Effects of camphor oil addition to diesel on the nanostructures and oxidative reactivity of combustion-generated soot

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Abstract

Less viscous and low cetane (LVLC) fuels have emerged as the promising alternative fuels or additives to fossil fuels. Camphor oil is one such potential LVLC fuel currently under consideration. However, it’s sooting propensity and subsequent effects on soot nanostructure, when blended with diesel, are not well understood. In this work, the effects of camphor and camphor oil addition to diesel on the sooting propensity, soot oxidative reactivity, and the chemical composition, structural disorders, and the morphology of soot particles are studied using a diffusion flame. The chemical and the microstructural changes in soot are investigated using several experimental techniques such as energy dispersive X-ray spectroscopy, high resolution transmission electron microscopy, Raman and electron energy loss spectroscopy, and powder X-ray diffraction, while the oxidative reactivity is studied using thermogravimetric analysis. The activation energy for O2-induced soot oxidation during the initiation stage shows a significant reduction in its value with the addition of camphor and camphor oil to diesel, which were 220 kJ/mol for diesel soot, 175 kJ/mol for 5% camphor/95% diesel soot, and 150 kJ/mol for 10% camphor oil/90% diesel soot. The blending of camphor and camphor oil with diesel results in soot with smaller fringe length and primary particle diameter, but increases the fringe tortuosity, the degree of crystal disorder, and the amounts of oxygen functionalities and aliphatics in soot. These physico-chemical changes in soot are used to explain the observed trend of oxidative reactivity. This study successfully demonstrates the potential of terpenoid keto compounds with characteristic bicyclic ring structure in improving
oxidative reactivity of combustion derived soots as desired in diesel particulate filter technologies.

Keywords: Soot; Nanostructure; Oxidative reactivity; Camphor oil; Diesel.

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1. Introduction

The majority of the ever-increasing demand of society for energy is fulfilled through the combustion of fossil fuels such as diesel and gasoline. Diesel, in particular, has served as one of the most important fuels for on-road and off-road transportation industry [1]. However, due to the technological limitations, its inefficient combustion results in the emission of gaseous pollutants such as CO\textsubscript{x}, NO\textsubscript{x}, and SO\textsubscript{x} and solid state pollutants such as soot [2]. Soot is composed of a series of polycyclic aromatic hydrocarbon (PAH) compounds, randomly stacked together in the form of disordered graphitic nano-crystallites [3]. Due to their physically and chemically stable structure, the constituent PAHs are classified as persistent organic pollutants (POPs) [4]. The PAH nano-crystallites are known to bio-magnify and bio-accumulate in the environment causing fatal diseases such as skin and pulmonary cancer [5]. The use of diesel particulate filters (DPF) has by far been the most practical solution to this problem, and is being implemented in developed countries. However, due to the high cost involved, such a technology does not find popularity among developing nations.

Biofuels produced from non-edible sources such as waste cooking oils, animal fats, algae, and seed oils such as Jatropha and Karanja that grow on waste lands [6–9] are being considered as promising alternative fuels or fuel blends to fossil fuels. Fuel modifications using these biodiesels, methyl esters in particular (obtained from the transesterification of biomass), have the capability to reduce soot emission, while simultaneously maintaining the required energy efficiency [10–12]. It has been reported that the presence of oxygenates in the fuel additives, their chemical structure, and their physical properties such as density, kinematic viscosity, and cetane number play a vital role in inducing increased oxidation rate of diesel fuel, thus reducing soot production in combustion engines [13,14]. Moreover, it is also desired to enhance the regeneration efficiency of DPF by altering soot nanostructures in a way that facilitates its faster oxidation once captured in DPF. For instance, Yehliu et al. [15] studied the nanostructure of soot obtained from diesel and biodiesel blends, and unravelled the presence of a large percentage of shell-core type nanostructure using high-resolution transmission electron microscopy (HRTEM) images. They attributed the high reactivity of soot from biodiesel blends
to this disordered shell-core structure of soot. A similar observation was made in other studies [16,17]. Several investigations have correlated the presence of high oxygen content and large surface area of the biodiesel-derived soot to its higher rate of oxidation [18,19]. Furthermore, the effect of dilution of diesel with the addition of biodiesel and the relative significance of oxygen content has been statistically studied by some research groups [20]. It has been validated that the oxygenated functional groups in biodiesel play a crucial role in reducing soot production. Moreover, a high oxygen content in soot leads to its high oxidative reactivity. However, there are also reports that contradict this observation. Yehliu et al. [15] found no significance of oxygen content in soot to its oxidative reactivity, and instead attributed the reactivity to the disordered structure of soot rather than the oxygen content. The effects of the amorphous nature of soot and the degree of lattice disorder have been evaluated using XRD and Raman analyses by several researchers, where these crystal defects have been shown to have a significant effect on the improved kinetics of soot oxidation [5,21,22]. Further studies signifying the influence of the chemical structure such as the carbon chain length of the methyl esters, the position of the C=C bond, and the presence of the number of methyl group substituent on aromatic rings are available in the literature [23–25]. The increase in the chain length in methyl esters enhances the sooting tendency, but reduces soot reactivity. The position of the C=C also influenced the soot reactivity, while the methyl substitution in aromatics decreases the soot oxidation rate. More recently, the studies on the oxidation of soots derived from diesel, biodiesel, and their blends over Ag/SiO$_2$, Ag/ZnO, and 3% Ag/CeO$_2$ have reported their excellent catalytic effects in oxidizing soots [26,27], and further research is being conducted to improve their efficiency for their possible application in DPF technology.

Since the above studies suggested that the blending of diesel with biodiesel is efficient in reducing the emitted pollutants while maintaining energy efficiency, a great deal of research focus has been devoted on the search for effective and economical methods to convert bio-waste into biodiesel over the past decade. Recently, the photocatalysis-driven synthesis of biodiesel using Cr/SiO$_2$ catalyst has shown promising results in this regard [28]. However, many challenges still remain to be addressed [29–31]. Recently, a new class of fuels, extracted from bio-based renewable sources, which can be utilized directly as neat fuels in combustion engines without their conversion into methyl esters, are being looked upon as a possible solution to the above problem [32]. The potential candidates for such fuels are the essential oils such as eucalyptus, pine, lemon, and orange oils [33,34]. Camphor oil is one such biofuel, which is currently being studied by some research groups [35,36]. It is an essential oil, traditionally extracted from the Cinnamon Camphora tree in most regions of Asia. It is
available in four grades: white, brown, yellow, and blue camphor oil. The white grade is the
most commonly available one, with primary chemical components as Camphor (~ 50%),
Linalool (~ 23%), 1,8 Cineol (~ 11.5%), and some minor terpene derivatives [37,38]. On the
other hand, solid camphor is obtained by recrystallization from camphor oil, which allows
exclusion of the above listed secondary components. In the fuel sector, it is classified into a
new class of fuels called less viscous and low cetane (LVLC) fuels. The LVLC fuels are
reported to improve fuel/air mixing, fuel evaporation, and fuel atomization in CI engines.
Moreover, these fuels offer suitable ignition delays, which in turn allows reduced fuel-to-air
ratio to improve combustion efficiency and reduces soot formation [34,39]. For instance, the
twin-cylinder compression ignition engine studies under full load condition with camphor oil-
blended diesel, conducted by Subramanian et al. [35], reported an improved brake thermal
efficiency, which is an indicative of the effective conversion of the chemical energy stored in
fuel into mechanical energy. The engine experiments, conducted by Kasiraman et al. [36],
involved the mixing of less viscous camphor oil with highly viscous cashew nut shell oil that
resulted in the improved physical properties of the blend and better engine performance.
Thiyagarajan et al. [40,41] investigated the blending of camphor oil with Karanja biodiesel and
found that the engine performance was improved with a reduction in CO₂ emission, although
NOx emission was elevated. Since camphor oil has a low cetane number, its blending with
other fuels such as diesel or biodiesel reduces the cetane number of the blend. In such cases,
cetane enhancers such as dimethyl ether, dimethyl carbonate have been used in conjunction to
improve engine performance [42,43]. While the research on improving the engine performance
and lowering pollutant emission with camphor oil blended fuels continues to grow, no studies
are available in the literature on the nanostructural properties and oxidation kinetics of soot
obtained from camphor oil-diesel blends. More importantly, camphor belongs to a special class
of naturally occurring organic compounds called the terpenoid ketone family with a
characteristic bicyclic ring structure and the presence of keto group, which can be utilized to
modify the soot nanostructure of the blended diesel fuels for improving its oxidative reactivity
as desired in diesel particulate filter technology. Such an investigation could open up new
doorways for tailored synthesis or isolation of naturally occurring terpenoids for large scale
utilization in designing advanced bio fuels. Since it is known that, in several combustion
systems such as diesel engines, wall wetting assisted, diffusion-driven processes govern soot
production rate, soot produced from a diffusion flame could serve as a suitable representative
of those processes, and the study of the changes in the physical and chemical characteristics
and the reactivity of soot particles could provide an insight into their formation and oxidative
removal efficiency in engines with DPF. This provided the motivation for the present study to
determine the sooting tendency of camphor oil–diesel blends, while simultaneously
investigating the modifications in soot nanostructures and its influence on the oxidative
reactivity of soot which can improve the efficiency of DPF technology.

2. Experimental details

The smoke points of diesel (D), camphor solid-diesel blends (CD), camphor oil (CO), and
camphor oil-diesel (COD) blended fuels were measured using an American Society for Testing
and Materials (ASTM) standard smoke point apparatus (model number RAP172, developed as
per ASTM D1322 standard [23]) at atmospheric pressure. The camphor oil (grade: white, >
99% purity) and camphor solid (> 99% purity) were purchased from Sigma Aldrich, while
diesel was obtained from a commercial diesel station in U.A.E. The smoke point apparatus is
made up of a metallic cylindrical fuel reservoir (length: 10.9 cm, internal diameter: 2.125 cm)
with a provision for air vent, a wick tube with a cotton wick of diameter 4.5 mm and a length
of 125 mm, a lamp body, and a stand that hold the burner fitted with a chimney of 13 cm height.
An adjustable glass window is mounted in front of it to assist in the flame height measurement
by a vertical scale fixed behind the flame. The cylindrical lamp body has a diameter of 81 mm,
which surrounds the burner and protects the flame from the environmental disturbances. A
screw mechanism allows the flame height to be varied by lowering or raising the fuel reservoir.
In order to collect soot from the tip of each fuel flame of 27 mm height, a filter assembly (from
Sierra Instruments, USA), with a vacuum pump provision, was mounted over the chimney. The
70 mm diameter, borosilicate microfiber filters, strengthened with woven glass (TX40H120-
WW, Pall Filters, USA), was used to collect soot samples from the flames of all the tested
fuels. The collected soot samples (approx. 20 mg) were then dried in a tube furnace (heating
rate of 20 °C/min up to a temperature of 400 °C) under N2 flow to remove water vapour and
any loosely-bound hydrocarbons.
These samples were then subjected to chemical and morphological characterization. The
nanostructural features of soot were found using FEI Tecnai G20 scanning HRTEM operated
at 200 kV with 0.24 nm point resolution, equipped with an energy-filtered camera. Soot
samples were ultra-sonicated in ethanol for 10 min to disperse soot and a drop of this solution
was applied to a lacey carbon-coated, 400 mesh copper grid followed by drying under an
infrared lamp. The electron energy loss spectra (EELS) were obtained using Gatan Image Filter
in the TEM mode with the energy resolution of 1 eV, and the spectra were recorded at a
dispersion of 5 eV per channel. For the oxidation of soot particles, a thermogravimetric
analyzer (TGA, NETZSCH STA 409PC-LUXX) was used. The dried soot samples were heated in air from 200 to 800 °C at the heating rates of 1, 3, 5, and 7 °C min⁻¹. The elemental composition of soot samples was analysed by surface coating the particles with Pd/Au for improved conductivity in a SEM-Carl Zeiss (JSM-5800LV) coupled with Ametex EDX PV6500 system. The quantitative information on the nanocrystallites present in soot were calculated from the XRD patterns found using Panalytical Empyrean powder X-ray diffractometer with Cu-Kα radiation (1.54 Å, 40 kV, 40 mA). The scan range was 10–80° with the scan step size of 0.02° and the scan speed of 10 s/step. The structural disorder in soot samples were further analysed using Raman spectrometer (Witech Alpha 300 RAS) equipped with 515 nm laser source and a dual purpose 50x objective lens.

3. Results and Discussion

3.1 Smoke point, threshold sooting index, and fuel consumption rate

The smoke points of diesel (D), camphor oil (CO), camphor oil-diesel blend (COD), and camphor solid-diesel blend (CD), as a function of blending percentage were measured by observing the flame height at which the sooting wings appear on both sides of the flame [23]. The comparative analysis of the variation in smoke point is presented in Figure 1a with a relative error of ±1 mm. It is observed that the blending of a small percentage of camphor oil to diesel results in the enhancement of the smoke point that increased from 19 for pure diesel to 23.3 and 21.3 mm for 10% camphor oil/90% diesel (10% COD) and 15% camphor oil/85% diesel (15% COD), respectively. The further increase in the blending percentage of camphor oil resulted in a gradual decrease in the smoke point enhancing the sooting tendency of the blend. When pure camphor oil was tested, a smoke point of 13.7±1 mm was measured, which is significantly below the smoke point of pure diesel. In order to further investigate the cause of such a significant decrease in the smoke point at higher camphor oil–diesel blending ratios, the smoke point measurements were performed by blending pure camphor solid with diesel, as it is soluble in diesel up to a saturation limit of 20%. This provided a way to eliminate the effect of secondary constituents present in the camphor oil and understand the sole influence of camphor on the sooting tendency of diesel. As seen in Figure 1a, the blending of camphor solid increases the smoke point of diesel to 22.3 (±1) with a small blending percentage of up to 5%, possibly due to the presence of tricyclic structure with oxygen atom in the camphor molecule that could enhance the fuel radical formation rate, leading to improved fuel oxidation and reduced soot formation [44,45]. The smoke point falls drastically with the further increase in
camphor content in diesel, consistent with the trend obtained in the case of higher blending percentage of camphor oil in diesel.

Since the smoke point is an instrument-dependent quantity, we further calculated an instrument-independent parameter, threshold sooting index (TSI), which is expressed as: \[ TSI = a(MW/SP) + b, \]
where \( SP \) = smoke point, \( MW \) = molecular weight of the fuel and, \( a=3.76 \) and \( b=-3.75 \) are instrumental constants determined using standard fuels with known TSI values [23]. The calculated TSI values for diesel (D), camphor oil (CO), 5% camphor solid/95% diesel blend (5% CD) and 10% COD are presented in Figure 1b. It is evident that both 5% CD and 10% COD blends are effective in reducing the sooting tendency of diesel.

![Figure 1: (a) Variation in smoke point (SP_{error} = ± 1 mm) with different blending percentages of camphor oil and camphor solid with diesel. (b) Calculated TSI for pure diesel (D), 5% CD, 10% COD, and pure camphor oil (CO).](image)

It is interesting to note that the relative decrease in the smoke point with increasing percentage of camphor solid in diesel is significantly more drastic as compared to that of COD blends. This could be attributed to the presence of a relatively greater amount of camphor in CD as compared to COD at the same blending percentage. The high sooting tendencies of pure camphor/camphor oil and of the blends containing their high percentages are explained as follows. It has been reported that, at low temperatures, camphor can undergo secondary reactions such as oxidative dimerization with its in-situ generated enoxy radical to form exo,exo’ or endo,endo’-3,3’-bicamphor, or its corresponding biisoborneol derivatives [46,47]. Although flame temperature is significantly high (>800 °C), many alkoxy and peroxy radicals are formed in flames via fuel pyrolysis and partial oxidation. Under low oxygen environments,
such initially formed alkoxy radicals may initiate dimeric nucleation similar to the case discussed above, and could enhance the cyclization reactions to form PAH-like structures. Subsequently, this may lead to the enhancement in soot formation with the increase in camphor blending percentage in diesel. When a high percentage of camphor oil is blended with diesel, the decrease in smoke point is relatively slower, which could be attributed to i) relatively lower percentage of camphor content (i.e. ~50% in pure oil) in the blend, and also ii) the presence of other oxygenated compounds such as linalool (~23%) and 1,8-cineol (~11.5%) that together increases the total oxygen content of the fuel to decrease sooting tendency.

For a given fuel, the change in the fuel mass flow rate in a flame can also affect significantly the mass production and physicochemical properties of soot. The fuel mass flow rate in diffusion flames is a simplified representation of the increase in the engine load that also results in an increased injection of fuel to the cylinder. In this study, the fuel mass flow rate was measured to ensure that the experimental results mimic diesel engine conditions, where soot emissions from various fuels are quantified at a constant fuel mass flow rate [48]. In our experiments, the fuel mass flow rates for the above fuels and their blends were calculated by continuously monitoring the weight loss of the fuel in the burner as a function of burning time for the fixed flame heights of 10, 20, and 30 mm. For each flame height, the weight loss of the fuel was plotted versus time (where, an outstanding linearity was obtained), and the slope of the plot depicted the fuel mass flow rate. A plot of fuel mass flow rate versus flame height was almost linear \( R^2 = 0.98 \), as shown in Figure SF1 (supplementary data file). It was found to be largely constant at a fixed flame height for all the fuels. The largest dissimilarity between the fuel mass flow rates at a fixed flame height is 7%, and in most of the cases it is below 4%. These small differences could mainly arise from the uncertainty of ±1 mm in flame height measurement in the smoke point apparatus. Even when a flame height is fixed, some fluctuation in the diffusion flame is experimentally seen that slightly changes the flow rate (as shown by the error bars in the plot). Thus, while it is theoretically viable to fix fuel mass flow rate and flame height, small fluctuations in the flame are always present. Since the differences are very small, it is justified to consider the fuel mass flow rates of different fuels to be same at a given flame height [49].

Since the measured average smoke points of D, 5% CD, and 10% COD fuels were 19, 23, and 24 (±1) mm, respectively, soots from all these fuels were collected at a fixed flame height of 27 mm (that was above the smoke points of all the tested fuels). The effect of camphor oil and camphor solid blending with diesel on the kinetics of their soot oxidation with \( \text{O}_2 \) was further
investigated using thermogravimetric analysis, and their results are presented in the next section.

3.2 Thermogravimetric analysis

Soot oxidation in engines and in DPFs is strongly dependent on its reactivity with O₂. Therefore, thermogravimetric analysis in air have been performed to investigate the kinetics of O₂-induced soot oxidation. The overall oxidation process can be generalised to produce either CO (g) or CO₂ (g) and other side products, without accounting for large complex reaction models. A typical soot oxidation in TGA involved heating known quantities of soot samples derived from D, 5% CD, and 10% COD blended fuels in air from 200-800 °C at different heating rates of 1, 3, 5, and 7 °C/min. The typical variation in soot conversion (α) and conversion rate (dα/dt) at varying heating rates in the case of soot obtained from 10% COD is presented in Figure 2a and 2b, respectively, while those for D and 5% CD soot can be found in Figure SF2 in the supplementary data file. The soot conversion (α) is defined as α = (Mo-Mt)/(Mo-Ml) with Mo being the initial soot mass, Mt being the mass of partially oxidized soot at temperature T, and Ml being the leftover mass (mostly ash). The reactivity of soot at different conversion levels is evaluated by calculating the activation energies for soot oxidation using Friedman method [50,51], wherein the rate of soot conversion (dα/dt) is equated to soot conversion (α) by Equation E1.
Figure 2: (a) soot conversion, $\alpha$, and (b) soot conversion rate ($d\alpha/dt$) measured in air at different heating rates. Comparative analysis of (c) soot conversion, $\alpha$ measured at a fixed heating rate of $3 \circ C \ min^{-1}$, and (d) variation in activation energy at different soot conversion levels measured in case of soots obtained from 5% CD, 10% COD and D.

$$\frac{d\alpha}{dt} = k(T)f(\alpha)$$  \hspace{1cm} (E1)

In the above equation, $k(T)$ is the rate constant (s$^{-1}$) of O$_2$-induced soot oxidation, expressed by the Arrhenius equation, $k=Ae^{-(Ea/RT)}$, where $A$ is the pre-exponential factor, $Ea$ is the activation energy of soot oxidation, and $T$ is the temperature [52]. The term, $f(\alpha)$ describes the reaction model or the conversion function. Since the concentration of O$_2$ is maintained constant throughout the TGA experiment, its expression in rate equation is omitted. For further details about the procedure and activation energy calculation, please refer to [25,53].

The calculated activation energies at different conversion levels and temperatures are presented in Figure 2d. It can be seen that, at low conversion level (i.e. $\alpha = 0.1$), the activation energies of the three soot samples differ significantly. The 10% COD soot has the lowest initial activation energy of 150 kJ/mol, followed by 175 kJ/mol for 5% CD soot, while the diesel soot had the highest initial activation energy of 220 kJ/mol. The higher oxygen content in soot...
increases the evaporation rate of loosely bound/volatile substances in soot and induces internal oxidation of the soot structure [54]. Therefore, the observed increase in the activation energy of soot oxidation could be attributed to the relative decrease in the amount of oxygen/oxygen functionalities on soot surface [54], in the order of 10% COD > 5% CD > D (which has been confirmed through elemental analysis, as shown later). The observed trend of activation energy is maintained up to a conversion level of 25%, beyond which, a reversal in activation energy trend between 5% CD and D soots is observed. The calculated value of the average activation energies at all conversion levels were 154, 165 and 170 kJ/mol for 10% COD, 5% CD, and D soots, respectively.

The activation energies for soot and carbon black are found in the range of 130-200 kJ/mol in the literature [55]. For commonly used palm oil-derived biodiesel blended fuels, which contain two oxygen atoms per molecule of methyl ester, (i.e., a high concentration of fuel-bound oxygen), the activation energy for soot oxidation is reported to be relatively lower in the range of 100-150 kJ/mol [54,56]. Such oxygen-rich fuel blends are reported to produce soots with high O/C ratio, and the subsequent decomposition of strongly bound oxygenated functional groups on soot to CO and CO$_2$ (especially above 400 oC) is known to enhance the soot oxidation rate [57,58]. The higher oxygen content in soot also leads to a better exposure of carbon atoms to ambient oxidants further accelerating the oxidation rate [59]. Since the concentration of fuel bound oxygen in camphor solid/camphor oil is relatively lower than biodiesel, the trend of average activation energies measured in this study for the 5% CD and 10% COD soot fall very much within the above expected range. However, it is to be noted that, some literature reports also have obtained contradictory results, wherein the soot oxidation rates have been found to be poorly dependent on the fuel-bound oxygen content/or the oxygen functionalities bonded to the soot surface [15]. Instead, the structural disorders in soot nano/microstructures have been reported to have much greater influence on the reactivity. The reversal in trend observed at the higher conversion levels in the present study, therefore, could have been influenced by a combination of parameters such as structural disorders, a larger amorphous content, a greater percentage of curvature in the lattice fringes, a decrease in the ratio of the number of edge to basal C atoms, and a higher amount of aliphatic hydrocarbons present in soot [60–62]. Therefore, all the soot samples were subjected to EDX elemental analysis, XRD, HRTEM, Raman, and EELS measurements to obtain more information about the average O/C ratio, microstructural disorder, PAH stack size, shape, and $\sigma/\pi$ bonding characteristics of soot, which will be discussed in the subsequent sections.
3.3 Elemental Analysis

To get an estimate of the relative distribution of carbon and oxygen atoms on the soot surface produced from D, 5% CD and 10% COD blended fuels, all soot samples were subjected to Energy Dispersive X-ray Spectroscopic (EDX) analysis using Ametex EDX PV6500 system, and the data was processed using eZAF Smart Quant analyser. The weight and the atomic percentages estimated for C and O (H atoms are not detected with this method) at three different portions of each soot sample are presented as inset in Figure 3.

Figure 3: EDX data of soot obtained from (a) D, (b) 5% CD, and (c) 10% COD, wherein A1, A2 and A3 represent elemental composition of three different regions of each soot sample.

Since element-specific calibration of the instrument was not performed, the results obtained are used as a qualitative indicator, but can be utilized to compare average O/C ratio of the three soot samples, as reported in [63]. As evident from Figure 3, the average O/C ratio increased from 0.224 (±0.002) for D soot to 0.239 (±0.004) and 0.264 (±0.003) for 5% CD and 10% COD soots, respectively. This indicates that the presence of higher amount of oxygen in the 10% COD blended fuel also induces relatively higher concentration of oxygen functionality in soot as compared to 5% CD and D, which correlates well with the initial oxidative reactivity of soot. Nonetheless, this could well be a coincidence and other structural disorders may have a significant role to play in inducing soot reactivity, as presented in the next section.
3.4 High Resolution Transmission Electron Microscopy (HRTEM)

Figure 4 represents the typical HRTEM images for D, 5% CD, and 10% COD soots at different resolutions of 5, 10 and 20 nm. It is evident from Figure 4(C,F,I) that all the soot samples are composed of agglomerated near-spherical particles, which are also called primary soot particles. These primary particles appear to be connected/fused to neighbouring particles due to the sintering effect. Each primary particle is composed of core-shell-like structure (see images A,D,G), where the core is composed of curved fringes of short length and the shell is composed of long range fringes, concentrically arranged along the periphery [6,64]. The short core fringes represent a high degree of structural disorder, which is due to the internal oxidation of soot particles, as they move up in the flame [6,64].

The comparative analysis of the average diameter of the primary particles of the three samples was performed using the Gatan software, where around 30-40 particles were identified from different HRTEM images at 20 nm resolution. The results are shown in Figure 5. It is evident that the average particle diameter is largest in the case of diesel soot (around 52 nm) as compared to the average diameters of 40 nm and 37 nm for 5% CD and 10% COD soots, respectively. The similarity in the particle diameters of 5% CD and 10 % COD soots indicates the role of camphor molecule in reducing the growth rate of soot nuclei, which is in agreement with the TSI measurements. It can be clearly seen in subfigure G that the fringe structure in the case of 10% COD soot is highly disordered with the existence of multiple nano core-shell structures within a single primary particle as compared to the other two soot samples.

The further quantitative analysis of fringe length and tortuosity was done using an in-house Matlab code (developed based on the algorithms of [65,66]), wherein the first step involves the negative transformation of the HRTEM image that converts it to grayscale. The multiple regions of interest (ROI) are then selected. On these ROI, several operations such as Gaussian filter, histogram equalization, and top-hat transformation are performed. In order to remove branches from fringes, firstly the all branch points are identified in the resulting image using the built-in Matlab “branchpoints” morphological operation. Starting from each of these branch points, the length of the fringe is noted in all directions. Once the smallest branch is identified, the connection to this branch is broken by setting the first pixel to this branch to zero. The newly constructed processed fringe structure is presented in Figure 6 (A,B,C) for diesel, 5% CD, and 10 % COD soot samples, respectively. The quantitative information about the fringe lengths and fringe curvature (i.e. tortuosity index) has been obtained by analysing all the fringe patterns in the above processed images, and the result obtained are presented as subfigures A1-A2, B1-B2, and C1-C2 in Figure 6 and Table 1.
Figure 4: HRTEM micrographs of (A-C) Diesel, (D-F) 5% CD, and (G-I) 10% COD soots at different resolutions of 5, 10 and 20 nm.

Figure 5: Average primary particle diameter for Diesel, 5% CD and 10% COD soot samples obtained using the HRTEM micrographs at 20 nm resolution.
Figure 6: Processed fringe nanostructure, fringe tortuosity and fringe length distribution in (i) diesel (A,A1,A2), (ii) 5% CD (B,B1,B2), and (iii) 10% COD (C,C1,C2) soots, respectively.

Table 1: Mean fringe length and mean fringe tortuosity of D, 5% CD and 10% COD soots.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean Fringe Length (nm)</th>
<th>Mean Fringe Tortuosity</th>
</tr>
</thead>
<tbody>
<tr>
<td>D</td>
<td>0.92</td>
<td>1.73</td>
</tr>
<tr>
<td>5% CD</td>
<td>0.86</td>
<td>1.82</td>
</tr>
<tr>
<td>10% COD</td>
<td>0.83</td>
<td>1.85</td>
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It is evident from Table 1 that diesel soot particles have the largest mean fringe length of 0.92 nm, which decreases to 0.86 nm and 0.83 nm in the cases of 5% CD and 10% COD.
respectively. Moreover, the analysis of the tortuosity index suggests that the short fringes in 10% COD soot also have the highest curvature (1.86) compared to diesel soot (1.73). Therefore, the shorter fringe length and the largely curved fringes together make the 5% CD and 10% COD soots highly disordered or amorphous, and hence, more prone to oxidation, as desired for the fast regeneration of DPF filters. The disorder in the crystal structure of these soot samples is further analysed using XRD and Raman spectroscopy.

3.5 X-Ray Diffraction analysis
The X-ray diffraction patterns of D, 5% CD, and 10% COD soots are presented in Figure 7, and the quantitative information about the PAH stacks is determined from their analyses. All the three soot samples displayed two typical distinctive peaks at 20 values of around 24.5° and 44°, respectively. A third minor peak around 80° of weaker intensity is observed due to the (110) plane, and is an indicator of the presence of non-graphitic carbon in the soot samples [67]. The observed broad peaks (24.5° and 44°) are indicative of the random alignment of nanocrystallites into a largely amorphous graphitic structure in soot. The peak at 20 value of 24.5° is assigned to the (002) plane, and it inculcates the information about the thickness of the PAH stack and the PAH interlayer spacing in it [25,68]. The peak at 20 value of 44° is assigned to the (100) plane and is indicative of the average PAH size in a soot sample. Further analysis of these peaks were carried out to obtain the structural parameters associated with the nanocrystallites present in soot samples, such as (i) PAH interlayer spacing (d_{002}) as per Bragg’s law (see equation E2), (ii) the nano-crystallite height (Lc), which gives the thickness of the PAH stack, using Scherrer formula (see Equation E3), and (iii) the nano-crystallite width (La) which provides the average PAH stack size using Equation E4 [69].

\[ d_{002} = \frac{\lambda}{2 \sin \theta_{002}} \]  
\[ L_c = \frac{0.9 \lambda}{B_{002} \cos \theta_{002}} \]  
\[ L_a = \frac{1.84 \lambda}{B_{100} \cos \theta_{100}} \]

In the above equations, \( \lambda \) is the wavelength of X-ray (1.54 Å for Cu-Kα), \( \theta_{002} \) and \( \theta_{001} \) are the Bragg’s angles, and \( B_{002} \) and \( B_{100} \) are the full width at half maximum (FWHM) for the two peaks, respectively. The Bragg’s angles and the FWHM were obtained via Gaussian fitting of the respective peaks using Matlab software. The obtained values are listed in Table 2.
Figure 7: XRD pattern of soot samples derived from 5, 5% CD and 10% COD.

Table 2: Crystallite parameters of soot samples obtained from XRD analysis

<table>
<thead>
<tr>
<th>Sample</th>
<th>d_{002} (Å)</th>
<th>L_c (Å)</th>
<th>L_a (Å)</th>
<th>L_c/d_{002}</th>
</tr>
</thead>
<tbody>
<tr>
<td>D</td>
<td>3.62</td>
<td>12.00</td>
<td>29.32</td>
<td>3.31</td>
</tr>
<tr>
<td>5% CD</td>
<td>3.63</td>
<td>11.78</td>
<td>27.35</td>
<td>3.24</td>
</tr>
<tr>
<td>10% COD</td>
<td>3.68</td>
<td>11.12</td>
<td>26.11</td>
<td>3.02</td>
</tr>
</tbody>
</table>

The calculated values of d_{002} for all three soot samples were in close range. The substantially greater values of d_{002} in soot compared to graphite (3.35 Å) indicates that all soot samples have a highly disordered structure. Moreover, the slightly greater d_{002} values in 10% COD soot as compared to the other two soot samples infer relatively weaker bonding interactions between the PAH stacks, which further enhances their oxidation rate in TGA experiments [70]. The estimated L_c value of 10% COD soot (11.12 Å) was found to be the lowest as compared to the other two soots, while the L_c/d_{002} ratio, which is a measure of the number of graphene layers in a coherent graphitic stack, decreases in the order, D > 5% CD > 10% COD. This suggests that the oxidation of camphor partially alters the oxidation mechanism of diesel in a way that favours face to face association over edge to edge coalescence of the graphene layers, as suggested by Hurt et al. [71]. The L_a values decrease substantially in the order D > 5% CD > 10% COD. The
low La value is an indicator of the small size of the PAH stack. Moreover, it has been well established that the smaller size of the PAH stack is associated with an increase in the edge carbon atoms as compared to the basal carbon atoms. Since the edge carbon atoms in the graphene sheet are more reactive compared to the basal carbon atoms, soot obtained from 10% COD are more easily oxidized by O$_2$ as compared to the other two soot samples [21].

3.6 Raman Spectroscopic analysis

The typical convoluted Raman bands, as observed in the first order region of 900-2000 cm$^{-1}$ of the three soot samples, is presented in Figure 8. The broad band near 1350 cm$^{-1}$, called the D band, is associated with the disordered structure of soot particles. The G band, observed near 1590 cm$^{-1}$, is relatively sharper, but slightly shifted above the expected value of 1580 cm$^{-1}$ for graphitic structure, further suggesting the high degree of disorder in the soot structure [72,73]. The five-curve deconvolution model was used to analyse all the Raman spectra, wherein Voigt function was used to fit the raw experimental data, as suggested in [74]. The corresponding five peaks were labelled as D1 = 1345 cm$^{-1}$, D2 = 1560 cm$^{-1}$, D3 = 1445 cm$^{-1}$, D4 = 1220 cm$^{-1}$, and G =1590 cm$^{-1}$. The D2 band corresponds to lattice vibrations, while the D3 band signifies the presence of highly amorphous carbon in soot. The genesis of D4 band is less understood, and is believed to be a result of C=C and C=C vibrations in polyene-like structures [72]. The relative intensity of the D and G bands serves as an indicator of the magnitude of discontinuities and disorientation of the graphene layers present in soot [74]. Therefore, the ratio of I$_D$/I$_G$ is obtained, and via an inverse relation, it is equated to the lattice width (La) using the Knight and White equation (E5), as suggested in [75]. The intensity of the D1 peak and a proportionality constant of 4.4 for the excitation wavelength of 515 nm is used, as suggested in [75,76]. The calculated values of I$_D$/I$_G$ and La are listed in Table 3.

$$L_a = 4.4 \left(\frac{I_{D1}}{I_G}\right)^{-1}$$  \hspace{1cm} (E5)

In Table 3, the I$_D$/I$_G$ ratio increases and the corresponding value of La decreases on moving from D soot to 10% COD soot. It is noted here that the calculated values of La are in close agreement with those obtained from XRD studies, which further confirms that the degree of lattice disorder increases due to the blending of camphor solid and camphor oil with diesel. Thus, both the XRD and the Raman studies very well compliment the TGA experiments, wherein the initial activation energy of soot oxidation by O$_2$ was found to decrease due to the blending of 5% camphor solid and 10% camphor oil with diesel in the early stages of oxidation process.
Figure 8: First order Raman spectra of A) Diesel, B) 5% CD and C) 10% COD soot samples.

Table 3: Structural parameters extracted from the Raman spectra of soot samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$I_{D1}/I_G$</th>
<th>$I_{la}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>D</td>
<td>1.45</td>
<td>29.7</td>
</tr>
<tr>
<td>5% CD</td>
<td>1.50</td>
<td>28.6</td>
</tr>
<tr>
<td>10% COD</td>
<td>1.59</td>
<td>26.8</td>
</tr>
</tbody>
</table>

3.7 Electron Energy Loss Spectroscopy (EELS)

The EELS technique serves as a very effective tool in investigating the ordered/disordered graphitic characteristics and aliphatic/aromatic structural composition of the soot samples based on the quantitative estimation of the peak structures and the relative intensity ratio of $I_{1s}$ to $\pi^*$ and $I_{1s}$ to $\sigma^*$ K edge transitions that occur on the soot surface [67]. When the EELS spectra of the three soot samples, presented in Figure 9, are compared, significant differences in the $\pi^*$ (285 eV) and $\sigma^*$ (292 eV) peaks are observed. The $\pi^*$ peak, which indicates the C=C
bond character due to presence of aromatics, is often referred to as the graphite peak, while the
σ* peak indicates the ordering of the graphitic layers [77]. Both the peaks are distinctly sharp
and well developed in the case of diesel soot. However, the two peaks are relatively less
developed and have broad hump-like structure in the case of soots collected from 5% CD and
10% COD, confirming that these soots have highly disordered graphene layers (in agreement
with the HRTEM, XRD, and Raman studies).

![Figure 9: Electron energy loss spectra of A) Diesel, B) 5% CD and C) 10% COD soots.](image)

The further analysis of the relative distribution of aromatics and aliphatics in soot is performed
by calculating the π*/σ* ratio using the EELS intensities of 285 and 292 eV peaks. The low
value of π*/σ* ratio indicates high aliphatic content in soot [68]. The obtained values of 0.767
(±0.019), 0.640 (±0.011), and 0.668 (±0.013) in the cases of D, 5% CD and 10% COD soots,
respectively, confirm that the 5% CD soot had the highest aliphatic content followed by 10%
COD and D soots. It is well known that aliphatic compounds are oxidized at a faster rate than
aromatics. This could explain the reversal in activation energy trend observed at the conversion
level of 30% in TGA, wherein 5% CD soot has the lowest activation energy. As the oxidation
progresses to higher conversion levels, the lack of smaller PAH crystallites and the loss of oxygenated species followed by the graphitization of soot structure could have resulted in a further increase in the activation energy for the oxidation of 5% CD soot. Since, the diesel soot already possesses relatively higher degree of graphitic character (as confirmed through EELS studies), the relative change in the graphitic character at higher conversion levels during its oxidation could be minimal, and hence, the increase in its activation energy of oxidation is relatively lower [67].

4 Conclusion
The effect of camphor solid and camphor oil addition to diesel on the sooting propensity and their subsequent effect on soot nanostructures were investigated. It was observed through smoke point and TSI measurements that the blending of both camphor solid (up to 5%) and camphor oil (up to 10%) could reduce soot formation rate possibly through the improved combustion of the fuel. The subsequent analysis of the produced soot particles through HRTEM, XRD, and Raman studies revealed that soots obtained from the above blends had smaller primary particle diameter with core-shell structure and a greater percentage of short lattice fringes with increased fringe tortuosity as compared to diesel soot. This resulted in an enhancement in the degree of disorder in the crystal structure, thus making it prone to easy attack by O\textsubscript{2} during oxidation. Moreover, the presence of higher oxygen and aliphatic contents in soot produced from the blends, as observed through EDX and EELS measurements, resulted in the enhancement of O\textsubscript{2}-induced soot oxidation kinetics as compared to diesel soot. All the above results therefore confirm that the blending of up to 10% camphor oil with diesel will not only improve the fuel combustion to reduce soot formation, but would also significantly improve the DPF regeneration efficiency by making soot more susceptible to oxidation.

Supporting Information
A plot showing the measure fuel flow rate as a function of flame height for different fuels and a plot on soot conversion and soot conversion rate at different temperatures and heating rates are provided in the supporting information file.

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