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Published in:
Carbon

Link to article, DOI:
10.1016/j.carbon.2018.10.072

Publication date:
2019

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Malmborg, V. B., Eriksson, A. C., Török, S., Zhang, Y., Kling, K. I., Martinsson, J., ... Pagels, J. (2019). Relating aerosol mass spectra to composition and nanostructure of soot particles. Carbon, 142, 535-546. https://doi.org/10.1016/j.carbon.2018.10.072

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Relating aerosol mass spectra to composition and nanostructure of soot particles

Vilhelm B. Malmborg a,*, Axel C. Eriksson a, Sandra Török b, Yilong Zhang c, Kirsten Kling d,*, Johan Martinsson e,2, Edward C. Fortner f, Louise Gren g, Sanghoon Kook c, Timothy B. Onasch f, Per-Erik Bengtsson b, Joakim Pagels a

a Division of Ergonomics and Aerosol Technology, Lund University, Box 118, SE-22100, Lund, Sweden
b Division of Combustion Physics, Lund University, Box 118, SE-221 00, Lund, Sweden
c School of Mechanical and Manufacturing Engineering, University of New South Wales, Sydney, NSW, 2052, Australia
d National Research Centre for the Working Environment, 2100, Copenhagen, Denmark
e Division of Nuclear Physics, Lund University, Box 118, SE-221 00, Lund, Sweden
f Center for Aerosol and Cloud Chemistry, Aerodyne Research, Inc., 45 Manning Rd., Billerica, MA, 01821, USA

A R T I C L E   I N F O

Article history:
Received 14 May 2018
Received in revised form 21 September 2018
Accepted 21 October 2018
Available online 22 October 2018

A B S T R A C T

The composition and carbon nanostructure of soot are important parameters influencing health and climate effects, and the efficacy of soot mitigation technologies. We used laser-vaporization, electron-ionization aerosol mass spectrometry (or SP-AMS) to systematically investigate relationships between aerosol mass spectra, carbon nanostructure (HRTEM), and composition (thermal-optical carbon analysis) for soot with varying physicochemical properties. SP-AMS refractory black carbon concentrations (based on C$_{2+}$/C$_{20}$ clusters) were correlated to elemental carbon ($r = 0.98$, $p < 10^{-3}$) and equivalent black carbon (aethalometer) concentrations. The SP-AMS large carbon (C$_{21}$, midcarbons and fullerene carbons) fraction was inversely correlated to fringe length ($r = 0.97$, $p = 0.028$) and linearly correlated to the fraction of refractory organic carbon that partially pyrolyze during heating ($r = 0.89$, $p < 10^{-4}$). This refractory organic carbon material was incompletely detected with conventional aerosol mass spectrometry (flash vaporization at 600 °C). This suggests that (SP-AMS) refractory carbon cluster analysis provides insight to chemical bonding and nanostructures in refractory carbon materials, lowcarbons (C$_{20}$) indicate mature soot and large carbons indicate refractory organic carbon and amorphous nanostructures related to C$_{5}$-components. These results have implications for assessments of soot particle mixing state and brown carbon absorption in the atmosphere and enable novel, on-line analysis of engineered carbon nano-materials and soot characteristics relevant for climate and health.

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1. Introduction

Atmospheric soot particles form under incomplete combustion conditions in internal combustion engines, gas turbine engines (e.g., airplanes), wild and prescribed burns, and domestic wood-stoves [1–3]. Soot is an effective absorber of incoming solar radiation [4] and causes adverse health effects in humans [5–8]. As a result, soot is an important short-term climate forcer and a major contributor to air pollution [9,10]. Soot particles are generally composed of light absorbing refractory carbonaceous components often referred to as black carbon (BC) [11], ash, and condensed non-refractory semi-volatile and low volatility organic material [12]. Both fuel and combustion conditions strongly influence physical and chemical properties of soot. Soot can exhibit large variations in, for example, the carbon nanostructure [13,14], hydrogen to carbon ratio [15], surface functional groups [16], and condensed polycyclic aromatic hydrocarbons (PAHs) [15,17]. These variations in soot properties, in turn, influence the applications and strategies to mitigate soot emissions, climate forcing, and human health relevant aspects of the soot.

High-resolution transmission electron microscopy (HRTEM) has been widely applied to characterize the soot nanostructure...
It is found in HRTEM images that the refractory part of soot particles are composed of carbon layers, appearing as fringes, whose physical dimensions and organization reveal important information regarding the maturity and graphitization level of the particles [18,25,26]. Soot maturity is a qualitative description of the optical and physicochemical characteristics of soot. Soot maturity is commonly used to relate soot properties with time/position inside a flame, for example at increasing heights above the burner in premixed flames. As soot matures, the hydrogen to carbon ratio decreases [27,28] and the structural order of the carbon nanostructure and size of carbon lamellae have been found to increase [29,30]. Immature soot particles exhibit amorphous carbon nanostructures characterized by short and often highly curved fringes. Mature soot particles, in general, have a core-shell internal structure with short fringe segments forming amorphous core regions that are concentrically surrounded by a shell of longer and straighter carbon layers [31,32]. Characterization using HRTEM has revealed strong connections between the carbon nanostructure and reactivity towards oxidation [13,14,16,22,24,33]. This relationship can influence the ability to efficiently remove soot, for example in engine exhaust using diesel particulate filters. Mature soot with longer and straighter fringes (i.e., large and flat carbon layers) is less reactive towards oxidation. The reduced reactivity with increasing soot maturity is explained by reduced curvature (i.e., reduced tortuosity) and a relative decrease in the number of reactive edge-site atoms as the carbon layers grow in size [13,33]. Moreover, soot maturity and variations in composition and nanostructure influence the mass absorption cross section (MAC) [34] and absorption Angstrom exponent [11,35–38] which to a large extent determine the soot climate impact.

Soot formation mechanisms in different environments are intriguingly complex [17] and the different environments in which soot formation occur can result in vastly different soot properties. Although soot formation mechanisms have long remained elusive [39], the hydrogen-abstraction-C2H2-addition (HACA) mechanism [40,41] has been important in describing the formation and growth of aromatics and soot particles. The HACA framework identifies stabilomer sequences in order for aromatic growth reactions to become irreversible, a critical element also with respect to the growth of other forms of carbon such as fullerenes [40]. PAHs have long been recognized as soot precursor molecules [17,42] and incipient soot particles have been considered to form by the stacking of PAHs into three-dimensional structures [31,43]. Reilly et al. [44] identified a transition from PAHs to fullerenes in the first, very nascent, soot particles in a flame. Johnson et al. [45] identified the importance of low molecular weight (<400 u) PAHs with pentagonal ring systems and fullerene formation in the very early stages of formation of a carbon black soot (engineered soot nanoparticles). In the latter experiment, fullerenes and fullerene nanostructures were incorporated into the soot matrix simultaneously with high molecular weight PAHs annealed into the soot shortly after soot formation. The fullerenes were incorporated into the soot but also destroyed upon further growth and oxidation [45]. More recently, Johansson et al. [46] provided strong evidence for soot formation occurring from resonance-stabilized C5-containing radical species and clustering of hydrocarbons by radical-chain reactions (CHRCR). The CHRCR mechanism proceeds through both aromatic and aliphatic soot precursors and can explain the typical core-shell nanostructure of mature soot particles.

Particulate matter from low temperature combustion processes often contain refractory organic carbon not removed by heating to 300 °C–650 °C [35,47,48]. This group of compounds is challenging to characterize due to its low volatility and thermal degradation due to pyrolysis. In fact, on-line state of the art techniques such as combinations of BC absorption monitors and conventional aerosol mass spectrometry may not be capable of detecting this group of species [35,49]. Yet, due to its aromatic character and enhanced UV-VIS (brown carbon) absorption, soot containing these refractory organic carbon compounds may induce additional health and climate effects. Furthermore, there may be relationships and overlap between disordered materials constituting immature soot and refractory organic carbon (e.g., [50]). Therefore, there is a need to compare methods used in different disciplines to decipher diverse aspects of these materials and to validate novel in-situ techniques to classify these groups of compounds.

An aerosol mass spectrometer capable of detecting refractory carbon components is the soot particle aerosol mass spectrometer (SP-AMS) [51]. In the SP-AMS, aerosol particles are focused into high vacuum (10⁻⁷ mbar) by an aerodynamic lens. Infrared light absorbing particles, such as BC-containing particles, vaporize (in-flight) when the focused beam of aerosol particles passes through an intracavity Nd:YAG IR laser (1064 nm). The vapors are ionized using electron ionization (70 ev) and subsequently detected in a high-resolution time-of-flight mass spectrometer. Molecules fragment upon vaporization and ionization in the SP-AMS. The SP-AMS refractory black carbon (rBC) concentration is determined from the elemental carbon cluster fragments C₅₋₆ (lowcarbons). Lowcarbon signal intensities are generally dominated by C₃ following by C₄, and have been shown to correlate with optically determined equivalent BC concentrations for mature soots of different sources [51–53]. Some soots (select flame soots, carbon blacks and biomass combustion emissions) can produce SP-AMS mass spectra with significant ion signals at higher carbon numbers [51–55] in the range of C₇₋₁₀ (midcarbons) and C₁₀₋₁₃ (fullerenes). These large carbons (C₇₋₁₀), observed for some soots in SP-AMS mass spectra, may have ring structures (midcarbons) and fullerene structures [52] due to the stability of these structures at large carbon numbers [56].

Both fullerenes and fullerene soot nanostructures incorporate five membered aromatic rings (C₅). Formation of C₅-containing nanostructures have been suggested to proceed through partial oxidation of benzene to the phenoxy radical and subsequent CO loss, resulting in C₅ [18]. Laser heating (at 1064 nm and other) can result in annealing of the soot nanostructure [57,58]. The degree of annealing depends on laser power (or heating rate) [58] and the initial order of the soot nanostructure [59]. Annealing of graphitizable and non-graphitizable carbon materials results in different end structures. C₅ structures were recently identified as the cause of non-graphitizability in non-graphitizable chars [60]. Whether midcarbons and fullerene carbons are existing structures in the soot or form upon annealing in the SP-AMS laser interaction (for example from C₅-containing precursors) prior to its complete vaporization has not yet been verified. However, these large carbons have been hypothesized to relate to variations in the initial soot carbon nanostructure and immature soot properties [51–54]. This study presents the first attempt to test this hypothesis by benchmarking the on-line SP-AMS mass spectra analysis against off-line HRTEM analysis of the soot carbon nanostructure and offline thermal-optical carbon (OC/EC) analysis of the soot composition. In this study, soot of varying maturity and diverse physicochemical properties were generated using a miniature combustion aerosol standard (miniCAST) soot generator. The results suggest strong relationships between the refractory carbon cluster (rBC) distribution in SP-AMS mass spectra, the soot carbon nanostructure, and particle composition.
2. Method

2.1. Experimental overview

The experimental set-up to examine the relationship between the soot carbon nanostructure, thermal-optical carbon fractions, and refractory carbon cluster \((rC^n)\) distributions in SP-AMS mass spectra is given in Fig. 1. The main set-up consisted of (a) a miniature combustion aerosol standard \((\text{miniCAST})\) soot generator (model 5201C; Jing ltd), (b) options to heat the aerosol in a thermodenuder \((250^\circ \text{C})\) and ceramic tube furnace \((500^\circ \text{C})\), (c) a dilution stage \((\text{approximate total dilution of 1:250})\), and (d) aerosol instruments. The set-up has previously been described in detail [35]. The mini-CAST was operated at five different operating points (OPs) \(1, 3, 5, 6, \) and \(7\) by altering the flow of oxidation air and by mixing the fuel gas with a nitrogen \((N_2)\) mixing flow (see Supplementary data and Table S1). For an increased OP number, an increased \(N_2\) dilution and reduced oxidation air flow was used that reduces the adiabatic flame temperature. This reduces the rate of the in-flame evolution of soot properties before the miniCAST flame is quenched and results in the generation of soot with different physiochemical properties and soot with less mature character [35,47]. The dilution was accomplished through two ejector diluters \((\text{Dekati}^\text{®} \text{ Diluter DL-1000})\) in series and supplied with air at room temperature that passed a HEPA-filter and an activated carbon filter. In addition to the main set-up, a more detailed heating experiment was conducted on soot from a single miniCAST setting \((\text{OPs using undiluted exhaust})\) by ramping the temperature of the ceramic tube furnace in steps from \(25^\circ \text{C}\) to \(925^\circ \text{C}\).

The diluted miniCAST soot aerosol was analyzed on-line using a soot particle aerosol mass spectrometer \((\text{SP-AMS}; \text{Aerodyne Inc., Billerica, USA})\), a scanning mobility particle sizer \((\text{SMPS}; \text{Electrostatic Classifier model 3071 TSI Inc, CPC model 3775 TSI Inc})\), and a 7 wavelength aethalometer \((\text{Model: AE33; Magee Scientific Corp., Berkeley, USA})\), Untreated \((\text{bypass})\) and heat-treated \((250^\circ \text{C} \text{ and } 500^\circ \text{C})\) miniCAST soot was also collected for off-line thermal-optical carbon analysis \((\text{i.e., OC/EC})\) and HRTEM analysis. The SP-AMS was used to probe the non-refractory organic aerosol \((\text{OA})\) and carbon clusters of refractory origin. In the experiments, positively charged carbon clusters were measured between mass-to-charge ratio \((m/z)\) 12 and \(m/z\) 708 \((C_i^1 - C_{29}^1)\) and divided in three classes: lowcarbons \((C_1^1 - C_{25}^1)\), midcarbons \((C_{26}^1 - C_{29}^1)\), and fullerene carbons \((C_{30}^1 - C_{59}^1)\). Large carbons \((C_{30}^1 - C_{59}^1)\) refer to the sum of midcarbons and fullerene carbons. The aethalometer provided real-time measurements of equivalent BC concentrations derived from light absorption at \(880 \text{ nm}\) and \(950 \text{ nm}\). Mass estimations from SMPS size distributions were derived as previously described in Török et al. [35]. The off-line thermal-optical carbon analysis was used to quantify the operationally defined organic carbon \((\text{OC}_{1-4})\), elemental carbon \((\text{EC})\), pyrolytic carbon \((\text{PC})\), and total carbon \((\text{TC})\) mass according to the Eusaar_2 protocol [61]. We use the standard definition for \(\text{OC}\) referring to the sum of \(\text{OC}_{1-4}\) and \(\text{PC}\). \(\text{OC}_{1-4}\) were evaluated in inert \(\text{He}\) and \(\text{EC}\) in \(\text{He}\) mixed with \(2\% \text{O}_2\). \(\text{PC}\) was determined from the thermograms and subtracted from the \(\text{EC}\) concentration. Increased light absorption may occur during the evaluation of \(\text{OC}_{1-4}\). \(\text{PC}\) mass is then defined by the material removed in the \(\text{He}\) mixed with \(2\% \text{O}_2\) evaluation of \(\text{EC}\) until reaching the initial light absorption. Further information on the \(\text{OC}/\text{EC}\) analysis is included in the Supplementary data. Information on the carbon nanostructure of the soot particles \((\text{fringe lengths and tortuosity, median and lognormal parameters})\) was derived from the HRTEM analysis and using image post-processing [62] previously used to characterize soot nanostructure [63–66]. A detailed description of the HRTEM analysis including a flowchart (Fig. S1) can be found in the Supplementary data.

2.2. Description of the SP-AMS set-up, calibration, and tuning

The SP-AMS [51] incorporates an infrared Nd:YAG \((\lambda = 1064 \text{ nm})\) intracavity laser into the Aerodyne aerosol mass spectrometer \((\text{AMS})\) [67]. The intracavity laser enables vaporization of highly refractory light absorbing \((\lambda = 1064 \text{ nm})\) particles, in particular refractory black carbon \((\text{rBC})\) components. The SP-AMS can be operated with the laser vaporizer on or off, while the tungsten vaporizer \((-600 \text{ °C})\), which vaporizes non-refractory components, is always heated. The vaporization is followed by electron ionization \((70 \text{ eV})\) of the vapors and detection of positively charged ions in a high resolution time-of-flight mass spectrometer. The mass spectrometer was set-up to record positive ions in the range \(m/z\) \(10\)–\(710\).

Mass calibration of non-refractory components was performed according to the standard procedure using \(300 \text{ nm}\) \((\text{mobility diameter})\) ammonium nitrate particles. The non-refractory organic aerosol \((\text{OA})\) include \(\text{PAHs}\) and was evaluated in the conventional AMS mode \((\text{SP-AMS laser off})\) and was assigned the default relative ionization efficiency \((\text{RIE})\) of 1.4 \((\text{relative to } \text{NO}_3)\). SP-AMS \((\text{laser on})\) mass calibration of refractory black carbon \((\text{rBC})\) was performed using \(300 \text{ nm}\) carbon black particles \((\text{Regal Black; Cabot Inc., Boston, MA, USA})\) and lowcarbon \((C_{<5}^1)\) intensities. The relative ionization efficiency of \(\text{rBC}\) to \(\text{NO}_3\) \((\text{RIE}_{\text{rBC}})\) was determined to \(0.29\). Non-refractory OA \((C_{30}^1 - C_{40}^1, C_{40}^1 - C_{20}^1)\) in the range \(m/z\) \(10\) to \(m/z\) \(120\) and \(\text{PAHs}\) in the range \(m/z\) \(202\)–\(300\). lowcarbons \((C_{<5}^1)\), and midcarbons \((C_{30}^1 - C_{39}^1)\) were derived from curve-fitting with high-resolution data in the PIKA analysis module \((\text{version 1.15D})\). Signal intensities from fullerenes carbons \((C_{30}^1 - C_{39}^1)\) were derived by integrating the SP-AMS signal at unit-mass resolution. Information on the SP-AMS quantification of \(\text{PAHs}\) and separation of midcarbon and \(\text{PAHs}\) ions at the same nominal \(m/z\) is found in the Supplementary data.

The influence of laser power on large carbons was evaluated by performing a laser intensity sweep on heat-treated \((250^\circ \text{C})\) miniCAST OP6 soot \((\text{section 2.1.2})\). The reported laser powers are the measured laser bleed-through of the output coupler mirror. We monitored laser power \((\text{mW})\) with an OPHIR power monitor and laser intensity \((\text{W/m}^2)\) with a CCD-camera. The camera laser intensity was calibrated against the power monitor. In this

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Fig. 1. Experimental set-up. The set-up included the soot extinction cell but is not part of the current analysis. Results on optical properties including the soot extinction cell have been published previously [35]. (A colour version of this figure can be viewed online.)
evaluation, the SP-AMS was set-up to record mass spectra including positively charged carbon cluster ions up to $C_{200}^+$ (m/z 10–2500). Above m/z 700, $C_{50}^+$ (m/z 720) and $C_{70}^+$ (m/z 840) were identified as two additionally strong fullerene carbon signals. However, including larger carbon clusters above m/z 700 did not significantly influence the ratio of large carbons to $rC_x^+$ (the intensity ratio of large carbons to the total refractory carbon clusters).

The tuning of the voltages in the SP-AMS mass spectrometer influenced the distribution of large carbons. Specifically, altering the tuning of the ion extractor between negative and positive voltages, with all other parameters constant, would significantly change the $rC_x^+$ mass spectrum. Different tunings could maximize or minimize large carbon cluster ion signals in a given SP-AMS $rC_x^+$ mass spectrum for soot particles that exhibited large carbon ion signals; changing tunings did not result in large carbon ion signals being observed for mature soot types. Thus, the results and conclusions presented here are not dependent upon the SP-AMS tuning, only the magnitude of the observed large carbon ion signals. A thorough investigation of SP-AMS factors influencing the detection of large carbons in the SP-AMS will be published elsewhere. Our current understanding advocates a tuning of voltages which optimizes large carbon ion signals and allows simultaneous and reproducible detection of lowcarbons, midcarbons, and fullerene carbons.

The OPs 5, 6, and 7 were all associated with strong signals from mid- and fullerene carbons. To estimate the mass concentrations of the material responsible for these large carbon signals, knowledge of their relative ionization efficiencies is crucial. The SP-AMS relative ionization efficiency for large carbons is currently unknown. If large carbons were associated with non-refractory aliphatic or aromatic hydrocarbons we would expect these ions to have relative ionization efficiencies similar to OA (RIEOA larger than 1). OA has traditionally been assigned a relative ionization efficiency of 1.4. Recent evaluations suggest an RIEOA of 1.6 ($\pm$0.3) for OA with carbon oxidation states commonly observed in ambient OA, and even higher RIE for reduced organic species [68]. Since large carbons appear to have originated from refractory components we have treated them as having the same relative ionization efficiency with respect to NO$_3$ as lowcarbons, hence their concentrations are given as Regal Black equivalent mass concentration (RIE$_{RB}$ = 0.29).

3. Results and discussion

3.1. SP-AMS analysis

3.1.1. Refractory carbon cluster ($rC_x^+$) analysis

Refractory carbon cluster ($rC_x^+$) distributions in SP-AMS mass spectra were notably different between the miniCAST operating points (OP). Fig. 2 shows the mass spectra of $rC_x^+$ from OP 3, 5, 6, and 7. A higher OP represents an increased N$_2$ dilution of the miniCAST propane fuel flow and reduced oxidation air flow rate, and thus reduced adiabatic flame temperatures. The mass spectrum from OP1 was similar to that of OP3 and is not shown. Lowcarbons ($C_1^–C_5^+$ with m/z 12–60) dominated the mass spectra of OP1 and OP3, while OPs 5, 6, and 7 showed increasing fractions of large carbons ($C_{29}^–rC_{14}^+$, m/z $\geq$ 72) with increasing N$_2$ fuel dilution. The large carbons consisted of midcarbons ($C_{30}^–C_{59}^+$, m/z 72–348) separated by 12 Da and fullerene carbons ($C_{50}^–C_{149}^+$, m/z 360–708) of mainly even carbon numbers separated by 24 Da.

Among individual large carbons, the strongest midcarbon cluster was $C_{11}^+$ (m/z 132) followed by $C_{10}^-C_{14}^+$, $C_{13}^+$ and $C_{14}^+$ (m/z 120, 180, 84, and 168). These carbon clusters contributed more than 10% each and their sum contributed more than two thirds of the total midcarbon signal. The strongest fullerene ions were $C_{36}^+$ (m/z 432), $C_{44}^+$ (m/z 528), and $C_{50}^+$ (m/z 600). The distributions of mid- and fullerene carbon cluster ions were similar between OPs 5, 6, and 7, although large carbon intensities increased relative to low carbons with increasing miniCAST OP.

3.1.2. Laser power influence on SP-AMS refractory $C_x^+$ mass spectra

Fig. 3 shows the laser power influence on signals from individual carbon clusters (Fig. 3a) and the total carbon ion signal when grouped into low-, mid-, and fullerene carbons (Fig. 3b). $C_x^+$ and lowcarbons, the most commonly used proxies for rBC, were approximately doubled by a threefold increase in laser power. Fig. 3a shows that the fullerene carbon cluster $C_{50}^+$ had a slightly elevated response to laser power compared to $C_{15}^+$. However, Fig. 3b shows that, when grouped, fullerene carbons ($C_{10}^-C_{59}^+$) had a similar response to laser power as did lowcarbons (C$_{1–5}$). The midcarbon cluster ions ($C_{6}^–C_{29}^+$) appeared to fragment more strongly with higher laser power than did lowcarbons or fullerene carbons (Fig. 3b). The midcarbons appeared to preferentially fragment into $C_7^-$. This preferential path for fragmentation of midcarbons resulted in $C_7^–$ to $C_{29}^+$ ratios greater than 1 at high laser powers (>15 mW in Fig. 3). Hence, high laser power and fragmentation of midcarbons may explain previous SP-AMS observations related to high $C_7^–$ to $C_{29}^+$ ratios and the simultaneous observation of large carbons [54]. This effect could potentially induce bias to the determination of especially low-, mid- and midcarbons when comparing instrument to instrument.

Thermal ionization of large carbons has been shown to occur in the SP-AMS laser vaporization at high laser fluences [52]. In addition, dissociation of $C_{50}^+$ (i.e., the expulsion of $C_5$ and other Cl clusters) may occur at similar energies to thermal ionization [69,70] and influence $rC_x^+$ distributions. Thermal ionization in the SP-AMS was shown to be of importance primarily for very large carbons ($C_{36}^–C_{59}^+$) [52]. The increased sensitivity to $C_{50}^+$ compared to $C_{15}^+$ with increasing laser power suggests a contribution from thermal ionization for this ion at the highest laser fluences. However, such a dependence was not seen for the full class of fullerene carbons in Fig. 3b. Thus our analysis shows that during the current experiments, laser induced thermal ionization and power dependent fragmentation had only a very minor impact on the carbon cluster distributions.

3.1.3. Non-refractory OA and $rC_x^+$ after heat-treatment to 250 °C and 500 °C

Fig. 4a shows the ratio of non-refractory organic aerosol (OA$_{AMS}$), including PAHs, derived in conventional AMS mode (laser off) to the total carbon mass (TC) derived from thermal-optical carbon analysis. Fig. 4b shows OA$_{AMS}$ to PAH$_{AMS}$ ratios. The ratios are shown for untreated soot (BP), after heating to 250 °C in the thermodenuder, and after heating to 500 °C in the ceramic furnace. MiniCAST OPs 1, 3 and 5 generated particles with low non-refractory OA$_{AMS}$ content (Fig. 4a, BP). Non-refractory OA$_{AMS}$ increased for OP6 and OP7 but remained low with respect to TC (OA$_{AMS}$ to TC ratios of 0.05 and 0.06 respectively). Approximately half of the OA$_{AMS}$ signal at OP6 and OP7 was attributed to PAHs (Fig. 4b). In the soot from miniCAST OP1 and OP3, four-membered PAHs at m/z 202 dominated the total particulate PAH mass (not shown). The mass fraction of PAHs at m/z 202 was low at OPs 5, 6, and 7, where larger 4, 5 and 6 membered PAHs (m/z 226–300) dominated the particle PAH mass. Heating the aerosol in the thermodenuder (250 °C) and ceramic furnace (500 °C) removed approximately 30% of the non-refractory OA$_{AMS}$ for OP7 (Fig. 4a) and up to 60% for OP6. Heating the aerosol in the ceramic furnace to 500 °C resulted in an almost complete evaporation of PAHs (PAH$_{AMS}$ to OA$_{AMS}$ ratios less than 0.15), OA to TC ratios will be further discussed in Fig. 6 (section 3.2).

Large carbon to $rC_x^+$ ratios and midcarbons and fullerene carbons to $rC_x^+$ ratios are shown in Fig. 5a and b, respectively, for
bypass (BP, 25 °C) and after heating to 250 °C in the thermodenuder and 500 °C in the ceramic furnace. Heating the soot to 250 °C had only minor influences on low carbons and mid carbons. Fullerene carbons increased slightly, which resulted in marginally higher large carbon to rC⁺ₓ ratios after heating to 250 °C. Further heating to 500 °C showed a moderate but significant decrease of mid carbons which resulted in lower large carbon to rC⁺ₓ ratios. Heating to 500 °C resulted in a minor decrease of the fullerene to rC⁺ₓ ratio for OP5, but did not influence this ratio in soot from OPs 6 and 7.

When comparing the SP-AMS laser-on mode and laser-off mode mass spectra, increases of fragments originating from aromatic or highly unsaturated aliphatic compounds (CₓHᵧₓ−ₓ) were observed especially for higher miniCAST operating point (not shown). A strong contribution of CₓHᵧₓ−ₓ fragments to OA mass spectra has previously been found in SP-AMS laser-on mode for immature diesel soot [53], and can thus be related to increased H:C ratios in immature soots [31]. Additionally, CO, CO₂ and CₓOᵧₓ increased for laser on (not shown). The relative abundance of these carbon and oxygen containing ions decreased with OP (increased N₂ fuel dilution) which likely reflects a less oxidized surface of the refractory material. However, the major difference in total soot aerosol mass spectra between the SP-AMS laser-on mode and laser-off mode were the refractory low-, mid- and fullerene carbon cluster ions (i.e., rC⁺ₓ).
In summary, Figs. 4 and 5 shows that heating the aerosol to 250 °C resulted in partial removal of non-refractory OAAMS and a minor increase of fullerene carbons. Increased temperature to 500 °C further reduced the non-refractory particle fraction and resulted in reduced midcarbon signal intensity. These results suggest that there existed no strong relationship between non-refractory OA (including PAHs) and the large carbons (i.e., midcarbons and fullerene carbons were not formed from material related to non-refractory OA on soot heated to 250 °C nor 500 °C). Thus, the origin of the observed large carbons appear to be more refractory than typical organic compounds, but less refractory than elemental carbon or graphitic carbon which are related to both SP-AMS low carbons and equivalent BC (aethalometer).

3.2. Relationship between SP-AMS mass spectra and thermal-optical carbon analysis

The SP-AMS low carbon intensity was linearly correlated to both the elemental carbon (EC) mass and the equivalent BC (aethalometer) concentration (Supplementary data, Fig. S2) when including data from all OPs. This supports the use of the low carbon (C_1-C_6) signal to derive refractory black carbon (rBC) mass concentrations from SP-AMS data, as has previously been suggested [52].

Table 1 shows the thermal-optical carbon analysis for the untreated (25 °C) miniCAST soot (for heat-treated samples, see Table S3 in Supplementary data). For soot from miniCAST OPs 1 and 3, EC dominated the total carbon (TC) mass while OC_1-4 were low (<5% of TC) and negligible amounts of pyrolytic carbon (PC) were formed upon heating. For the miniCAST OPs 5, 6, and 7, OC (the sum of OC_1-4 and PC) fractions increased with increasing OP up to more than 50% of the TC mass for OP 6 and 7.

Fig. 6 presents trends of the contribution to the total carbon mass (TC) from OC (sum of OC_1-4 + PC), non-refractory OAAMS, and the sum of non-refractory OAAMS and large carbon (C_+6) concentration (RIE_{EB} equivalent). Comparing the OAAMS to TC ratios and the OC to TC ratios in Fig. 6, OAAMS (i.e., laser off) appears to represent OC for OPs 1 and 3, but only represent a small fraction of OC for OPs 5, 6, and 7. Including both OAAMS and C_+6 strongly improved the agreement with OC/TC ratios for all OPs, with a fairly strong and significant correlation (r = 0.89, p < 10^-4). The comparison in Fig. 6 is qualitative, as the RIE of the refractory material generating large carbons requires further evaluation, yet informative, and illustrates a strong correlation and a possible linear relationship between SP-AMS refractory large carbons and refractory OC (including PC).

In addition to Fig. 6, Fig. S3 (supplementary data) compares the non-refractory OAAMS mass and OC (including PC) mass while Fig. S4 compares the sum of non-refractory OAAMS and large carbon (RIE_{EB} equivalent) concentration versus OC mass. Including both non-refractory OAAMS and large carbons significantly improved the correlation and reduced the difference in derived concentrations.
between the SP-AMS and thermal-optical carbon analysis (see Fig. S3 and Fig. S4). The reported OAAMS includes mass contribution from hydrogen and other heteroatoms. We have not attempted to remove this contribution for comparison with OC. For these flame soots this would only result in minor adjustments of the ratios and was neglected in this simplified comparison.

From these results, we hypothesize that refractory large carbons \( (C_{1-4} + PC) \) derive mainly from material quantified as OC3, OC4, and PC in the thermal-optical carbon analysis. However, we anticipate that the classification of these components according to OC1-4, PC, and EC may depend on several factors (e.g., sampled mass on filters, relative carbon mass from semi-volatile organic components, analysis protocol, compound molecular weight, etc.) and the observations in this study may not be generalized to all soot samples. We therefore expect that components giving rise to the large carbon signal can be evaluated in OC3, OC4, and PC, but also in EC1 and EC2 depending on the conditions.

The organic carbon classes (OC1-4 and PC) in thermal-optical carbon analysis are related to the volatility of organic compounds. A higher OC class is synonymous to compounds of lower volatility. PC represents the least volatile organic carbon fraction [71]. In the EUSAAR_2 protocol, OC3 is evaluated at 450°C and OC4 is evaluated at 650°C in He. Components that pyrolyze upon heating to 450°C and 650°C in He may be forming similar pyrolysis products on the AMS tungsten vaporizer (when used in the conventional AMS mode with the SP-AMS laser off). It can therefore be expected that soot from these higher miniCAST OPs include material classified as OC1-4 and PC that cannot be vaporized on time-scales and temperatures associated with the AMS flash vaporization (~5 s, 600°C, and high vacuum) [72].

### 3.3. Soot nanostructure analysis by HRTEM and relation to SP-AMS mass spectra

High resolution images of the carbon nanostructure were obtained by high-resolution transmission electron microscopy (HRTEM). Fig. 7 shows HRTEM images of representative soot particles from the miniCAST OPs 3, 5, 6 and 7. SMPS particle size distributions show soot particle size decreasing from approximately 300 nm to 80 nm when changing from OP3 to 7 [35]. In Fig. 7, a prominent change in the physical appearance of soot particles is apparent with increased miniCAST OP (increased N\(_2\) flow, reduced oxidation air flow, and altered flame characteristics).

To analyze the carbon nanostructure, semi-automated image processing of carbon fringe length and tortuosity was conducted on images of non-overlapping parts of soot particles from miniCAST OPs 3, 5, 6, and 7. Fringe lengths decreased strongly with higher miniCAST OP (Table 2). The geometric mean fringe lengths \((l_g)\) decreased from 0.61 nm to 0.51 nm for OP3 to OP7, the fringe length distributions became narrower (decreased geometric standard deviation, \(\sigma_l\)), and the proportion of fringes larger than 1 nm decreased for higher OP. In contrast, the curvature of the fringes (their tortuosity) increased with higher operating point. The geometric mean fringe tortuosity \((t_g)\) increased from 1.18 at OP3 to 1.23 at OP7.
mature), to less graphitized (less mature) soot. C₅ species in carbon lamellae introduce curvature (tortuosity) in the soot nanostructure [26,33,73] and prevents the stacking of parallel layers [74,75], resulting in amorphous soot nanostructures. The decrease in fringe length and increase in tortuosity is a result of decreasing adiabatic flame temperatures and altered flame characteristics with increasing miniCAST OP. Similar relationships between combustion temperatures and soot nanostructures have been found in, for example, diesel engines. High curvature structures have been linked to reduced combustion temperatures in diesel engines applying exhaust gas recirculation [76] and fringe lengths of in-cylinder soot have been shown to increase from the early to late combustion phase [77] and engine out [23]. The mean fringe length derived for OP3 is comparable to the fringe lengths observed in diesel engine exhaust soot during combustion with a fatty acid methyl ester [13]. The more amorphous soot from OP6 and OP7 have mean fringe lengths more similar to soot from a wildfire [73].

Fig. 8 shows the geometric mean fringe length (left axis, note the reversed scale), the large carbon to rC₅₊ ratio (right axis) and sum of OC₃-₄ and PC to TC ratio (right axis) for OP 3, 5, 6, and 7. The large carbon to rC₅₊ ratio was negatively correlated to the geometric mean fringe length (Pearson correlation \( r = -0.97; p = 0.028 \)), and positively correlated to the mean fringe tortuosity (\( r = 0.92; p = 0.079 \)). These correlations suggest a strong relationship between the SP-AMS carbon cluster distribution and the soot nanostructure. A similar correlation was also found between the average fringe lengths and the OC₃-₄ and PC to TC ratio (\( r = -0.93 \) and \( p = 0.066 \)).

Fullerenes have C₅ rings incorporated into their structure. The SP-AMS detection of fullerene carbons can thus be a result of C₅-containing components present in the short carbon lamellae. While graphitic (C₆-containing) soot nanostructures almost exclusively fragment to low carbons, C₅-containing carbon lamellae may fragment into larger, and energetically favorable, fullerene structures. Midcarbons were exclusively detected simultaneously with fullerene carbons, making it plausible that they form from similar principles as fullerene carbons in the SP-AMS. Hence, the correlation between large carbons in SP-AMS soot mass spectra and shorter fringe lengths (and increased tortuosity) can represent an increased amount of C₅-containing species incorporated into the soot nanostructure. As a result, changes in the SP-AMS rC₅₊ distribution may possibly provide a real-time measurement of key parameters controlling soot reactivity.

### 3.4. SP-AMS measured refractory BC during evaporation and oxidation

The relative intensities of low-, mid-, and fullerene carbons changed upon heat treating the soot (Fig. 5). In addition to the

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**Table 2**

| MiniCAST Operating point | N₂ dilution of fuel (l/min) | Fringe length \( l_m, l_g, \sigma_l \) | Fringe tortuosity \( \tau_m, \tau_g, \sigma_\tau \) | Fraction of total fringes > 1 nm | Number of analyzed fringes (n) |
|--------------------------|---------------------------|-------------------------------------|-------------------------------------|---------------------------------|-----------------------------|
| 3                        | 0.100                     | 0.56, 0.61, (±0.02)                 | 0.61, 0.61                         | 1.13, 1.18                      | 35%, 2387                   |
| 5                        | 0.200                     | 0.52, 0.56, (±0.01)                 | 0.56, 0.56                         | 1.15, 1.20                      | 25%, 3388                   |
| 6                        | 0.250                     | 0.50, 0.54, (±0.02)                 | 0.52, 0.52                         | 1.15, 1.21                      | 21%, 839                    |
| 7                        | 0.300                     | 0.47, 0.51, (±0.001)                | 0.49, 0.49                         | 1.18, 1.23                      | 15%, 4019                   |

**Fig. 7.** Images from high-resolution transmission electron microscopy (HRTEM) of representative miniCAST soot particles (top) and soot carbon nanostructure (bottom).
bypass and thermodenuder experiments, a more detailed heating study where miniCAST OP5 soot was heated in small steps up to 925 °C in the ceramic tube furnace was carried out. Fig. 9 shows the evolution with temperature of SMPS derived mass, equivalent BC (aethalometer), and SP-AMS lowcarbons, midcarbons, fullerene carbons. The evolution of particle size distribution (electrical mobility) during the heating experiment is available in Supplementary data (Fig. S5). A general decrease in concentrations was observed with increasing temperature in the range 25 °C–250 °C and may partly be explained by evaporation of volatile material, but mainly due to increased thermophoretic losses in the cooling section after the thermodenuder. Fullerene carbons initially increased relative to low- and midcarbons when the soot was heated to between 150 °C – 400 °C. At higher temperatures (400 °C – 600 °C), midcarbons and fullerene carbons decreased rapidly relative to lowcarbons. The equivalent BC concentration, SMPS derived mass, and the SP-AMS lowcarbon concentration all decreased sharply at temperatures higher than 625 °C, at similar temperatures to the oxidation onset of mature diesel soot [78]. The equivalent BC and lowcarbon soot components were almost completely oxidized at 925 °C.

The evaporation/oxidation experiment shows that lowcarbons and equivalent BC were associated with the same material. Particle components from which the midcarbons and fullerenes originated were removed at temperatures approximately 200 °C lower than components associated with lowcarbons and equivalent BC. The particle mass derived from mobility measurements with the SMPS was largely unaffected between 400 °C–625 °C. This suggest that mobility sizes were insignificantly affected by mass loss from components associated with SP-AMS large carbons. Hence, the loss of large carbon associated material may have been related to internal oxidation patterns similar to those observed for some diesel engine soot at reduced combustion temperatures [14]. Alternatively, the loss of large carbon associated material may have been related to the pyrolysis of such components and a complete restructuring to refractory BC material. An increase of lowcarbons was not observed simultaneously to the decrease of large carbons, strengthening the hypothesis that the large carbon associated material was oxidized/evaporated rather than pyrolyzed.

3.5. SP-AMS rC_4^+ analysis and Raman spectroscopy

Changes in the nanostructure upon heating of soot from a miniCAST soot generator (model 5201C; Jing ltd) were previously studied using Raman spectroscopy by Ess et al. [79]. We identified similarities in the behavior of our SP-AMS lowcarbon to rC_4^+ ratio (the complement ratio to the large carbon to rC_4^+ ratio) and their Raman spectroscopy ID/IG ratio analysis. Changes in the soot nanostructure were studied upon heating (in air) by evaluating the ratio between the intensity of the D peak to the intensity of the G peak (ID/IG) in Raman spectra [79]. The ID/IG ratio is related to the

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**Fig. 8.** Geometric mean fringe lengths ($l_p$) (left axis, note the reversed scale), large carbon to rC_4^+ ratios (right axis), and OC_3+PC to TC ratios (right axis) for the untreated soot from miniCAST OPs 3, 5, 6, and 7. Error bars represent (95%) confidence intervals for average fringe lengths and large carbon to rC_4^+ ratios, and estimated errors for the OC/EC analysis. The average fringe length was inversely correlated to the large carbon to rC_4^+ ratio ($r = -0.97, p = 0.028$) and sum of OC_3+ and PC to TC ratio ($r = -0.93, p = 0.066$). (A colour version of this figure can be viewed online.)

**Fig. 9.** Soot evaporation and oxidation experiment. SMPS mass estimations (using effective densities from non-heated samples), equivalent BC (aethalometer), and SP-AMS concentrations of low-, mid-, and fullerene-carbons normalized to their concentrations at room temperature (25 °C, the starting point) during temperature ramping of miniCAST OP5 soot (raw exhaust, low oxygen content) from 25 °C–925 °C in a thermodenuder (25 °C–250 °C) and ceramic furnace (250 °C–925 °C).
carbon nanostructure and suggested to be proportional to fringe length [80,81]. In order to produce soot with higher OC content, Ess et al. [79] reduced only the oxidation air flow of the miniCAST (no N2 was added). Thus, their “set point” and our “operating point” values represent different miniCAST conditions.

On the right axis, Fig. 10 shows the results from Ess et al. [79] of the ID/IG ratio for soot from set point 1 and 2 during a temperature ramp from 25 °C to 600 °C. For set point 1 soot, they observed an ID/IG ratio close to 1.2 and negligible changes upon temperature ramping. For set point 2 (shown) and set point 3 (not shown) soot, they instead observed a small but significant decrease in the ID/IG ratio from heating up to 300 °C, indicating a decreasing structural order of the soot nanostructure. Further heating (300 °C–600 °C) strongly increased the ID/IG ratio up to ~1.2 and was interpreted as an ordering of the nanostructure [79].

On the left axis, Fig. 10 shows the lowcarbon to rC+C ratio for miniCAST soot from OP 1, 3, and 5, as well as for diesel soot sampled from an experimental heavy duty engine [53]. Diesel soot and the mature miniCAST soot at OP 1 and 3 had high lowcarbon to rC+C ratios (>0.95) which were not influenced by heating to 500 °C. For soot from OP5, the lowcarbon to rC+C ratio was constant between 25 °C–150 °C but showed a slight decrease upon heating between 150 °C–325 °C. Heating above 400 °C sharply increased the lowcarbon to rC+C ratio, from 0.65 at 325 °C to 0.93 at 550 °C. The ratio remained nearly constant and above 0.9 upon further heating from 550 °C to 925 °C.

The increase of large carbons between 150 °C to 325 °C for soot from OP5 occurred at similar temperatures as the decrease in ID/IG ratio for soot from set point 2 [79]. Similarly, the sharp increase in lowcarbon to rC+C ratio between 400 °C–550 °C coincided with the sharp increase in ID/IG ratio of the set point 2 soot. The similarities between the SP-AMS rC+C analysis and Raman spectroscopy ID/IG analysis provide further evidence for a strong relationship between SP-AMS rC+C distributions and the soot nanostructure. The increase in fullerene carbons from heating of the soot between 150 °C–325 °C thus appears related to a partial restructuring to less ordered carbon nanostructures, or the formation of such structures from charring of OA. Consequently, we have interpreted the sharp decay of large carbons at 400 °C as related to the oxidation (removal) of particle components associated with the SP-AMS large carbon clusters, and increased graphitization of the carbon nanostructure.

4. Conclusions

This study explored relationships between refractory carbon cluster (rC+C) distributions in SP-AMS mass spectra, soot carbon nanostructure, and thermal/oxidation properties of soot components. A linear correlation was found between the SP-AMS large carbon to rC+C ratio and decreasing average fringe length (r = −0.97, p = 0.028). The comparison between the thermal-optical carbon analysis and the SP-AMS rC+C mass spectra indicated that low-carbons derive from EC components while large carbons (mid- and fullerene carbons) originated from refractory organic carbon components classified as OC3−4 and components undergoing pyrolysis as OC evolves in the thermal-optical carbon analyzer (i.e., PC). Material related to large carbon ions exhibited different oxidation behavior than that related to lowcarbons and equivalent BC. At temperatures ~200 °C lower than the oxidation of lowcarbons and equivalent BC, the large carbon clusters (C+C) were no longer detected in SP-AMS mass spectra. Therefore, large carbons (both midcarbons and fullerene carbons) most likely represent particle components chemically separated from and with optical properties different from BC.

In addition to C6 rings also found in graphitic carbon, fullerenes incorporate C5 rings. Soot with high tortuosity (curved lamellae) and more reactive nanostructures have been hypothesized to form by partial oxidation of benzene and C5 chemistry in flames [18,26]. The large carbons in SP-AMS mass spectra may thus provide information on refractory soot composition and nanostructures related to soot reactivity. In modern diesel engines, soot forms in large amounts shortly after ignition and is oxidized in the late combustion cycle. As a result, only a small fraction of the soot initially formed in the combustion exits the engine into the tailpipe [53,82,83]. Identifying soot components which are easily oxidized (removed) may therefore aid in developing new soot mitigation strategies for internal combustion engines. Absorption Angstrom exponents were previously reported to increase with higher miniCAST OP [35]. This suggests that soot particle components responsible for SP-AMS large carbons can have absorption spectra more similar to brown carbon (with absorption Angstrom exponents > 1).

This study also identifies major uncertainties related to the characterization of mass and composition for immature soots. We
may conclude that for the low temperature combustion conditions and immature soots sampled in this study, including only equivalent BC (or EC) and non-refractory OA$_{AMS}$ (conventional AMS measurement, laser off) significantly underestimated the total particulate carbon mass. The additional carbon mass consisted of refractory organic carbon components correlated to large carbons in the SP-AMS (laser on) mass spectra. The incapacity to evaporate the refractory organic carbon material at 500 °C in the furnace, the persistent large carbons in SP-AMS mass spectra, and the stability of the soot nanostructures under the electron beam during HRTEM, suggest that these refractory organic carbon components were constituents incorporated into and part of the internal soot carbon nanostructure (see also e.g., [47]). This may challenge atmospheric BC and OA internal mixing state models assuming stratified particles consisting of a BC core and a coating of condensed low volatility organic constituents (e.g., [48, 84]). It also suggests that relationships between equivalent BC to non-refractory OA ratios and the absorptivity of OA may be biased if refractory organic components are present in the soot. SP-AMS large carbons have been observed in soot mass spectra from a number of combustion sources including biomass combustion and in low temperature diesel combustion [51,52,54,55,85]. We therefore suggest a need to reevaluate the internal mixing state model (BC core and organic coating) for soot emissions from such combustion appliances, especially in relation to the underlying causes of increased absorption Angstrom exponents and toxicity.

The results in this study form a basis for interpreting large carbons (C$_{18}^+$) in aerosol mass spectra from soot and BC-containing materials. This interpretation can aid the design of novel on-line investigations related to refractory organic carbon and C$_{5}$-containing structures in engineered carbon nanomaterials and particulate emissions from low temperature combustion. Moreover, it can aid in deciphering the optical, cloud forming, and health relevant properties of such components in ambient soot. Future studies should identify suitable calibrants and develop a mass calibration procedure for SP-AMS large carbons, as well as improve the accuracy with which rC$_{18}^+$ analysis can provide information on variations in the soot nanostructure.

Acknowledgements

The authors gratefully acknowledge the financial support from The Swedish Research Council FORMAS (research grant 2013-453), The Swedish Research Council VR (research grant 2013-5021), and Nanolund at Lund University (p08-2014). Part of the HRTEM work was conducted under and financed by the Danish Center for Nanosafety. We acknowledge technical support by Dr. Jens Kling (Technical University of Denmark), and Dr Natalia Ivleva (Technical University of Munich) for sharing Raman spectroscopy data.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.carbon.2018.10.072.

References

[1] P. Parent, C. Laffon, I. Marhaba, D. Ferry, T.Z. Regier, I.K. Ortega, et al., Nano-scale characterization of aircraft soot: a high-resolution transmission electron microscopy, Raman spectroscopy, X-ray photoelectron and near-edge X-ray absorption spectroscopy study, Carbon 101 (2016) 86–100.
[2] I.E. Nielsen, A.C. Eriksson, R. Lindgren, J. Martinsson, R. Nyström, E.Z. Ristovski, B. Miljevic, N.C. Surawski, L. Morawska, K.M. Fong, F. Goh, et al., Respiratory health effects of diesel particulate matter, Respilology 17 (2) (2012) 201–212.
[3] T.C. Bond, S.J. Doherty, D.W. Fahey, P.M. Forster, T. Berntsen, B.J. DeAngelo, et al., Binding the role of black carbon in the climate system: a scientific assessment. J. Geophys. Res. Atmos. 119 (11) (2014) 5380–5552.
[4] S. Steiner, C. Bisig, A. Petri-Fink, B. Rothen-Rutishauser, Diesel exhaust: current knowledge of adverse effects and underlying cellular mechanisms, Arch. Toxicol. 90 (2016) 1541–1553.
[5] N. Matti Maricq, Chemical characterization of particulate emissions from diesel engines: a review, J. Aerosol Sci. 38 (11) (2007) 1079–1118.
[6] F. Stanzione, A. Tregrossi, A. Ciajolo, Probing structures of soot formed in premixed flames of methane, ethylene and benzene, Proc. Combust. Inst. 34 (1) (2013) 1885–1902.
[7] W. Wang, C. Song, J. Song, G. Lv, H. Fang, W. Zhang, Aliphatic C–H and oxygenated surface functional groups of diesel in-cylinder soot: characterizations and impact on soot oxidation behavior, Proc. Combust. Inst. 34 (2) (2013) 3099–3106.
[8] H. Wang, Formation of nascent soot and other condensed-phase materials in flames, Proc. Combust. Inst. 33 (1) (2011) 41–67.
[9] W.J. Greico, J.B. Howard, R.L. Vander Wal, Soot processes in compression ignition engines, Prog. Energy Combust. Sci. 13 (3) (2007) 379–402.
[10] W.J. Greico, J.B. Howard, Soot oxidation: dependence upon initial oxidative reactivity of diesel engine soot, Combust. Flame 155 (4) (2008) 675–695.
[11] C. Russo, M. Alfè, J.-N. Rouzaud, F. Stanzione, A. Tregrossi, A. Ciajolo, Probing structures of soot formed in premixed flames of methane, ethylene and benzene, Proc. Combust. Inst. 34 (1) (2013) 1885–1902.
[12] B. Rehalla, C. Bae, Morphology and nano-structure of soot in diesel spray and engine exhaust, Fuel 203 (2017) 47–56.
[13] Y. Ying, D. Liu, Effects of butanol isomers additions on soot nanostructure and reactivity in normal and inverse ethylene diffusion flames, Fuel 205 (2017) 272–309.
[14] R.L. Vander Wal, A. Strzelec, T. Toops, C. Stuart Daw, C.L. Genzale, Forensics of soot: C5-related nanostructure as a diagnostic of in-cylinder chemistry, Fuel 113 (2013) 522–526.
[15] R.L. Vander Wal, A. Strzelec, T. Toops, C. Stuart Daw, C.L. Genzale, Forensics of soot: C5-related nanostructure as a diagnostic of in-cylinder chemistry, Fuel 113 (2013) 522–526.
