# Impacts of the July 2012 Siberian Fire Plume on Air Quality in the Pacific Northwest

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Abstract. Biomass burning emissions emit a significant amount of trace gases and aerosols and can affect atmospheric chemistry and radiative forcing for hundreds or thousands of kilometers downwind. They can also contribute to exceedances of air quality standards and have negative impacts on human health. We present a case study of an intense wildfire plume from Siberia that affected the air quality across the Pacific Northwest on July 6-10, 2012. Using satellite measurements (MODIS True Colour RGB imagery and MODIS AOD), we track the wildfire smoke plume from its origin in Siberia to the Pacific Northwest where subsidence ahead of a subtropical Pacific High made the plume settle over the region. The

- 20 normalized enhancement ratios of  $O_3$  and  $PM_1$  relative to CO of 0.26 and 0.08 are consistent with a plume aged 6-10 days. The aerosol mass in the plume was mainly submicron in diameter ( $PM_1/PM_{2.5} = 0.96$ ) and the part of the plume sampled at the Whistler High Elevation Monitoring Site (2182 m ASL) was 88% organic material. Stable atmospheric conditions along the coast limited the initial entrainment of the plume and caused local anthropogenic emissions to buildup. A synthesis of air quality from the regional surface monitoring networks describes changes in ambient  $O_3$  and  $PM_{2.5}$  during the event and
- 25 contrasts them to baseline air quality estimates from the AURAMS chemical transport model without wildfire emissions. Overall, the smoke plume contributed significantly to the exceedances in  $O_3$  and  $PM_{2.5}$  air quality standards and objectives that occurred at several communities in the region during the event. Peak enhancements in 8-hr  $O_3$  of 34-44 ppbv and 24-hr  $PM_{2.5}$  of 10-32 µg/m<sup>3</sup> were attributed to the effects of the smoke plume across the Interior of British Columbia and at the Whistler Peak High Elevation Site. Lesser enhancements of 10-12 ppbv for 8-hr  $O_3$  and of 4-9 µg/m<sup>3</sup> for 24-hr  $PM_{2.5}$
- 30 occurred at Whistler Peak and across coastal British Columbia and Washington State. The findings suggest that the large air quality impacts seen during this event were a combination of the efficient transport of the plume across the Pacific, favorable entrainment conditions across the BC interior and the large scale of the Siberian wildfire emissions. A warming climate increases the risk of increased wildfire activity and events of this scale re-occurring under appropriate meteorological conditions.

#### **1** Introduction

Wildfires emit significant amounts of primary pollutants including fine particulate matter ( $PM_{2.5}$ ), carbon monoxide (CO), 5 nitrogen oxides ( $NO_x$ ), and non-methane hydrocarbons (NMHCs). These pollutants can react to form ozone ( $O_3$ ) and

- secondary organic aerosols (SOA), which can affect downwind air quality and climate over a wide range of scales from local (Akagi et al., 2011, 2012, 2013; Smolyakov et al., 2014), to regional (Wigder et al., 2013), to inter-continental (Jaffe et al. 2001, 2004; Bertschi et al., 2004; Bertschi and Jaffe, 2005; Lapina et al., 2006; Pfister et al., 2006; Weiss-Penzias et al., 2007; Bein et al., 2008; Strada et al., 2012; Jaffe and Wigder, 2012). Short-term exposures to air pollution from biomass
- <sup>10</sup> burning have been found to be associated with a range of health impacts, including respiratory symptoms, increased hospital admissions and emergency room visits and premature mortality (Henderson and Johnston, 2012). The rate of  $O_3$  and SOA formation in wildfire plumes, and hence the potential for adverse air quality and health impacts, can vary considerably due to several factors, including biomass fuel type, combustion characteristics, atmospheric conditions, and downwind distance (Akagi et al., 2011; Jaffe and Wigder, 2012). The  $O_3$  production in biomass burning is driven by
- 15 photochemical reactions of NO<sub>x</sub> and NMOCs and typically occurs in a NO<sub>x</sub> limited environment. O<sub>3</sub> enhancements relative to CO, also known as the normalized enhancement ratio (NER;  $\Delta O_3/\Delta CO$ ), for temperate and boreal wildfires generally increase as the plume ages and is typically in the range of 0.1 to 0.7 (Jaffe and Wigder, 2012). A similar trend is present for the NER of PM<sub>1</sub> ( $\Delta PM_1/\Delta CO$ ) for plumes aged <2 days; yet for older plumes, SOA formation can be overshadowed by PM loss due to wet and dry deposition, cloud processing and/or evaporation (Wigder et al., 2013). Further study is still required
- 20 to better understand the interactions of the above factors on downwind air quality for long-range transport (LRT) events. The Russian boreal forests represent the largest forested region on Earth with an aerial coverage of approximately 8 million km<sup>2</sup> (Stocks, 2004). About 1% of this region is damaged from 10,000 to 35,000 forest fires annually (Isaev et al., 2002). Jaffe et al. (2004) estimated that the active Siberian wildfire season in 2003 caused summertime O<sub>3</sub> enhancements of 9-17 ppbv, which led to episodic O<sub>3</sub> exceedances in western North America. Other studies have also demonstrated the LRT of
- 25 wildfire smoke from Siberia can affect air quality over Asia (Jeong et al., 2008; Jung et al., 2016) and contribute to artic haze (Brock et al., 2011).

This study examines an episodic LRT event of Siberian wildfire smoke to the Pacific Northwest that had significant and widespread air quality impacts on July 6-10, 2012. Record temperatures and dry weather across Siberia in 2012 led to increased wildfire activity. Over 17,000 wildfires were detected in July and August 2012 alone. The Fire Inventory from

30 NCAR model (FINN; Wiedinmyer et al., 2011) estimated that 48 Tg of CO emissions occurred by August 2012 which was

already more than double the biomass burning emission for the entire 2010 season and second only to the 72 Tg from the 2003 season (NASA, 2012).

Cottle et al. (2014) examined the impact of this event in the Lower Fraser Valley (LFV) in British Columbia, Canada using lidar measurements taken at the University of British Columbia. HYSPLIT trajectory analysis confirmed that aerosol layers

- 5 subsiding over the LFV had largely originated from over wildfire source areas in Siberia at least 6 days earlier. Aerosol backscatter measurement and low depolarization volume ratios during the event showed the progressive entrainment of smoke into the LFV through July 10<sup>th</sup>, 2012. Their findings suggest that the high PM<sub>2.5</sub> observed by the region's fixed air quality monitoring network was associated with the progressive entrainment of aerosols into the LFV. This study expands on the Cottle et al. (2014) work in a number of significant ways: it analyzes potential air quality impacts over a much greater
- 10 geographical area encompassing large parts of British Columbia and Washington State; it uses detailed air quality measurements at a high elevation background site to provide insight into plume chemistry; and it makes use of photochemical modelling to establish baseline air quality conditions in the absence of any wildfire emissions to better quantify the impacts of the plume on O<sub>3</sub> and PM<sub>2.5</sub> levels.
- The objectives of this study are 1) to characterize the trans-Pacific advection of the 2012 Siberian fire plume and its eventual entrainment into the local valleys of British Columbia and northwest Washington State, 2) to determine its contribution to degraded air quality and exceedances of regional air quality objectives and national standards and 3) to provide information on the chemical composition of the smoke plume.

# 2. Methods

## 20 2.1 Monitoring Data

#### 2.1.1 Ambient Air Quality Monitoring Data

Ambient  $O_3$  and  $PM_{2.5}$  data were obtained from several air quality monitoring networks (Fig. 1 c and d), including 35 stations from the British Columbia Ministry of Environment (BCMoE) network (BC MoE, 2014a), 22 air quality monitors within the Lower Fraser Valley (LFV) Air Quality Monitoring Network (Metro Vancouver, 2014), and 15 stations from the

- 25 Washington State Monitoring Network (WSMN) (WA ECY, 2012). The air quality monitoring network across British Columbia is, in part, co-managed by the National Air Pollution Surveillance Network (NAPS) (Dabek-Zlotorzynska et al., 2011). Study sites also included two marine background sites (Amphitrite Point, Ucluelet, site C, 18 m above sea level (ASL)) on Vancouver Island (McKendry et al., 2014); and Cheeka Peak, site D, 480 m ASL in Washington State (Weiss-Penzias et al., 2004)) and two elevated background sites (Whistler Peak, site 1, 2182 m ASL, located 120 km north of
- 30 Vancouver (Takahama et al., 2011; Macdonald et al., 2011); and Mt. Rainier Jackson Visitor Centre, site F, 1782 m ASL,

located in Pierce County, Washington State). A description of the data analyzed and sampling methodologies for the various networks and sampling sites is summarized in Table 1.

# 2.1.2 Lidar Sites

Two lidars were used in this study: one located on the University of British Columbia Point Grey campus in Vancouver (site

- 5 B, 80 m ASL) and a second located at the north end of the Whistler village (site A, 650 m ASL), approximately 8 km north of the Whistler Peak High Elevation (WHI) (Gallagher et al., 2012; Strawbridge, 2013). The lidars use a Continuum Inlite III laser with an approximate output power of 1.5W at 10 Hz and operates simultaneously at wavelengths of 532 and 1064 nm. The system measures the return signals through 3 detection channels: one at 1064 nm, and two at 532 nm (one for each polarization). Backscatter information provided by the upward pointing setup has a vertical resolution of 3 m from near the
- 10 ground up to 15 km and uses a 10s averaging period. Additional details on the lidar systems can be found in Strawbridge (2013).

#### 2.1.3 Whistler Peak High Elevation site

Air quality measurements were collected at the Whistler Peak High Elevation site (see Table 1 for details). CO was measured with Thermo Environmental Instruments Inc., Model 48C-Trace Level analyzer and  $O_3$  was measured with a Thermo Environmental Instruments Inc., UV absorption monitor (TECO 49C). Particle chemistry was provided both 15 through integrated filter samples and continuously with an Aerosol Chemical Speciation Monitor (ACSM). Inorganic ions (Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, and Ca<sup>2+</sup>) were analyzed on 48-hour filter samples by ion chromatography (IC) and ICP-MS. OC, POC, and EC measurements were also obtain on a Sunset lab OC-EC aerosol analyzer (Sunset Laboratory Inc., 2016) using the EnCan Total-900 thermal method (Huang et al., 2006) with an 8-day sampling period (July 3-11) split between day and night. The ACSM provided particle chemical composition for  $SO_4^{2^-}$ ,  $NO_3^-$ ,  $CI^-$ ,  $NH_4^+$ , organics on a 30-min 20 time resolution. Optical measurements of black carbon (rBC) were acquired using the Single Particle Soot Photometer (SP2). During the study, the SP2 instrument had a steady 6% loss in its laser power; however internal diagnostics suggest that the power loss is unlikely to significantly affect the readings because heating aerosol during intake was unaffected. SP2 readings were also affected by a leak in the inlet plumbing noticed after the event. Particle size distributions were measured by an optical particle counter (OPC) over the (0.25 to 32 um) for the entire period and also from a Scanning Mobility 25

Particle Sizer (SMPS) (14-572 nm diameter) from July 9 onwards. Details of the trace gas and particle measurements at Whistler Peak are discussed by Macdonald et al. (2011) and Takahama et al. (2011).

#### 2.2 Trajectory and Dispersion Modelling

A detailed analysis by Cottle et al. (2014) using HYSPLIT (Hybrid Single-Particle Lagrangian Integrated Trajectory) 30 trajectories and high resolution smoke dispersion models generated with the Bluesky framework (Sullivan et al., 2008) concluded that wildfire sources within North America did not significantly influence air quality within the LFV during the July 6-10, 2012 period.

A similar analysis was done using the CMC Trajectory Model (D'Amours et al., 2015) with meteorological fields from the

- 5 GEM-LAM 15 km model at 15 minute intervals for nearby North America wildfires (Long Draw, Oregon and Waldo Canyon, Colorado). Forward trajectories were integrated for a 72 hour period at release heights of 10 m, 100 m, and 1000 m above ground level (AGL). Fig. S1 shows the forward trajectories for the Long Draw and Waldo Canyon wildfires released at 00 UTC on July 9<sup>th</sup>, 2012 and confirmed that smoke from these sources were unlikely to have contributed to the degraded ambient air quality observed over the larger geographical extent in this study. Trajectory modelling was not done for the few
- 10 small wildfires within the Interior of British Columbia. Overall, the effects of these sources are expected to be localized and with only minor air quality enhancements until after July 9<sup>th</sup>, 2012.

# 2.3 Analysis of O<sub>3</sub> and PM<sub>2.5</sub> impacts

Based on measurements collected at over 70 air quality monitoring stations in the Pacific Northwest, the air quality impacts of the Siberian wildfire plume were assessed by examining the  $O_3$  and  $PM_{2.5}$  concentrations at various averaging periods of

- 15 1-hr, 8-hr and 24-hr, depending on the pollutant. For each monitoring station, severe air quality episodes were identified using the regional objective and national standards, applicable at the time of the event (Table 2), as thresholds, and were compared to the average historical July daily maxima values from 2000 to 2010. It should be noted that although some of these standards, for example the Canada Wide Standard (CWS) for  $O_3$  (CCME, 2014), are based on multi-year statistics, all standards were compared to hourly observations of their respective averaging periods.
- 20 To further characterize the timing and spatial extent of these impacts, baseline concentrations of  $O_3$  and  $PM_{2.5}$  over the Pacific Northwest, in the absence of any wildfire emissions from either Siberia or from within North America, were simulated using the AURAMS (A Unified Regional Air-quality Modelling System) air quality transport model (Gong et al., 2006 with updates from Kelly et al., 2012) and were compared with observations.

# 2.3.1 AURAMS baseline simulation

- AURAMS was run over a 12 day period (July 5-15, 2012) using a nested configuration of 12 and 4 km grid spacing with the inner (4 km) domain covering southern British Columbia and northern Washington State (Fig. 1 a and c). Meteorology for the simulations was provided by Environment and Climate Change Canada's (ECCC) Global Environmental Multiscale (GEM) Limited-Area Model (LAM) weather forecast model (Côté et al., 1998) run at a 2.5 km resolution and then interpolated to the AURAMS domains. GEM-LAM was run in a series of 30-hour simulations starting from 00Z with the
- 30 first six hours of each simulation discarded as "spin-up" in order to allow model fields to reach steady-state. The model was run with anthropogenic emissions based on the 2010 Canadian and 2008 U.S. emission databases. Emission totals for both databases were adjusted to 2012 levels using Metro Vancouver forecasted and backcasted LFV emission estimates (GVRD,

2007). The model used biogenic emissions that calculated using the Biogenic Emissions Inventory System (BEIS) version 3.0.9 emissions algorithms and Biogenic Emissions Landuse Database 3 (BELD3) data. AURAMS employed the ADOM-II gas-phase chemical mechanism (Stockwell et al., 1989) and used a chemically-speciated 12-bin sectional distribution to characterize particulate matter from 0.01 to 40.96 µm in diameter.

- 5 Lateral boundary conditions to the AURAMS model along its outer (12 km) domain were supplied by seasonally representative climatological values for some species and for O<sub>3</sub> an additional adjustment was made to the climatological boundary conditions in response to the local tropopause height (Makar et al. 2010). Lateral boundary conditions to the inner 4 km domain were supplied from the 12 km model output. AURAMS runs did not include any wildfire emissions from either Siberia or from within North America.
- The reliability of the AURAMS baseline simulation (Table S2) was examined by comparing the 1-hr observed and modelled  $O_3$  and  $PM_{2.5}$  during non-event days (July 5 and from July 12-15). Within the 4 km domain, the model bias in 1-hr  $O_3$  and  $PM_{2.5}$  is relatively low (2.35 ppbv and 1.14 µg/m<sup>3</sup>, respectively). Over the 12 km domain, the  $O_3$  and  $PM_{2.5}$  performance degraded, with  $O_3$  bias and RMSE of 14.3 ppbv and 20.86 ppbv, respectively, and  $PM_{2.5}$  bias and RMSE of 5.84 µg/m<sup>3</sup> and 10.21 µg/m<sup>3</sup>, respectively. However, it should be noted that the majority of the study sites fall within the 4 km domain.
- 15 Overall, the 1-hr model performance of the baseline simulation is comparable to the assessed reliability of the AURAMS model from other studies (Makar et al., 2014).

#### 2.3.2 LRT enhancements analysis

LRT enhancements were identified when ambient air quality conditions in the Pacific Northwest departed from the baseline conditions as modelled by AURAMS, beyond a reasonable margin of error. As the model bias can affect the magnitude of the LRT enhancement estimates, the uncertainty in these enhancements was quantified in a conservative manner in this

20 the LRT enhancement estimates, the uncertainty in these enhancements was quantified in a conservative manner in this study. This was done by examining the range of differences between the 8-hr and 24-hr observed and modelled values for O<sub>3</sub> and PM<sub>2.5</sub>, respectively, during non-event days (July 5 and from July 12-15) at each monitoring location. Table S1 provides uncertainty estimates in 8-hr O<sub>3</sub> and 24-hr PM<sub>2.5</sub> for selected sites. For reporting purposes, enhancements for multiple monitoring locations were aggregated, when appropriate, to reflect the average contribution of the Siberian plume across the different regions in the Pacific Northwest.

#### 3. Results and Discussion

# 3.1 July 2012 smoke event overview

On June 29<sup>th</sup>, 2012, smoke originating from wildfires over Eastern Russia, as noted by the NASA Earth Observatory group, drifted eastward across the Sea of Japan, the Sea of Okhotsk, and Kamchatka (Fig. 1 b). Moderate Resolution Imaging

30 Spectroradiometer (MODIS) Aerosol Optical Depth (AOD) imagery between July 1<sup>st</sup> and July 6<sup>th</sup>, shown in Fig. S2 (a to f),

illustrates the progression of the aerosol plume (AOD > 1) across the Pacific to western North America. Long-range transport of aerosols was aided by strong zonal flow established as a developing storm tracked from the Bearing Sea to the Gulf of Alaska. Subsidence ahead of the subtropical Pacific high is thought to have contributed to the descent of the plume over the study area. In western North America, subsidence and mountain wave activity are found to be important factors in

5 bringing mid-tropospheric dust layers in range of boundary layer entrainment process (McKendry et al., 2001; Hacker et al., 2001).

On July 7<sup>th</sup>, the MODIS True Colour imagery (Fig. 2) shows the smoke plume oriented northeast to southwest across the southern British Columbia and Washington State. At this time, degraded air quality conditions were observed inland over the Interior of British Columbia, where both the 8-hr  $O_3$  and 24-hr  $PM_{2.5}$  exceedances occurred at Kamloops (site 8). From

- 10 July 8-9, poor air quality conditions continued to spread across the Interior, coinciding with the northward and inland progression of the plume as collaborated by AOD imagery (Fig. S2 g), and caused air quality exceedances at several communities (see Table 3). Over in the coastal region, elevated  $O_3$  and  $PM_{2.5}$  also occurred on July 8<sup>th</sup>, with isolated 8-hr  $O_3$ exceedances at Chilliwack (site 3) and Enumclaw (site 5). MODIS AOD on July 10<sup>th</sup> (Fig. S2 h) show the plume remnant had shifted southeastward and out of the Pacific Northwest domain.
- 15 Figure 3 provides an overview of the maximal enhancements to 8-hr O<sub>3</sub> and 24-hr PM<sub>2.5</sub> estimated based on the differences between the ambient air quality observation and the baseline AURAMS air quality modelling for the event. Overall, the baseline modelling suggests that the greatest air quality impacts from the Siberian wildfire smoke were in the Interior of British Columbia and at the Whistler High Elevation monitoring site with peak increases of 34-44 ppbv for 8-hr O<sub>3</sub> and 10-32 µg/m<sup>3</sup> for 24-hr PM<sub>2.5</sub>. The coastal portions of British Columbia and Washington State saw lesser enhancements of 10-14
- 20 ppbv and 4-9 ug/m<sup>3</sup>, respectively. The regional disparity in the enhancement may be attributable to difference in subsidence across the region. Coastal regions had stable atmospheric conditions, leading to stagnant conditions under a strong thermal inversion between 500 and 1000 m ASL (Fig. 4 a). Stable atmospheric conditions likely limited the entrainment of the plume over the coastal area and caused a buildup of local anthropogenic emissions. Over the Interior of British Columbia, dry and less stable conditions existed with daytime mixing heights over 3 km ASL (Fig. 4 b). The variations in 8-hr O<sub>3</sub> and 24-hr
- 25 PM<sub>2.5</sub> enhancements (with uncertainty estimates) across the Pacific Northwest are listed in Table 4 and are discussed in more detail in the following sections.

# 3.2 Plume chemistry at Whistler High Elevation Monitoring site

The Whistler Peak High Elevation air chemistry monitoring site is situated to monitor free tropospheric background air pollutants and to study long-range transport events across the Pacific (Leaitch et al., 2009; Macdonald et al, 2011). On July

30 6<sup>th</sup>, significant increases in 1-hr O<sub>3</sub>, CO, and PM<sub>2.5</sub> mark the onset of the smoke event at Whistler Peak (Fig. 5) and coincide with the timing of elevated aerosol layers detected by the Whistler lidar. Aerosol backscatter ratios show that the plume persisted over Whistler from July 6 14:00 PST (WHI1) to July 8 06:00 PST (WHI2) at an elevation of 2-3.6 km ASL. Backscatter ratios also indicate the entrainment of the aerosols deeper into the Whistler Valley as hourly observed O<sub>3</sub>,  $PM_{2.5}$ , and CO reached peaks of 86 ppbv, 24  $\mu$ g/m<sup>3</sup>, 276 ppbv, respectively. The maximum 1-hr O<sub>3</sub> and CO values were approximately 50 and 167 ppbv above the average July background levels as reported in Macdonald et al. (2011) for 2002-2006. O<sub>3</sub> conditions prior to the event were consistent with July background levels.

- The ratio of  $O_3$  to CO enhancements ( $\Delta O_3/\Delta CO$ ) provides some insight into the relative photochemical production of  $O_3$  in 5 wildfire plumes (Jaffe and Wigder, 2012) and is strongly influenced by the travel time between the fire and measurement location (Mauzerall et al., 1998). For this study, the plume age is estimated between 7-10 days old based on a Siberian origin of June 29<sup>th</sup>, 2012. The  $\Delta O_3/\Delta CO$  ratio was estimated using observations from WHI1 to WHI2 and excluding cases where CO was lower than the July background (110 ppbv). The regression slope of O<sub>3</sub> and CO for the event ranged from 0.15 to 0.24 with a mean value of 0.26 (n= 20, r<sup>2</sup>= 0.80) and is comparable to other similarly aged Siberian plumes with 10 excess ratios of 0.22-0.36 (6-10 days; Bertschi et al., 2004) and 0.15-0.84 (7-10 days, n=5; Bertschi and Jaffe, 2005).
- As shown in Figure 5d, during the smoke event (WHI1 to WHI2), the PM<sub>2.5</sub> mass was dominated by submicron aerosol with an average PM<sub>1</sub>/PM<sub>2.5</sub> ratio of 0.96. The particle composition as determined from the ACSM (Fig. 6) during the event was on average approximately 87.6% organic mass, 3.7% NO<sub>3</sub><sup>-</sup>, 5.9% SO<sub>4</sub><sup>2-</sup>, 2.5% NH<sub>4</sub><sup>+</sup>, 0.3% Cl<sup>-</sup>. PM<sub>1</sub> mass is estimated from the ACSM, since the size distribution data from 10 nm to 10 µm, measured by OPC, indicate that most of the mass is below 0.7
- 15  $\mu$ m; in Figure S3, an example is shown for July 9<sup>th</sup>, 2012 when the SMPS became operational again. Further, a comparison of the mass concentrations estimated from the SMPS, based on the assumptions of spherical particles and a particle mass density of 1.2 g/cm<sup>3</sup> as a result of the dominant organic composition during that period (Fig. 6), with the ACSM for July 9-31 is illustrated in Figure S4. The comparison is strong over the 518 1-hr averaged data : r<sup>2</sup> = 0.95; slope = 1.10. As in the example in Figure S3, the PM<sub>2.5</sub> mass is estimated with the addition of the OPC data.
- 20 The evolution of PM concentrations in wildfire plumes is largely affected by the rate of secondary organic aerosol (SOA) formation (Akagi et al., 2012). Using the regression methods described in Wigder et al. (2013), ΔPM<sub>1</sub>/ΔCO is estimated at 0.08 (n=20, r<sup>2</sup>=0.94) during the WHI1 to WHI2 period. This ratio is comparable to those of long-range transported plume (>740 km) of 0.03 and 0.06 found in Weiss-Penzias et al. (2007) and Wigder et al. (2013), respectively. Compared to the median ratios of 0.17, 0.29, 0.29, and 0.19, corresponding to distances of <140, 140-340, 340-540, 540-740 km, respectively.</p>
- as reported by Wigder et al. (2013), the ratio found in this study is relatively low, suggesting that the net  $PM_1$  loss exceeds SOA formation during transport as the plume ages. A recent wildfire study (Jolleys et al., 2012) found little SOA production in aged biomass burning plumes; however, it was noted that other aircraft campaigns had found a large variability in OA enhancements.

The OC/EC ratio is also considered as its increase may indicate a contribution from SOA formation to the OC as a smoke

30 plume ages. From the 8-day integrated  $PM_1$  quartz filter samples, separated by day and night, the value of OC/EC was 3.8 (day) and 3.4 (night) at Whistler Peak for the period July 3-11. During the WHI1-WHI2 event, an OC/EC ratio of 3.7 was also estimated by scaling the average ACSM OM and rBC to the 8-day integrated OC and EC sample, respectively. In the week prior to July 1<sup>st</sup>, 2012, a large variation in OC/EC (approximately 1.2 to 7) was observed in the suburb of Novosibrisk, approximately 400-800km south to southwest of the main sources of the Siberian wildfire (Smolyakov et al., 2014). Due to

this large variation in OC/EC close to the source as attributed to variations in burning phase and fuel types, no further inferences of SOA formation in the plume were made. The ratios of m/z44 and m/z43 to OM derived from the ACSM, indicate that the organic aerosol during the height of the BB plume (WHI1 to WHI2) was more oxygenated than at any other time during July 5-11 (Fig. S5). If SOA production during the plume transport was insignificant, then this observation implies either a relatively high level of oxygenation near the source, heterogeneous oxidation of the OM during plume

- 5 implies either a relatively high level of oxygenation near the source, heterogeneous oxidation of the OM during plume transport, or the loss of less oxygenated components of the OM during transport.
   The overall contribution of the July 6-10 Siberian Plume event to the air quality standards for O<sub>3</sub> and PM<sub>2.5</sub> at Whistler Peak (site 1) were estimated using AURAMS baseline modelling (also shown in Fig. 5). Generally, the model depicts relatively clean air quality conditions (approximately 30 ppbv and 1.0 µg/m<sup>3</sup> for O<sub>3</sub> and PM<sub>2.5</sub>, respectively) on July 5<sup>th</sup> with an
- 10 increase trend in baseline  $O_3$  (between 5 to 10 ppbv from July 5-8). It is estimated that the Siberian wildfires contributed an additional 44 ppbv to the daily maximum 8-hr  $O_3$  and 10  $\mu$ g/m<sup>3</sup> to the 24-hr PM<sub>2.5</sub> concentration (Table 4).

#### 3.3 Lower Fraser Valley (LFV)

The UBC lidar (site B) data indicated a plume arrival time of 12:00 PST on July 6<sup>th</sup> (LFV1, Fig. 7a) and a plume elevation of between 2 – 3.6 km ASL over Metro Vancouver which is similar to that found by the Whistler lidar (site A) but with generally weaker aerosol backscatter ratio readings (Fig. 7a). The plume had a shorter overall persistence over Metro Vancouver with the plume's trailing edge shifting northward by 03:00 PST on July 8<sup>th</sup> (LFV2). It appears that the conditions favourable for the transport of Siberian smoke to LFV were also conducive to O<sub>3</sub> production (e.g. light winds, clear skies, and high temperatures). This makes the attribution of Siberian smoke impacts on local air quality slightly more challenging in the LFV than at Whistler. Figure 7 (b,c) depicts the increasing O<sub>3</sub> and PM<sub>2.5</sub> across the LFV during July 6-10 with the

- 20 network average 1-hr O<sub>3</sub> and 1-hr PM<sub>2.5</sub> rises to 54 ppbv and 18 µg/m<sup>3</sup>, respectively, on July 8<sup>th</sup>. Peak O<sub>3</sub> condition of 79 ppbv observed at Hope Airport (site 4) and PM<sub>2.5</sub> of 26 µg/m<sup>3</sup> at Port Moody Rocky Point (site 2) arose as inland temperature reached a maximum of 31<sup>o</sup>C at Chilliwack (site 3) on July 8<sup>th</sup>. Inland temperatures and O<sub>3</sub> conditions moderated subsequently on July 9<sup>th</sup> following a brief period of elevated convection in the early morning hours that affected parts of Washington State and LFV. High O<sub>3</sub> concentrations were reported over the high elevation (>1000m ASL) site in North
- 25 Vancouver, located close to the coast over the norther portion of the LFV, by a separate field campaign (Bart et al., 2014). The AURAMS model was used to assess the underlying baseline air quality (Fig. 7 b and c) and to assess the local impacts through a site-by-site analysis of differences with LFV monitoring network data (Fig. 8 a and b). For July 6-8, the AURAMS baseline simulation without wildfire emissions captured the rise in daytime O<sub>3</sub> on average across the LFV network (Fig. 7 b) and on a site-by-site basis (Fig. 8 a). In contrast, the baseline simulation was consistently lacking in nighttime O<sub>3</sub> titration.
- 30 Previous air quality studies have shown that these discrepancies do not significantly affect the rest of the baseline AQ simulation (Makar et al, 2014). As such, the estimates of  $O_3$  enhancement within the LFV are based on daytime hours only. The 1-hr  $O_3$  enhancements occurred on July 9-10 appears, based on the lidar data, to be about 18 27 hours after the plume advected north of the LFV (Fig. 8 a). The average enhancement of peak 8-hr maximum  $O_3$  across the LFV was estimated to

be 10 ppbv (Table 4; hourly variability shown as shaded regions in Fig. 8a). Spatial maps of 8-hr  $O_3$  (Fig. S6) show widespread enhancements across the LFV on July 9<sup>th</sup> lingered in the western portion on July 10<sup>th</sup>. The lack of  $O_3$  enhancement estimated throughout the LFV on July 8<sup>th</sup> (Fig. 8a and Fig. S6) suggests the exceedance-level  $O_3$  episode in Chilliwack (Fig. 8c) may have been due to local  $O_3$  production and not the Siberian wildfire.

- 5 The  $PM_{2.5}$  enhancements associated with the LRT event developed in the morning of July 8<sup>th</sup>, much earlier than with any O<sub>3</sub> enhancements, and then persisted through July 10<sup>th</sup>. The Siberian plume is estimated to have enhanced the 24-hr  $PM_{2.5}$  by a maximum of 9 µg/m<sup>3</sup> for sites across the LFV. Spatially, the largest enhancements to  $PM_{2.5}$  appear to be concentrated over the northern portion of the LFV (Fig. 3 and by day in Fig. S6). This suggests that entrainment of the plume may have been facilitated by slope flows along the northern mountainous terrain, yet appears not to have reached the floor of the LFV, as
- 10 evidenced by  $O_3$  data from the fixed monitoring network.

#### 3.4 Washington State

In general, the air quality concentrations in Washington State (WA) during the episode were similar to those of the LFV. The average hourly  $PM_{2.5}$  across WA rose to a maximum of ~15 µg/m<sup>3</sup> on July 9<sup>th</sup> and the peak 8-hr O<sub>3</sub> of 72 ppbv

- 15 (Fig. S7) occurred at Enumclaw on July 8<sup>th</sup>. The LRT smoke enhancements in WA (Table 4) were similar to those in the LFV for  $O_3$  with a 10 ppbv increase to the 8-hr  $O_3$  and a weaker contribution to the 24-hr  $PM_{2.5}$  of 4 µg/m<sup>3</sup> (Fig. 3 and by day in Fig. S6). However, the timing and geographical extent of the impacts of  $PM_{2.5}$  and  $O_3$  differed considerably. In particular, the impacts on  $PM_{2.5}$  were minimal over WA until July 10<sup>th</sup>, while  $O_3$  underwent some modest enhancement on July 9<sup>th</sup> at a limited number of monitoring locations (Fig. S7). It is likely that some of the discrepancies (between the
- 20 Canadian and US portions of the LFV in the estimated plume impacts) arise from model errors stemming from different methodologies used to create the 2010 Canadian and 2008 US emissions inventories.

#### 3.5 The Interior of British Columbia

The character of the Siberian plume and its impacts over the Interior of British Columbia differed considerably from the enhancements seen over the LFV and WA. The air quality for two communities (Fig. 9), Kamloops (site 8) and Williams Lake (site 9), reflect the dramatic response seen across the Interior. The arrival of the plume at these communities, 19:00 PST on July 6 for Kamloops (site 8) and 12:00 PST on July 8<sup>th</sup> for Williams Lake (site 9), resulted in a sudden shift in hourly PM<sub>2.5</sub> and elevated O<sub>3</sub> on the subsequent day. Based on the AURAMS modelling, it is estimated that the plume contributed an additional 18 and 38 ppbv at Kamloops (site 8) and Williams Lake (site 9), respectively, to the daily maximum 8-hr O<sub>3</sub>, and an additional 32 μg/m<sup>3</sup> and 30 μg/m<sup>3</sup>, respectively, to the 24-hr PM<sub>2.5</sub> (Table 4). The suddenness and magnitude of these

30 impacts suggests that the plume became well mixed into the boundary layer after its transition across the Coast Mountains, which also corroborates with the entrainment signatures noted on the Whistler lidar that shows the plume mixed down to the base of the Whistler Valley (Fig. 5). Unfortunately, the 12 km model study domain did not extend north sufficiently to provide baseline estimates of the air quality conditions to Quesnel (site 10), Prince George (site 11), and the Northern Interior (site 12 to 14), which also experienced conditions that exceeded national and provincial air quality standards. Instead, a baseline air quality level was inferred from the observations on July 6<sup>th</sup> and the enhancements at these sites (listed in Table 4) were found to be consistent

5 with those experienced at Williams Lake (site 9; based on a July  $6^{th}$  baseline, as well as the 12km domain) except for slightly weaker 24-hr PM<sub>2.5</sub> enhancements of 22-24  $\mu$ g/m<sup>3</sup>.

On July 10<sup>th</sup>, as the bulk of the Siberian plume shifted northeastward out of British Columbia, remnants of the plume shifted over the Southern Interior and Whistler; however, attribution of the plume's impacts was impeded due to local wildfire activity (Matthew Creek; approx. 100 km west of Kelowna; 155 acres). Overall, the maximal air quality enhancements due

10 to smoke from the Siberian wildfires in the Southern Interior was approximately 15 ppbv and 15  $\mu$ g/m<sup>3</sup> for 8-hr O<sub>3</sub> and 24-hr PM<sub>2.5</sub>, respectively, based on the impact analysis conducted at Vernon (site 7) for the July 6-9 period (Fig. S10).

# 4. Conclusions

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The extreme 2012 Russian boreal fire season resulted in the trans-Pacific transport of a large smoke plume across the Pacific in early July which had substantial, yet varied, impacts across large part of British Columbia and Washington State. Air quality concentrations in excess of the Canada Wide Standard (65 ppbv for 8-hr  $O_3$ ) and the British Columbia Air Quality

- Objective (25  $\mu$ g/m<sup>3</sup>) of 24-hr PM<sub>2.5</sub> were experienced on July 6-10 at several municipalities across British Columbia. The aged smoke plume (6-10 days old), as supported from data from Whistler Peak, had a mean normalized enhancement ratio ( $\Delta O_3/\Delta CO$ ) of 0.26 and a PM<sub>1</sub>/CO ratio of 0.08, consistent with other long-range transport smoke studies of similar origin. Fine organic aerosol mass (~88% based on the ACSM measurements) accounted for the majority of the 1-hr and 24-
- 20 hr PM<sub>2.5</sub> enhancements of  $23 \ \mu g/m^3$  and  $10 \ \mu g/m^3$ , respectively. Analysis of OC/EC did not provide evidence for significant SOA production during transport as the plume ages. However, analysis of the ratios of m/z44 and m/z43 to OM derived from the ACSM, the level of oxygenation of organic material in this plume was much higher than during flanking times. If SOA formation during transport was minimal, then the relatively high level of oxygenation may have resulted from three factors: the OM near the source was relatively oxygenated; heterogeneous oxidation of the OM during plume transport; or
- 25 removal of less oxygenated components of the OM during transport. Baseline air quality modelling provided by the AURAMS model was used to delineate the scope of the LRT enhancements to air quality under varying atmospheric conditions. Generally, the largest enhancements to 8-hr O<sub>3</sub> of 34-44 ppbv and 24-hr PM<sub>2.5</sub> of 10-32 µg/m<sup>3</sup> occurred at Whistler Peak (site 1), Kamloops (site 8) and across the Central and Northern Interior of British Columbia (site 9 to 14). In contrast, lesser LRT enhancements of approximately 10 ppbv to 8-hr O<sub>3</sub> and 4-9 µg/m<sup>3</sup> to
- 30 24-hr PM<sub>2.5</sub> were estimated for coastal British Columbia and Washington State. Stable atmospheric conditions along coastal area of British Columbia and Washington State likely limited the initial entrainment of the plume. The AURAMS modelling

approach accounted for enhancements in anthropogenic air quality due to the smoke plume in the majority of cases with the exception of July 8<sup>th</sup> at Chilliwack (site 3) and Enumclaw (site 5), where elevated levels may have been due to other factors. Long range transport smoke events, such as the 2012 Siberian forest fire plume, highlight the far reaching impacts of wildfires and how air quality impacts can vary with downwind meteorology, plume age and entrainment pathways. This

5 demonstrates the need to include wildfire emissions within chemical transport model, and to derive effective parameterizations for the lofting and subsequent reactions in modelling ambient air quality conditions. Lateral boundary for chemistry used in regional air quality modeling application should be responsive to the long-range transport of pollutants. Wildfires sources pose an increasing concern as global climate change is expected to increase wildfire frequency and severity in many regions (IPCC, 2007; Gillett, 2004; Liu et al., 2013). As such, wildfire smoke is expected to contribute

10 more to the ambient levels for O<sub>3</sub>, PM<sub>2.5</sub>, CO, and other pollutants and can potentially increase the frequency of extended periods of degraded air quality.

## Data availability

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Data sources used in this paper are listed in the supplemental material (Table S3). For ECCC data not readily accessible, please contact the corresponding author (Andrew.Teakles@canada.ca) for assistance.

#### Supplementary material related to this article is available online at:

#### **Author contribution**

- A. Teakles designed the study, developed the methodology and prepared the manuscript with contributions from co-authors.
- R. So consolidated and analyzed the data and provided contributions to the methodology.
- 20 J. Baik provided assistance in developing R code to generate figures.
  - B. Ainslie generated the AURAMS model run.
  - S. Hanna, A. Bertram and L. Huang provided data and their expertise on data and instrumentation comparability.
  - K. Strawbridge provided the lidar data and his expertise in interpreting this data.
  - AM. Macdonald and R. Leaitch provided and analyzed data collected at Whistler Peak High Elevation site.
- 25 D. Toom assisted with the ACSM data and supplementation analysis on OA oxygenation.
  - I. McKendry provided his expertise on the LFV analysis.
  - R. Vingarzan reviewed and provided contributions to the methodology and manuscript composition.
  - C. Schiller reviewed the manuscript and provided her chemistry expertise to the data interpretation.
  - R. Nissen performed the synoptic analysis.

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# 15 References

Akagi, S.K., Yokelson, R.J., Burling, I.R., Meinardi, S., Simpson, I., Blake, D.R., McMeeking, G.R., Sullivan, A., Lee, T., Kreidenweis, S., Urbanski, S., Reardon, J., D. W. T. Griffith, Johnson, T.J., Weise, D.R., 2013. Measurements of reactive trace gases and variable O3 formation rates in some South Carolina biomass burning plumes. Atmospheric Chemistry and Physics 13, 1141-1165.

20 Akagi, S.K., Craven, J.S., Taylor, J.W., McMeeking, G.R., Yokelson, R.J., Burling, I.R., Urbanski, S.P., Wold, C.E., Seinfeld, J.H., Coe, H., Alvarado, M.J., Weise, D.R., 2012. Evolution of trace gases and particles emitted by a chaparral fire in California. Atmospheric Chemistry and Physics 12, 1397-1421.

Akagi, S.K., Yokelson, R.J., Wiedinmyer, C., Alvarado, M.J., Reid, J.S., Karl, T., Crounse, J.D., Wennberg, P.O., 2011. Emission factors for open and domestic biomass burning for use in atmospheric models. Atmospheric Chemistry and Physics

25 11, 4039-4072.

Bart, M., Williams, D.,E., Ainslie, B., McKendry, I., Salmond, J., Grange, S.,K., Alavi-Shoshtari, M., Steyn, D., Henshaw, G.,S., 2014. High density ozone monitoring using gas sensitive semi-conductor sensors in the Lower Fraser Valley, British Columbia. Environmental science & technology 48, 3970-7.

BC MOE (British Columbia Ministry of Environment), 2014a. BC air quality - network description. Accessed 06/19 2015.

30 http://www.bcairquality.ca/assessment/network-description.html BC MOE (British Columbia Ministry of Environment), 2014b. Ambient air quality objectives. Environmental Standards Branch. Accessed 06/19 2015. <u>http://www.bcairquality.ca/reports/pdfs/aqotable.pdf</u> Bein, K.J., Zhao, Y., Johnston, M.V., Wexler, A.S., 2008. Interactions between boreal wildfire and urban emissions. Journal of Geophysical Research - Atmospheres 113, D07304.

Bertschi, I.T., Jaffe, D.A., Jaeglé, L., Price, H.U., Dennison, J.B., 2004. PHOBEA/ITCT 2002 airborne observations of transpacific transport of ozone, CO, volatile organic compounds, and aerosols to the northeast pacific: Impacts of Asian anthropogenic and Siberian boreal fire emissions. Journal of Geophysical Research - Atmospheres 109, D23S12.

Bertschi, I.T. and Jaffe, D.A., 2005. Long-range transport of ozone, carbon monoxide, and aerosols to the NE pacific troposphere during the summer of 2003: Observations of smoke plumes from Asian boreal fires. Journal of Geophysical Research - Atmospheres 110, D05303.

5

Brock, C.A., Cozic, J., Bahreini, R., Froyd, K. D., Middlebrook, A. M., McComiskey, A., Brioude, J., 2011. Characteristics,

10 sources, and transport of aerosols measured in spring 2008 during the aerosol, radiation, and cloud processes affecting Arctic Climate (ARCPAC) Project. Atmospheric Chemistry and Physics 11, 2423-2453. CCME (Canadian Council of Ministers of the Environment), 2014. Canada-wide standards for particulate matter and ozone -2012 final report. Canadian Council of Ministers of the Environment. Accessed 06/19 2015.

http://www.ccme.ca/files/Resources/air/pm ozone/PN 1526 2012 CWS for PM and Ozone Final Report.pdf

- 15 Côté, J., Gravel, S., Methot, A., Patoine, A., 1998. The operational CMC-MRB global environmental multiscale (GEM) model. part I: Design considerations and formulation. Monthly Weather Review 126, 1373-1395. Cottle, P., Strawbridge, K., McKendry, I., 2014. Long-range transport of Siberian wildfire smoke to British Columbia: Lidar
- observations and air quality impacts. Atmospheric Environment 90, 71. Dabek-Zlotorzynska, E., Dann, T.F., Kalyani Martinelango, P., Celo, V., Brook, J.R., Mathieu, D., Ding, L., Austin, C.C.,
- 2011. Canadian national air pollution surveillance (NAPS) PM2.5 speciation program: Methodology and PM2.5 chemical composition for the years 2003–2008. Atmospheric Environment 45, 673-686.
   D'Amours R, Malo A, Flesch T, Wilson J, Gauthier J-P, Servranckx R (2015) The Canadian Meteorological Center's Atmospheric transport and dispersion modelling suite. Atmos Ocean. doi:10.1080/07055900.2014.1000260
   Environment Canada, 2013. 10 years of data from the National Air Pollution Surveillance (NAPS) network data summary
- 25 from 1999 to 2008. En49-2/7-40-PDF. Accessed 01/28 2016. https://www.ec.gc.ca/rnspa-naps/77FECF05-E241-4BED-8375-5A2A1DF3688C/NAPS\_Annual\_Report\_August2013\_E.pdf Gallagher, J., McKendry, I., Cottle, P., Macdonald, A., Leaitch, R., Strawbridge, K., 2012. Application of lidar data to assist airmass discrimination at the whistler mountaintop observatory. Journal of Applied Meteorology and Climatology 51, 1733-1739.
- 30 Gillett, N.P., Weaver, A.J., Zwiers, F.W., Flannigan, M.D., 2004. Detecting the effect of climate change on Canadian forest fires. Geophysical Research Letters 31, L18211.

Gong, W., Gong, S., Dastoor, A.P., Bouchet, V.S., Makar, P.A., Moran, M.D., Pabla, B., Ménard, S., Crevier, L., Cousineau, S., Venkatesh, S., 2006. Cloud processing of gases and aerosols in a regional air quality model (AURAMS). Atmospheric Research 82, 248-275.

Greater Vancouver Regional District (GVRD), 2007. 2005 Lower Fraser Valley air emissions inventory & forecast and backcast. technical report. <u>http://www.metrovancouver.org/services/air/emissions/Pages/default.aspx</u> Grimm Technologies Inc., 2010. Grimm model 1.109 - Portable aerosol spectrometer. Accessed 01/28 2016. http://www.dustmonitor.com/Research/1109.htm

- Hacker, J. P., McKendry, I. G., Stull, R. B., 2001. Modeled downward transport of Asian dust over Western North America during April 1998. J. Appl. Meteorol. 40, 1617–1628.
   Health Canada, 1999. National ambient air quality objectives for ground-level ozone summary science assessment document. federal provincial working group on air quality objectives and guidelines. July 1999. Accessed 06/19 2015. http://www.hc-sc.gc.ca/ewh-semt/alt\_formats/hecs-sesc/pdf/pubs/air/naaqo-onqaa/ground\_level-ozone-
- 10 tropospherique/summary-sommaire/ozone-summary-sommaire-eng.pdf

Henderson, S.B., Johnston, F.H., 2012. Measures of forest fire smoke exposure and their associations with respiratory health outcomes. Current Opinion in Allergy and Clinical Immunology 12, 221-227.

Huang, L., Brook, J.R., Zhang, W., Li, S.M., Graham, L., Ernst, D., Chivulescu, A., Lu, G., 2006. Stable isotope measurements of carbon fractions (OC/EC) in airborne particulate: A new dimension for source characterization and apportionment. Atmospheric Environment 40, 2690-2705.

IPCC, 2007. Climate Change 2007: Impacts, Adaptation and Vulnerability: Contribution of Working Group II to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change, M.L. Parry, O.F. Canziani, J.P. Palutikof, P.J. Van Der Linden and C.E. Hanson, Eds. Cambridge University Press, Cambridge, UK, 976.

Isaev, A.S., Korovin, G.N., Bartalev, S.A., Ershov, D. V., Janetos, A., Kasischke, E. S., Shugart, H. H., French, N. H. F.,
Orlick, B. E., Murphy, T.L., 2002. Using Remote Sensing to Assess Russian Forest Fire Carbon Emissions. Climatic Change

55, 235-249.

Jaffe, D. and Wigder, N., 2012. Ozone production from wildfires: A critical review. Atmospheric Environment 51, 1-10. Jaffe, D., Bertschi, I., Jaeglé, L., Novelli, P., Reid, J.S., Tanimoto, H., Vingarzan, R., Westphal, D.L., 2004. Long-range transport of Siberian biomass burning emissions and impact on surface ozone in western North America. Geophysical

25 Research Letters 31, L16106.

15

30

Jaffe, D., Anderson, T., Covert, D., Trost, B., Danielson, J., Simpson, W., Blake, D., Harris, J., Streets, D., 2001. Observations of ozone and related species in the northeast pacific during the PHOBEA campaigns 1. ground-based observations at Cheeka Peak. Journal of Geophysical Research 106, 7449-7461.

Jeong, J.I., Park, R.J., Youn, D.,2008. Effects of Siberian forest fires on air quality in East Asia during May 2003 and its climate implication. Atmospheric Environment 42, 8910-8922.

Jolleys, M.D., Coe, H., McFiggans, G., Capes, G., Allan, J.D., Crosier, J., Williams, P.I., Allen, G., Bower, K.N., Jimenez, J.L., Russell, L.M., Grutter, M., Baumgardner, D., 2012. Characterizing the aging of biomass burning organic aerosol by use of mixing ratios: a meta-analysis of four regions. Environmental Science & Technology 46, 13093-13102.

Jung, J., Lyu, Y., Lee, M., Hwang, T., Lee, S., Oh, S., 2016. Impact of Siberian forest fires on the atmosphere over the Korean Peninsula during summer 2014. Atmospheric Chemistry and Physics 16, 6757-6770.Kelly, J., Makar, P.A., Plummer, D.A., 2012. Projections of mid-century summer air-quality for North America: Effects of

5 Lapina, K., Honrath, R.E., Owen, R.C., Martín, M.V., Pfister, G., 2006. Evidence of significant large-scale impacts of boreal fires on ozone levels in the midlatitude northern hemisphere free troposphere. Geophysical Research Letters 33, L10815. Leaitch, W.R., Macdonald, A.M., Anlauf, K.G., Liu, P.S.K., Toom-Sauntry, D., Li, S.-., Liggio, J., Hayden, K., Wasey, M.A., Russell, L.M., Takahama, S., Liu, S., van Donkelaar, A., Duck, T., Martin, R.V., Zhang, Q., Sun, Y., McKendry, I., Shantz, N.C., Cubison, M., 2009. Evidence for asian dust effects from aerosol plume measurements during INTEX-B 2006

changes in climate and precursor emissions. Atmospheric Chemistry and Physics 12, 5367-5390.

20

- near whistler, BC. Atmospheric Chemistry and Physics 9, 3523-3546.
   Liu, Y., Goodrick, S.L., Stanturf, J.A., 2013. Future U.S. wildfire potential trends projected using a dynamically downscaled climate change scenario. Forest Ecology and Management 294, 120.
   Macdonald, A.M., Anlauf, K.G., Leaitch, W.R., Chan, E., Tarasick, D.W., 2011. Interannual variability of ozone and carbon monoxide at the Whistler high elevation site: 2002-2006. Atmospheric Chemistry and Physics 11, 11431-11446.
- 15 Makar, P.A., Nissen, R., Teakles, A., Zhang, J., Zheng, Q., Moran, M.D., Yau, H., diCenzo, C., 2014. Turbulent transport, emissions and the role of compensating errors in chemical transport models. Geoscientific Model Development 7, 1001-1024.

Makar, P.A., Gong, W., Mooney, C., Zhang, J., Davignon, D., Samaali, M., Moran, M.D., He, H., Tarasick, D.W., Sills, D., Chen, J., 2010. Dynamic adjustment of climatological ozone boundary conditions for air-quality forecasts. Atmospheric Chemistry and Physics 10, 8997-9015.

Mauzerall, D.L., Logan, J.A., Jacob, D.J., Anderson, B.E., Blake, D.R., Bradshaw, J.D., Heikes, B., Sachse, G.W., Singh, H., Talbot, B., 1998. Photochemistry in biomass burning plumes and implications for tropospheric ozone over the tropical South Atlantic. Journal of Geophysical Research 103, 8401-8423.

- 25 McKendry, I. G., Hacker, J. P., Stull, R., Sakiyama, S., Mignacca, D., Reid, K., 2001. Long range transport of Asian dust to the Lower Fraser Valley, British Columbia, Canada. J. Geophys. Res. 106(D16), 18 361–18 370. McKendry, I., Christensen, E., Schiller, C., Vingarzan, R., Macdonald, A.M., Li, Y., 2014. Low ozone episodes at Amphitrite point marine boundary layer observatory, British Columbia, Canada. Atmosphere-Ocean 52, 271-280. McKendry, I., Macdonald, A.M., Leaitch, W.R., van Donkelaar, A., Zhang, Q., Duck, T., Martin, R.V., 2008. Trans-pacific
- 30 dust events observed at Whistler, British Columbia during INTEX-B. Atmospheric Chemistry and Physics 8, 6297-6307. Metro Vancouver, 2014. Integrated air quality and greenhouse gas management plan progress report. Accessed 06/19 2015. http://www.metrovancouver.org/services/air-quality/AirOualityPublications/2014IAOGGMPProgressReport.pdf NASA, 2012. Wildfires in Siberia (September 13. 2012). Accessed 06/19 2014. http://earthobservatory.nasa.gov/IOTD/view.php?id=79161

Pfister, G.G., Emmons, L.K., Hess, P.G., Honrath, R., Lamarque, J.-., Martin, M.V., Owen, R.C., Avery, M.A., Browell, E.V., Holloway, J.S., Nedelec, P., Purvis, R., Ryerson, T.B., Sachse, G.W., Schlager, H., 2006. Ozone production from the 2004 North American boreal fires. Journal of Geophysical Research - Atmospheres 111, D24S07.

Puget Sound Clean Air Agency, 2013. 2012 air quality data summary & appendix. Accessed 08/19 2014. http://www.pscleanair.org/airq/reports.aspx

Smolyakov, B.S., Makarov, V.I., Shinkorenko, M.P., Popova, S.A., Bizin, M.A., 2014. Effects of Siberian wildfires on the chemical composition and acidity of atmospheric aerosols of remote urban, rural and background territories. Environmental pollution 188, 8-16.

Stocks, B.J., 2004. Forest fires in the boreal zone: climate change and carbon implications. International Forest Fire News 10 (IFFN) 31, July – December 2004, 122-131.

Stockwell, W., Lurmann, F., 1989. Intercomparison of the ADOM and chemical mechanisms. Electrical Power Research Institute Topical Report EPRI 3412, 254 pp.

Strada, S., Mari, C., Filippi, J.-B., Bosseur, F., 2012. Wildfire and the atmosphere: Modelling the chemical and dynamic interactions at the regional scale. Atmospheric Environment 51, 234-249.

15 Strawbridge, K.B., 2013. Developing a portable, autonomous aerosol backscatter lidar for network or remote operations. Atmospheric Measurement Techniques 6, 801-816.

Sullivan, D.C., Raffuse, S.M., Pryden, D.A., Craig, K.J., Reid, S.B., Wheeler, N.J.M., Chinkin, L.R., Larkin, N.K., Solomon, R., Strand, T., 2008. Development and applications of systems for modeling emissions and smoke from fires: the BlueSky smoke modeling framework and SMARTFIRE. In: 17th International Emissions Inventory Conference, June 2 – 5, 2008,

20 Portland, OR.

5

Sunset Laboratory Inc., 2016. Sunset Laboratory Inc. - Cutting Edge Carbon Aerosol Particulate Analysis Instruments. Assessed 07/21 2016. http://www.sunlab.com/

Takahama, S., Schwartz, R.E., Russell, L.M., Macdonald, A.M., Sharma, S., Leaitch, W.R., 2011. Organic functional groups in aerosol particles from burning and non-burning forest emissions at a high-elevation mountain site. Atmospheric Chemistry

25 and Physics 11, 6367-6386.

U.S. EPA (Environmental Protection Agency), 2014. National ambient air quality standards (NAAQS). Accessed 06/19 2015. <u>www.epa.gov/air/criteria.html</u>

WA ECY (Washington State Department of Ecology), 2008. Ozone monitoring procedure. 95-201G (rev. 4/08) Accessed 06/19 2015. https://fortress.wa.gov/ecy/publications/documents/95201g.pdf

30 WA ECY (Washington State Department of Ecology), 2012. 2012 ambient air monitoring network report. Accessed 06/19 2015. https://fortress.wa.gov/ecy/publications/publications/1202008.pdf Weiss-Penzias, P., Jaffe, D., Swartzendruber, P., Hafner, W., Chand, D., Prestbo, E., 2007. Quantifying Asian and biomass burning sources of mercury using the Hg/CO ratio in pollution plumes observed at the Mount Bachelor Observatory. Atmospheric Environment 41, 4366-4379.

Weiss-Penzias, P., Jaffe, D.A., Jaeglé, L., Liang, Q., 2004. Influence of long-range-transported pollution on the annual and diurnal cycles of carbon monoxide and ozone at Cheeka Peak Observatory. Journal of Geophysical Research - Atmospheres 109, D23S14.

Wiedinmyer, C., Akagi, S., Yokelson, R., Emmons, L., Al-Saadi, J., Orlando, J., 2011. The fire INventory from NCAR

5 (FINN): A high resolution global model to estimate the emissions from open burning. Geoscientific Model Development 4, 625-641.

Wigder, N.L., Jaffe, D.A., Saketa, F.A., 2013. Ozone and particulate matter enhancements from regional wildfires observed at Mount Bachelor during 2004-2011. Atmospheric Environment 75, 24-31.

Network/ Site	Measured Parameters	Instrument(s)	Reference
British Columbia			
British Columbia Ministry of Environment (BCMoE)	O <sub>3</sub>	Thermo Environmental Instruments Inc., UV absorption monitor (TECO 49C)	
network & Lauren Franze Vallari	PM <sub>2.5</sub>	Thermo Scientific TEOM 1400ab with sample equilibration system	Environment Canada (2013); Metro Vancouver (2014)
Air Quality Monitoring Network		Met One BAM-1020 Thermo Scientific Model 5030 SHARP	
Amphitrite Point, Ucluelet (site C)	O <sub>3</sub> PM <sub>2.5</sub>	Thermo 49i O <sub>3</sub> analyzer Thermo Sharp 5030 particulate monitor	McKendry et al. (2014)
UBC Lidar (site B) Whistler Village Lidar (site A)	aerosol backscatter	Lidar	Strawbridge (2013)
Whistler Peak High Elevation site (site 1)	СО	Thermo Environmental Instruments Inc., Model 48C-Trace Level analyzer	Macdonald et al. (2011)
	O <sub>3</sub>	Thermo Environmental Instruments Inc., UV absorption monitor (TECO 49C)	Macdonald et al. (2011)
	Refractory Black Carbon	Droplet Measurement Technologies Inc., Single Particle Soot Photometer (SP2)	Takahama et al. (2011)
	Particle Chemistry < 1 micron (SO <sub>4</sub> <sup>2-</sup> , NO <sub>3</sub> <sup>-</sup> , Cl <sup>-</sup> , NH <sub>4</sub> <sup>+</sup> , Organics)	Aerodyne Aerosol Chemical Speciation Monitor	Takahama et al. (2011)
	Particle size distributions from 0.25 to 32 µm diameter	GRIMM portable aerosol spectrometer 1.109	Grimm Technologies Inc. (2010)
	Particle size distributions in range ~ 14 nm to 572 nm diameter	TSI 3081 Scanning Mobility Particle Sizer (SMPS) with a TSI 3775 Condensation Particle Counter (CPC) operated at a low flow setting	McKendry et al. (2008)
	Integrated Inorganic Particle Chemistry (in this study: SO <sub>4</sub> <sup>2-</sup> )	Teflon filters; 48-hr samples	McKendry et al. (2008)
	Integrated Organic Chemistry (OC and EC)	Sunset lab OC-EC aerosol analyzer (EnCan Total -900 Thermal method); 8-days sample	Sunset Laboratory Inc. (2016); Huang et al. (2006)
Washington State			
Washington State	$O_3$	Teledyne-API T400	WA ECV (2008 & 2012)
(WSMN) Network	PM <sub>2.5</sub>	Nephelometer	WAECY (2008 & 2012)

# Table 1. Instrument list for various monitoring networks used in the study.

Cheeka Peak (site D)	O <sub>3</sub>	Teledyne-API T400	WA ECY (2008 & 2012)
	PM <sub>2.5</sub>	Radiance Research M903	
Mt. Rainier Jackson Visitor Center (site F)	O <sub>3</sub>	Teledyne-API T400	WA ECY (2008 & 2012)

Standard/ Objective	Pollutant	Averaging period	Value	Reference
Canada NAAQS	O <sub>3</sub>	1-hr	82 ppbv	Health Canada (1999)
Canada CWS			65 ppbv	CCME (2014)
BC provincial and Metro	$O_2$	8-hr	65 pphy	BC MOE (2014b)
Vancouver	03	0-111	05 pp0 (	Metro Vancouver (2014)
U.S. EPA NAAQS			75 ppbv	U.S. EPA (2014)
Canada CWS			$30 \mu\text{g/m}^3$	CCME (2014)
BC provincial and Metro	PM.	24-hr	$25 \ \mu g/m^3$	BC MOE (2014b)
Vancouver	1112.5			Metro Vancouver (2014)
U.S. EPA NAAQS			$35 \ \mu g/m^3$	U.S. EPA (2014)

Table 2: 2012 Ambient air quality standards and objective for O<sub>3</sub> and PM<sub>2.5</sub>.

Table 3. Summary of ambient  $O_3$  and  $PM_{2.5}$  conditions for stations that had severe (in bold) air quality degradation (exceeding one of the values listed in Table 2) between July 6th and July 11th. Values indicated reflect the maxima of 1, 8, and 24-hr averaging periods. Values in bracket denote historical July average daily maxima of 1 and 8-hr average periods and the historical July average of 24-hr running mean.

Station ()	Station(s)	O <sub>3</sub> (pr	obv)	$PM_{2.5} (\mu g/m^3)$		
Station(s)	ID	8-hr	1-hr	24-hr	1-hr	
Whistler						
Whistler Peak High Elevation Site	1	83	86	12	24	
Lower Fraser Valley		15 (20)	<b></b>	10 (7)	26 (12)	
Port Moody Rocky Point Park	2	45 (28)	55 (34)	19 (7)	26 (13)	
Chilliwack Airport	3	67 (35)	74 (41)	14 (6)	21 (12)	
Hope Airport <sup>a</sup>	4	64.8 (39)	79 (43)	18 (6)	21 (11)	
Washington State <sup>b</sup>						
Enumclaw	5	72	80	NA	NA	
Southern Interior						
Kelowna College	6	74 (40)	78 (44)	15 (5)	30 (13)	
Vernon Science Centre	7	67 (32)	73 (36)	19 (5)	30 (12)	
Kamloops Fire Station	8	65 (39)	72 (43)	36 (5)	44 (12)	
Central Interior						
Williams Lake <sup>c</sup>	9	80 (34)	84 (37)	34 (5)	46 (16)	
Quesnel <sup>c</sup>	10	76 (31)	92 (35)	31 (6)	57 (20)	
Prince George <sup>c</sup>	11	NA	NA	31 (6)	49 (17)	
Northern Interior (N. INT)						
Burns Lake Fire Centre	12	NA	NA	28 (3)	38 (9)	
Houston Firehall	13	NA	NA	27 (3)	38 (9)	
Telkwa	14	NA	NA	26 (3)	36 (11)	

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<sup>a</sup> air quality conditions at Hope Airport are noted here as it nearly exceeded the regional 8-hr ozone objective <sup>b</sup> encompasses coastal stations in northwestern Washington State as shown in Figure 1

<sup>c</sup> multiple stations in the area of interest

Table 4. Maximum daily enhancement (observed – model) of daily maximum 8-hr rolling average  $O_3$  and 24-hr rolling average  $PM_{2.5}$  based on various air quality baselines (4km and 12km AURAMS and historical values) from July 6<sup>th</sup> to July 11<sup>th</sup>, 2012.

			O <sub>3</sub> (ppbv)			PM <sub>2.5</sub> (μg/m	3)
Area of Interest	Baseline	No. of Sites	enhancement	Uncertainty range	No. of Sites	enhancement	Uncertainty range
Whistler Peak		1	43.7	(15.8, 49.2)	1	10.3	(7.7, 10.6)
$\mathrm{LFV}^{\mathrm{a,b}}$		20	9.9	(1.6, 22.0)	17	8.7	(6.0, 12.8)
Vancouver Island	AURAMS (4km)	6	13.7	(1.7, 27.0)	9	9.4	(7.1, 13.1)
Ucluelet	(4811)	1	12.4	(0.1, 14.9)	1	6.6	(0.9, 5.8)
U.S. <sup>a,b,c</sup>		10	10.4	(0, 23.4)	10	3.8	(4.1, 12.4)
Kelowna <sup>a</sup>		1	5.6	(5.5, 19.6)	1	7.7	(9.6, 11.0)
Vernon <sup>a</sup>	AURAMS	1	14.6	(16.7, 32.6)	1	14.7	(9.8, 13.1)
Kamloops <sup>a</sup>	(12km)	1	18.2	(22.0, 33.3)	1	32.1	(29.2, 33.1)
Williams Lake		1	37.5	(20.3, 57.0)	2	30.3	(23.9, 31.3)
Williams Lake		1	38		2	31	
Quesnel	Historical	1	43		3	24	
Prince George	(July 6th)	NA	NA	-	2	23	-
Northern Interior		1	34		4	22	

<sup>a</sup> estimated enhancement excluding July 10<sup>th</sup> and July 11<sup>th</sup>

5  ${}^{b}O_{3}$  enhancements were based on daytime values

<sup>c</sup> encompass northwest coastal stations in Washington State as shown in Figure 1

Station	Station #	Elevation (m)	Lat.	Long.	8-hr O <sub>3</sub> uncertainty <sup>*</sup>	24-hr PM <sub>2.5</sub> uncertainty <sup>*</sup>
Whistler						
Whistler Peak High Elevation Site	1	2182	50.06	-122.96	(-5.6, 27.9)	(-0.3, 2.5)
Whistler Lidar	А	660	50.13	-122.95		
Lower Fraser Valley						
Port Moody Rocky Point Park	2	17	49.28	-122.85	(-11.6, 7.9)	(-3.3, 5.3)
Chilliwack Airport	3	12	49.16	-121.94	(-22.6, 1.4)	(1.2, 2.4)
Hope Airport	4	39	49.37	-121.50	(-18.2, 8.8)	(-4.0, 10.0)
UBC Lidar	В	80	49.26	-123.25		
Vancouver Island						
Ucluelet Amphitrite Point	С	14	48.92	-125.54	(-2.6, 12.3)	(0.9, 5.7)
Washington State						
Enumclaw	5	402	47.14	-121.93	(-22.8, 7.3)	
Cheeka Peak	D	480	48.30	-124.62	(-3.1, 4.9)	(-11.5, 2.4)
Quillayute	E	59	47.94	-124.56		
Mt Rainier Jackson Ctr	F	1782	46.78	-121.74	(-21.5, 12.7)	
Southern Interior						
Kelowna College	6	347	49.86	-119.48	(-14.0, 0.1)	(-3.3, -1.9)
Vernon Science Centre	7	476	50.23	-119.28	(-18.1, -2.2)	(1.6, 4.9)
Kamloops Fire Station	8	381	50.70	-120.39	(-15.2, -3.8)	(-1.0, 2.9)
Kelowna Airport	G	344	49.96	-119.38		
Central Interior						
Williams Lake Columneetza School	9	631	52.14	-122.15	(-19.5, 17.2)	(-1.1, 6.4)
Williams Lake CRD Library	,	609	52.13	-122.14		(-0.9, 6.5)
Quesnel Maple Drive		614	52.96	-122.50		
Quesnel Senior Secondary	10	490	52.98	-122.49		
Quesnel West Correlieu School		478	52.97	-122.52		
Prince George Gladstone School	11	617	53.86	-122.76		
Prince George Plaza 400	11	588	53.91	-122.74		
Norther Interior						
Burns Lake Fire Centre	12	710	54.23	-125.76		
Houston Firehall	13	602	54.40	-126.65		
Telkwa	14	515	54.69	-127.05		

Table S1. Station information and modelling uncer	rtainty for select sites used in this study.
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\*AURAMS model uncertainty (observed-model) during non-event days

	y-intercept	Slope	r	Bias	MAE	RMSE
Current Study						
O <sub>3</sub> (4km)	8.5	0.66	0.69	2.35	7.57	9.55
PM <sub>2.5</sub> (4km)	5.68	0.35	0.35	1.14	3.71	4.97
O <sub>3</sub> (12km)	16.46	0.89	0.6	14.3	16.32	20.86
PM <sub>2.5</sub> (12km)	9.75	0.44	0.24	5.84	7.79	10.21
Makar et al. (2014)						
O <sub>3</sub> (All; Table 4a)	15.32	0.7	0.64	8.48	12.53	16.17
PM <sub>2.5</sub> (All; Table 4b)	3.36	0.45	0.28	1.2	5.28	7.72
PM <sub>2.5</sub> (LFV sites; Table 5)	7.95	0.24	0.23	2.82	6.77	10.53

Table S2. 1-hr AURAMS model performance for  $\mathrm{O}_3$  and  $\mathrm{PM}_{2.5}$  at 4 and 12km domain.

# Table S3. Source information and data availability

Site	Data	Source
British Columbia		
BCMoE network	O <sub>3</sub> , PM <sub>2.5</sub>	BCMoE (http://envistaweb.env.gov.bc.ca/)
Lower Fraser Valley Air Quality Monitoring Network	O <sub>3</sub> , PM <sub>2.5</sub>	BCMoE (http://envistaweb.env.gov.bc.ca/)
Amphitrite Point, Ucluelet	O <sub>3</sub> , PM <sub>2.5</sub>	ECCC ,contact info: Andrew.Teakles@canada.ca
UBC & WHI Lidar	Lidar	ECCC, contact info: Andrew.Teakles@canada.ca
Whistler Peak High Elevation Site	Particle size distribution , O <sub>3</sub> , CO, Filter packs IC (tSO4), Black Carbon, TOT EC & OC, ACSM, Lidar	ECCC, contact info: Andrew.Teakles@canada.ca
BC surface air quality stations	O <sub>3</sub> , PM <sub>2.5</sub> (climatology)	ECCC NAPS website (annual summaries: http://maps-cartes.ec.gc.ca/rnspa-naps/data.aspx)
Washington State		
WSMN	O <sub>3</sub> , PM <sub>2.5</sub> (neph)	Puget Sound Clean Air website
		(http://airgraphing.pscleanair.org/)
Cheeka Peak	O <sub>3</sub> , PM <sub>2.5</sub>	Department of Ecology, State of Washington, https://fortress.wa.gov/ecy/enviwa/
Mt. Rainier Jackson Visitor	0.	Department of Ecology, State of Washington,
Center	03	https://fortress.wa.gov/ecy/enviwa/
Air Quality Modelling		
All sites	AURAMS O <sub>3</sub> , PM <sub>2.5</sub>	ECCC, contact info: Andrew.Teakles@canada.ca
Satellite		
World	MODIS AOD	http://modis.gsfc.nasa.gov/data/
Pacific	MODIS true colour	http://visibleearth.nasa.gov/view.php?id=78406
Others		
Quillayute & Kelowna	Radiosondes	http://weather.uwyo.edu/upperair/sounding.html
Elevation data (for fig 1)	GLOBE Elevation	http://www.ngdc.noaa.gov/mgg/topo/globe.html



Fig. 1. Map of the monitoring network data and AURAMS modelling domain used in the study. The pink and blue regions in (a) highlights the two geographical regions of interest, British Columbia and Washington state, respectively. The black outlined area in (a) highlights the geographical location of the MODIS true colour Image on June 29<sup>th</sup>, 2012 from the NASA earth Observatory site (b). The blue and red outlined areas in either (a) and/or (c) highlight the 12 km and 4 km AURAMS grid domains, respectively. The green outline in (c) highlights the Northern Interior sites that fall outside the gridded domain. The Lower Fraser Valley (LFV; black outline in c) inset (d) displays the Metro Vancouver air quality monitoring network and the location of the UBC lidar. Abbreviations used in this figure are defined in Table S1 and throughout the text. Topography shaded in panels (c) and (d).



Fig. 2. MODIS true colour Image composite maps for July 7, 8, and 9, 2012 overlaid with daily maximum of 8-hr  $O_3$  (a, b, and c, respectively) and of 24-hr  $PM_{2.5}$  (d, e, f, respectively) for the study sites. The locations where observed concentration exceed the national or regional air quality standards for either  $O_3$  or  $PM_{2.5}$  are indicated in red

5 triangles in the appropriate panel and described in Table 3. The light blue circle on the MODIS source imagery indicates the location of the Kelowna Airport (site G) on all panels. Abbreviations used in this figure are defined in Table S1 and throughout the text.



Fig. 3. Overview of the maximal enhancements to 8-hr O<sub>3</sub> and 24-hr PM<sub>2.5</sub> estimated for the study area (a and b, respectively) and the LFV (c and d, respectively) based on the differences between the ambient air quality
observation and the AURAMS baseline modelling for the July 6-10, 2012 smoke event. Abbreviations used in this figure are defined in Table S1 and throughout the text.



Fig. 4. Skew-T atmospheric radiosondes for Quillayute (site E) and Kelowna Airport (site G) at 00Z on July 8<sup>th</sup>, 2012. The atmospheric dry bulb temperature (T) and dewpoint temperature (Td) are shown from the surface up to 100 mb. Black dashed line is a trace of a surface parcel lift to estimate the mixed layer depth at the height where it intersects the environment dry bulb temperature.



Fig. 5. Whistler lidar (site A, 650 m ASL) measured backscatter ratio of the 1064 nm channel to that of clear air for the July 2012 Siberian smoke plume (a) and observed (solid) 1-hr  $O_3$  (b), 1-hr CO (c), and 1-hr  $PM_1$  (based on the ACSM) and  $PM_{2.5}$ . (d) at Whistler Peak High Elevation station (site 1, 2182 m ASL, indicated as a horizontal white line in panel a). AURAMS baseline (dashed) 1-hr  $O_3$  and 1-hr  $PM_{2.5}$  are shown in panel b) and d), respectively. Elevated aerosol backscatter in panel (a) are noted from July 6 14:00 PST (WHI1) to July 8 06:00 PST (WHI2) by red dashed lines. The shaded regions indicate night time hours between 18:00 PST to 07:00 PST.



Fig. 6. Aerosol chemistry measurements taken at Whistler Peak High Elevation site (site 1) between July 6<sup>th</sup> 2012 and July 12<sup>th</sup>, 2012. Hourly Organics (Org),  $NO_3$ ,  $NH_4^+$ ,  $SO_4^{-2}$  measured by the ACSM (a, b), EC and OC as sampled over a 8-day period (July 3-11, with day and night split) using the EnCan Total-900 thermal method (T900) (a), SO4 as a 3-day integrated sample by quartz filter pack (FP) (b), and rBC as acquired using the Single Particle Soot Photometer (SP2) (c). The shaded regions indicate night time hours between 18:00 PST to 07:00 PST.



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Fig. 7. UBC lidar (site 2) observations and air quality conditions across the Lower Fraser Valley network between July 6<sup>th</sup> and 12<sup>th</sup>, 2012. Panel (a) shows the ratio of measured backscatter of the 1064 nm channel to that of clear air. Panel (b) and (c) indicates the mean observed (solid) and AURAMS baseline without wildfire emissions (dashed) 1-hr O<sub>3</sub> (ppbv) and 1-hr PM<sub>2.5</sub> ( $\mu g/m^3$ ) for 19 and 17 sites, respectively, across the LFV air quality monitoring network. The light and dark grey shading represents the range and inter-quartile range, respectively, across the network at a particular hour. The vertical shaded regions indicate night time hours between 18:00 PST to 07:00 PST.



Fig. 8. Contributions of the Siberian wildfire smoke plume based on the differences of observed 1-hr  $O_3$  (a) and  $PM_{2.5}$  (b) and the AURAMS baseline without wildfire emissions on a site-by-site basis for the LFV. For (a) and (b), the average contribution is labelled by the blue solid line. The light grey and dark grey shading represents the range and inter-quartile range of the contribution across the network at a particular hour. Panel (c) shows the AURAMS baseline and observed  $O_3$  timeseries for Chilliwack, BC (site 3). The light blue shaded regions indicate night time hours between 18:00 PST to 07:00 PST.



Fig. 9. The hourly observed (blue) and baseline AURAMS (red, dashed) without wildfire emissions  $O_3$  (top) and  $PM_{2.5}$  (bottom) for Kamloops (site 8) (a) and Williams Lake (site 9) (b). The shaded regions indicate night time hours between 18:00 PST to 07:00 PST.



Fig. S1. Five day forward trajectories using the CMC Trajectory model from the Long Draw, Oregon (a) and Waldo Canyon, Colorado wildfires (b) released at 00UTC on July 9<sup>th</sup>, 2012 at heights of 10 m, 100 m, and 1 km AGL.



Fig. S2. The daily average MODIS Aerosol Optical Depth product for July 1-6 (a to f) illustrates the eastward progression of the Siberian Fire plume in early July onto the Western Pacific prior to its arrival off the coast of 5 Washington and Oregon States on July 6<sup>th</sup>, 2012 (f). Enhanced AOD values spread across South and Central British Columbia by July 8<sup>th</sup>, 2012 (g) then shift southeastward out of the Pacific Northwest domain on July 10<sup>th</sup>, 2012 (h). Mean sea level pressure is contour (in solid black) on all panels.



Fig. S3. Comparison of SMPS and GRIMM aerosol number and volume distributions for July 9<sup>th</sup>, 2012 16h30 -20h30 PST.



Fig. S4. Timeseries of SMPS PM<sub>1</sub> and ACSM readings between July 9<sup>th</sup> and July 31<sup>st</sup>, 2012. The ACSM data have 5 been adjusted for a collection efficiency of 0.5.



Fig. S5. The m/z44 and m/z43 components of the organic aerosol spectrum derived from the ACSM spectra. The ratio of organic signal at m/z44 (Org43) vs. m/z43 (Org44) is compared to show that the organic aerosol was more oxygenated during the LRT smoke event (July 6<sup>th</sup> 14:00 PST to July 8<sup>th</sup> 06:00 PST) than at any other times during the 5 6 day period from July 5<sup>th</sup> 12:00 to 11<sup>th</sup> 18:00 PST. Dashed black lines mark the 1:1 and 2.5 ratio lines for reference.



Fig. S6. Spatial map of the differences of the AURAMS baseline without wildfire emissions to daytime averaged 8-hr  $O_3$  (ppbv, panels a, b, c, d) and of the daily averaged  $PM_{2.5}$  ( $\mu g/m^3$ , e, f, g, h) from July 7<sup>th</sup>, 2012 to July 10<sup>th</sup>, 2012.



Fig. S7. Hourly average observed (solid) and AURAMS baseline (dashed) without wildfire emissions: hourly  $O_3$  (a) and  $PM_{2.5}$  (b) for sites across the Washington State air quality monitoring network. The light grey and dark grey shading represents the range and inter-quartile range (IQR) across the network at a particular hour. The shaded regions indicate night time hours between 18:00 PST to 07:00 PST.



Fig. S8. The observed and modelled baseline hourly  $O_3$  (a) and  $PM_{2.5}$  (b) for Vernon (site 7). The shaded regions indicate night time hours between 18:00 PST to 07:00 PST.