¹ Inter-comparison of Elemental and Organic Carbon

- ² Mass Measurements from Three North American
- ³ National Long-term Monitoring Networks at a co-

⁴ located Site

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- 20 Keywords
- 21 Black carbon, thermal evolution, air pollution, carbonaceous aerosol, IMPROVE, CAPMON, CABM

22 Abstract

23 Carbonaceous aerosol is a major contributor to the total aerosol load and being monitored by 24 diverse measurement approaches. Here, ten years (2005-2015) of continuous carbonaceous aerosol 25 measurements collected at the Centre of Atmospheric Research Experiments (CARE) in Egbert, Ontario, Canada on quartz-fiber filters by three independent networks (Interagency Monitoring of PROtected 26 27 Visual Environments (IMPROVE), Canadian Air and Precipitation Monitoring Network (CAPMoN), and 28 Canadian Aerosol Baseline Measurement (CABM)) were compared. Specifically, the study evaluated 29 how differences in sample collection and analysis affected the concentrations of total carbon (TC), 30 organic carbon (OC), and elemental carbon (EC). Results show that different carbonaceous fractions 31 measured by various networks were consistent and comparable in general among the three networks 32 over the ten years period, even with different sampling systems/frequencies, analytical protocols and 33 artifact corrections. The CAPMON TC, OC, and EC obtained from the DRI-TOR method were lower than 34 those determined from the IMPROVE A TOR method by 17%, 14% and 18%, respectively. When using 35 transmittance for charring correction, the corresponding carbonaceous fractions obtained from the 36 Sunset-TOT were lower by as much as 30%, 15%, and 75%, respectively. In comparison, the CABM TC, 37 OC, and EC obtained from a thermal method (ECT9) were higher than the corresponding fractions from 38 IMPROVE_A TOR by 20-30%, 0-15% and 60-80%, respectively. Ambient OC and EC concentrations were 39 found to elevate when ambient temperature exceeded 10 °C. These increased ambient concentrations 40 of OC during summer were possibly attributed to the secondary organic aerosol (SOA) formation and 41 forest fire emissions, while elevated EC concentrations were potentially influenced by forest fire 42 emissions and increased vehicle emissions. Results also show that the pyrolyzed organic carbon (POC) 43 obtained from the EnCan-Total-900 (ECT9) protocol could provide additional information on SOA 44 although more research is still needed.

45 Introduction

Carbonaceous aerosols, including elemental carbon (EC), which is often referred to as black carbon (BC) and organic carbon (OC), make up a large fraction of the atmospheric fine particulate matter (PM) mass (Heintzenberg, 1989). Atmospheric OC and BC particles that are emitted directly into the atmosphere have both natural (e.g., biomass burning or forest fires) and anthropogenic (e.g., internal combustion engines) sources. A significant amount of the particulate OC is also formed in the atmosphere through oxidation and condensation of volatile organic compounds (e.g., isoprene and terpenes), which are emitted directly from vegetation. BC is a by-product of incomplete combustion of

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hydrocarbon fuels, generated mainly from fossil fuel combustion and biomass burning. Atmospheric
particles have direct and indirect influences on climate, visibility, air quality, ecosystems, and adverse
human health effects (Bond et al., 2013; Japar et al 1986; Lesins et al., 2002; Watson, 2002).

Atmospheric BC absorbs solar radiation while OC primarily scatters (Schulz et al., 2006). However, BC and OC co-exist in atmospheric particles and the net radiative forcing of the aerosol particles depends on the particle size, composition, and the mixing state of the particles, while all of these variables also change as aerosol particles age (Fuller et al., 1999; Lesins et al., 2002).

60 Black carbon is a generic term in the literature and it is often interchanged with other terms 61 such as EC, soot, refractory BC, light absorbing carbon, or equivalent BC (Petzold et al., 2013). Although 62 BC is highly relevant to climate research, there is no universally agreed and clearly defined terminology 63 concerning the metrics of carbonaceous aerosol. The use of different terminology is linked to the 64 different methodologies used to measure different physical or chemical properties of BC. The scientific 65 community generally accepts the definitions that BC particles possess the following properties: (1) 66 strongly absorbing in the visual spectrum with an inverse wavelength (λ) dependence (i.e., λ^{-1}) (Bond 67 and Bergstrom 2006), (2) refractory in nature with a vaporization temperature near 4000 K (Schwarz et 68 al., 2006), (3) insoluble in water and common organic solvents (Fung, 1990), (4) fractal-like aggregates of 69 small carbon spherules (Kittelson, 1998), (5) containing a large fraction of graphite-like sp²-bonded carbon atoms (Bond et al., 2013; Petzold et al., 2013), and (6) chemically inertness in the atmosphere 70 71 (Bond et al. 2013). In this article, the recommendation from Petzold et al. (2013) is adopted as the 72 definition of BC whenever the context of climate effects impacted by strong light-absorption 73 carbonaceous substance is mentioned. EC is referred to as the carbon mass determined from the 74 thermal evolution analysis (TEA) or thermal optical analysis (TOA) of carbonaceous materials at the 75 highest temperature set point (e.g., >550 °C) under an oxygenated environment. It is also assumed that 76 ambient EC and BC concentrations time series correlate with each other.

TOA and TEA have been applied in many long-term monitoring networks with various protocols to quantify OC and EC concentrations from aerosol deposits on quartz-fiber filters (Birch and Cary, 1996; Cachier et al., 1989; Cavalli et al., 2010; Chow et al., 1993; Huang et al., 2006; Huntzicker et al., 1982) due to the simplicity in filter sample collection and the analytical procedures. TOA and TEA provide a direct measurement of the carbon mass in the collected PM mass. One of the limitations of TOA and TEA is the need for sufficient sampling time to accumulate enough mass for precise measurements (i.e., ensuring a high signal to noise ratio) which constrains the temporal resolution of such samples. In addition, EC and OC are defined differently in different protocols and could affect the absolute mass
values measured. Generally, OC is quantified under a pure helium (He) atmosphere at a low heating
temperature whereas EC is quantified under an oxygen (O₂)/He atmosphere at high temperatures.
Estimates of total carbon (TC=OC+EC) derived from different TOA and TEA methods are generally
consistent, whereby the differences in OC and EC estimates could vary from 20 to 90%, and often larger
differences are found for EC, owing to its smaller contribution to TC (Cavalli et al., 2010; Chow et al.,
1993; 2001; 2005; Countess 1990; Watson et al., 2005; Hand et al., 2012).

91 During thermal analysis, some of the OC chars to form pyrolyzed organic carbon (POC) when 92 heated in the inert He atmosphere, darkening the filter (Chow et al., 2004; Watson et al. 2005). When 93 O₂ is added, POC combusts to EC, resulting in an overestimation of EC of the filter. The formation of 94 POC depends on the nature of the organic materials, the amount of the oxygenated compounds in the 95 collected particles, the rate, duration, and temperature of the heating, and the supply of O_2 in the 96 carrier gas (Cachier et al. 1989; Chan et al., 2010; Han et al. 2007; Yang and Yu, 2002). POC in TOA is 97 estimated by monitoring reflectance and/or transmittance of a 633-650 nm laser beam, which is termed 98 thermal optical reflectance (TOR) or thermal optical transmittance (TOT) method, respectively. When 99 the reflected or transmitted laser signal returns to its initial intensity at the start of the analysis (i.e., at 100 OC/EC split point), it is assumed that artifact POC has left the sample and the remaining carbon belongs 101 to EC. The carbon mass before the split point is defined as OC whereas that after the split point is 102 defined as EC. POC is defined as the mass determined between the time when O₂ is introduced and the 103 OC/EC split point. Different from TOA, the TEA used in this study applies a different approach for POC 104 determination (see below).

105 Quartz-fiber filters adsorb organic vapors (Chow et al., 2009; Turpin et al., 1994; Viana et al., 106 2006; Watson et al., 2010), resulting in non-PM contributions to OC and charring enhancement within 107 the filter. These vapors are adsorbed passively when the filter is exposed to air and more so as air is 108 drawn through the filter during PM sampling. Sampling at low filter face velocities for long period of 109 time could lead to more adsorption (McDow and Huntzicker, 1990), while using high filter face velocities 110 for longer sample durations may result in evaporation of semi-volatile compounds as negative artifact 111 (Khalek, 2008; Sutter et al., 2010; Yang et al., 2011). The positive OC artifact from adsorption usually 112 exceeds the negative evaporation artifact, especially at low temperatures, resulting in OC 113 overestimation (Watson et al., 2009; WMO, 2016). This can be corrected by subtracting the OC

concentration from field blanks or backup filters located downstream of a Teflon-membrane or quartz fiber filter (Chow et al., 2010; Watson et al., 2005; 2010).

Previous studies further suggested that TOT could over-estimate the POC mass more than TOR, resulting in higher POC (and lower EC) because of the charring of the adsorbed organic vapors within the filter (Chow et al 2004; Countess 1990). Since only a portion (0.5-1.5 cm²) of the filter is analyzed, inhomogeneous PM deposits add to measurement uncertainty when OC and EC are normalized to the entire filter deposit area. Deposits that are light or too dark can cause unstable laser signals that affect the OC/EC split (Watson et al., 2005).

122 The short lifetime of atmospheric aerosols (in days to weeks) and the different chemical and 123 microphysical processing that occur in the atmosphere result in high spatial and temporal variations of 124 aerosol properties. To facilitate the determination of the trends in emission changes and evaluation of 125 the effectiveness of emission mitigation policies (Chen et al. 2012), long-term consistent atmospheric 126 measurements are required, including aerosol carbon fractions. The emission sources of OC and EC at a 127 regional and global scales are often constrained through the use of atmospheric transport models in 128 conjunction of long-term OC and EC measurements (Collaud Coen et al., 2013; Huang et al., 2018). 129 Usually an integration of datasets from different networks is necessary for sufficient spatial coverage. 130 The objective of this study is to conduct an inter-comparison study for evaluating the comparability and 131 consistency of ten years co-located carbonaceous aerosol measurements at Egbert made by three North 132 American networks (Interagency Monitoring of PROtected Visual Environments, Canadian Air and 133 Precipitation Monitoring Network, and Canadian Aerosol Baseline Measurement), all of which use 134 different sampling instruments, frequencies, durations, analytical methods, and artifact corrections. 135 This inter-comparison study is also expected to provide some suggestions/recommendations for 136 improving the compatibility and consistency of long-term measurements.

137 Sampling and Measurements

138 Sampling Site

The sampling station is the Center for Atmospheric Research Experiments (CARE) located near Egbert, Ontario (44°12′ N, 79°48′ W, 251 m a.s.l.), Canada. This station is owned and operated by Environment and Climate Change Canada (ECCC), and is located 70 km NNW of the city of Toronto. There are no major local anthropogenic sources within about 10 km of the site. Air that reaches this site from southern Ontario and the northeastern United States typically carries urban or anthropogenic 144 combustion pollutants that were emitted within last two days (Rupakheti et al. 2005; Chan and

145 Mozurkewich 2007; Chan et al., 2010). Air from the north generally contains biogenic emissions and is

often accompanied with the presence of SOA during summer (Chan et al., 2010; Slowik et al., 2010).

147 Table 1 compares the instrument and analytical specifications among the three networks.

148 *The Interagency Monitoring of PROtected Visual Environment Network*

149 The IMPROVE network, established in 1987, includes regional-scale monitoring stations for 150 detecting visibility trends, understanding long-term trends, and evaluating atmospheric processes 151 (Malm 1989; Malm et al., 1994; Yu et al., 2004). IMPROVE operates about 150 sites and provides long-152 term records of PM₁₀ and PM_{2.5} (particles with aerodynamic diameter less than 10 and 2.5 microns, 153 respectively) mass as well as PM_{2.5} composition, including anions (i.e., chloride, nitrate, and sulfate), and 154 carbon (OC and EC). IMPROVE 24-hour samples at Egbert were acquired once every third day from 2005 155 to 2015. The sampling period was from 08:00 to 08:00 local standard time (LST) except for August 16, 156 2006 through October 24, 2008 (from 00:00 to 00:00 LST). Module C of the IMPROVE sampler uses a 157 modified air-industrial hygiene laboratory (AIHL) cyclone with a 2.5 μ m cut point at a flow rate of 22.8 158 liters per minute (L/min). PM samples were collected onto a 25 mm diameter quartz-fiber filter (Tissue 159 quartz, Pall Life Sciences, Ann Arbor, MI, USA), which were pre-fired at 900°C for four hours. Once 160 sampled, filters were stored in freezer until they were ready to be analyzed in the DRI laboratory in 161 Reno. All samples were analyzed by the IMPROVE_A thermal/optical reflectance protocol (Figure S1a; 162 Supplementary information) (Chow et al., 2007) as shown in Table S1 (Supplementary information). The 163 IMPROVE data (denoted as IMPROVE_A TOR) were obtained from the website 164 http://vista.cira.colostate.edu/IMPROVE (Malm et al., 1994; IMPROVE, 2017).

165 **The Canadian Air and Precipitation Monitoring Network**

CAPMON was established in 1983 to understand the source impacts of acid rain-related
 pollutants from long-range transport to the Canadian soil and atmosphere. The network operates 30
 regionally representatives sites (as of 2015) across Canada with most located in Ontario and Quebec.
 Measurements include PM, trace gases, mercury (both in air and precipitation), tropospheric ozone, and
 multiple inorganic ions in air and precipitation. In addition, a few number of sites include carbon (OC
 and EC) meausrements (https://www.canada.ca/en/environment-climate-change/services/air pollution/monitoring-networks-data/canadian-air-precipitation.html).

173 Twenty-four-hour samples (08:00 to 08:00 LST) were acquired every third day from 2005 to 174 2015 using the Modified Rupprecht and Patashnick (R&P) Model 2300 PM_{2.5} Speciation Sampler with 175 ChemComb cartridges and PM_{2.5} impactor plates with impactor foam to direct particles onto a 47 mm 176 diameter tissue quartz-fiber filter (Thermo Scientific, Waltham, MA, USA) operated at 10 L/min. 177 Samples were made on the same date when the IMPROVE samples were collected. A second parallel 178 cartridge is configured with a 47 mm front Teflon-membrane filter and a quartz-fiber backup filter to 179 estimate vapor adsorption artifact. All guartz-fiber filters were pre-fired at either 800°C or 900°C for 180 over two hours and cooled at 105°C overnight and stored in freezer (-15 °C) before being used for 181 sampling. All sampled filters were shipped cold and stored in freezer until they are ready to be analyzed 182 in the CAPMoN laboratory in Toronto.

183 Carbon was determined using the Sunset laboratory-based carbon analyzer (Sunset Laboratory 184 Inc., OR, USA; http://www.sunlab.com/) following the IMPROVE-TOT protocol from 2005 to 2007 185 (denoted as Sunset-TOT), then by DRI Model 2001 Thermal/Optical Carbon Analyzer following the 186 IMPROVE-TOR protocol (denoted as DRI-TOR) from 2008 to 2015 (Chow et al., 1993). As shown in Table 187 S1, the temperature settings for IMPROVE protocol (i.e., DRI-TOR) for CAPMoN samples are lower than 188 those of IMPROVE_A TOR protocol for IMPROVE samples by 20°C to 40°C (Figure S1b). Overall, Chow et 189 al. (2007) found that the small difference in the temperature-ramp between these protocols results in 190 correlated but different OC, EC, and TC mass.

191 The Canadian Aerosol Baseline Measurement Network

192 The Climate Chemistry Measurements and Research (CCMR) Section in the Climate Research 193 Division of ECCC has operated the Canadian Aerosol Baseline Measurement (CABM) network since 2005 194 to acquire data relevant to climate change (https://www.canada.ca/en/environment-climate-195 change/services/climate-change/science-research-data/greenhouse-gases-aerosols-196 monitoring/canadian-aerosol-baseline-measurement-program.html). The CABM network includes 6 197 sites (as of 2016) for aerosol chemical, physical, and optical measurements that cover ecosystems at 198 coastal, interior urban/rural areas, boreal forests, and the Arctic. Measurements are intended to 199 elucidate influences from various emission sources on regional background air, including biogenic 200 emissions, biomass burning as well as anthropogenic contributions from industrial/urban areas. 201 The CABM filter pack system uses a PM_{2.5} stainless steel cyclone (URG-2000-30EHS) operated at

202 16.7 L/min for sampling from 2006 to 2015 with an operator manually changing the 47 mm quartz-fiber

203 filter on a weekly basis. All quartz-fiber filters were pre-fired at 900°C overnight prior being sampled. 204 Once sampled, filters were shipped cold and then stored in freezer until they were ready to be analyzed 205 in the CCMR laboratory in Toronto. A TEA method, EnCan-Total-900 (ECT9), developed by Huang et al. 206 (2006) and refined later (Chan et al., 2010), was used to analyze the OC, POC, and EC on the guartz-fiber 207 filters using a Sunset laboratory-based carbon analyzer. The ECT9 protocol was developed to permit 208 stable carbon isotope (^{13}C) analysis of the OC and EC masses without causing isotope fractionation, as it 209 was demonstrated by Huang et al. (2006). This method first heats the filter at 550°C and 870°C for 600 s 210 each in the He atmosphere to determine OC and POC (including carbonate carbon; CC), respectively, and 211 then combusts the sample at 900°C under 2% O_2 and 98% He atmosphere for 420 s to determine EC 212 (Figure S1c and Table S1). The ECT9 POC definition (released as CO₂ at 870 °C) includes the charred OC, 213 and some calcium carbonate (CaCO₃) that decomposes at 830°C, as well as any refractory OC that is not 214 combusted at 550°C. Chan et al. (2010) found that POC determined by ECT9 was proportional to the 215 oxygenated compounds (e.g., aged aerosol from atmospheric photochemical reaction) and possibly 216 humic-like materials. Consistent with the IMPROVE_A TOR protocol (Chow et al., 2007), OC is defined as 217 the sum of OC and POC, as CC is usually negligible in $PM_{2.5}$.

CABM sites are also equipped with Particle Soot Absorption Photometer (PSAP; Radiance Research, Seattle, WA, USA) that continuously monitor aerosol light absorption at 1 minute time resolution, as changes in the amount of light transmitted through a quartz-fiber filter. Assuming the mass absorption coefficient (MAC) for aerosol is constant at Egbert, the one minute PSAP absorption measurements are linearly proportional to the BC or EC concentrations. In this study, five years of PSAP data (2010-2015) collected at Egbert was used to assess the impact of different sampling duration on the derived monthly averages EC values.

225 Differences in Sampling and Analysis among Networks

226 Depending on the sharpness (i.e., slope) of the inlet sampling effectiveness curve (Watson et al., 227 1983), different size-selective inlets may introduce measurement uncertainties. CAPMON uses 228 impactors whereas CABM and IMPROVE use cyclones. Impactor may have larger pressure drops across 229 the inlet that might enhance semi-volatile PM evaporation. Larger solid particles might bounce off when 230 in contact with the impactor and be re-entrained in the PM_{2.5} samples if the impactor is overloaded 231 (Flagan and Seinfeld, 1998; Hinds, 1999). Atmospheric mass size distributions typically peak at about 10 232 μ m with a minimum near 2.5 μ m, therefore, the difference in mass collected with different impactors or 233 cyclones among the three networks is not expected to be large (Watson and Chow, 2011). Analyzing OC

and EC content by TEA or TOA also subject to a number of artifacts, including adsorption of volatile
 organic compound (VOC) gases by quartz-fiber filter, leading to positive artifact, and evaporation of
 particles, leading to negative artifact (Malm et al., 2011).

The small filter disc (25 mm diameter) and high flow rate (22.8 L/min) in the IMPROVE sampler result in a 5- to 7-fold higher filter face velocity (i.e., 107.7 cm/s) than that for the CAPMoN and CABM samplers (16-20 cm/s). McDow and Huntzicker (1990) assert that higher filter face velocity may reduce sampling artifacts. However, very high face velocity (>100 cm/s) may enhance OC volatilization (Khalek 2008).

Both IMPROVE and CAPMoN networks correct for vapor adsorption, while CABM network does not. For CAPMoN measurements, the organic artifact derived from each 24-hour backup quartz-filter was subtracted from the corresponding OC measurement. For IMPROVE measurements (up until 2015), monthly median OC value obtained from the backup quartz-filters from 13 sites (not including Egbert) was subtracted from all samples collected in the corresponding month. Monthly averaged OC values were then derived from the 24-hour artifact corrected measurements.

248 Multiple studies show that using the same TOA protocol on both DRI and Sunset carbon 249 analyzers can produce comparable TC concentrations (Chow et al., 2005; Watson et al., 2005). 250 However, large differences in EC are found between the reflectance and transmittance POC correction 251 (Chow et al., 2004; 2005; Watson et al., 2005). Difference in OC and EC definitions among different TOA 252 and TEA protocols introduces measurement uncertainties. Among the TOA methods, how POC is 253 determined from the laser signals at different temperatures in the inert He atmosphere introduce 254 uncertainties. Large uncertainties in laser transmittance were found for lightly- and heavily-loaded 255 samples (Birch and Cary, 1996). For the CABM samples, the POC determined at 870 °C by ECT9 256 represents different OC properties and does not equal the charred OC obtained by Sunset-TOT, DRI-TOR, 257 or IMPROVE_A TOR.

Both IMPROVE and CAPMON data sets are once every third day 24-hour measurements collected on the same date while the CABM data is weekly integrated samples. A comparison between the integrated weekly samples and 24-hour samples have already been done by Yang et al. (2011) and therefore will not be repeated here. Based on two years of Egbert measurements (2005-2007), Yang et al. (2011) suggested that integrated weekly samples might experience reduced vapor adsorption but increased losses of semi-volatile organics leading to lower OC measurements. Weekly EC values were higher than those from 24-hour samples, which were attributed to the higher analytical uncertainties for
the lower loadings on the 24-hr samples (Yang et al., 2011).

266 Five years (2010-2015) of real-time (1 min average) PSAP particle light absorption 267 measurements (at 567 nm) was used here as a proxy common EC data set to assess the effect of 268 different sample duration on monthly average EC concentrations. First, the 1 min PSAP data was 269 averaged to 24-hour once in every three day samples and integrated weekly samples, respectively, and 270 the comparison of the two data sets are compared in Figure 1a. The results demonstrate that both data 271 sets capture the variations adequately. Monthly averages derived from the two sets of measurements 272 show highly correlated results (r=0.78; Figure 1b) and a slope of 0.96 (Figure 1c). Assuming the 273 variations in light absorption can represent the variations in EC, these results suggest that monthly 274 averaged EC based on integrated weekly sampling is about 4% lower than the monthly averaged EC 275 based on 24 hour sampling.

276 Results and Discussions

277 NIST urban dust standard comparison

278 The National Institute of Standards and Technology (NIST) Urban Dust Standard Reference 279 Material (SRM) 8785 Air Particulate Matter on Filter Media is intended primarily for use to evaluate 280 analytical methods used to characterize the carbon composition of atmospheric fine PM (Cavanagh and 281 Watters, 2005; Klouda et al., 2005). These samples were produced by resuspension of the original SRM 282 1649a urban dust sample, followed by collection of the fine fraction (PM_{2.5}) on quartz-fiber filters 283 (Klouda et al., 2005; May and Trahey, 2001). Past studies on SRM 1649a and SRM 8785 have shown 284 consistent composition and both samples were supplied with certified values for OC and EC (Currie et 285 al., 2002; Klouda et al., 2005). The consistency between the ECT9 and the IMPROVE_A TOR analytical 286 methods was assessed by analyzing NIST SRM 8785 filters. Four SRM 8785 filters with mass loading of 287 624-2262 μg were analyzed following the ECT9 method by the ECCC laboratory and the IMPROVE A TOR 288 protocol by the DRI laboratory during 2009-2010.

The values in the SRM 8785 certificate were reported in grams of OC or EC per grams of PM mass, which are average mass ratios based on analysis of a small numbers of randomly selected samples. Figure 2a-c shows that measurements by IMPROVE_A TOR protocol were within uncertainties of the certificate values. Ratios measured with ECT9 were greater, but not significantly different from the certificate values. When fitting the ECT9 measurements to the IMPROVE_A TOR measurements using a linear regression (Figure 3a-c), good correlations (r=0.9-0.99) were observed with 21-25% higher
in values by the ECT9 method than the IMPROVE_A TOR.

The parameter EC/TC, calculated based on the reported certificate values, were compared with the average EC/TC values determined from the inter-comparison study (ICP) by the DRI group (using IMPROVE_A TOR) and the ECCC group (using ECT9) (Figure 2d). These results show that EC/TC reported by both analytical methods were statistically the same as the certificate value.

Finally, the EC/TC value was further verified by analyzing SRM 1649a samples with the ECT9 method. The combusted CO₂ from OC, EC, and TC were analyzed for the isotope ratios (i.e., ${}^{14}C/{}^{12}C$) expressed as a fraction of modern carbon (i.e., FM_i is the ratio of ${}^{14}C/{}^{12}C$ in the sample i, relative to a modern carbon standard) for individual mass fractions (i.e., FM_{TC}, FM_{OC}, and FM_{EC}). Using isotopic mass balance, the EC/TC ratio can be derived from Eq. [1]:

305
$$FM_{TC} = FM_{OC} \times \left(1 - \frac{EC}{TC}\right) + FM_{EC} \times \frac{EC}{TC}$$
[1]

The ${}^{14}C/{}^{12}C$ ratio were determined by off-line combustion method at the Keck Carbon Cycle accelerator 306 307 mass spectrometry (KCCAMS) Facility at University of California Irvine. A FM_{TC} value of 0.512 was 308 obtained, which is close to certificate values that range from 0.505 to 0.61 (Currie et al., 2002). Average 309 measured values of FM_{OC} and FM_{EC} for the SRM 1649a via ECT9 were 0.634 (n=3) and 0.349 (n=3), 310 respectively. This yields an EC/TC ratio of 0.425, which is comparable to the ECT9 value of 0.44, and 311 close to the reported certificate value of 0.49 and the IMPROVE A TOR value of 0.47 (Figure 3d), 312 reconfirming a good separation of OC from EC using the ECT9 method. This analysis also confirms the consistency between the IMPROVE_A TOR and ECT9 methods. 313

314 Vapor Adsorption Corrections

315 Figure 4 shows the monthly averaged carbon concentration time series with and without the 316 artifact correction for CAPMoN samples over the period from 2005 to 2015. Vapor adsorption 317 contributes to a large amount of the measured OC (Figure 4a), but a negligibly amount to EC (Figure 4b) 318 and POC after 2008 (Figure 4c). The median vapor adsorption artifact was 0.79 μ g/m³ from 2008 to 319 2015 for DRI-TOR, representing about 50.9% of the uncorrected OC, compared to $0.92 \,\mu$ g/m³ (43.3% of 320 uncorrected OC) using the Sunset-TOT before 2008 (Supplemental Figure S2). Linear least square 321 regressions between corrected and uncorrected carbon in Figure 5 shows a slope of 0.52 for OC and 322 0.56 for TC with good correlations (r=0.93-0.94). Sunset-TOT measurements acquired prior 2008 are

mostly scattered around the regression line, with higher concentrations. On average, about 48% of the uncorrected OC ($0.84 \mu g/m^3$) can be attributed to vapor adsorption. The low filter face velocity (15.5 cm/s) in CAPMoN samples could be one of the contributing factors.

Figure 5c indicates that artifact corrected EC concentrations are 7.8% (0.02 µg/m³) lower than the uncorrected values. The artifact magnitude is close to the detection limit of 0.022 µg/m³ (0.197 µg/cm²) and within analytical uncertainties (Chow et al., 1993). Some Sunset-TOT EC measurements are scattered from the regression line, indicating a more accurate and consistent adsorption correction for DRI-TOR (Figure 5b). Although not expected to impact EC concentration, vapor adsorption directly affects POC correction and thus influences EC mass determination.

332 Figure 5d shows that 4.3% (0.01 μ g/m³) of POC was caused by vapor adsorption using the DRI-333 TOR protocol. For Sunset-TOT, however, up to 21.1% (0.17 μ g/m³) of the POC was detected on the 334 backup filter. Note that POC is part of OC and is a charring correction in the DRI-TOR and Sunset-TOT 335 protocols. Results show that filter transmittance is influenced by both surface and within filter charring 336 and EC from different sources have been observed to have different filter penetration depths (Chen et 337 al., 2004; Chow et al., 2004). Based on the available information from this study, an optical correction 338 by reflectance appears to be more appropriate and give more consistent results when POC 339 concentration is relatively large compared to EC (Chen et al., 2004). Regardless, the absolute POC and 340 EC concentrations were much lower than OC and the adsorption correction on TC is mostly attributed to 341 the OC artifact.

342 Since the IMPROVE aerosol samples were acquired at a higher filter face velocity (107.7 cm/s), it 343 is expected that the magnitude of the vapor adsorption correction would be smaller for the IMPROVE 344 samples. This is supported by the observations from Watson et al. (2009) at six anchor IMPROVE sites (i.e., Mount Rainier National Park, Yosemite National Park, Hance Camp at Grand Canyon National Park, 345 346 Chiricahua National Monument, Shenandoah National Park, and Okefenokee National Wildlife Refuge), 347 suggesting that vapor adsorption obtained from backup quartz filters represented about 23% of the 348 uncorrected OC values. Filter fibers are saturated over a long sampling interval (Khalek, 2008; Watson 349 et al., 2009), thus, artifacts for the CABM samples are expected to be lower relatively.

350 Comparison among IMPROVE, CAPMoN, and CABM Measurements

Figure 6 shows the temporal variations of the monthly averaged IMPROVE_A TOR, CAPMON
 Sunset-TOT, DRI-TOR, and CABM ECT9 measurements. Also included in the figure are the monthly

averaged temperature and the wind direction and speed (expressed in wind barbs). It is evident that
better correlations of TC, EC and OC were found between the protocols that use same POC correction
method (DRI-TOR and IMPROVE_A TOR) than between Sunset-TOT (which uses transmittance for POC
correction) and IMPROVE_A TOR (Table 2). Especially correlation of EC between Sunset-TOT and
IMPROVE_A TOR was poor.

Comparisons of the monthly averaged carbonaceous measurements among different networks are summarized in Figure 7. When fitting the monthly averaged DRI-TOR and Sunset-TOT measurements to IMPROVE_A TOR measurements using a linear regression fit through the origin (i.e., Regression 1) typically yields less than unity slopes (0.64-0.97; Table 2), suggesting that the carbonaceous masses reported by CAPMoN were in general lower than those by IMPROVE. Fitting the measurements allowing an intercept (i.e., Regression 2) typically yields least square slopes close to unity (>0.92) with small intercepts.

The effect of using transmittance or reflectance for POC determination is apparent. The Sunset-TOT POC correction is larger because transmittance is affected by the charred OC within the filter. This is consistent with the larger regression slopes in POC (Regression 1: 1.8) between Sunset-TOT and IMPROVE_A TOR protocol than the slope in POC (1.0) between the DRI-TOR and IMPROVE_A TOR protocol.

The ECT9 versus IMPROVE_A TOR via Regression 1 slopes are equal to or greater than unity,
ranging from 1.0 to 1.8 (Table 2). Linear regression with intercept (i.e., Regression 2) yields lower slopes
(0.6-1.7) with positive intercepts (0.06-0.18 µg/m³), signifying higher TC and EC concentrations for ECT9
samples. Higher intercepts (0.12-0.18 µg/m³) for TC, OC, and POC are consistent with ECT9
measurements uncorrected for vapor adsorption. However, the systematically higher TC, OC and EC by
21-25% via ECT9 relative to those via IMPROVE_A TOR in SRM 8785 could not be simply attributed to
the uncorrected vapor adsorption.

In specific, ECT9 OC concentrations are 15% higher than the IMPROVE_A TOR measurements
(Table 2) with good correlation (r=0.87; Table S2). the ECT9 method yielded 66-83% higher EC than
IMPROVE_A TOR, with moderate correlation (r=0.74). Differences in combustion temperatures for
OC/EC split determination could contribute to these discrepancies. Heating under an oxidative
environment at a constant temperature of 900 °C in the ECT9 protocol could combust more highly
refractory carbon than the IMPROVE_A TOR protocol, which only heats progressively from 580 °C to 840

°C. Another minor factor could include inhomogeneous deposition of mass loading on the filter spot.
When plotted on different scales, Figure S3 shows that the two EC data sets track well, capturing both
long-term trends and seasonal variations.

386 A slope approaching unity (1.00) was obtained when fitting the ECT9 POC to IMPROVE A TOR 387 POC through the origin (Figure 7d). Refitting the data allowing an intercept leads to a slope of 0.62 with 388 a y-intercept (0.12; Table 2), comparable in magnitude to the vapor adsorption artifact. The correlation 389 coefficient between ECT9 POC and IMPOVE A TOR POC is low (r=0.46; Table S3). However, correlation 390 between IMPROVE_A TOR POC and IMPROVE_A TOR OC is much higher (r=0.91), and even to a lesser 391 extent between IMPROVE_A TOR POC and IMPROVE_A TOR EC (r=0.71). In comparison, ECT9 POC has 392 weak correlation with ECT9 OC (r=0.65) and ECT9 EC (r=0.37). These observations show that the POC 393 definition in ECT9 is not dominated by charred OC correction and likely include the characterization of 394 other oxygenated organic materials as observed in Chan et al. (2010). Additional research is needed to 395 verify if ECT9 POC is proportional to SOA formation.

396 Seasonality in Carbon Concentration and Possible Origination

Figure 6 shows elevated carbon during summer, consistent with the observations from Yang et al. (2011) and Healy et al. (2017). A sigmoid function was applied here to characterize the relationship between ambient carbon concentration and ambient temperature. The Sigmoid function has a characteristic "S" shape and represents an integral of a Gaussian function. Relationships between carbon concentrations and ambient temperatures are illustrated in Figure S5. Apparent increases in OC and TC concentrations are found when ambient temperatures exceed about 10 °C; a phenomenon not as apparent in EC. EC from the week-long CABM samples are more scattered.

404 The TC, OC, and EC from all measurements are averaged and shown in Figure 8 with the 405 following best-fitted sigmoid functions:

406
$$TC = 1.053 + \left\{ \frac{3.558}{1 + exp\left(\frac{23.081 - T}{3.760}\right)} \right\}$$
[2]

407
$$OC = 0.780 + \left\{ \frac{1.838}{1 + exp\left(\frac{20.089 - T}{2.978}\right)} \right\}$$
[3]

408
$$EC = 0.239 + \left\{ \frac{1.446}{1 + exp\left(\frac{34.776 - T}{8.404}\right)} \right\}$$
[4]

Equations [2]-[4] show that lower limits of the observed TC, OC, and EC concentrations are 1.05, 0.78,
and 0.24 μgC/m³, with the half way of the maximum growth curve occurring at about 23 °C, 20 °C, and
35 °C, respectively. The predicted maximum concentrations for TC, OC, and EC are 4.61, 2.62, and 1.69
µgC/m³, respectively.

413 Preliminary analysis based on simple wind roses and Lagrangian particle dispersion transport 414 model (FLEXible PARTicle dispersion model) (Stohl et al., 2005) was conducted (Supporting Materials). 415 Results from the analysis appear to suggest that human activities (e.g., local transportation, residential 416 heating, and industrial activities), biogenic emissions (e.g., monoterpenes) from the boreal forest, SOA 417 formation, biomass burning, and transboundary transport could contribute to the variations of OC and 418 EC at Egbert in a complicated way (Ding et al., 2014; Chan et al., 2010; Leaitch et al., 2011; Passonen et 419 al., 2013; Tunved et al., 2006; Lavoué et al 2000; Healy et al. 2017), which requires additional research 420 to confirm. At Egbert, increasing ambient temperature from 10 °C to 20 °C leads to higher OC 421 concentrations from 0.84 to 1.61 µgC/m³ (91.7% increase) and EC concentration from 0.31 to 0.45 422 μ gC/m³ (45.2% increase). The temperature dependency of OC and EC suggests a potential climate 423 feedback mechanism consistent with the observations from Leaitch at al. (2011) and Passonen et al. 424 (2013).

425 Chan et al. (2010) showed that ECT9 POC possesses a positive relationship with oxygenated 426 organics and aged aerosol particles. The seasonality in ECT9 POC is compared with the average OC and 427 EC seasonality observed at Egbert (Figure 8d). Interestingly, the ECT9 POC concentration does not show 428 a gradual exponential shape of function as for OC and EC. Instead, it shows a small but obvious two-step 429 function when plotted against ambient temperature. The ECT9 POC temperature dependent results 430 (Figure 8d) suggest constant sources of background emissions of possible oxygenated organic 431 compounds that is independent from the measured OC, with additional secondary organic compound 432 (SOA) formation at higher temperatures (e.g., >15 °C). Future study is needed to verify this.

433 Summary of the Inter-comparison Study

Ten years of OC and EC measurements at Egbert were obtained from three independent
networks (IMPROVE, CAPMON, CABM) and observable differences in carbon concentrations were
attributed to different sampling methods, analytical protocols, sampling time, and filter artifact
corrections. Vapor adsorption did not affect EC values but contributed 20-50% of the measured OC,
depending on the sampling filter face velocity. The higher TC and OC concentration of the CABM

measurements by 20-30% and 15%, respectively, compared to the IMPROVE measurements could be
partially due to the absence of vapor adsorption correction. These results are consistent with other
inter-comparison study before data adjustments (Hand et al., 2012). The differences in analytical
protocol also play a role in causing higher carbon values, supported by the higher TC, OC and EC values
from the SRM8785 analysis obtained by the ECT9 method compared to those by IMPROVE_A TOR
method. Pyrolyzed OC (POC) from ECT9 is shown to be more than a charring correction and more
research is needed to develop its relationship with SOA.

Important observations from the inter-comparison study are: (1) CAPMON DRI-TOR TC, OC, and
EC are 5-17%, 7-16%, and 7-18% lower than the corresponding masses from IMPROVE_A TOR. (2)
CAPMON Sunset-TOT TC, OC, and EC are lower than the IMPROVE_A TOR values by up to 30%, 15%, and
75%. (3) CABM TC, OC and EC by ECT9 are higher than the IMPROVE_A TOR values by 20-30%, 0-15%,
and 60-80%, respectively.

451 Carbon concentrations observed from all three networks exhibited a non-linear positive 452 dependency with ambient temperature, which can be characterized by a sigmoid function. Although 453 further research is needed, preliminary observations suggested that increased anthropogenic activities, 454 urban emissions, SOA formation, forest fire emissions, and long range transport could have an impact on 455 the observed OC and EC at Egbert. The increase in OC concentration with temperature is consistent 456 with the climate feedback mechanisms reported from various studies. The different characteristic 457 temperature dependency of the ECT9 POC suggests the need for future investigation, which could 458 provide additional insights of SOA formation from acquired carbonaceous measurements.

459 Suggestions Going Forward

460 Long-term measurements play important roles for detecting the trends in atmospheric 461 compositions, constraining their emission changes, and allow for assessing the effectiveness of emission 462 mitigation policies at regional scales (WMO, 2016; 2003), provided that the measurements are 463 consistent and comparable across different networks. Recognizing the absence of a universally 464 accepted carbonaceous standard, long-term inter-comparison studies become challenging and even 465 more important. Echo the recommendations from the World Meteorological Organization (WMO) 466 guidelines and recommendations for long-term aerosol measurements (WMO, 2016; 2003), this study 467 illustrates the importance of measurement consistency (e.g., sampling method/procedures, analytical 468 instrument/method/protocols and data processing, quality assurance and quality control protocols)

- 469 within a network over a long period of time. As indicated in the guidelines, regular inter-comparison of
- 470 filter samples should be encouraged. These activities include analyzing exchanged common filter
- 471 samples and co-located filter samples. In addition, there is a need to develop proper reference
- 472 materials for assessing comparability and consistency, and incorporating the use of such reference as
- 473 part of the inter-comparison effort.
- 474 Nomenclature

475	AIHL	Air-industrial hygiene laboratory
476	AMS	Accelerator mass spectrometry
477	BC	Black carbon
478	CABM	Canadian Aerosol Baseline Measurement
479	CAPMoN	Canadian Air and Precipitation Monitoring Network
480	CARE	Center for Atmospheric Research Experiment
481	CCMR	Climate Chemistry Measurements and Research
482	DRI	Desert Research Institute
483	DRI-TOR	CAPMoN measurements using IMPROVE on DRI analyzer with TOR correction
484	EC	Elemental carbon
485	ECCC	Environment and Climate Change Canada
486	ECT9	EnCan-Total-900 protocol
487	FID	Flame ionization detector
488	FLEXPART	FLEXible PARTicle dispersion model
489	ICP	Inter-comparison study
490	IMPROVE	Interagency Monitoring PROtected Visual Environments
491	IMPROVE_A TOR	IMPROVE_A TOR protocol on DRI analyzer
492	KCCAMS	Keck Carbon Cycle accelerator mass spectrometry
493	MAC	Mass absorption coefficient
494	NIST	National Institute of Standard and Technology
495	OC	Organic carbon
496	PM	Particulate matter
497	POC	Pyrolyzed organic carbon
498	PSAP	Particle Soot Absorption Photometer
499	SOA	Secondary organic aerosol
500	SRM	Standard Reference Material
501	Sunset-TOT	IMPROVE TOT protocol on Sunset analyzer
502	TC	Total carbon
503	TEA	Thermal evolution analysis
504	TOA	Thermal optical analysis
505	TOR	Thermal optical reflectance
506	TOT	Thermal optical transmittance
507	UCI	University of California Irvine
508	WMO	World Meteorological Organization
509		

511 Acknowledgements

- 512 Authors would like to acknowledge Elton Chan and Douglas Chan of ECCC for providing the FLEXPART
- 513 model results and providing technical advice. IMPROVE measurements were obtained directly from the
- 514 IMPROVE web site (http://vista.cira.colostate.edu/IMPROVE/Data/QA_QC/Advisory.htm). IMPROVE is a
- 515 collaborative association of state, tribal, and federal agencies, and international partners. U.S.
- 516 Environmental Protection Agency is the primary funding source, with contracting and research support
- 517 from the National Park Service. IMPROVE carbon analysis was provided by Desert Research Institute.
- 518 Funding of this study was initiated by Climate Change Technology and Innovation Initiative (CCTI)
- 519 program, operated through Natural Resources Canada (NRCan), and supported by Clean Air Regulatory
- 520 Agenda (CARA) initiative and ECCC internal federal funding.

521 Supplementary Information:

- 522 The supplement related to this article, which includes additional details on thermal/optical analysis, the
- 523 experimental parameters used in different temperature protocols (IMPROVE, IMPROVE_A, ECT9),
- radiocarbon analysis, vapor adsorption uncertainty, seasonality of carbonaceous measurements, wind
- rose analysis, and FLEXPART back trajectory analysis at Egbert are available online.

526 Author Contributions:

- 527 TWC and LH wrote the paper, with KB, JW, JCC, CIC, GMS, KJ provide contributions to the article. All
- 528 authors commented on the manuscript.

529 **Competing interest**:

530 The authors declare that they have no conflict of interest.

531 References:

Beverly, R.K., Beaumont, W., Tauz, D., Ormsby, K.M., von Reden, K.F., Santos, G.M. and Southon, J.R.:
The Keck Carbon Cycle AMS Laboratory, University of California, Irvine: Status report, Radiocarbon 52,
301-309, 2010.

535 Birch, M. E. and Cary, R. A.: Elemental carbon-based method for monitoring occupational exposures to 536 particulate diesel exhaust, Aerosol Sci. Technol., 25, 221-241, 1996.

Bond, T. C. and Bergstrom, R. W.: Light absorption by carbonaceous particles: An investigative review,
Aerosol Sci. Technol., 40, 27-67, 2006.

- Bond, T. C., Doherty, S. J., Fahey, D. W., Forster, P. M., Berntsen, T., DeAngelo, B. J., Flanner, M. G.,
- 540 Ghan, S., Karcher, B., Koch, D., Kinne, S., Kondo, Y., Quinn, P. K., Sarofim, M. C., Schultz, M. G., Schulz,
- 541 M., Venkataraman, C., Zhang, H., Zhang, S., Bellouin, N., Guttikunda, S. K., Hopke, P. K., Jacobson, M. Z.,
- 542 Kaiser, J. W., Klimont, Z., Lohmann, U., Schwarz, J. P., Shindell, D., Storelvmo, T., Warren, S. G., and
- 543 Zender, C. S.: Bounding the role of black carbon in the climate system: A scientific assessment, J.
- 544 Geophys. Res. Atmos., 118, 5380-5552, 2013.
- 545 Cachier, H., Bremond, M. P., and Buat-Ménard, P.: Thermal separation of soot carbon, Aerosol Sci.
 546 Technol., 10, 358-364, 1989.
- 547 Cavalli, F., Viana, M., Yttri, K. E., Genberg, J., and Putaud, J. P.: Toward a standardized thermal-optical
 548 protocol for measuring atmospheric organic and elemental carbon: The EUSAAR protocol, Atmos. Meas.
 549 Tech., 3, 79-89, 2010.
- 550 Cavanagh, R. R. and Watters, Jr., R. L.: National Institute of Standards and Technology: Report of
- 551 Investigation Reference Material 8785: Air particulate matter on filter media (A fine fraction of SRM
- 552 1649a urban dust on quartz-fiber filter), 2005.
- 553 Chan, T. W. and Mozurkewich, M.: Application of absolute principal component analysis to size 554 distribution data: identification of particle origins, Atmos. Chem. Phys, 7, 887-897, 2007.
- 555 Chan, T. W., Huang, L., Leaitch, W. R., Sharma, S., Brook, J. R., Slowik, J. G., Abbatt, J. P. D., Brickell, P. C.,
- Liggio, J., Li, S. M., and Moosmüller, H.: Observations of OM/OC and specific attenuation coefficients
 (SAC) in ambient fine PM at a rural site in central Ontario, Canada, Atmos. Chem. Phys, 10, 2393-2411,
 2010.
- 559 Chen, L.-W. A., Chow, J. C., Watson, J. G., Moosmüller, H., and Arnott, W. P.: Modeling reflectance and
 560 transmittance of quartz-fiber filter samples containing elemental carbon particles: Implications for
 561 thermal/optical analysis, J. Aerosol Sci., 35, 765-780, 2004.
- 562 Chen, L.-W. A., Chow, J. C., Watson, J. G., and Schichtel, B. A.: Consistency of long-term elemental
 563 carbon trends from thermal and optical measurements in the IMPROVE network, Atmos. Meas. Tech., 5,
 564 2329-2338, 2012.
- 565 Chow, J. C., Watson, J. G., Pritchett, L. C., Pierson, W. R., Frazier, C. A., and Purcell, R. G.: The DRI
- Thermal/Optical Reflectance carbon analysis system: Description, evaluation and applications in U.S. air
 quality studies, Atmos. Environ., 27A, 1185-1201, 1993.
- Chow, J. C., Watson, J. G., Crow, D., Lowenthal, D. H., and Merrifield, T. M.: Comparison of IMPROVE and
 NIOSH carbon measurements, Aerosol Sci. Technol., 34, 23-34, 2001.
- 570 Chow, J. C., Watson, J. G., Chen, L.-W. A., Arnott, W. P., Moosmüller, H., and Fung, K. K.: Equivalence of 571 elemental carbon by Thermal/Optical Reflectance and Transmittance with different temperature
- 572 protocols, Environ. Sci. Technol., 38, 4414-4422, 2004.
- 573 Chow, J. C., Watson, J. G., Louie, P. K. K., Chen, L.-W. A., and Sin, D.: Comparison of PM_{2.5} carbon 574 measurement methods in Hong Kong, China, Environ. Poll, 137, 334-344, 2005.
- 575 Chow, J. C., Watson, J. G., Chen, L.-W. A., Chang, M.-C. O., Robinson, N. F., Trimble, D. L., and Kohl, S. D.:
- 576 The IMPROVE_A temperature protocol for thermal/optical carbon analysis: Maintaining consistency
- with a long-term database, J. Air Waste Manage. Assoc., 57, 1014-1023, 2007.

- 578 Chow, J. C., Watson, J. G., Lowenthal, D. H., and Chen, L.-W. A.: Climate change Characterization of
- 579 black carbon and organic carbon air pollution emissions and evaluation of measurement methods Phase
- 580 II: Characterization of black carbon and organic carbon source emissions, Desert Research Institute,
- 581 Reno, NVDRI 04-307, 2009.
- 582 Chow, J. C., Bachmann, J. D., Kinsman, J. D., Legge, A. H., Watson, J. G., Hidy, G. M., and Pennell, W. R.:
- 583 Multipollutant air quality management: Critical review discussion, J. Air Waste Manage. Assoc., 60, 584 1151-1164, 2010.
- 585 Collaud-Coen, M. C., Andrews, E., Asmi, A., Baltensperger, U., Bukowiecki, N., Day, D., Fiebig, M.,
- 586 Fjaeraa, A. M., Flentje, H., Hyvarinen, A., Jefferson, A., Jennings, S. G., Kouvarakis, G., Lihavainen, H.,
- 587 Myhre, C. L., Malm, W. C., Mihapopoulos, N., Molenar, J. V., O'Dowd, C., Ogren, J. A., Schichtel, B. A.,
- 588 Sheridan, P., Virkkula, A., Weingartner, E., Weller, R., and Laj, P.: Aerosol decadal trends Part 1: In-situ
- optical measurements at GAW and IMPROVE stations, Atmos. Chem. Phys, 13, 869-894, 2013.
- 590 Countess, R. J.: Interlaboratory analyses of carbonaceous aerosol samples, Aerosol Sci. Technol., 12, 114-591 121, 1990.
- 592 Currie, L. A., Benner, B. A., Jr., Cachier, H., Cary, R., Chow, J. C., Druffel, E. R. M., Eglinton, T. I.,
- 593 Gustafsson, Ö., Hartmann, P. C., Hedges, J. I., Kessler, J. D., Kirchstetter, T. W., Klinedinst, D. B., Klouda,
- G. A., Marolf, J. V., Masiello, C. A., Novakov, T., Pearson, A., Prentice, K. M., Puxbaum, H., Quinn, J. G.,
- 595 Reddy, C. M., Schmid, H., Slater, J. F., Watson, J. G., and Wise, S. A.: A critical evaluation of
- 596 interlaboratory data on total, elemental, and isotopic carbon in the carbonaceous particle reference
- 597 material, NIST SRM 1649a, Journal of Research of the National Institute of Standards and Technology,
- 598 107, 279-298, 2002.
- 599 Ding, L., Chan, T. W., Ke, F. and Wang, D. K. W.: Characterization of chemical composition and
- 600 concentration of fine particulate matter during a transit strike in Ottawa, Canada, Atmos. Environ., 89,601 433-442, 2014.
- Flagan, R. C. and Seinfeld, J. H.: Fundamentals of Air Pollution Engineering, Prentice Hall, EnglewoodCliffs, NJ, 1988.
- Fuller, K. A., Malm, W. C., and Kreidenweis, S. M.: Effects of mixing on extinction by carbonaceous
 particles, Journal of Geophysical Research, 104, 15941-15954, 1999.
- Fung, K. K.: Particulate carbon speciation by MnO2 oxidation, Aerosol Sci. Technol., 12, 122-127, 1990.
- Han, Y. M., Cao, J. J., An, Z., Chow, J. C., Watson, J. G., Jin, Z. D., Fung, K. K., and Liu, S.: Evaluation of the
 thermal/optical reflectance method for quantification of elemental carbon in sediments, Chemosphere,
 69, 526-533, 2007.
- Hand, J.L., Schichtel, B.A., Pitchford, M., Malm, W.C., and Frank, N.H. (2012): Seasonal composition of
 remote and urban fine particulate matter in the United States, Journal of Geophysical Research, 117,
 D05209, doi:10.1029/2011JD017122..
- Healy, R. M., Sofowote, U., Su, Y., Debosz, J., Noble, M., Jeong, C. H., Wang, J. M., Hilker, N., Evans, G. J.,
- Doerksen, G., Jones, K., and Munoz, A.: Ambient measurements and source apportionment of fossil fuel
 and biomass burning black carbon in Ontario, Atmos. Environ., 161, 34-47, 2017.
- Heintzenberg, J. (1989). Fine particles in the global troposphere: A review, Tellus, 41, 149-160.

- 617 Hinds, W. C.: Straight-line acceleration and curvilinear particle motion. In: Aerosol Technology.
- 618 Properties, Behavior, and Measurement of airborne Particles, 2nd Ed., John Wiley & Sons, Inc., New 619 York, 1999.
- 620 Huang, L., Brook, J. R., Zhang, W., Li, S. M., Graham, L., Ernst, D., Chivulescu, A., and Lu, G.: Stable
- 621 isotope measurements of carbon fractions (OC/EC) in airborne particulate: A new dimension for source 622 characterization and apportionment, Atmos. Environ., 40, 2690-2705, 2006.
- 623
- Huang, L.: The issue of harmonizing the methodologies for emission inventories of GHGs with those of 624 SLCFs (in terms of measurement perspective), IPCC Expert Meeting on Short Lived Climate Forcers,
- 625 Geneva, May 28-31, 2019, https://www.ipcc-nggip.iges.or.jp/public/mtdocs/1805_Geneva.html
- 626 Huntzicker, J. J., Johnson, R. L., Shah, J. J., and Cary, R. A.: Analysis of organic and elemental carbon in
- 627 ambient aerosols by a thermal-optical method. In: Particulate Carbon: Atmospheric Life Cycle, Wolff, G. 628 T. and Klimisch, R. L. (Eds.), Plenum Press, New York, NY, 1982.
- 629 IMPROVE: Interagency Monitoring of Protected Visual Environments, National Park Service, Ft. Collins, 630 CO, 2017.
- 631 Japar, S. M., Brachaczek, W. W., Gorse, R. A., Jr., Norbeck, J. H., and Pierson, W. R.: The contribution of 632 elemental carbon to the optical properties of rural atmospheric aerosols, Atmos. Environ., 20, 1281-
- 633 1289, 1986.
- 634 Khalek, I. A.: 2007 diesel particulate measurement research, Coordinating Research Council, Alpharetta, 635 GA, 2008.
- 636 Kittelson, D. B.: Engines and nanoparticles: A review, J. Aerosol Sci., 29, 575-588, 1998.
- 637 Klouda, G. A., Filliben, J. J., Parish, H. J., Chow, J. C., Watson, J. G., and Cary, R. A.: Reference material 638 8785: Air particulate matter on filter media, Aerosol Sci. Technol., 39, 173-183, 2005.
- 639 Lavoué, D., Liousse, C., Cachier, H., Stocks, B. J., and Goldammer, J. G.: Modeling of carbonaceous 640 particles emitted by boreal and temperate wildfires at northern latitudes, J. Geophys. Res. Atmos., 105, 641 26871-26890, 2000.
- 642 Leaitch, W. R., MacDonald, A. M., Brickell, P. C., Liggio, J., Sjostedt, S. J., Vlasenko, A., Bottenheim, J. W.,
- 643 Huang, L., Li, S. M., Liu, P. S. K., Toom-Sauntry, D., Hayden, K. A., Sharma, S., Shantz, N. C., Wiebe, H. A.,
- 644 Zhang, W., Abbatt, J. P. D., Slowik, J. G., Chang, R. Y. W., Russell, L. M., Schwartz, R. E., Takahama, S.,
- 645 Jayne, J. T., Ng, N. L.: Temperature response of the submicron organic aerosol from temperate forests, 646 Atmos. Environ., 45, 6696-6704, 2011.
- 647 Lesins, G., Chylek, P., and Lohmann, U.: A study of internal and external mixing scenarios and its effect
- 648 on aerosol optical properties and direct radiative forcing, J. Geophys. Res., 107, 4904,
- 649 10.1029/2001JD000973, 2002.
- 650 Malm, W. C.: Atmospheric haze: Its sources and effects on visibility in rural areas of the continental 651 United States, Env. Mon. Ass., 12, 203-225, 1989.
- 652 Malm, W. C., Sisler, J. F., Huffman, D., Eldred, R.A., and Cahill, T. A.: Spatial and seasonal trends in 653 particle concentration and optical extinction in the United States, J. Geophys. Res., 99, 1347-1370, 1994.

- Malm, W.C., Schichtel, B.A., and Pitchford, M.L.: Uncertainties in PM_{2.5} gravimetric and speciation
 measurements and what we can learn from them, J. Air & Waste Manage. Assoc., 61, 1131-1149, 2011.
- May, W. E. and Trahey, N. M.: National Institute of Standards and Technology: Certificate of Analysis
 Standard Reference Material 1649a: Urban dust, 2001.
- 658 McDow, S. R. and Huntzicker, J. J.: Vapor adsorption artifact in the sampling of organic aerosol: Face 659 velocity effects, Atmos. Environ., 24A, 2563-2571, 1990.
- Paasonen, P., Asmi, A., Petäjä, T., Kajos, M. K., Äijälä, M., Junninen, H., Holst, T., Abbatt, J. P. D., Arneth,
 A., Birmili, W., van der Gon, H. D., Hamed, A., Hoffer, A., Laakso, L., Laaksonen, A., Leaitch, W. R., PlassDülmer, C., Pryor, S. C., Räisänen, P., Swietlicki, E., Wiedensohler, A., Worsnop, D. R., Kerminen, V. M.,
 and Kulmala, M.: Warming-induced increase in aerosol number concentration likely to moderate climate
- 664 change, Nature Geoscience, 6, 438-442, 2013.
- 665 Petzold, A., Ogren, J. A., Fiebig, M., Laj, P., Li, S. M., Baltensperger, U., Holzer-Popp, T., Kinne, S.,
- 666 Pappalardo, G., Sugimoto, N., Wehrli, C., Wiedensohler, A., and Zhang, X. Y.: Recommendations for 667 reporting "black carbon" measurements, Atmos. Chem. Phys, 13, 8365-8379, 2013.
- 668 Rupakheti, M., Leaitch, W. R., Lohmann, U., Hayden, K., Brickell, P., Lu, G., Li, S. M., Toom-Sauntry, D.,
- 669 Bottenheim, J. W., Brook, J. R., Vet, R., Jayne, J. T., and Worsnop, D. R.: An intensive study of the size and 670 composition of submicron atmospheric aerosols at a rural site in Ontario, Canada, Aerosol Sci. Technol.,
- 671 *39,* 722-736*,* 2005.
- Santos, G.M., Moore, R., Southon, J., Griffin, S., Hinger, E., Zhang, D.: AMS 14C preparation at the
 KCCAMS/UCI Facility: Status report and performance of small samples. Radiocarbon, 49, 255-269, 2007.
- 674 Schulz, M., Textor, C., Kinne, S., Balkanski, Y., Bauer, S., Berntsen, T., Berglen, T., Boucher, O., Dentener,
- 675 F., Guibert, S., Isaksen, I. S. A., Iversen, T., Koch, D., Kirkevag, A., Liu, X., Montanaro, V., Myhre, G.,
- Penner, J. E., Pitari, G., Reddy, S., Seland, O., Stier, P., and Takemura, T.: Radiatives forcing by aerosols as
- derived from the AeroCom present-day and pre-industrial simulations, Atmos. Chem. Phys, 6, 5225-5246, 2006.
- ,
 - 679 Schwarz, J. P., Gao, R. S., Fahey, D. W., Thomson, D. S., Watts, L. A., Wilson, J. C., Reeves, J. M.,
 - Darbeheshti, M., Baumgardner, D. G., Kok, G. L., Chung, S. H., Schulz, M., Hendricks, J., Lauer, A.,
 - Kärcher, B., Slowik, J. G., Rosenlof, K. H., Thompson, T. L., Langford, A. O., Loewenstein, M., and Aikin,
 - 682 K.C.: Single-particle measurements of midlatitude black carbon and light-scattering aerosols from the
 - boundary layer to the lower stratosphere, J. Geophys. Res., 111, D16207, doi:10.1029/2006JD007076,
 - 684 2006.
 - Slowik, J. G., Stroud, C., Bottenheim, J. W., Brickell, P. C., Chang, R. Y. W., Liggio, J., Makar, P. A., Martin,
 - 686 R. V., Moran, M. D., Shantz, N. C., Sjostedt, S. J., van Donkelaar, A., Vlasenko, A., Wiebe, H. A., Xia, A. G.,
- Zhang, J., Leaitch, W. R., and Abbatt, J. P. D.: Characterization of a large biogenic secondary organic
 aerosol event from eastern Canadian forests, Atmos. Chem. Phys, 10, 2825-2845, 2010.
- 689 Stohl, A., Forster, C., Frank, A., Seibert, P., and Wotawa, G.: Technical note: The Lagrangian particle 690 dispersion model FLEXPART version 6.2, Atmos. Chem. Phys, 5, 2461-2474, 2005.
- Sutter, B., Bemer, D., Appert-Collin, J. C., Thomas, D., and Midoux, N.: Evaporation of liquid semi-volatile
 aerosols collected on fibrous filters, Aerosol Sci. Technol., 44, 395-404, 2010.

- Tunved, P., Hansson, H. C., Kerminen, V. M., Strom, J., Dal Maso, M., Lihavainen, H., Viisanen, Y., Aalto,
- P. P., Komppula, M., and Kulmala, M.: High natural aerosol loading over boreal forests, Science, 312,261-263, 2006.
- Turpin, B. J., Huntzicker, J. J., and Hering, S. V.: Investigation of organic aerosol sampling artifacts in the
 Los Angeles Basin, Atmos. Environ., 28, 3061-3071, 1994.
- Viana, M., Chi, X., Maenhaut, W., Cafmeyer, J., Querol, X., Alastuey, A., Mikuska, P., and Vecera, Z.:
- Influence of sampling artefacts on measured PM, OC, and EC levels in carbonaceous aerosols in an urban
 area, Aerosol Sci. Technol., 40, 107-117, 2006.
- Watson, J. G., Chow, J. C., Shah, J. J., and Pace, T. G.: The effect of sampling inlets on the PM₁₀ and PM₁₅
 to TSP concentration ratios, J. Air Pollut. Control Assoc., 33, 114-119, 1983.
- Watson, J. G.: Critical review: Visibility: Science and regulation, J. Air Waste Manage. Assoc., 52, 628-713, 2002.
- Watson, J. G., Chow, J. C., and Chen, L.-W. A.: Summary of organic and elemental carbon/black carbon
 analysis methods and intercomparisons, Aerosol Air Qual. Res., 5, 65-102, 2005.
- Watson, J. G., Chow, J. C., Chen, L. W. A., and Frank, N. H.: Methods to assess carbonaceous aerosol
 sampling artifacts for IMPROVE and other long-term networks, J. Air & Waste Manage. Assoc., 59, 898911, 2009.
- Watson, J. G., Chow, J. C., Chen, L.-W. A., and Wang, X. L.: Measurement system evaluation for fugitive
 dust emissions detection and quantification, Desert Research Institute, Reno, NV, 2010.
- 712 Watson, J. G., and Chow, J. C.: Ambient aerosol sampling in: Aerosol Measurement: Principles,
- techniques and applications, Third Edition, edited by Kulkarni, P., Baron, P.A., and Willeke, K., pp. 591614, Hoboken, NJ, USA: Wiley, 2011.
- 715 WMO/GAW aerosol measurement procedure: Guidelines and recommendations, WMO TD No. 1178,716 2003.
- WMO/GAW aerosol measurement procedures: Guidelines and recommendations, 2nd ed., WMO-No.
 1177, 2016.
- Yang, F., Huang, L., Sharma, S., Brook, J. R., Zhang, W., Li, S. M., and Tan, J. H.: Two-year observations of
 fine carbonaceous particles in variable sampling intervals, Atmos. Environ., 45, 2418-2426, 2011.
- 721 Yang, H. and Yu, J. Z.: Uncertainties in charring correction in the analysis of elemental and organic
- carbon in atmospheric particles by thermal/optical methods, Environ. Sci. Technol., 36, 5199-5204,
- 723 2002.
- Yu, S. C., Dennis, R. L., Bhave, P. V., and Eder, B. K.: Primary and secondary organic aerosols over the
- 725 United States: Estimates on the basis of observed organic carbon (OC) and elemental carbon (EC), and
- air quality modeled primary OC/EC ratios, Atmos. Environ., 38, 5257-5268, 2004.

Table 1 Specifications for the filter sampling systems and analytical instruments/methods used by the three networks.

	IMPROVE	CAI	CABM	
Data coverage period	2005-2015	2005-2007	2008-2015	2005-2015
Analytical instrument	DRI	Sunset	DRI	Sunset
Thermal/optical protocol	IMPROVE_A	IMPROVE	IMPROVE	ECT9
Pyrolyzed organic carbon detection	Reflect.	Transmit.	Reflect. & Transmit.	Retention time
Particle size selection method	Cyclone	Impactor plates	Impactor plates	Cyclone
Particle size cut off diameter (nm)	2.5	2.5	2.5	2.5
Sampling flow rate (L/min)	22.8	10.0	10.0	16.7
Filter media model	2500QAT-UP	2500QAT-UP	2500QAT-UP	2500QAT-UP
Quartz filter diameter (mm)	25	47	47	47
Filter deposition exposure area (cm ²)	3.53	10.75	10.75	13.85
Filter face velocity (cm/s)	107.65	15.50	15.50	20.09
Sampling frequency	Daily every 3 days	Daily every 3 days	Daily every 3 days	Integrated weekly
Daily sampled air volume (L/day)	31680	14400	14400	24048
Air volume per sample (m ³)	31.68	14.4	14.4	168.3
Positive artifact correction	Yes	Yes	Yes	No
Filter blank correction	Yes	No	No	Yes
Number of 24-h sample	1228	254	907	-
Number of weekly sample	-	-	-	476
Number of monthly averaged sample	124	28	93	117

730 **Table 2** Regression results (slope, correlation coefficient, and total number of points) obtained when fitting various CABM (ECT9) and CAPMoN

731 (Sunset-TOT & DRI-TOR) carbonaceous mass concentration time series against IMPROVE (IMPROVE_A TOR) measurements. IMPROVE_A TOR

and ECT9 measurements cover the period from 2005 to 2015. Sunset-TOT and DRI-TOR measurements cover the periods for 2005-2008 and

733 2008-2015, respectively. Regression 1 indicates the best-fitted slope through the origin. Regression 2 is the best-fitted slope with intercept (in

734 brackets).

	Regression 1	Regression 2	R	Ν
Sunset-TOT TC vs IMPROVE_A TOR TC	0.888±0.033	0.713±0.112 (0.301±0.186)	0.78	28
Sunset-TOT OC vs IMPROVE_A TOR OC	0.967±0.041	0.873±0.135 (0.125±0.170)	0.79	28
Sunset-TOT EC vs IMPROVE_A TOR EC	0.639±0.042	0.233±0.130 (0.171±0.053)	0.33	28
Sunset-TOT POC vs IMPROVE_A TOR POC	1.769±0.091	1.776±0.351 (-0.003±0.127)	0.70	28
DRI-TOR TC vs IMPROVE_A TOR TC	0.832±0.015	0.946±0.044 (-0.164±0.059)	0.91	93
DRI-TOR OC vs IMPROVE_A TOR OC	0.835±0.017	0.934±0.046 (-0.116±0.050)	0.90	93
DRI-TOR EC vs IMPROVE_A TOR EC	0.818±0.019	0.929±0.072 (-0.032±0.020)	0.81	93
DRI-TOR POC vs IMPROVE_A TOR POC	0.986±0.028	1.230±0.080 (-0.073±0.023)	0.85	93
ECT9 TC vs IMPROVE_A TOR TC	1.304±0.022	1.197±0.065 (0.164±0.093)	0.88	107
ECT9 OC vs IMPROVE_A TOR OC	1.149±0.021	1.004±0.056 (0.179±0.064)	0.87	107
ECT9 EC vs IMPROVE_A TOR EC	1.834±0.046	1.661±0.149 (0.056±0.046)	0.74	107
ECT9 POC vs IMPROVE_A TOR POC	0.998±0.031	0.615±0.082 (0.124±0.025)	0.59	107

735

- 737 **Figure 1** (a) Real-time Particle Soot Absorption Photometer (PSAP) measurements averaged to match
- the corresponding sampling frequencies used in different networks. (b) Monthly PSAP measurements
- derived from (a). (c) Comparison of the different sets of measurements from (b) with the 1:1 line shown
- 740 in red.



- 744 Figure 2 Comparison of the TC, OC, and EC measurements of the NIST SRM samples reported by the
- 745 ECCC and DRI groups during the inter-comparison study (ICP) conducted between 2009 and 2010.
- 746 "Reported" represent the published value in the NIST SRM certificate (Cavanagh and Watters, 2005).
- 747 Error bars represent uncertainties covering 95% confidence interval. In (d), the ECT9 value (in green)
- 748 represents the calculated EC/TC ratio determined based on stable carbon isotope measurement
- 749 obtained from the SRM 1649a sample (Currie et al., 2002).



- 752 **Figure 3** Comparison of: (a) TC, (b) OC, and (c) EC concentrations obtained from the same NIST SRM
- 8785 filters reported by ECCC following the TEA (ECT9) method and by DRI following the IMPROVE_A
 TOR protocol during the inter-comparison study in 2009/2010.







Figure 4 Monthly averaged CAPMoN (a) OC, (b) EC, and (c) POC mass concentration time series with and
 without vapor adsorption correction. Note that the y-axes in Figures 4b and 4c are on different scale.

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- 767 **Figure 5** Relationship between the monthly averaged CAPMoN vapor adsorption corrected and
- 768 uncorrected measurements for (a) TC, (b) OC, (c) EC, and (d) POC. Black solid markers represent the TOR
- 769 measurements (2008-2015) analyzed by the DRI analyzer (i.e., DRI-TOR). Red open markers represent
- the TOT measurements before 2008 analyzed by the Sunset analyzer (i.e., Sunset-TOT). The red line
- represents the best-fitted linear regression of all the DRI-TOR measurements through the origin. All the
- corresponding statistics (i.e., best-fitted slope, correlation coefficient, total number of measurement
- points) are included in the legend.





- **Figure 6** Monthly averaged (a) TC, (b) OC, (c) EC, and (d) POC concentration time series obtained from
- three different networks at Egbert. CAPMoN measurements before 2008 were obtained using Sunset-
- TOT method (in green) while measurements starting 2008 were obtained using DRI-TOR method (in
- 780 orange).





Figure 7 Comparison of the monthly averaged carbonaceous mass concentrations from the DRI-TOR (red
circles and orange triangles) and ECT9 (black squares) protocols against IMPROVE_A TOR protocol. The
different straight lines represent the linear regression best fitted line through the origin (i.e., Regression
1). The fitted parameters for all corresponding data sets with (Regression 2) and without (Regression 1)
the y-intercept are summarized in Table 2.



793 Figure 8 Figure shows the relationship of averaged (a) TC, (b) OC, and (c) EC concentrations from all

794 networks as a function of ambient temperature. Each data point represent the average value of all

795 network measurements within a 3°C temperature range. Uncertainties are standard deviations of the

- 796 measurements. Red curve represents the best-fitted Sigmoid function. Figure 10(d) shows the
- 797 seasonality of ECT9 POC compared to the average OC and EC seasonality. Black solid curve represents
- 798 the best-fitted Sigmoid function on all ECT9 POC measurements.

