO, estimated from the back-trajectories, ranged from 3-35 hours.
In between these two large regional BB events there was a time period that was influenced by
230 Siberian wildfires (solid box in Figure 1). During August there were intense forest fires around
Lake Baikal in Siberia, peaking on 8/8/2015 with a total fire area of 681 km^2 , and an estimated
CO and BC emissions of 3.22×10^8 and 1.33×10^6 kg/day, respectively (*Fire INventory from*
NCAR (FINN) data) [Wiedinmyer *et al.*, 2011]. Transport times from northeastern Asia to MBO
during these events ranged from 4-10 days. NASA MODIS aqua and terra images show the
235 eastward transport of smoke from the Lake Baikal fires during this time period
[<https://worldview.earthdata.nasa.gov/>, 2016]. We used V3.30 aerosol classification products
from the Cloud-Aerosol Lidar with Orthogonal Polarization (CALIOP) instrument on the Cloud-
Aerosol Lidar Infrared Pathfinder Satellite Observation (CALIPSO) satellite to confirm the

transport of plumes of smoke from the Siberian fires to North America [<http://www-calipso.larc.nasa.gov/>, 2016; Winker et al., 2010; Winker et al., 2009]. Aerosol plumes are identified as one of six types: dust, polluted continental, polluted dust, smoke (biomass burning), clean continental or clean marine aerosols [Omar et al., 2009].

3.2. Overview of Summer 2015 BB Events

Table 1 provides an overview of the 19 BB events from MBO during the summer of 2015. We calculated water vapor enhancement (ΔWV) to indicate the origin of the event air mass. Positive ΔWV suggest the air mass ascended from the boundary layer (BL) to MBO, while near zero or negative values mean the air mass is relatively dry and likely descended or arrived from the free troposphere (FT) [Baylon et al., 2015; Wigder et al., 2013]. All of the regional BB events have ΔWV values ≥ 1.00 g/kg, while all of the Siberian-influenced events have ΔWV values near zero or negative. The precision and total uncertainties for all of the parameters derived from optical measurements are provided for these events in Table S1.

We found the $\Delta\sigma_{\text{scat}}/\Delta\text{CO}$ (σ_{scat} at STP) enhancement ratio to range from 0.48-1.29 Mm^{-1} ppbv^{-1} , with the majority of events being between 0.8 and 1.25 Mm^{-1} ppbv^{-1} . We found $\Delta\text{PM1}/\Delta\text{CO}$ (PM1 at STP) to range from 0.18-0.43 $\mu\text{g cm}^{-3}$ ppbv^{-1} . These values are in the same range as BB plumes seen previously at MBO [Baylon et al., 2015; Wigder et al., 2013].

In 2015 many fires were burning throughout the northwestern U.S. So in contrast to previous work at MBO, we were not able to calculate transport time for any of the regional BB events observed as they were influenced by multiple fires with various transport times. Figure 2 provides an example of this and exemplifies the impossibility of determining an exact transport time.

3.3. Optical Properties of the BB Aerosol at MBO

We observed significant differences in the optical properties of regional and Siberian-influenced BB events. The Siberian-influenced events had higher absorption coefficients relative to other measurements. This resulted in higher $\Delta\sigma_{\text{abs}}/\Delta\text{CO}$, higher MAE ($\Delta\sigma_{\text{abs}}/\Delta\text{PM1}$), and lower ω ($\sigma_{\text{scat}}/(\sigma_{\text{scat}} + \sigma_{\text{abs}})$) compared to regional BB events (Figures 3 and 4). We found no significant differences for $\Delta\sigma_{\text{scat}}/\Delta\text{CO}$ or MSE ($\Delta\sigma_{\text{scat}}/\Delta\text{PM1}$) between regional and Siberian events. Back-trajectories for the Siberian events (events 10-15) originated at high elevation over Siberia,

suggesting that the BB emissions were lofted to altitudes of 4-10 km (Figure 5). The Siberian events at MBO were observed over the course of a week (8/17/2015-8/23/2015); therefore the back-trajectories in Figure 5 represent a sustained meteorological pattern that consistently transported Siberian smoke to North America throughout the week. Aerosol vertical profiles measured by CALIOP corroborate the transport of BB plumes from the Siberian fires across the Pacific at altitudes of 4-10 km. Large BB plumes were identified over Northeast Asian and the North Pacific consisting primarily of BB smoke and some polluted dust over the Northern Pacific from 8/8/2015-8/17/2015. Figures S1-S4 show selected CALIPSO transects from 8/13/2015-8/16/2015 over the Pacific. The location and altitude of these plumes match the back-trajectories calculated from MBO for the Siberian events (Figure 5), verifying that events 10-15 are heavily influenced by the Siberian fires.

We suggest that the Siberian BB events observed at MBO represent hotter, more flaming portions of the fires which have higher BC emissions and thus higher absorption enhancements compared to the regional BB events. The hotter parts of the fires have more pyro-convective energy to loft the plume high into the atmosphere where it can then undergo long-range transport. During the ARCTAS-A flight campaign in Alaska, Siberian fire plumes were found to have a much larger BC/CO ratio ($8.5 \pm 5.4 \text{ ng m}^{-3} \text{ ppbv}^{-1}$) than North American fire plumes ($2.3 \pm 2.2 \text{ ng m}^{-3} \text{ ppbv}^{-1}$) [Kondo *et al.*, 2011]. This difference was attributed to the Siberian fires having a higher modified combustion efficiency (MCE). In addition, for the Siberian BB plumes they found MCE to increase with altitude. Jolleys *et al.* [2015] correspondingly found higher $\Delta\text{BC}/\Delta\text{OA}$ (Δ black carbon/ Δ organic aerosol) ratios to increase with altitude in eastern Canadian BB plumes. Intense, flaming fire plumes have higher injection heights into the atmosphere due to enhanced pyro-convection, whereas smoldering fires have low thermal convective energy and are mostly contained within the boundary layer. BB aerosol lofted to the free troposphere via pyro-convection is less likely to be removed and can have a longer atmospheric lifetime of up to 40 days [Bond *et al.*, 2013]. The back-trajectories for the Siberian events corroborate this idea. They were all relatively dry (water vapor mixing ratio $< 5 \text{ g kg}^{-1}$) with little precipitation during transport, suggesting the aerosol in the Siberian events was subjected to very limited wet deposition, which is the main removal mechanism from the atmosphere. Flaming conditions produce more BC and less OA generally, which leads to amplified absorption [Vakkari *et al.*, 2014; Yokelson *et al.*, 2009]. Flaming conditions are associated with high MCE values [Reid *et*

al., 2005a]. Unfortunately, we were not able to calculate MCE values for the Siberian events at
300 MBO due to extensive dilution and boundary layer mixing during transport [Yokelson et al.,
2013a].

While the ω values for the Siberian events are significantly lower relative to the regional
events, they are all high (> 0.95) compared to typical flaming conditions measured in the
laboratory or near-field measurements [S Liu et al., 2014a; Vakkari et al., 2014]. S Liu et al.
305 [2014a] found a robust relationship between ω and MCE in laboratory BB emissions where MCE
was negatively correlated with ω . However, observations have found that ω increases
significantly hours after emission in BB plumes [Reid et al., 2005a; Vakkari et al., 2014]. A
previous study at MBO found that well-aged BB plumes do not follow the S Liu et al. [2014a]
parametrization [Briggs et al., 2016]. All of the BB plumes observed by Briggs et al. [2016] had
310 $\omega > 0.91$ despite MCE values as high as 0.98, and no relationship was found between ω and MCE.
The high ω values typical of aged BB plumes are most likely due to SOA formation and
increased scattering efficiency as the particles ages and increases in size through coagulation and
condensation. Given this we believe the ω 's seen in these Siberian plumes are different and
significantly higher than the ω 's directly after emission and are therefore cannot be equated to an
315 MCE value.

We found AAE values for the Siberian events to be significantly lower than regional BB
events (Figures 3 and 4). High AAE values are indicative of the presence of brown carbon (BrC).
Brown carbon is a fraction of OA that selectively absorbs short wavelengths [Andreae and
Gelencser, 2006; Chen and Bond, 2010; Kirchstetter et al., 2004]. There are two possible
320 explanations for the difference in AAE values. The first is that the flaming conditions that
produced the Siberian events seen at MBO had higher BC and lower OA emissions, which
inherently have lower AAE as total absorption is dominated by BC and less BrC is initially
produced. Laboratory and field studies have corroborated this and observed an inverse
relationship between MCE and AAE [Holder et al., 2016; S Liu et al., 2014a; McMeeking et al.,
325 2014]. The other explanation is that BrC is lost during transport through photobleaching,
volatilization, and aerosol-phase reactions. Forrister et al. [2015] determined that BrC decreased
with transport with a half-life of 9 hours and that AAE decreases from ~ 4.0 to ~ 2.5 24 hours after
emission. All of the regional BB events were influenced by multiple fires that had transport times

330 varying from 3- 35 hours. With each event being influenced by at least one fire with a transport time ≤ 6 hours, this short transport time is consistent with the higher AAE values we observed.

3.4. Mass Scattering Efficiency

Mass scattering efficiencies (MSEs) are important for calculating the radiative forcing effects of aerosols in global climate and chemical transport models. Estimates of MSE are used to convert aerosol mass measurements to aerosol optical properties [Briggs *et al.*, 2016; Hand and Malm, 2007; Pitchford *et al.*, 2007]. MSE is dependent on particle composition, which determines the particle's refractive index and hygroscopicity, and aerosol size distribution [Hand and Malm, 2007]. We calculated MSE as the slope of the RMA regression of σ_{scat} and PM1 ($\Delta\sigma_{\text{scat}}/\Delta\text{PM1}$). R^2 values were >0.94 for all events. We found MSE values ranged from 2.50-4.76 $\text{m}^2 \text{g}^{-1}$, which are consistent with previously measured values.

340 During 2013 at MBO, MSE values estimated using AMS organic matter (OM) data and the σ_{scat} for four wildfire plumes ranged from 2.8-4.8 $\text{m}^2 \text{g}^{-1}$ (mean: 3.7 $\text{m}^2 \text{g}^{-1}$) [Briggs *et al.*, 2016]. Levin *et al.* [2010] calculated MSE values for fresh BB smoke from a variety of fuels to range from 1.5–5.7 $\text{m}^2 \text{g}^{-1}$, with most of the values falling between 2.0 and 4.5 $\text{m}^2 \text{g}^{-1}$. Reid *et al.* [2005a] reviewed MSE values from BB events and found a range between 3.2 and 4.2 $\text{m}^2 \text{g}^{-1}$ (mean: 3.8 $\text{m}^2 \text{g}^{-1}$) for temperate and boreal fresh smoke, and larger values for aged smoke (3.5-4.6 $\text{m}^2 \text{g}^{-1}$; mean: 4.3 $\text{m}^2 \text{g}^{-1}$). MSE values upwards of $\sim 6 \text{m}^2 \text{g}^{-1}$ have been observed for aged BB plumes [Hand and Malm, 2007; McMeeking *et al.*, 2005]. Due to the large variation in MSE values for BB events, assigning an average MSE value to convert aerosol mass measurements to aerosol optical properties or vice versa introduces significant uncertainties.

350 We investigated the cause for the variation in the MSE values that we observed. We found MSE's for BB events to be positively correlated with D_{pm} ($R^2 = 0.73$) (Figure 6a). If two D_{pm} values associated with bimodal size distributions are removed, the correlation increases substantially ($R^2 = 0.88$). A positive correlation between MSE and mean particle diameter has previously been observed in ambient data [Lowenthal and Kumar, 2004] and laboratory studies [McMeeking *et al.*, 2005]. Theoretically according to Mie theory, MSE will increase as the average particle diameter grows, through coagulation and condensation, toward the measurement wavelength (550 nm) [Seinfeld and Pandis, 2006].

3.5. BB Size Distributions

Figure 7 shows the BB aerosol number size distributions for the regional events (solid lines) and
360 Siberian events (dashed lines) we observed at MBO in $dN/d\log D_p$. We found D_{pm} and σ_g of the
number distributions to range from 138-229 nm and 1.53-1.89, respectively. The size
distributions observed at MBO are similar to *Janhäll et al.* [2010], who compiled aged BB size
distributions. They found the accumulation mode mean diameter to range from 175-300 nm with
geometric standard deviations of 1.3–1.7. No dependence was found in D_{pm} in plumes of
365 regional or Siberian origins. Similarly during the ARCTAS-B flight campaign, aged BB plumes
of western Canadian and Asian origins were found to have similar size distributions (Canadian:
 $D_{pm} = 224 \pm 14$ nm, $\sigma_g = 1.31 \pm 0.05$; Asian: $D_{pm} = 238 \pm 11$ nm, $\sigma_g = 1.31 \pm 0.03$) [*Kondo et al.*,
2011]. The BORTAS-B flight campaign in Eastern Canada observed aged BB plumes with
median diameters of 180-240 nm [*Sakamoto et al.*, 2015].

370 We observed clear bimodal distributions with an accumulation mode (100-500 nm) and
Aitken mode (20-100 nm) for five events (2, 3, 11, 14 and 15). The Aitken mode in these size
distributions most likely represents a secondary source from within the boundary layer. A
prominent “tail” consisting of higher than expected number concentrations of small-diameter
particles (30-90 nm) was observed for most of the unimodal events at MBO. It would be
375 expected that particles in this size range would grow to larger particles through coagulation
relatively quickly. *Sakamoto et al.* [2015] observed a similar elevation in the number
concentration of small particles during the BORTAS-B campaign. They attempted to account for
the existence of the tail with a Lagrangian box model of coagulation and dilution but were unable
to do so. Coagulation should cause a significant decrease in Aitken mode particles in a matter of
380 hours; and nucleation and condensation growth rates would have to be unreasonably high to
maintain these small particles.

We observed no clear distinction between the size distributions from regional and
Siberian events. These results are consistent with previous studies that have not observed a
dependence from plume age, transport time, or source location on the BB size distribution.
385 *Kondo et al.* [2011] found little difference between the D_{pm} of Siberian and Canadian BB plumes
despite different chemical composition, optical properties, and transport times. Similarly,
Sakamoto et al. [2015] found no trend in size distribution with plume transport distance. In a

study performed in the Front Range of Colorado, *Carrico et al.* [2016] found no significant difference between the size distribution of an hours old and a days old fire plume.

390 As previously stated, we found MSE's for BB events to be positively correlated with D_{pm} . This makes physical sense due to increased light scattering efficiency of larger particles closer to the wavelength of light (550 nm). In addition, we found event integrated D_{pm} to be correlated with event integrated σ_{scat} ($R^2 = 0.65$) and PM1 mass ($R^2 = 0.72$), and moderately correlated with CO ($R^2 = 0.41$) (Figure 6b,c,d). D_{pm} was not found to be correlated with any
395 normalized enhancement ratio ($\Delta\sigma_{scat}/\Delta CO$, $\Delta PM1/\Delta CO$). CO, σ_{scat} , and PM1 can be thought of as surrogates for plume concentration. The correlation between these proxies of plume concentration and D_{pm} indicates that in general, the more concentrated BB plumes have larger size distributions.

In a related study, *Sakamoto et al.* [2016] selected subsets of the MBO BB regional
400 events presented here and tested them against parameterizations of the aged size distribution. The parameterizations calculate D_{pm} and σ_g from inputs that can be derived from emissions-inventory and meteorological parameters. The seven inputs are: emission median dry diameter, emission distribution modal width, mass emissions flux, fire area, mean boundary-layer wind speed, plume mixing depth, and time/distance since emission. We identified eleven plumes from
405 regional events that had consistent transport to known regional fires. The simple fits captured over half of the variability in observed D_{pm} and modal width, even though the freshly emitted D_{pm} and modal widths were unknown. The results demonstrate that the parameterizations presented in *Sakamoto et al.* [2016] section 3.4 can be successfully used to estimate aged BB size distributions in regional BB plumes with transport times up to 35 hours. Using these
410 parameterizations to estimate BB plume size distribution in global and regional aerosol models is a significant improvement to assuming fixed values for size-distribution parameters.

The *Sakamoto et al.* [2016] parameterizations were particularly sensitive to mass emissions flux and fire area, as well as wind speed and transport time. If mass emissions flux is interpreted as surrogate for plume concentration, this agrees with our conclusion that increased
415 plume concentration results in a larger size distribution.

4. Conclusions

We characterized the physical and optical properties of 19 aged biomass burning events observed at the Mt. Bachelor Observatory in the summer of 2015. Regional (Northern California and Southwestern Oregon) and Siberian events were observed. Our main conclusions were:

- 420 - $\Delta\sigma_{\text{scat}}/\Delta\text{CO}$ (σ_{scat} at STP) enhancement ratio ranged from 0.48-1.29 Mm^{-1} ppbv⁻¹, with the majority of events being between 0.8 and 1.25 Mm^{-1} ppbv⁻¹.
- Siberian-influenced events had significantly higher $\Delta\sigma_{\text{abs}}/\Delta\text{CO}$ and MAE, and lower ω compared to regional events. We propose this is due to MBO sampling the portion on Siberian smoke that has been lofted to higher elevation through pyro-convection, thereby
- 425 preferentially sampling emissions of strong flaming combustion conditions. In general flaming conditions produce more BC, which would explain the amplified absorption in the Siberian events.
- Absorption Ångström exponent values were significantly lower for the Siberian events than regional events, which indicates lack of BrC produced by the Siberian fires or loss of
- 430 BrC during transport through photobleaching, volatilization, and aerosol-phase reactions.
- Mass scattering efficiencies ranged from 2.50-4.76 $\text{m}^2 \text{g}^{-1}$. MSE was positively correlated with D_{pm} ($R^2 = 0.73$), which agrees with Mie Theory.
- Aerosol number size distribution D_{pm} and σ_{g} ranged from 138-229 nm and 1.53-1.89, respectively. Five of the nineteen events had bimodal distributions, the rest being
- 435 unimodal. The unimodal distributions had a prominent “tail” of small-diameter particles (30-90 nm). No distinction could be made between regional and Siberian size distributions.

4. Author Contribution

James R. Laing performed the data analysis and prepared the manuscript with assistance

440 from all co-authors.

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Table 1. Identified BB plumes at MBO during the summer of 2015. All enhancement ratios are obtained by taking the slope of a RMA linear regression between the two species. ND (“no data”) indicates missing data. WC in the MAE column signifies a weak correlation ($R^2 < 0.60$).

Event number	Event date and time (UTC)	Event duration (hours)	Source fire location	ΔWV (g/kg)	$\Delta\sigma_{\text{scat}}/\Delta CO$ (Mm^{-1} ppbv $^{-1}$)	$\Delta\sigma_{\text{abs}}/\Delta CO$ (Mm^{-1} ppbv $^{-1}$)	MSE (m^2 g $^{-1}$)	MAE (m^2 g $^{-1}$)	AAE (467-660 nm)	ω (528 nm)	D_{pm} (nm)	σ_g
1	7/31/15 15:35-17:10	1.58	OR	0.16	1.13	0.036	ND	ND	3.15	0.97	164	1.72
2	8/9/15 2:55-8:55	6	CA, OR	1.62	0.89	WC	3.17	0.085	3.45	0.98	138	1.82
3	8/9/15 13:35-8/10/15 0:00	10.42	CA, OR	2.07	1.24	0.033	3.29	0.087	3.72	0.98	156	1.7
4	8/10/15 1:10-5:55	4.75	CA, OR	1.86	1.05	0.03	3.78	0.108	3.86	0.97	182	1.54
5	8/10/15 6:05-11:40	5.58	CA, OR	1.25	1.09	0.034	3.44	0.106	4.02	0.97	183	1.61
6	8/10/15 11:45-14:35	2.83	CA, OR	1.32	0.94	WC	3.27	WC	4.12	0.99	177	1.61
7	8/10/15 14:40- 8/11/15 6:15	15.58	CA, OR	1.83	1.17	0.032	3.64	0.098	3.52	0.98	186	1.62
8	8/11/15 14:20-18:45	4.42	CA, OR	1.11	1.07	0.029	2.5	0.066	2.74	0.98	160	1.78
9	8/14/15 10:00-15:35	5.58	OR	1.12	0.48	0.007	2.75	0.042	3.06	0.99	165	1.67
10	8/17/15 0:05-3:55	3.83	Siberia	-0.87	1.39	0.078	ND	ND	2.48	0.95	176	1.57
11	8/17/15 17:15- 8/18/15 7:00	13.75	Siberia	-0.22	1.06	0.060	ND	ND	2.5	0.95	179	1.69
12	8/18/15 16:05 - 8/19/15 16:40	24.58	Siberia	0.56	1.29	0.075	ND	ND	2.3	0.95	196	1.64
13	8/19/15 17:40 - 8/20/15 3:05	9.42	Siberia	0.6	1.12	0.052	ND	ND	2.25	0.96	175	1.76
14	8/22/15 15:30-18:05	2.58	Siberia	-3.1	1.97	0.078	4.76	0.188	3.59	0.96	229	1.73
15	8/23/15 3:55-7:00	3.08	Siberia	-2.45	1.09	0.059	2.84	0.156	2.51	0.96	162	1.89
16	8/23/15 9:50 - 8/25/15 6:50	45	CA, OR	1	1.13	0.029	4.06	0.107	3.15	0.98	205	1.58
17	8/25/15 12:45 - 8/26/15 19:00	30.25	CA, OR	1.67	0.88	0.027	3.75	0.111	3.12	0.98	181	1.6
18	8/26/15 7:15 - 8/28/15 11:15	40	CA, OR	1.35	0.89	0.031	3.7	0.128	3.48	0.97	191	1.53
19	8/28/15 17:40 - 8/29/15 6:15	12.58	CA, OR	1.54	0.69	ND	2.94	ND	ND	ND	164	1.58
Regional BB events (mean \pm stdev)				1.38 \pm 0.49	0.97 \pm 0.21	0.03 \pm 0.01	3.36 \pm 1.03	0.09 \pm 0.04	3.45 \pm 1.04	5.71 \pm 1.65	170 \pm 15.7	1.67 \pm 0.08
Siberian BB events (mean \pm stdev)				-0.91 \pm 1.56	1.32 \pm 0.34	0.07 \pm 0.01	3.8 \pm 2.05	0.17 \pm 0.09	2.61 \pm 0.49	4.16 \pm 0.6	181 \pm 19.7	1.77 \pm 0.1

ΔWV is water vapor enhancement, calculated for each event by subtracting the average WV for the summer sampling period from the WV value at the time when maximum CO was observed.

Aerosol scattering σ_{scat} (550 nm) and absorption σ_{abs} (528 nm) measurements were converted to STP.

MSE and MAE calculated as the $\Delta\sigma_{\text{scat}}/\Delta PM1$ and $\Delta\sigma_{\text{abs}}/\Delta PM1$ enhancement ratios, respectively.

D_{pm} is the geometric mean diameter and σ_g is the geometric standard deviation of the SMPS aerosol size distribution.

WC indicates a weak correlation in the MAE column ($R^2 < 0.60$).

ND indicates missing data. PM data was not available for events 1 and 10-13; absorption data was not available for events 19 and 20.

Figures:

Figure 1. Time series of CO, PM1, and aerosol scattering (σ_{scat}) at MBO during August. Threshold values (dashed black lines) used for BB event criteria are displayed for CO (150 ppbv) and scattering (20 Mm^{-1}). The dotted boxes represent multi-day periods of regional BB and encompasses events 2-8 and 16-19, respectively. The solid box represents the period influenced by Siberian BB and encompasses events 10-15.

Figure 2. Hysplit Back-trajectories for Event 19. The blue back-trajectories have a starting height of 1700 m amgl (above model ground level), the green a starting height of 1500 m amgl, and the red a starting height of 1300 m amgl. The squares are MODIS fire spots from 8/27/2015-8/29/2015 and are colored based on their fire radiative power (FRP).

Figure 3. Scatter plots of (a) mass absorption efficiency (MAE) and (b) absorption enhancement ratio $\Delta\sigma_{\text{abs}}/\Delta\text{CO}$ versus single scattering albedo (ω). MAE values were not calculated for four of the six Siberian-influenced events due to missing PM1 data.

Figure 4. Boxplots of (a) $\Delta\sigma_{\text{abs}}/\Delta\text{CO}$, (b) single scattering albedo (ω) measured at 528 nm, and (c) Absorption Ångström exponent (AAE) for absorption measurements at 467 and 660 nm for regional BB events and Siberian influenced events. N indicates the number of events for each box. Lower and upper whiskers represent the minimum and maximum values, respectively. Lower and upper lines of the box represent the 25th and 75th percentiles, respectively. The red line in the middle of the box represents the median, and the red plus mark represents outliers.

Figure 5. Most of the HYSPLIT back-trajectories for Siberian events (events 10-15) plotted as a function of altitude. Roughly 10% of the back-trajectories that did not follow the main transport track were not plotted. Forest fires from 8/7/2015 to 8/16/2015 identified by the Fire INventory from NCAR (FINN) fires are marked by red squares. These transects are not sequential and do not track one plume of Siberian smoke, but rather illustrate the extensive eastward transport of Siberian smoke over the course of the week. The four black boxes represent the locations of smoke plumes identified by CALIPSO cross sections detailed in Figures S1-S4.

Figure 6. Scatter plots of (a) MSE, (b) σ_{scat} , (c) PM1, and (d) CO versus D_{pm} for the BB events at MBO in the summer of 2015.

Figure 7. Event integrated aerosol number size distributions (corrected to STP) in dN/dlogDp ($\# \text{ cm}^{-1}$). The regional BB events have solid colored lines and the Siberian BB events have dashed black lines.