

PHYSICOCHEMICAL AND  
TOXICOLOGICAL ASSESSMENT OF THE  
SEMI-VOLATILE AND NON-VOLATILE  
FRACTIONS OF PM FROM HEAVY- DUTY  
VEHICLES OPERATING WITH AND  
WITHOUT EMISSIONS CONTROL  
TECHNOLOGIES

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Final report

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## Table of Contents

|   |    |
|---|----|
| <b>List of Figures</b> .....  | 4  |
| List of Tables .....  | 5  |
| Abstract.....   | 6  |
| 1. Executive Summary.....   | 7  |
| 1.1 Background .....  | 7  |
| 1.2 Methods.....  | 7  |
| 1.3 Results.....  | 7  |
| 1.4 Conclusions .....   | 8  |
| 2. Introduction .....   | 10 |
| 2.1 Statement of Significance .....   | 10 |
| 2.2 Background .....  | 11 |
| 2.3 Study Objectives.....   | 12 |
| 2.4 Major Components of the Study.....  | 12 |
| 3. Experimental Methods.....  | 14 |
| 3.1 Dynamometer Setup.....  | 14 |
| 3.2 Vehicles .....  | 15 |
| 3.3 Equipment and Instruments .....   | 17 |
| 4. Physical Characteristics of the Exhaust Particles (Semi-volatile and Non-volatile) from Retrofitted Vehicles ..... | 19 |
| 4.1 Introduction .....  | 19 |
| 4.2 Data Reduction.....   | 19 |
| 4.3 Results.....  | 21 |
| 4.4 Conclusions .....   | 32 |
| 5. Chemical Characteristics of Exhaust Particles from Retrofitted Vehicles.....                                       | 34 |
| 5.1 Introduction .....  | 34 |
| 5.2 Sample Analysis.....  | 34 |
| 5.3 Results.....  | 35 |
| 5.4 Conclusions .....   | 50 |
| 6. Toxicological Characteristics of Semi-volatile and Non-volatile Fraction of DEPs from Retrofitted Vehicles .....   | 51 |

|     |   |    |
|-----|---|----|
| 6.1 | Introduction .....                        | 51 |
| 6.2 | Sample Analysis.....                      | 52 |
| 6.3 | Results .....                             | 53 |
| 6.4 | Conclusions .....                         | 63 |
| 7.  | List of Publications from this Study..... | 64 |
| 8.  | Major Conclusions.....                    | 65 |
| 9.  | Acknowledgments.....                      | 66 |
| 10. | References .....                          | 67 |

## List of Figures

|  |    |
|--|----|
| Figure 3-1 Dynamometer set-up.....   | 15 |
| Figure 4-1 Particle number size distributions .....  | 23 |
| Figure 4-2 Number (DMS) and mass emission factors (nano-MOUDI).....  | 27 |
| Figure 4-3 Reduced Variable 1 (R): Ratio of total particle to solid or thermo-denuded particle count. ....   | 28 |
| Figure 4-4 Size distribution of thermo denuded aerosols for cruise mode.....   | 29 |
| Figure 4-5 Mass fractions remaining at different thermodenuder temperature settings. ....  | 29 |
| Figure 4-6 Reduced Variable 2: Surface rated diameter.....   | 30 |
| Figure 4-7 Time series of surface diameter (Ds), surface concentration (EAD signal and total particle concentration (DMS count) for a) DPX; b) Baseline- UDDS run.....   | 31 |
| Figure 4-8 Reduced Variable 3: Mass specific surface concentration. ....   | 32 |
| Figure 5-1 Emission factors of integrated chemical species measured by MOUDI-NanoMOUDI. a). Cruise 50 mph b). Transient UDDS.....  | 36 |
| Figure 5-2 EC, WSOC and WIOC emission rates at a) Cruise 50mph and b) Transient UDDS c) Proportion of WSOC to OC. ....   | 38 |
| Figure 5-3 (a) Emission factors of PAHs, hopanes and steranes of baseline and control device-equipped vehicles in UDDS cycle. (b) Emission factors of n-alkanes and acids of baseline and control device-equipped vehicles in UDDS cycle. ....   | 40 |
| Figure 5-4 (a) Emission factors of PAHs, hopanes and steranes of baseline and control device-equipped vehicles in cruise cycle. (b) Emission factors of n-alkanes and acids of baseline and control device-equipped vehicles in cruise cycle.....  | 42 |
| Figure 5-5 Comparison of the ratios of PAHs, hopanes and steranes to organic carbon emission factors in $\mu\text{g/g}$ from baseline and controlled vehicles. (a) running on UDDS cycle. (b) cruise cycle. (c) Ratios of PAHs, hopanes and steranes to organic carbon emission factors in $\mu\text{g/g}$ as reported by Riddle et al., (2007)..... | 44 |
| Figure 5-6 Emission rates of size-resolved PM chemical species. a) Cruise 50 mph, b) Transient UDDS c) Baseline vehicle.....   | 46 |
| Figure 5-7 a, b: Size-resolved EC/OC mass distribution at a) Cruise mode b) UDDS mode.....   | 49 |
| Figure 6-1 a, b, c: DTT consumption in n-moles $\text{min}^{-1} \mu\text{g}^{-1}$ of PM of thermo-denuded and undenuded PM.....  | 55 |
| Figure 6-2 Relationship between oxidative activity and semi-volatile PM fraction at $150^\circ\text{C}$ .....  | 56 |
| Figure 6-3a,b,c: DTT consumption per unit distance ( in n-moles $\text{min}^{-1} \text{km}^{-1}$ ) traveled by vehicles for Cruise and UDDS cycles and per hour( in n-moles $\text{min}^{-1} \text{hr}^{-1}$ ) for Idle.....   | 58 |
| Figure 6-4 Reactive Oxygen Species (ROS) activity of PM from the tested vehicles, expressed as (a) per mass of PM, and (b) per km (or hr for idle) of vehicle driven.....  | 59 |
| Figure 6-5 Percent removal of ROS activity in relation to that of aggregate water soluble metals after Chelex treatment of the exhaust PM samples from test vehicle-configurations under different driving cycles.....   | 60 |

## List of Tables

|  |    |
|--|----|
| Table 3-1 Details of test fleet.....   | 9  |
| Table 4-1 Size fractionated mass EFs (mg km <sup>-1</sup> for Cruise and UDDS, mg hr <sup>-1</sup> for Idle).....  | 26 |
| Table 6-1 Correlation coefficient (R) and significance level (P) for DTT activity and selected chemical species.....                                       | 61 |
| Table 6-2 Correlation coefficients and associated levels of significance for the regression between ROS activity and major water-soluble metals of PM..... | 62 |

## Abstract

This final report presents the investigation results of the CARB vehicle emissions study, with focus on the physicochemical and toxicological properties of the semi-volatile and non-volatile fractions of PM from heavy duty diesel vehicles, operating with and without emissions control technologies. A wide variety of diesel fuelled vehicles, representing the current in-use fleet, have been tested in the California Air Resources Board's (CARB) heavy-duty diesel emission testing laboratory (HDETL) in downtown Los Angeles, including a 1998 Kenworth truck, a diesel hybrid electric bus, a school bus, and a Caltrans truck on three driving cycles, i.e. steady state cruise (50mph), transient [EPA urban dynamometer driving schedule (UDDS)] and idle to simulate various real-world driving conditions. The tested emission control devices include a Continuously Regenerating Technology (CRT); CRT in combination with a selective catalytic reduction system (Zeolite or vanadium based SCRTs) etc. Detailed physico-chemical and toxicological characteristics of PM were measured for each vehicle and driving cycle, including physical properties (e.g. PM mass and size distribution), chemical (EC, OC, organic compounds, trace elements, inorganic ions) and toxicological [dithiothreitol (DTT) and macrophage reactive oxygen species (ROS) assays] characterization of the collected PM samples.

Substantial reduction in PM mass emissions (>90%) was accomplished for the tested vehicles operating with advanced emission control technologies, while such a reduction was not observed for particle number concentrations under cruise conditions, with the exceptions of the Hybrid-CCRT and EPF vehicles. Significant reductions in the emission of major chemical constituents (TC, OC, EC, and organic compounds) were achieved by the introduction of retrofits. Sulfate dominated the PM composition in vehicle configurations (V-SCRT-UDDS, Z-SCRT-Cruise, CRT, DPX) with considerable nucleation mode and total carbon was dominant for the configurations with less (ZSCRT-UDDS) or insignificant (CCRT, Horizon) nucleation. V-SCRT and Z-SCRT effectively reduced PAHs, hopanes and steranes, n-alkanes and acids by more than 99%, and often to levels below detection limits for both cruise and UDDS cycles. The CRT technology also showed similar reductions with SCRT for medium and high molecular weight PAHs, acids, but with slightly lower removal efficiencies for other organic compounds. Despite an increase in the intrinsic activity (both DTT and ROS, per mass basis) of exhaust PM with use of most control technologies, the overall activity (expressed per km or per hr) was substantially reduced for retrofitted configurations compared to the baseline vehicle. Significant reduction in DTT activity (by 50-100%) was observed for thermally-denuded PM from vehicles with retrofitted technologies (PM with significant semi-volatile fraction). On the other hand, Chelex treatment of PM samples removed a substantial ( $\geq 70$  %) fraction of the ROS activity, corroborating the effect of transition metals on this activity. Correlation analysis performed between measured activity and the chemical constituents showed that DTT activity is strongly associated ( $R=0.94$ ) with the water soluble organic carbon (WSOC), while the ROS activity was mostly driven by Fe content of the PM samples.

## 1. Executive Summary

### 1.1 Background

The objective of this project was to enhance a planned ARB vehicle emissions study with the research component to determine the physicochemical and toxicological properties of particulate matter (PM) from heavy duty vehicles operating with and without emissions control technologies. The key objective of this comprehensive 4-year project was to determine the physicochemical and toxicological properties of the semi-volatile and non-volatile fractions of PM from heavy duty diesel vehicles operating with and without emissions control technologies.

### 1.2 Methods

In this study, we assessed the PM-related oxidative activity from a wide variety of vehicles to represent the in-use fleet, including diesel vehicles with and without advanced PM emission control technologies. We investigated different driving cycles, since engine operation is known to affect the concentration, relative amounts and chemical composition of the nucleation and accumulation PM modes emitted. Experiments were carried out at the California Air Resources Board's (CARB) heavy-duty diesel emission testing laboratory (HDETL) in downtown Los Angeles. Ayala et al., (2002) described the dynamometer specifications in details. Three driving cycles, i.e. steady state cruise (50mph), transient [EPA urban dynamometer driving schedule (UDDS)] and idle were tested to simulate various real-world driving conditions. The fuel used to run the engines was CARB ultra-low sulfur diesel (ULSD) with sulfur content less than 15 ppm. Tunnel blank levels were measured and vehicles were conditioned (warmed up) every day before the start of official runs. The CVS was cleaned prior to starting the project. The test fleet comprised of four heavy-duty diesel vehicles in seven configurations. A 1998 Kenworth truck served as a baseline vehicle, without any emission control technology. The same Kenworth truck was also tested with three different control technologies: a Continuously Regenerating Technology [CRT], consisting of a diesel oxidation catalyst (DOC) followed by an uncatalyzed trap; CRT in combination with a selective catalytic reduction system [Zeolite or vanadium based SCRTs]. The other three test vehicles were a diesel hybrid electric bus, a school bus, and a Caltrans truck. Detailed physico-chemical and toxicological characteristics of PM were measured for each vehicle and driving cycle, including physical properties (e.g. PM mass and size distribution), chemical (EC, OC, organic compounds, trace elements, inorganic ions) and toxicological [dithiothreitol (DTT) and macrophage reactive oxygen species (ROS) assays] characterization of the collected PM samples.

### 1.3 Results

Substantial reduction in PM mass emissions (>90%) was accomplished for the HDDV operating with advanced emission control technologies. This reduction was not observed for particle number concentrations under cruise conditions, with the exceptions of the Hybrid-CCRT and EPF vehicles, which were efficient in controlling both - mass and number emissions. In general, significant nucleation mode particles (<50nm) were formed during cruise cycles in comparison with the UDDS cycles, which emit higher PM mass in the accumulation mode. The nucleation mode particles (<50nm) were mainly internally mixed, and evaporated considerably between 150 to 230°C.

Significant reductions in the emission of major chemical constituents (TC, OC, EC, and organic compounds) were achieved by the introduction of retrofits. V-SCRT and Z-SCRT effectively reduced PAHs, hopanes and steranes, n-alkanes and acids by more than 99%, and often to levels below detection

limits for both cruise and UDDS cycles. The CRT technology also showed similar reductions with SCRT for medium and high molecular weight PAHs, acids, but with slightly lower removal efficiencies for other organic compounds. Sulfate dominated the PM composition in vehicle configurations (V-SCRT-UDDS, Z-SCRT-Cruise, CRT, DPX) with considerable nucleation mode and TC was dominant for the configurations with less (ZSCRT-UDDS) or insignificant (CCRT, Horizon) nucleation.

Despite an increase in the intrinsic activity (both DTT and ROS, per PM mass basis) of exhaust PM with use of most control technologies, the overall activity (expressed per km or per hr) was substantially reduced for retrofitted configurations compared to the baseline vehicle. Significant reduction in DTT activity (by 50-100%) was observed for thermally-denuded PM from vehicles with retrofitted technologies (PM with significant semi-volatile fraction). On the other hand, Chelex treatment of undenuded PM samples removed a substantial ( $\geq 70\%$ ) fraction of the ROS activity. Correlation analysis performed between measured activity and the chemical constituents showed that DTT activity is strongly associated ( $R=0.94$ ) with the water soluble organic carbon (WSOC), while the ROS activity was mostly driven by the Fe content of the PM samples.

## 1.4 Conclusions

The newer diesel engines with emission control devices are very efficient in reducing the mass emission of particulate matter. However, enhanced formation of nucleation mode particles is observed in the exhausts of some retrofitted configurations. The vast majority of these nucleation mode particles are semi-volatile in nature.

The total emissions (per distance of vehicle traveled) of the major chemical species (e.g. elemental and organic carbon) are substantially reduced in the retrofitted vehicles compared to the baseline vehicle. However, sulfate emissions increase for the configurations with enhanced nucleation mode particles in their exhaust. Although the retrofitted vehicles emit less water soluble organic carbon (WSOC) per mile of vehicle driven, the per PM mass water soluble fraction of the organic carbon (WSOC/OC) is increased for most configurations.

With the introduction of control devices, the individual ratios of speciated organic compounds to OC are reduced significantly for PAHs, while the reduction was more modest for hopanes and steranes, implying that fuel and lubricating oil have substantially different contributions to the OC emitted by vehicles operating with control devices compared to the baseline vehicle. We hypothesize that PAHs can form in combustion processes and/or originate from diesel fuel, whereas hopanes and steranes come from lubricating oils.

Despite an increase in the intrinsic oxidative activity (both DTT and ROS, per mass basis) of exhaust PM with the use of control technologies for most configurations, the overall activity (expressed per km or per hr) was substantially reduced for retrofitted configurations compared to the baseline vehicle.

The semi-volatile fraction of the exhaust particles was observed to be highly oxidative in nature as demonstrated by a significant reduction in DTT activity (by 50-100%) observed for thermally-denuded PM. However, non-volatile species - particularly transition metals, are also responsible in cellular oxidative stress, as indicated by a substantial removal ( $\geq 70\%$ ) of the ROS activity after Chelex treatment of the PM samples.

An important caveat of the toxicological findings of this study is that they are all based on molecular or cellular assays that examine the toxicity of the PM suspension collected from a given vehicle and driving

configuration based on bulk PM mass. By their nature and design, these investigations did not take into account important parameters determining the toxicity and overall health effects attributable to the inhalation of an aerosol, such as particle size. The substantial reduction in the overall particle size distribution of newer vehicles creates an aerosol with a much higher lung deposition fraction than the baseline vehicle, and with considerably different toxicokinetics inside the human body once inhaled. Such important investigations can only be addressed by in vivo inhalation exposure studies to these aerosols, whether using animal models or human volunteers (or both), and are greatly needed in order to provide a more complete perspective to the results of this study.

## 2. Introduction

### 2.1 Statement of Significance

This study addresses the well-publicized and significant effects of particulate matter (PM) emissions from heavy-duty diesel vehicles on health and the environment. Although PM originating from these sources is known to be responsible for a myriad of adverse health outcomes, ranging from cancer to cardiopulmonary disease, and environmental problems, ranging from global warming to acid rain, the fields of combustion, atmospheric science, and public health are sufficiently separated that a unified plan of action is presently not feasible. Epidemiological and toxicological studies have demonstrated strong links between ambient particulate matter mass exposure and adverse health outcomes (NRC, 2004). However, it is not clear which physical or chemical properties of particulate matter (PM) pose the greatest health risk.

The objective of this project was to enhance a planned ARB vehicle emissions study with the research component to determine the physicochemical and toxicological properties of particulate matter (PM) from heavy duty vehicles operating with and without emissions control technologies. Heavy-duty diesel trucks constitute only a small fraction of the total fleet in California but have an important contribution to the emissions of fine and ultrafine particles. The outcomes of this study will be very helpful in assessing the health consequences of population exposure to diesel traffic sources in Los Angeles, where on average individuals spend 2-3 hours per day commuting in heavily congested freeways. The Los Angeles Basin (LAB) is home to more than 15 million individuals and it has been described as the most polluted airshed in the US, with a complex, persistent and unique PM pollution problem. Despite considerable improvements in air quality over the past two decades, Los Angeles continues to exhibit the most severe ozone and PM air quality problem in the U.S. because of the continuing increase in population and traffic. The LAB covers approximately 12,000 square miles and this region has experienced faster population growth than the rest of the nation since World War II. Growth is expected to increase significantly towards the end of the century and through 2020. The population is expected to increase 1.5 times by the year 2020 from 1990 levels. By 2020 the south coast air basin is expected to have a population of 21,000,000 as compared to 3,800,000 in 1950. The region has tens of thousands of heavy duty trucks transporting goods from the LA Harbor (the busiest in the US) to the rest of the US, all putting primary particles and reactive gases that act as particle precursors into the atmosphere.

The topography and climate of the LAB are some of the reasons for this area having high air pollution potential. Several air quality studies including our own, have shown that the LAB is known for distinct areas of different meteorological conditions (microclimates), which result in enormous spatial variations of PM, including species of toxicological interest, i.e., organic and elemental carbon, (OC, EC), nitrate, metals, polycyclic aromatic hydrocarbons (PAH) and gaseous co-pollutants. This is in contrast to metropolitan areas of the eastern US, where a large fraction of PM pollution is regional, and in which PM<sub>2.5</sub> and their components are more uniformly distributed.

A major objective of this comprehensive 4-year project was to determine the physicochemical and toxicological properties of the semi-volatile and non-volatile fractions of PM from heavy duty diesel vehicles operating with and without emissions control technologies. With the exception of uncontrolled diesel, the majority of PM emitted by these vehicles is semi volatile in nature, formed by condensation of super saturated vapors as they cool in the ambient atmosphere. Measurements of the relative toxicity of these particles compared to the more refractory (non volatile “accumulation” mode) PM are valuable in terms of assessing the need for additional control strategies.

As part of this study, we assessed the PM-related oxidative activity from a wide variety of vehicles to represent the in-use fleet, including diesel vehicles with and without advanced PM emission control technologies. We investigated different driving cycles, since engine operation is known to affect the concentration, relative amounts and chemical composition of the nucleation and accumulation PM modes emitted. This study highlights the need of promulgating effective vehicular emission control strategies aiming at reducing public risk in terms of acute as well as chronic health outcomes.

The semi-volatile PM fraction of vehicle emissions is extremely important in terms of its contribution to human exposure. Current emission control technologies remove effectively the non-volatile fraction, but are less effective at controlling the volatile fraction. Removal of the non-volatile PM fraction can increase the concentration of the volatile aerosol fraction by enhancing nucleation of condensing organic vapors (Bagley et al., 1998; Kittelson, 1998). Knowing how the toxicity of vehicular PM varies with particle component volatility will be vital in directing the design of emission control technologies that will be more effective in protecting the public health. The discussion on the treatment of the solid (refractory) and semi-volatile fractions in emission tests is still ongoing. While it is becoming obvious that the PM mass measurement alone is no longer adequate for emissions certification, the best alternative is still under discussion. The recommendation by European Union (EU) is to measure the number concentration of solid particles in addition to mass, maintaining the test cycles and the primary dilution system (CVS full exhaust flow tunnel) used now. EU claims that “this is a pragmatic approach, which allows a fast realization” (Burtscher, 2005). The unequivocal demonstration of the significant toxicity of the semi-volatile nucleation PM mode from heavy duty engines in this study raises significant concerns on the validity, effectiveness and overall wisdom of such a standard. The research described in this study generates invaluable information to the CARB, who is responsible for developing statewide strategies to reduce the emission of smog-forming pollutants and toxics by mobile sources.

## 2.2 Background

Measurements of particle emissions from motor vehicles have been accomplished via dynamometer source testing (Chase et al., 2000; Kwon et al., 2003; Sakurai et al., 2003a; Sakurai et al., 2003b; Schauer et al., 1999; Schauer et al., 2002), roadway tunnel sampling (Fraser et al., 1998), on-road chase experiments (Shah et al., 2004a; Vogt et al., 2003) and roadside measurements (Harrison et al., 2003; Sturm et al., 2003; Zhu et al., 2002a; Zhu et al., 2004; Zhu et al., 2002b). In general, particles directly emitted from diesel engines are in the size range from 20 nm to 130 nm in aerodynamic diameter for (Morawska et al., 1998). Many of the organic chemical constituents of these particles are semi-volatile, existing simultaneously in the gas and particle phases at equilibrium (Schauer et al., 1999; Schauer et al., 2002). Changes in ambient temperature and gas phase concentrations of these components can affect the measured particle size distributions due to evaporation or condensation. It has been shown that the particle size distributions determined by dynamometer testing are dependent on the dilution ratios and

dilution air conditions of the sampling apparatus (Holmen and Qu, 2004; Lyyranen et al., 2004; Vaaraslahti et al., 2004).

Recent emissions testing in either dynamometer or on-road testing facilities have shown that particles emitted from these diesel vehicles are externally mixed, i.e. different particles of the same size can have different chemical compositions. Depending on vehicle type, age and ambient conditions, between 70-90% of the particles by number and 10-30% by mass may consist of more volatile material than others (known as semi-volatile), and partially or completely evaporate upon heating (Sakurai et al., 2003b).

These results were further corroborated by the modeling study of Zhang et al., (2004), as well as the experimental field studies of Kuhn et al., (2005b), who demonstrated that condensation, evaporation, and dilution were the major factors affecting aerosol size distributions in the first 250 m downwind from freeways. The high concentrations of particle numbers in the proximity of freeways raise concerns for population exposure during commute. The particle volatility, which causes the dynamically shifting size distributions of these freshly emitted particles, needs to be better characterized to accurately assess population exposure to the physical and chemical properties of PM. For example, the volatility of these particles explains the more rapid decay in their concentration with respect to distance from a roadway, compared to that of non-labile PM species (such as EC) or gaseous co-pollutants such as CO and NO<sub>x</sub>, the concentration decrease of which would be affected mostly by atmospheric dilution. The increased dilution with distance to roadway decreases the concentration of vapors in equilibrium with these particles, which enhances evaporation of these vapors from the particle phase to re-establish equilibrium (Zhang et al., 2004). The exposure and health implications of these findings have not yet been investigated. Considering that the majority of people's exposure during commute will be dominated (at least based on particle numbers) by these particles, it would be useful to know whether the non-volatile or semi-volatile material is more toxic.

## 2.3 Study Objectives

The main goal of this research is to provide information on the physico-chemical characteristics as well as relative toxicity of semi-volatile fraction of the diesel exhaust particles compared to the larger, non-volatile (refractory), mostly carbonaceous fraction, collected from a variety of different engines, each operating under different driving conditions in dynamometer facilities. CARB plans an extensive emissions research program at its heavy-duty emissions laboratories and our study complements and enhances CARB's research efforts and leverages the significant capacity offered by our group. This was a collaborative study led by the Research Division of the California Air Resources Board and the Aerosol Laboratory of the University of Southern California with support from University of California Los Angeles (UCLA) and University of Wisconsin Madison (UWM) for chemical and toxicological analysis.

## 2.4 Major Components of the Study

The major components of this study can be summarized as below:

- 1) Conduct dynamometer experiments to collect the total and size fractionated diesel exhaust particles on appropriate substrates for physical, chemical and toxicological analysis, along with real-time data collection (particle number concentration and size distribution)

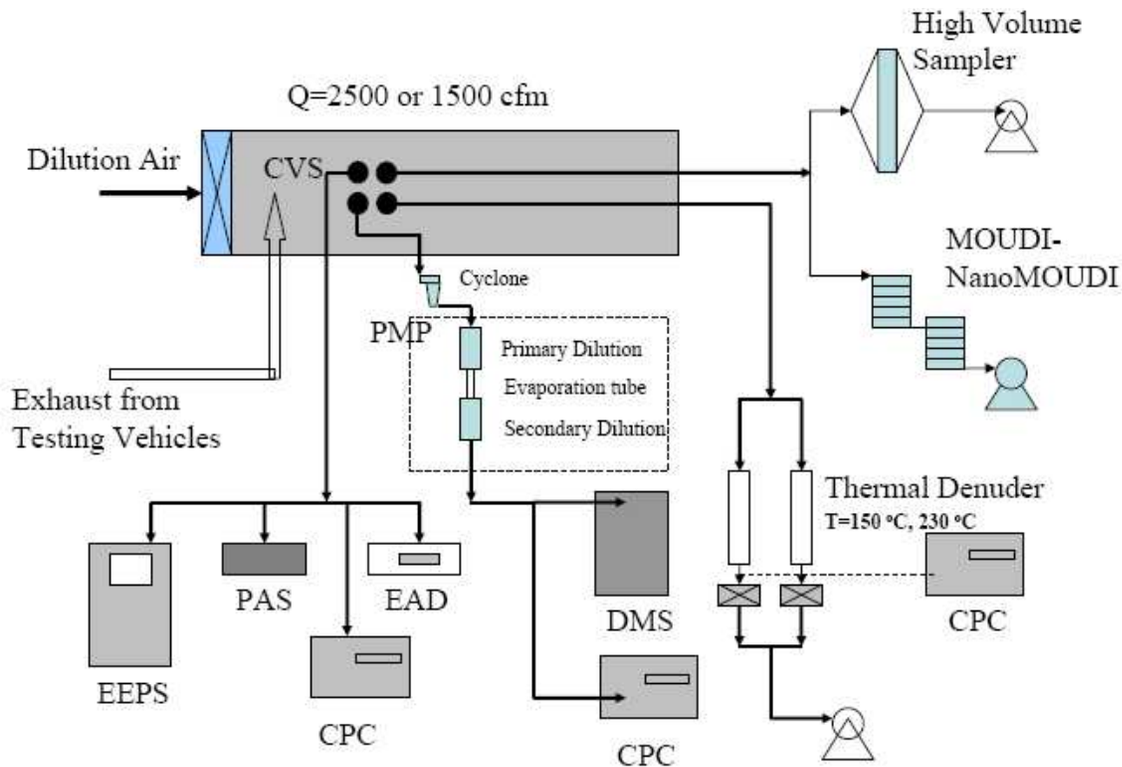
- 2) Conduct the physical (e.g. PM mass and size distribution), chemical (EC, OC, organic compounds, trace elements, inorganic ions) and toxicological [dithiothreitol (DTT) and macrophage reactive oxygen species (ROS) assays] characterization of the collected PM samples.
- 3) Conduct data analysis, and prepare manuscripts for publication in the peer-reviewed literature.
- 4) Prepare quarterly progress reports for CARB.
- 5) Prepare and submit the final report to CARB at the end of the project.

## 3. Experimental Methods

### 3.1 Dynamometer Setup

Experiments were carried out at the California Air Resources Board's (CARB) heavy-duty diesel emission testing laboratory (HDETL) in downtown Los Angeles. Ayala et al., (2002) described the dynamometer specifications in details. Figure 3.1 shows the schematic of the experimental setup. The sampling train includes heavy-duty chassis dynamometer, constant volume sampling (CVS) dilution tunnel and aerosol samplers. Diesel vehicle exhausts were transported by a stainless steel hose pipe and diluted with filtered air through the CVS. Measurements were taken 18 diameter lengths downstream of the exhaust introduction in the CVS. Three driving cycles, i.e. steady state cruise (50mph), transient [EPA urban dynamometer driving schedule (UDDS)] and idle were tested to simulate various real-world driving conditions. The fuel used to run the engines was CARB ultra-low sulfur diesel (ULSD) with sulfur content less than 15 ppm. Tunnel blank levels were measured and vehicles were conditioned (warmed up) every day before the start of official runs. The CVS was cleaned prior to starting the project.

The dilution air flow rate at the CVS was 2600 cfm ( $74 \text{ m}^3 \text{ min}^{-1}$ ) for cruise and UDDS cycles, and 1600 cfm ( $45.4 \text{ m}^3 \text{ min}^{-1}$ ) for the idle, respectively. For EPF and Hybrid-CCRT, the flow rates were maintained at 1600 cfm for all cycles. These flow-rates result in approximate dilution ratios of 6-9 for cruise, 5-80 for UDDS, and 15-25 for idle.



**Figure 3-1 Dynamometer set-up**

### 3.2 Vehicles

The test fleet comprised of four heavy-duty diesel vehicles in seven configurations (Table 3.1). A 1998 Kenworth truck served as a baseline vehicle, without any emission control technology. The other three test vehicles were a diesel hybrid electric bus, a school bus, and a Caltrans truck.

The diesel hybrid electric bus (San Joaquin Valley RTD) is equipped with a catalyzed continuously regenerative trap, or CCRT, consisting of a diesel oxidation catalyst (DOC) followed by a catalyzed trap, which was virtually brand-new, with only 1000 miles on the odometer.

The Kenworth truck was tested with three different control technologies: a Continuously Regenerating Technology [CRT], consisting of a diesel oxidation catalyst (DOC) followed by an uncatalyzed trap; CRT in combination with a selective catalytic reduction system [Zeolite or vanadium based SCRTs]. The two SCRT technologies consist of a wall-flow particulate trap (CRT) followed by a SCR section. The CRT was the same in each configuration. The difference between them lies in the choice of catalysts (vanadium or Zeolite) for the SCR to control oxides of nitrogen (NO<sub>x</sub>).

The Caltrans truck, with a smaller engine (7.6 L) than the Kenworth truck (11 L, Table 3.1), is retrofitted with an Engelhard DPX filter. The DPX filter is comprised of a diesel particulate trap with a catalytic wash-coat.

The last test vehicle was an Elk Grove school bus, equipped with an electric particle filter (EPF-Horizon). The EPF consists of a non-catalyzed silicon carbide substrate for PM control, coupled with an electric heating element and a small blower. The trap is regenerated periodically using electricity from the grid (plug in configuration) during non-operational periods—mostly at night.

Hereafter, the test fleet is referred as baseline, CRT, V-SCRT, Z-SCRT, DPX, hybrid/CCRT and School Bus/Horizon/EPF.

**Table 3-1 Details of test fleet**

| Vehicle                          |                        |      |        |                  |           |                | Engine                 |          | After treatment (AT)                 |                         | Dilution                          |
|----------------------------------|------------------------|------|--------|------------------|-----------|----------------|------------------------|----------|--------------------------------------|-------------------------|-----------------------------------|
| Make                             | Nomenclature           | Year | Miles  | Curb Weight (lb) | GVWR (lb) | Tested Wt (lb) | Model                  | Size [L] | Type                                 | Miles on AT             | Approximate Dilution Ratio in CVS |
| Kenworth                         | V-SCRT                 | 1998 | 360000 | 26,640           | 80,000    | 53,320         | Cummins M11, reflashed | 11       | Vanadium based SCRT                  | 50,000                  | 9.2 Cruise 6-30 UDDS 14 Idle      |
| Kenworth                         | Z-SCRT                 | 1998 | 360000 | 26,640           | 80,000    | 53,320         | Cummins M11, reflashed | 11       | Zeolite based SCRT                   | 0 on SCR, 50,000 on CRT | 9.2 Cruise 6-30 UDDS 14 Idle      |
| International                    | DPX                    | 1999 | 40,000 | 15,030           | 27,500    | 20,920         | International DT466E   | 7.6      | Engelhard DPX                        | 30,000                  | 6.2 Cruise 5-25 UDDS 22 Idle      |
| Gillig (35ft) with Alison Hybrid | Hybrid/CCRT            | 2007 | 1000   | NA               | NA        | NA             | Cummins                | 5.9      | CCRT                                 | 1000                    | 5-50 UDDS                         |
| Thompson-School Bus              | School Bus/EPF/Horizon | 1988 | 325000 | NA               | NA        | NA             | Cummins                | 5.9      | Cleaire - Horizon                    | 32,000                  | 8.3 Cruise 8-80 UDDS 25 Idle      |
| Kenworth                         | CRT                    | 1998 | NA     | 26,640           | 80,000    | 53,320         | Cummins M11, reflashed | 11       | Continuously Regenerating Technology | 64,000                  | 9.2 Cruise 6-30 UDDS 14 Idle      |
| Kenworth                         | Baseline               | 1998 | 374000 | 26,640           | 80,000    | 53,320         | Cummins M11, reflashed | 11       | None                                 | NA.                     | 9.2 Cruise 6-30 UDDS 14 Idle      |

### 3.3 Equipment and Instruments

Specific descriptions of some of the instruments and equipment used for this study are provided below.

#### ***Nano-MOUDI***

Size-resolved samples were collected using a micro-orifice uniform deposited impactor (MOUDI) upstream of a nano-MOUDI (MSP Corporation, Minneapolis, MN) loaded with pre-cleaned aluminum foil substrates. Particles were classified in the following aerodynamic size ranges: 10-18 nm, 18-32 nm, 32-56 nm, 56-100 nm, 100-180 nm, 180 nm-2.5  $\mu\text{m}$  and  $>2.5 \mu\text{m}$ . The MOUDI-Nano MOUDI tandem was operated for multiple runs in order to accumulate sufficient mass for chemical analysis for each vehicle and driving cycle.

#### ***DMS/ EEPS***

Size distribution of engine exhausts from the CVS was monitored every second by two multiple channel differential mobility spectrometers: a DMS500 (Cambustion) and an engine exhaust particle sizer (EEPS 3090, TSI Inc.). Both DMS and EEPS classify particles on the basis of their electrical mobility diameter. The cut-off size ranges of EEPS and DMS are 5.6-523 nm and 4.5- 1000 nm, respectively. With high time resolutions, they are both capable of tracking transient particle behavior, especially during UDDS cycles. With few exceptions (V-SCRT, Z-SCRT only), in which the DMS was placed downstream of the particle measurement program (PMP) sampler, both instruments were connected directly to CVS.

#### ***PMP***

A particle measurement program (PMP) protocol was developed in Europe to measure the solid particle emissions from light duty vehicles. The sampling train of PMP contains a volatile particle remover (VPR) and a particle counter ( $>23\text{nm}$ ). The VPR provides two stage dilution connected by an evaporation tube (ET). The temperature for the primary dilution, ET and secondary dilution is 150, 300 and  $\sim 35^\circ\text{C}$ , respectively. Detailed information for the PMP can be seen in Herner et al., (2007a).

#### ***Thermodenuder***

Particle volatility was determined by two thermodenuders (Model ELA-230, Dekati Ltd) sampling in parallel, each heating the entering aerosol to 150 and 230  $^\circ\text{C}$ , respectively. The thermodenuder consists of a heating section, followed by an adsorption/cooling unit. As aerosol stream was drawn from the CVS and passed through the heating tube, part of its volatile/semi-volatile components was sheared off. These labile species adsorb onto a layer of activated charcoal placed on the walls of the thermodenuder, leaving the non-volatile PM fraction to be collected on Teflon filters (47mm, PTFE, Gelman) placed downstream of the thermodenuders. Multiple runs were integrated to achieve desired sample mass loadings on these filters to perform various chemical and toxicological analyses. Solid particle number concentrations and size distributions were monitored intermittently by a condensation particle counter (CPC 3022 A, TSI Inc., MN) and a differential mobility particle sizer (DMA, TSI 3085).

#### ***Electrical Aerosol Detector (EAD)***

Several instruments are currently in use to assess PM surface related properties. In this study, a diffusion charger, the TSI EAD 3070A, was used to characterize the transient as well as steady state

behavior of aerosols. The EAD consists of a unipolar diffusion charger and an electrometer. Particles are charged by diffusion and then drawn through the electrometer which records total current (I) carried by the aerosol stream. Studies have shown that the response of EAD is a function of  $D_p^{1.13-1.16}$  (Jung and Kittelson, 2005; Wilson et al., 2007; Woo et al., 2001). This response is particularly a useful PM metric because it corresponds to the actual particle surface area exposed to the environment and can be used to quantify, for example, the area available for adsorption of gaseous species or for interaction with the epithelial tissue in the lungs (Wilson et al., 2007). Because the EAD signal is closely related to aerosol diameter (Jung and Kittelson, 2005) the manufacturer has marketed this instrument to indicate total aerosol length in  $\text{mm}/\text{cm}^3$ .

### ***High Volume Sampler***

The high volume sampler is a specially designed filter holder [Hi-Q Environmental Products Co., CA; (Misra et al., 2002)] to collect particles on 20x25 cm filters at a relatively high flow rate (450 l/min). Integrated PM samples from the CVS were collected using this sampler on Teflon coated glass fiber filters (Pallflex, Fiberfilm T60A20 – 8x10 inch, Pall Corp., East Hills, NY).

This section reports the physical properties, including size distribution, volatility (in terms of number and mass), surface diameter and agglomeration of particles emitted from heavy-duty diesel vehicles (HDDV) retrofitted with advanced emission control devices. Substantial reduction in PM mass emissions (>90%) was accomplished for the HDDV operating with advanced emission control technologies. This reduction was not observed for particle number concentrations under cruise conditions, with the exceptions of the Hybrid-CCRT and EPF vehicles, which were efficient in controlling both - mass and number emissions. In general, significant nucleation mode particles (<50nm) were formed during cruise cycles in comparison with the UDDS cycles, which emit higher PM mass in the accumulation mode. The nucleation mode particles (<50nm) were mainly internally mixed, and evaporated considerably between 150 to 230°C. Compared to the baseline vehicle, particles from vehicles with controls (except of the Hybrid-CCRT) had a higher mass specific surface area.

## 4. Physical Characteristics of the Exhaust Particles (Semi-volatile and Non-volatile) from Retrofitted Vehicles

### 4.1 Introduction

Diesel exhaust particles (DEPs) are normally agglomerates of hundreds of volatile/semi-volatile species adsorbed onto its refractory carbonaceous core (Bayona et al., 1988). Various on-road as well as dynamometer experiments have shown that these particles are predominantly externally mixed (except particles <20nm). Particle volatility is strongly dependent on gas to particle phase partitioning and is extremely sensitive to dilution and temperature conditions (Abdul-Khalek et al., 1999; Biswas et al., 2007; Kuhn et al., 2005a; Liu et al., 2007a; Wei et al., 2001). Particle volatility plays key roles in shaping the particle size spectrum and eventually determining the level of human exposure to different aerosol components (volatile or non-volatile) originating from traffic emissions.

In addition to volatility, particle surface characteristics are important physical parameters in determining PM toxicity. Some researchers have argued that particle

surface area is a better metric to predict health endpoints than particle mass or number and should be included as an essential element while considering new regulatory tools (Maynard, 2006; Nygaard et al., 2004; Oberdorster et al., 2005). This is because the availability of reaction sites to cause cell damage is more likely to be directly proportional to surface area available to lungs (Maynard, 2006). Surface properties will also be relevant in future as the control technologies are reported to generate considerable number of small particles (Bagley et al., 1998; Geller et al., 2005) leading to a net increase in surface area per unit mass of PM.

In this section we focus on the PM physical properties, i.e. size distribution, volatility and surface characteristics for diesel vehicles retrofitted with state-of-the-art after-treatment devices. Comparisons among different vehicle types and driving cycles and also with respect to a baseline vehicle (without any control technology) are discussed.

### 4.2 Data Reduction

The physical properties of aerosols from the various vehicles and driving cycles were reduced to three variables, originally introduced by Ntziachristos and Samaras (2006), intended to serve as means to discriminate the effects of fuel, driving condition and vehicle control technology on particle emissions in dynamometer studies. These variables are:

**Reduced Variable 1: Volatility Ratio**

$$R = \frac{N_{Exhaust}}{N_{TD}} \quad (1)$$

Where

$N_{Exhaust}$  = Total dilution corrected DMS or EEPS number concentration at the CVS ( $D_p \geq 7\text{nm}$ )

$N_{TD}$  = Number concentration measured by CPC after the thermodenuders ( $D_p \geq 7\text{nm}$ ).

This is a measure of particle volatility in terms of number concentration.

**Reduced Variable 2: Surface Rated Diameter**

The EAD electrometer current is given by

$$I = N_p N_t e Q \quad (2) \quad (\text{Woo et al., 2001})$$

Where,  $I$  = Total current measured by EAD (fA)

$N_p$  = Charge (charge/particle) attachment =  $0.0181 D_s^{1.13}$  (Jung and Kittelson, 2005)

$D_s$  = Surface rated diameter, nm

$e$  = Elementary charge,  $1.6 \times 10^{-19}$  coulomb

$Q$  = Aerosol flow rate through EAD, 1.5 lpm (1.5 lpm aerosol, 1 lpm sheath flow)

$N_t$  = Total exhaust particle number, 10-1000nm from DMS; 10-523nm for EEPS.

*Note: Particle number in the size range of 523-1000nm is insignificant.*

Now substituting in equation 2:

$$D_s = \left( \frac{I}{0.0181 N_t e Q} \right)^{1/1.13} \quad (3)$$

**Reduced Variable 3: Mass Specific Surface Area**

$$A = \frac{S}{PM} \quad (4)$$

A = Mass specific surface area ( $\text{m}^2 \text{g}^{-1}$ )

S = Surface concentration, in  $\text{m}^2 \text{km}^{-1}$

For the calculation of A, the EAD current was converted to particle surface area using the conversion factor of  $65 \mu\text{m}^2/\text{pA}$  ( $R^2=0.9$ , Alveolar deposition) reported by Wilson et al., (2007).

PM = Total particle mass collected between 10nm-2.5 $\mu\text{m}$  nano MOUDI stages, in  $\text{mg km}^{-1}$ .

The nano-MOUDI substrate mass ( $\leq 2.5\mu\text{m}$ ) measurements were in excellent agreement with parallel CARB reference filter measurements ( $R^2=0.99$ ; Slope =0.95, Intercept:  $2.4 \text{ mg km}^{-1}$ ) and was used to calculate mass based parameters. Filter mass was used only for V-SCRT-Cruise (nano-MOUDI data not available)

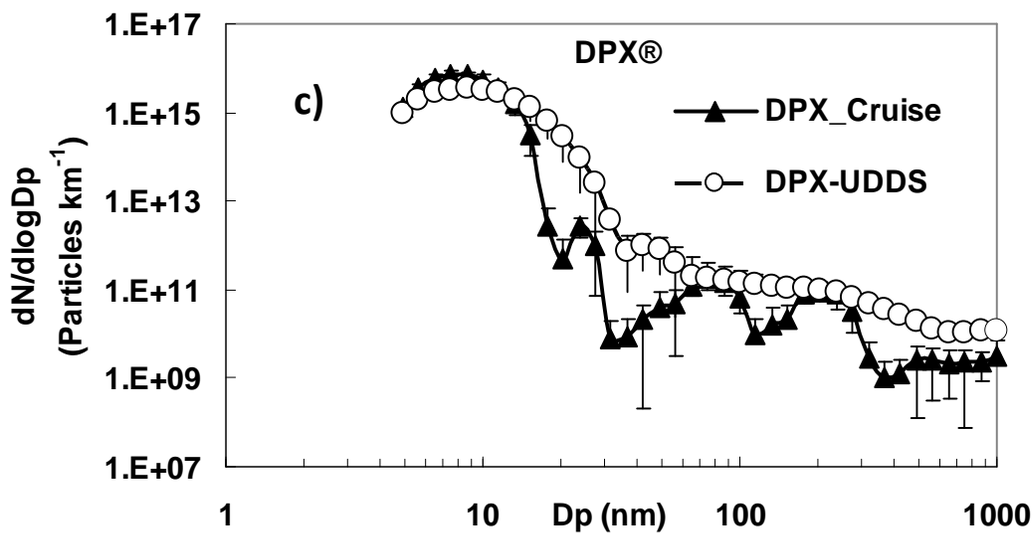
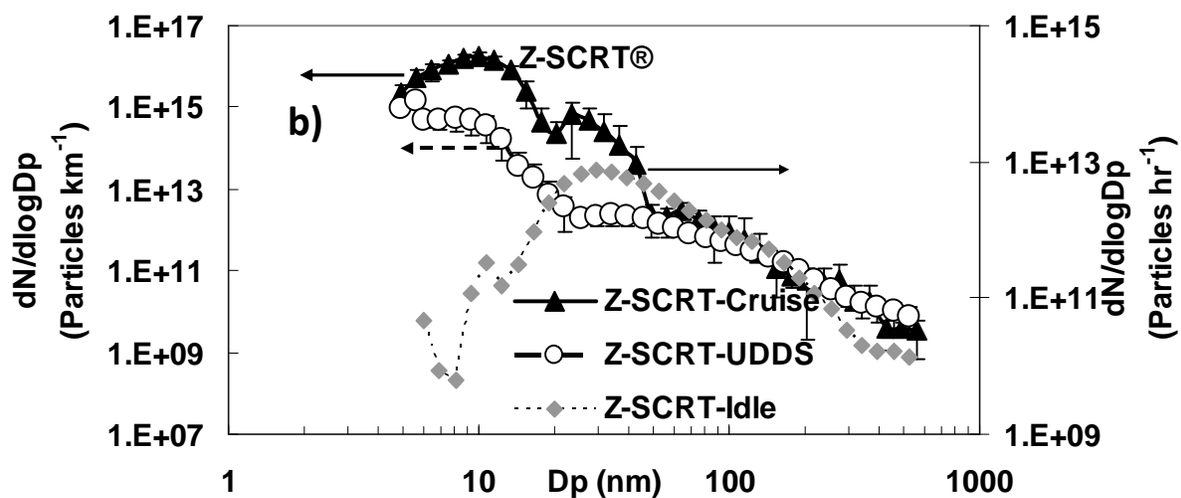
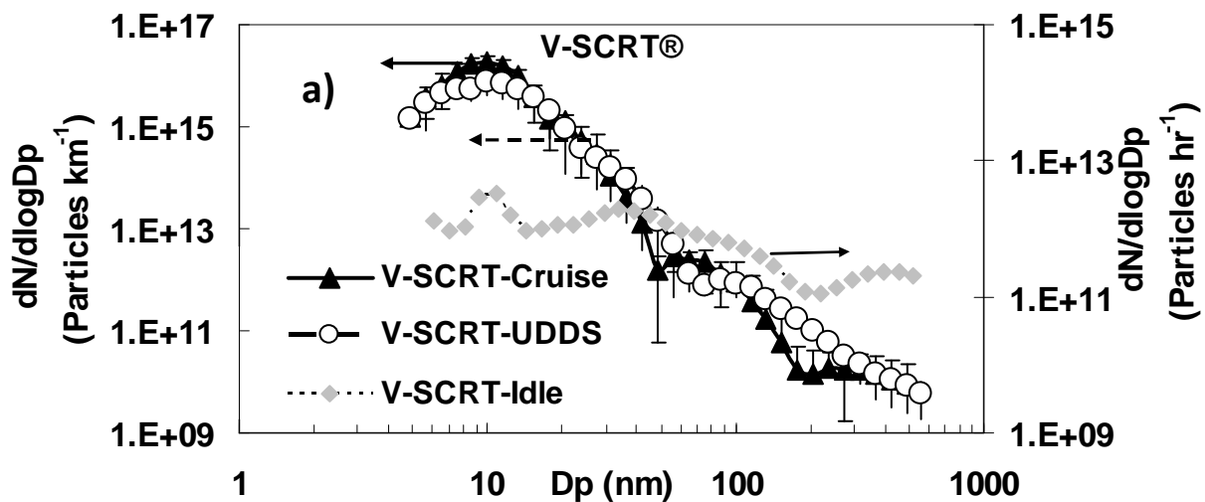
It is a measure of particle agglomeration, which increases with decreasing A .

## 4.3 Results

### Size Distribution

Figure 4.1 presents tunnel blank- subtracted mean size distributions for vehicles at different operating conditions. These distributions derived from DMS/EEPS measurements are grand averages of multiple runs for each driving cycle. The data are reported in terms of number per vehicle kilometer traveled for cruise and UDDS cycles, and number per hour for idle.

While performing preliminary quality assurance – quality control (QA/QC) of real time data for SCRTs (V-cruise, UDDS, Z-Cruise), we noticed that few EEPS sizes channels (10-20nm) were almost always saturated due to particle over-loadings. These size bins were subsequently replaced with corresponding secondary dilution-corrected DMS data (the DMS was placed downstream of the PMP secondary dilution with the ET off) to obtain more accurate size distributions. For the rest of test fleets [except CRT], we used data from the DMS, which was directly connected to the CVS. We have added a tunnel blank distribution in Figure 4.1f (Baseline vehicle).



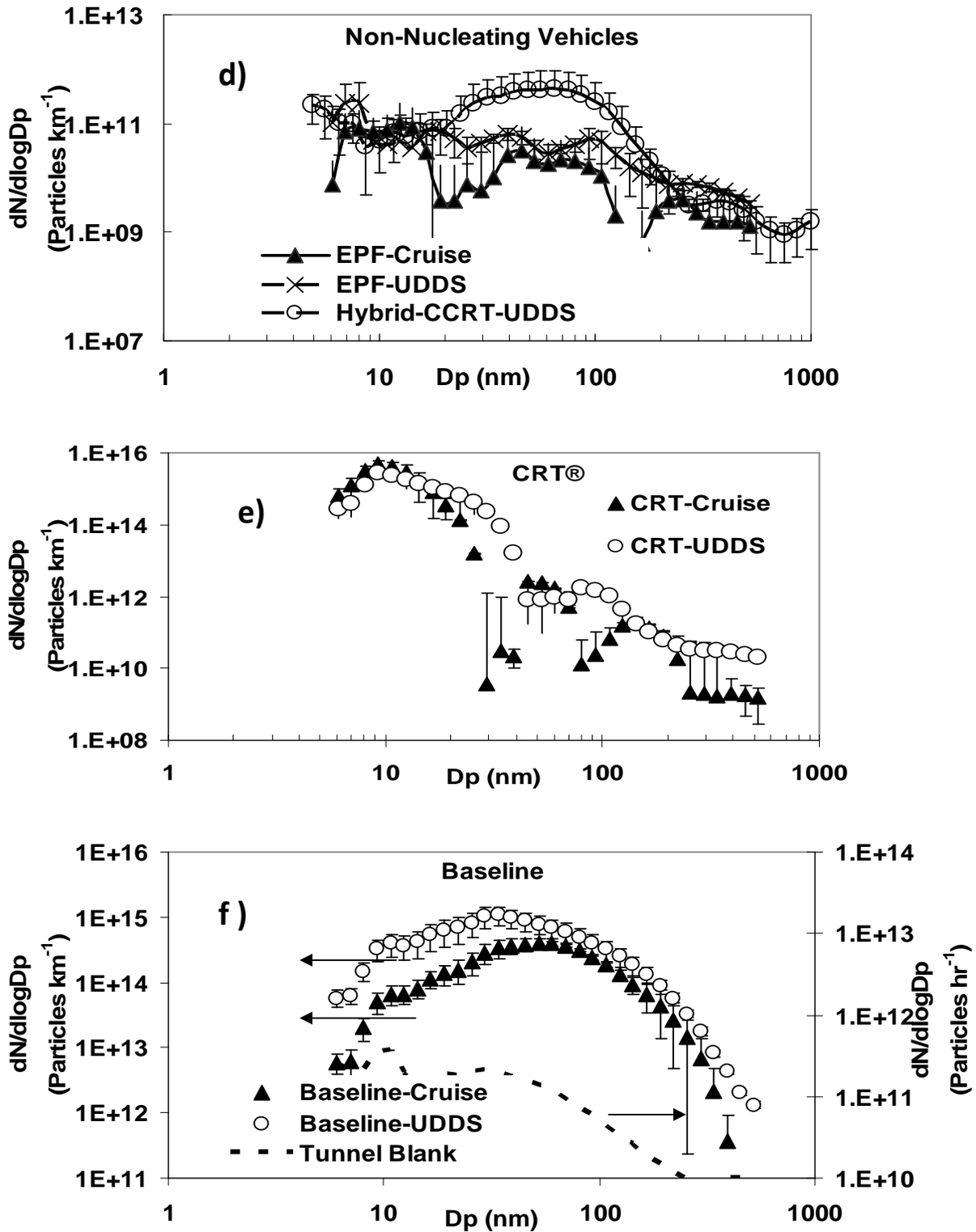


Figure 4-1 a, b, c, d, e, f Particle number size distributions

Size distribution patterns for both SCRTs are quite similar with sharp modes (Figure 4.1 a,b) at ~ 10nm. The distinguishable feature is the less prominent nucleation mode for UDDS runs especially for the Z-SCRT-UDDS cycle. This may be due to the fact that the Zeolite-SCRT requires much higher temperature to trigger and sustain nucleation than the vanadium based SCRT catalysts (Herner et al., 2007b). Moreover, in general the Zeolite catalysts have lot more catalytic surface area than vanadium catalysts and the Z-SCRT system used for this study is completely new. These provide higher storage sites for sulfate generated by the upstream DOC and DPF during transient and low temperature testing.

Although the main purpose of SCR technologies is to reduce NO<sub>x</sub> by ammonia, at elevated temperatures, their in-built catalysts may encourage the formation of sulfate, an important component acting potentially as seed aerosol for particle formation by condensation of semi-volatile organic vapors. For a brief sampling period, we bypassed the SCR portion from exhaust after-treatment system (SCRT) just to investigate the impact of the SCR catalysts. Although, this modification did not result in visible alteration of the shape (on log scale) of the distributions (CRT, Figure 4.1e), number concentration decreased by a factor of 2-3 from the V-SCRT and Z-SCRT cruise cycles—suggesting SCR catalysts' role on nucleation. Unlike cruise or UDDS cycles, idle runs are characterized with remarkably low particulate number emission rates, coupled with broad size distributions (Figure 4.1a, b). The second test engine, Engelherd-DPX (Figure 4.1c) displays a dominant nucleation mode, almost identical to the SCRTs/CRT. It has been hypothesized that the catalyst wash-coat on DPF (if saturated) may be enhancing the conversion of SO<sub>2</sub> to SO<sub>3</sub>/sulfate and partially stimulates the nucleation process (Hansen et al., 2001).

Contrary to the general notion that particulate filters augment nucleation, the hybrid vehicles [with a CCRT] and the school bus [with an EPF] were found to be highly efficient in suppressing if not eliminating this PM mode. The Hybrid-CCRT vehicle (Figure 4.1d) resulted in concentrations (CVS) in the range of ~10<sup>4</sup> particles cm<sup>-3</sup>, thus a 1,000-fold improvement over the previously tested vehicles (>10<sup>7</sup> particles.cm<sup>-3</sup>). We hypothesize that the initial capacity of its relatively new trap (with only 1000 miles on it) to store sulfur has significantly suppressed the formation of nuclei mode particles. Once all the storage sites are saturated, nucleated sulfate particles are expected. For this vehicle only a few odd large particles are left downstream (Figure 4.1d). The school bus (Figure 4.1d), however, was the cleanest amongst the entire test fleet with number emissions less than 1500 particles.cm<sup>-3</sup> measured in the CVS. It is important to note here that this vehicle is equipped only with an uncatalyzed filter (EPF) which is least likely to enhance the nucleation process. Thus, nucleation is not only control device specific but also a function of age and operating conditions e.g. temperature of the catalysts.

The baseline truck, on the other hand, represents the older genre of vehicles and was found to emit substantial amounts of larger particles (Figure 4.1f) with modes in the 60-100 nm range. Because of their large surface area, these accumulation mode particles act as adsorption sites and thus perfect sinks for organic vapors, leading to suppression of nucleation mode (Liu et al., 2007b).

### ***Size Segregated Mass Emission Factors***

Size fractionated mass emissions factors (in mg km<sup>-1</sup> or mg hr<sup>-1</sup>) are calculated based on the loadings on the MOUDI-nano-MOUDI impaction plates (Table 4.1). The mass loadings on the individual substrates are generally low for retrofitted vehicles and depending on size ranges, variation in the order of 20-40% (*Standard Deviation/Mean*; from few duplicate measurements) are observed.

Although a direct correspondence of mobility and aerodynamic diameters is not accurate without establishing some conversion factors, we can utilize the information from Table 4.1 to complement the

mobility size distributions (Figure 4.1) described before. Number and mass based size distributions are found to be in reasonable agreement with each other in terms of their trends.

The emission factors are remarkably low ( $\sim 1-12 \text{ mg km}^{-1}$ ) for the fleet operated with control technologies compared to the baseline vehicle ( $\sim 80-316 \text{ mg km}^{-1}$ ). While the majority (>95%) of PM mass is concentrated between 100nm-2.5 $\mu\text{m}$  for the baseline truck, nuclei modes are clearly visible for vehicles retrofitted with control devices. Significant reduction (>90%) of the mass is achieved for vehicles retrofitted with control devices.

Some general trends and inferences can be drawn from Figure 4.1 and Table 4.1. Consistent with a previous study (Vaaraslahti et al., 2004), the majority of the control technologies evaluated here have promoted bulk production of nano-size (nucleation) particles during steady state and high speed segment of transient running cycles. The cruise cycles on average generate higher nucleation and lesser accumulation mode particles than the UDDS cycles. The differences between these driving cycles are even more pronounced in mass distributions (Table 4.1): significant shifts towards larger sizes are apparent in UDDS runs due to increased emission of accumulation mode particles during the acceleration processes (Polidori et al., 2008). Aerosol formation mechanism, poorly understood till date, seems to be a function of vehicle type and driving conditions and retrofit design.

**Table 4-1 Size fractionated mass EFs (mg km<sup>-1</sup> for Cruise and UDDS, mg hr<sup>-1</sup> for Idle)**

| Size<br>(nm)    | V-SCRT |       | Z-SCRT |      |      | DPX    |      |       | Hybrid<br>CCRT | CRT    |      | EPF    |      |      | Baseline |      |        |
|-----------------|--------|-------|--------|------|------|--------|------|-------|----------------|--------|------|--------|------|------|----------|------|--------|
|                 | UDDS   | Idle  | Cruise | UDDS | Idle | Cruise | UDDS | Idle  | UDDS           | Cruise | UDDS | Cruise | UDDS | Idle | Cruise   | UDDS | Idle   |
| <b>10-18</b>    | 0.73   | 3.18  | 0.90   | 0.17 | 1.06 | 0.65   | 0.56 | 4.91  | 0.26           | 1.35   | 1.25 | 0.04   | 0.05 | 5.92 | 1.02     | 1.39 | 19.363 |
| <b>18-32</b>    | 1.44   | 2.48  | 1.48   | 0.34 | 3.54 | 0.29   | 0.46 | 1.89  | 0.12           | 0.86   | 1.52 | 0.22   | 0.08 | 3.48 | 1.96     | 2.54 | 60.241 |
| <b>32-56</b>    | 2.46   | 1.77  | 1.60   | 0.32 | 1.42 | 0.30   | 1.25 | 12.45 | 0.22           | 0.98   | 3.19 | 0.12   | 0.16 | 5.92 | 5.96     | 10.7 | 565.83 |
| <b>56-100</b>   | 1.36   | 4.25  | 1.15   | 0.17 | 0.35 | 0.11   | 0.60 | 3.40  | 0.33           | 0.49   | 0.55 | 0.14   | 0.10 | 4.53 | 9.98     | 19.6 | 600.26 |
| <b>100-180</b>  | 0.86   | 12.03 | 0.53   | 0.62 | 3.54 | 0.27   | 0.41 | 7.92  | 0.17           | 0.57   | 1.52 | 0.07   | 0.05 | 3.13 | 29.4     | 86.8 | 2327.9 |
| <b>180-2500</b> | 1.07   | 7.78  | 0.97   | 0.55 | 0.00 | 0.21   | 0.20 | 3.02  | 0.57           | 1.15   | 2.63 | 0.15   | 0.29 | 8.71 | 31.1     | 192  | 6049.9 |
| <b>&gt;2500</b> | 0.83   | 6.37  | 0.50   | 0.66 | 3.18 | 0.20   | 0.35 | 5.66  | 0.35           | 0.37   | 0.83 | 0.12   | 0.17 | 7.66 | 1.03     | 3.41 | 346.39 |

Figure 4.2 is an informative graph showing a plot of number EFs vs. Mass EFs (nano-MOUDI, except V-SCRT-Cruise) for each vehicle and driving cycle. Each data point corresponds to a cruise or an UDDS cycle for a given vehicle. The graph shows that for several of the after-treatment devices tested, particle number emissions increase with reduced mass emissions. There are some outliers or exceptions observed for Z-SCRT at UDDS cycle, Hybrid-CCRT, EPF and Idle (not shown) for which both number and mass EFs are relatively low, possibly due to the lack of catalytic formation of sulfate: EPF has no catalyst; the Hybrid CCRT was not yet aged, and therefore the sulfur storage sites were not fully saturated, thereby preventing or suppressing the formation of sulfur. Moreover, the idling cycle does not lead to high enough catalyst temperatures that would convert sulfur to sulfate. Thus, most of the new after-treatment devices appear to be highly efficient in reducing mass emissions but some are not effective in controlling number concentrations due to the formation of nuclei mode particles. These findings are potentially very important if number based standards are considered in the future.

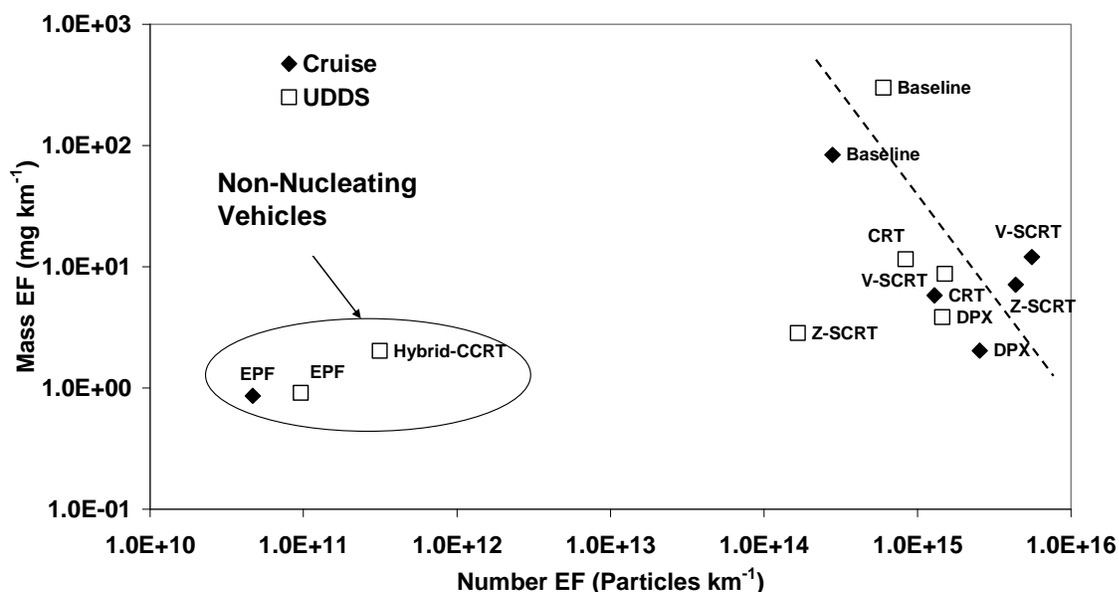


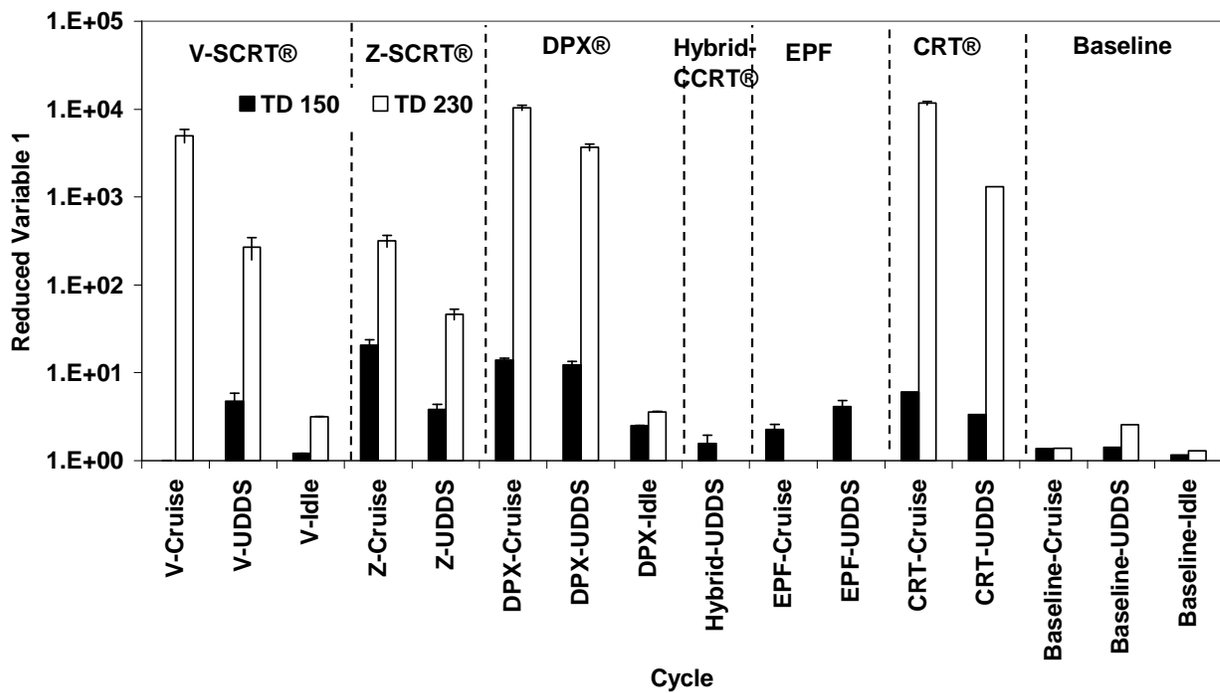
Figure 4-2 Number (DMS) and mass emission factors (nano-MOUDI).

### Particle Volatility

The extent of particle volatility in terms of total count is illustrated quantitatively by the reduced variable 1 (R) plotted in Figure 4.3 for various vehicles and cycles. R was calculated at 150 and 230°C except in few occasions (Hybrid-CCRT, EPF) when both the thermodenuders were operated at 150°C to maximize our capability to collect mass of non-volatile particles. Theoretically, this ratio (R) should be greater or equal to unity and its increase corresponds to higher particle volatility. For the majority of the cases, R at 230°C is at least an order of magnitude higher than their counterparts at 150°C, indicating complete disappearance of large fraction of particles within this temperature window.

Particle volatility was also observed to be somewhat sensitive to driving conditions and type of control technology used. For instance, cruise mode particles are extremely volatile with maximum R

values (~ 4000-10000) at 230°C for V-SCRT, DPX and CRT. The fact that less than ~0.1% of particles persisted at 230°C for most of the fleet with control technologies can logically be explained by the presence of unstable (volatile) fresh nano particles. Although the same Kenworth truck is used to evaluate various after-treatment devices (V-SCRT, Z-SCRT, CRT), the Zeolite catalysts, especially during UDDS runs, seem to emit particles with a slightly higher heat resilience, which is consistent with the fact that that nucleation mode particle were in lower concentrations for that vehicle and cycle (Figure 4.1b). As expected, the baseline vehicle had the highest fraction of non-volatile particles in all cycles. Lower volatility is also observed during all idle runs, EPF and Hybrid-CCRT vehicles. While the baseline truck emits large amounts of refractory elemental carbon, lower number loss for other vehicles/cycles is attributed to their insignificant particle emission rate and/or absence of the nucleation mode particle formation.



**Figure 4-3 Reduced Variable 1 (R): Ratio of total particle to solid or thermo-denuded particle count.**

The significant particle number loss between 150 to 230°C, especially for cruise cycles, may be better elucidated by the evaporation profiles shown in Figure 4.4. The majority of particles in the 7-20nm size range for DPX and CRT have disappeared as the aerosol stream is heated from 150°C to 230°C. This is in total contrast to the baseline vehicle, where no noticeable shift in size spectrum is observed. Matter et al., (1999) reported very similar thermal desorption trends between 172 and 204°C for particles sampled downstream of a DPF.

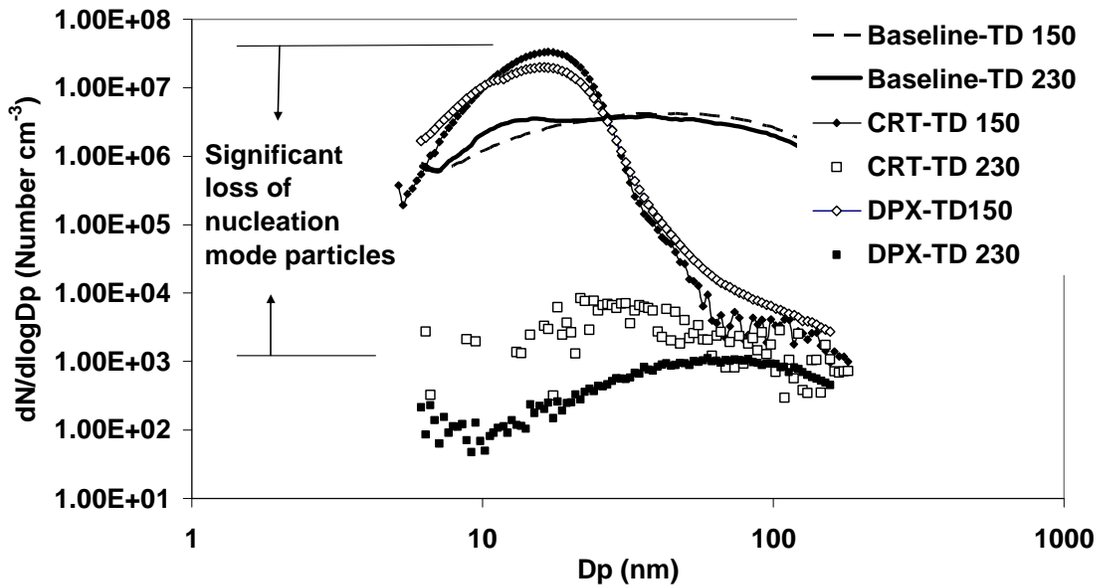


Figure 4-4 Size distribution of thermo denuded aerosols for cruise mode.

Note: Available only for DPX, CRT and Baseline

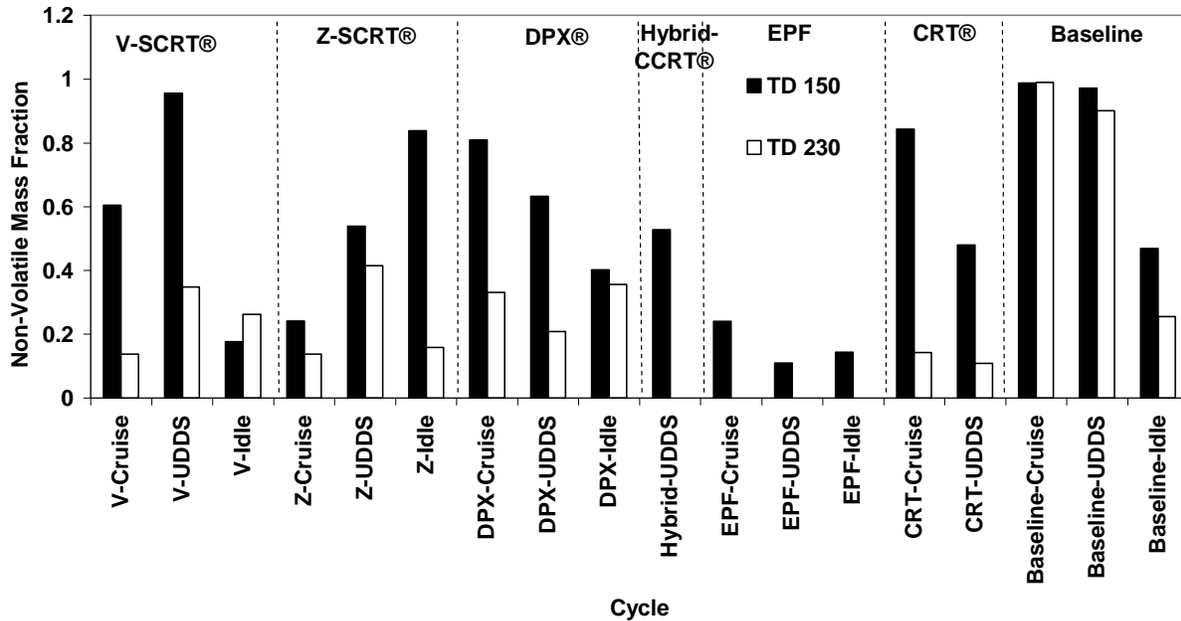


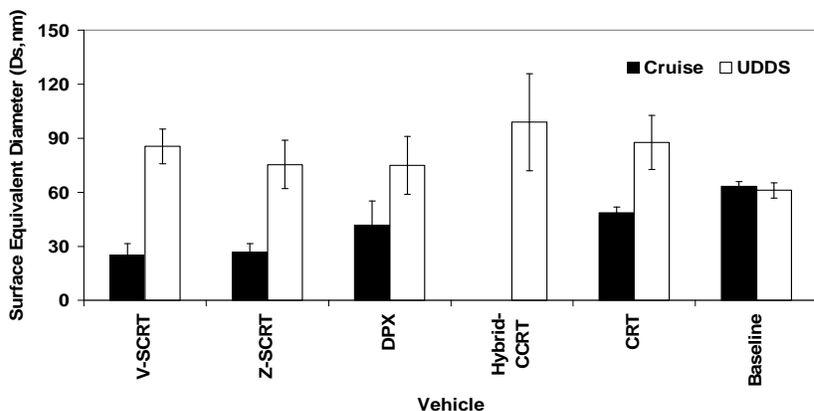
Figure 4-5 Mass fractions remaining at different thermodenuder temperature settings.

Figure 4.5 presents the amount of mass fraction retained (non-volatiles) at 150 and 230°C. We obtained these ratios from gravimetric filter measurement of denuded and undenuded samples. Particle number is not a good surrogate of its mass. Thus, except in the case of baseline vehicle, number

loss (Figure 4.3, 4.4) has not linearly translated into equivalent particle mass reduction. While more than 99% of the particles completely disappear at 230°C for SCRTs, CRT and DPX at cruise and UDDS runs, an appreciable fraction of mass (20-40%) remains intact. Quite a reverse trend is observed for idling, EPF and Hybrid-CCRT (low number emission scenarios with insignificant nucleation mode)—relatively low number volatility is accompanied with significant mass loss. A study (Biswas et al., 2007) conducted in close proximity of a freeway with the highest percentage of diesel fleet (~18%) in the US demonstrated that a substantial amount of particle mass could be lost from larger ultrafine particles (>40nm) due to heating without losing a significant amount of their total counts. The mode of the size distribution shifted to lower size ranges, as a result of the depletion of the aerosol mass, but overall these particles simply decreased in size but did not disappear during heating, unlike the much smaller nucleation mode PM.

**Particle Surface Properties: Surface Rated Diameter (Reduced Variable 2)**

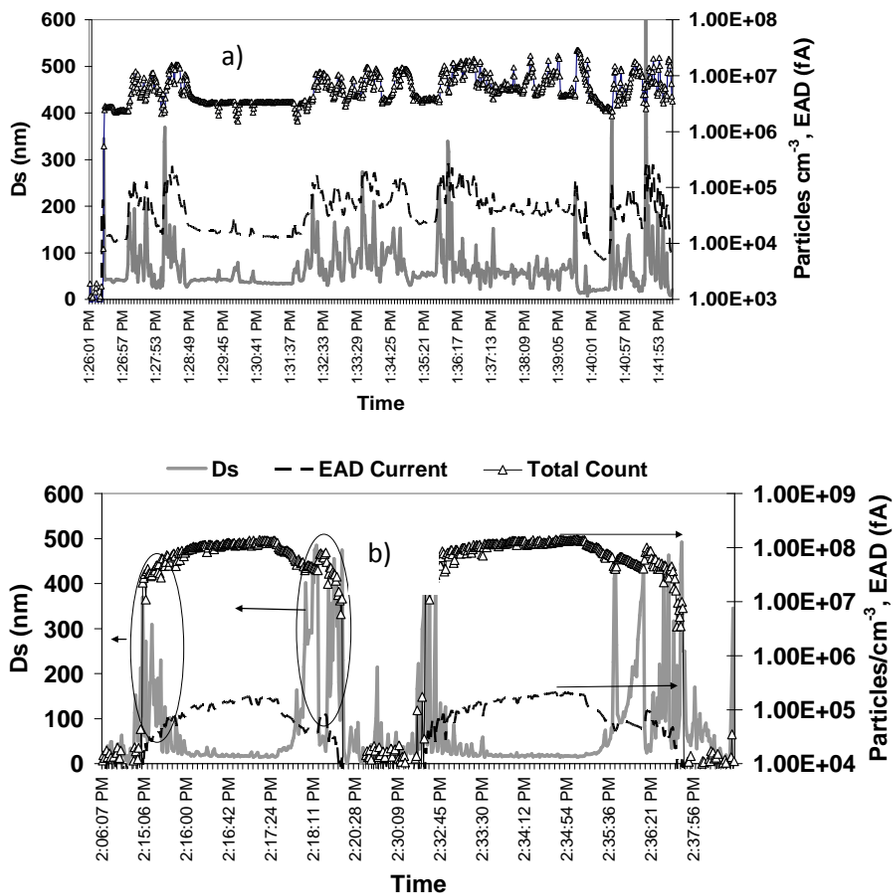
The surface rated diameters (Ds) calculated from EAD surface concentrations are illustrated in Figure 4.6. Ideally, Ds should be very close, if not higher than, the arithmetic mean particle diameter. Average surface rated diameters are calculated for all the vehicles except for school bus (EPF) where EAD signals are below the instruments detection limit. It is evident from the plot that cruise mode Ds, irrespective of types of control technologies, are smaller (~25-40nm) compared to the transient driving cycles (~75-100nm). This is consistent with the increased formation of elevated level of nuclei mode particles (< 50nm) during the steady state and larger amount of accumulation mode particles in UDDS cycles. Considering negligible variations in the accumulation modes (Harris and Maricq, 2001), also shown in our Figure 4.1 and Table 4.1, we can thus use Ds as a good indicator for the presence or absence of freshly nucleated particles.



**Figure 4-6 Reduced Variable 2: Surface rated diameter.**

During a typical transient run (i.e. UDDS cycles), an engine experiences acceleration, deceleration, steady state and idling phases, which result in frequent changes in exhaust particle characteristics in terms of their size distribution, total count and surface concentration. Acceleration and high speed cruise modes produce significant number of small particles, followed by their absence during deceleration and idling. Thus, rapid variation in Ds coupled with higher accumulation modes (Figure 4.1, Table 4.1) eventually translates into higher mean surface diameter for UDDS runs. This phenomenon can

be elucidated succinctly by time series plots (Figure 4.7 a,b) of particle number, EAD response (fA) and corresponding mean surface diameter (Ds) for a selected DPX UDDS run (Figure 4.7b). EAD signals and total counts track each other well; however, the Ds plot follows a reverse trend. Surface diameter goes up and down (50-500nm) at the initiation of acceleration or deceleration when the total count is still low, or drops down drastically. The lowest surface diameter (~20-30nm) was recorded during the later stages of acceleration, or at high load conditions (steady state) as small particles predominate the size distribution.

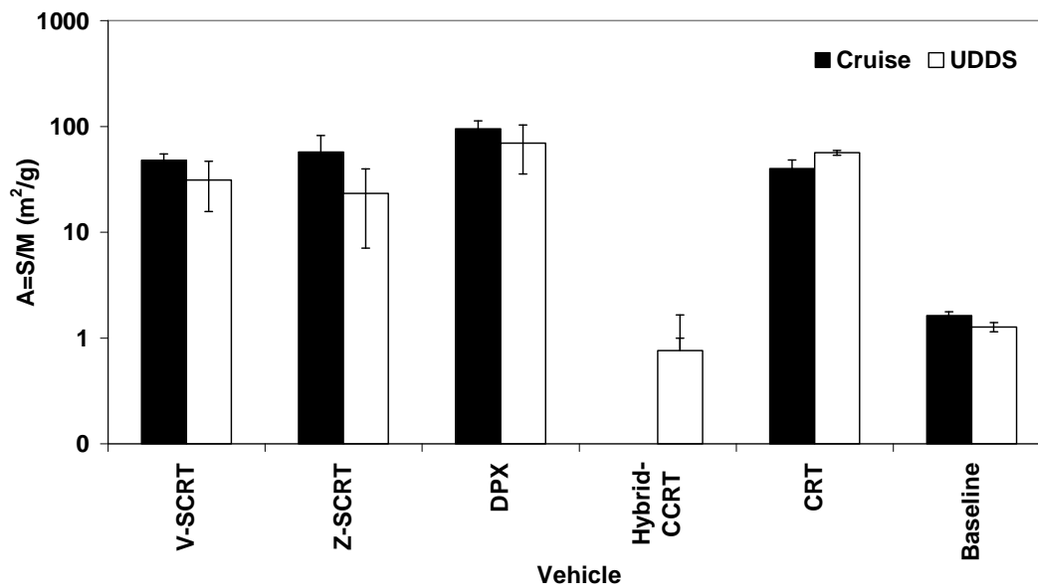


**Figure 4-7 a, b: Time series of surface diameter (Ds), surface concentration (EAD signal and total particle concentration (DMS count) for: a) Baseline and: b) DPX- UDDS run.**

Unlike the other vehicles (where similar behavior observed as that of Figure 4.7b), the baseline truck maintains a high particle number concentration ( $>10^6$  particle  $\text{cm}^{-3}$ ) over the entire span of the UDDS runs, which greatly dampens the fluctuation in Ds (Figure 4.7a). This explains its resultant lower mean surface diameter (~60nm) (Figure 4.6) despite the significant contribution of accumulation mode particles. Also, the lower difference in Ds for the cruise and UDDS can be attributed to the relatively higher number of nucleation mode particles in UDDS than cruise unlike other vehicles.

**Particle Surface Properties: Mass Specific Surface Concentration (Reduced Variable 3)**

Figure 4.8 shows mass specific surface concentrations for the various tested vehicles and driving cycles. This PM property is inversely proportional to the mean particle size of DEPs, thereby providing an indirect measure of the extent of particle agglomeration. Therefore, the decrease of 'A' implies an increased contribution of coagulated or accumulation mode particles. The accumulation mode particles scavenge nucleation precursor species (Liu et al., 2007b). Thus, particle agglomeration is an important parameter to indicate suppression of nuclei mode particles.



**Figure 4-8 Reduced Variable 3: Mass specific surface concentration.**

Cruise cycles on average have slightly higher values (5-10%) of 'A' compared to their respective UDDS cycles of a given vehicle, indicating their less agglomerated particle structures. Although the shape and size of accumulation modes are quite similar, if not identical, for both cycles (Figure 4.1), steady state operations are associated with higher nucleation mode. As EAD signal is weighed towards smaller particles (Jung and Kittelson, 2005), cruise cycles are likely to trigger more EAD response than transient cycles. On the other hand, accumulation modes prevail over nucleation modes (Figure 4.1) for Hybrid-CCRT at UDDS runs and for the baseline vehicle, resulting in lower mass specific surface concentrations ( $\sim 1 \text{ m}^2\text{g}^{-1}$ ). Higher fractal agglomerates emitted by the Baseline vehicle may result in a lower value of A.

## 4.4 Conclusions

This section presented some of the first detailed particle characterization for advanced NOx and PM retrofits for heavy duty diesel vehicles. Remarkable reductions in PM mass emissions (>90%) were found for the test fleet compared to the baseline vehicle. However, enhanced nucleation mode particles were observed for some of the vehicles especially during cruise cycles. Comparing to cruise cycles, the UDDS cycles emit higher particle mass in the accumulation mode. Idle runs are characterized with remarkably low particle number emission rates, coupled with fairly broad size distributions. The Hybrid-CCRT and EPF vehicles were efficient in controlling both mass and number emissions. The majority of particles by number evaporated by heating the aerosol to 150 - 230 °C, suggesting the nucleation mode particles are predominantly internally mixed and consist of semi-volatile compounds. Particles from the test fleet

(except Hybrid-CCRT) have shown about 100-fold higher active surface area per unit mass than the baseline vehicle.

This section focuses on PM chemical characteristics [Total carbon (TC), Elemental carbon (EC), Organic Carbon (OC), speciated organic compounds, inorganic ions, and water-soluble organic carbon (WSOC) from the tested vehicles. Significant reductions in the emission of major chemical constituents (TC, OC, EC, and organic compounds) were achieved by the introduction of retrofits. V-SCRT and Z-SCRT effectively reduced PAHs, hopanes and steranes, n-alkanes and acids by more than 99%, and often to levels below detection limits for both cruise and UDDS cycles. The CRT technology also showed similar reductions with SCRT for medium and high molecular weight PAHs, acids, but with slightly lower removal efficiencies for other organic compounds. Sulfate dominated the PM composition in vehicle configurations (V-SCRT-UDDS, Z-SCRT-Cruise, CRT, DPX) with considerable nucleation mode and TC was dominant for the configurations with less (ZSCRT-UDDS) or insignificant (CCRT, Horizon) nucleation.

The transient operation increases EC emissions, consistent with its higher accumulation PM mode content. In general, solubility of organic carbon is higher (average ~ 5 times) for retrofitted vehicles than the baseline vehicle. The retrofitted vehicles with catalyzed filters (DPX, CCRT) had decreased OC solubility (WSOC/OC: 8-25%) unlike those with uncatalyzed filters (SCRTs, Horizon; WSOC/OC~ 60-100%).

The particle bound organics-to-OC ratios ( $\mu\text{g/g}$ ) was lower for retrofitted vehicles compared to the baseline truck. However, the reduction in the ratio was considerable for PAHs, while it was insignificant for hopanes and steranes, implying that fuel and lubricating oil have substantially different contributions to the total OC emitted by vehicles operating with after-treatment control devices.

## 5.

### hemical Characteristics of Exhaust Particles from Retrofitted Vehicles

#### 5.1 Introduction

This section focuses on the chemical components (EC, OC, WSOC, inorganic ions, and speciated organic compounds) of PM from heavy-duty diesel vehicles (HDDV) tested during the experimental phase. Comparisons are made with previous studies to illustrate the effects of control technologies on the chemical composition of exhaust PM samples. The effect of driving cycles is also discussed by comparing steady-state cruise and transient driving cycles.

#### 5.2 Sample Analysis

Aluminum substrates from MOUDI-Nano MOUDI stages were pre and post-weighed during the sampling to determine the mass loadings. The elemental and organic carbon (EC, OC) content of deposited PM was analyzed by thermal-optical method (Kleeman et al., 1999). It should be noted here that for Al substrates, EC/OC split is not precisely defined; only relative fractions of EC and OC can be estimated. The ion concentrations were derived by ion chromatography (Zhang et al., 2008) technique. The water-soluble organic carbon (WSOC) content extracted from Teflon coated GF filters were analyzed by Shimadzu TOC-5000A liquid analyzer (Decesari et al., 2000).

The speciated organic compounds, including polycyclic aromatic hydrocarbons (PAH), hopanes and steranes, n-alkanes and organic acids, were quantified on the HiVol PM samples by gas chromatography-mass spectrometry (GC/MS). Methods for the quantification of individual compounds were based on already established solvent extraction procedures reported earlier (Schauer et al., 1999). Samples were extracted in dichloromethane and methanol and were combined and reduced in volume to approximately 1 mL by rotary evaporation, followed by

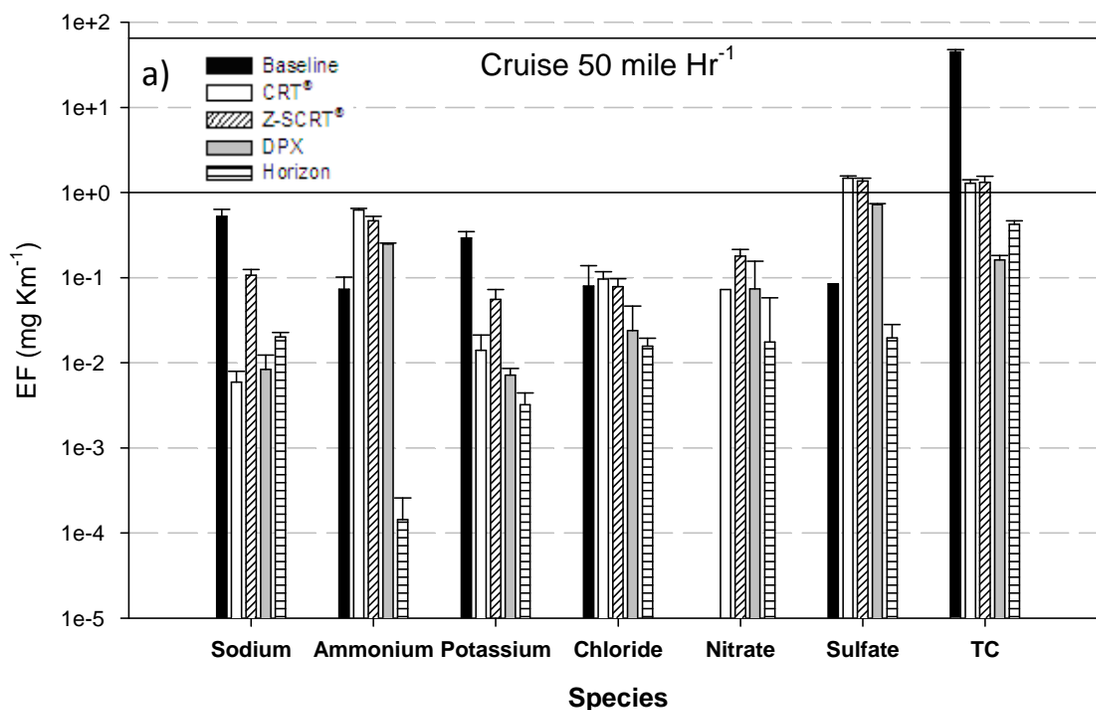
pure nitrogen evaporation. The underivatized samples were analyzed by auto injection into a GC/MSD system (GC model 5890, MSD model 5973, Agilent). A 30 m × 0.25 mm DB-5MS capillary column (Agilent) was used with a split-less injection. Along with the samples, a set of authentic quantification standard solutions were also injected and used to determine response factors for the compounds of interest. All the results were blank-corrected prior to data analysis and converted to mass emitted per distance traveled.

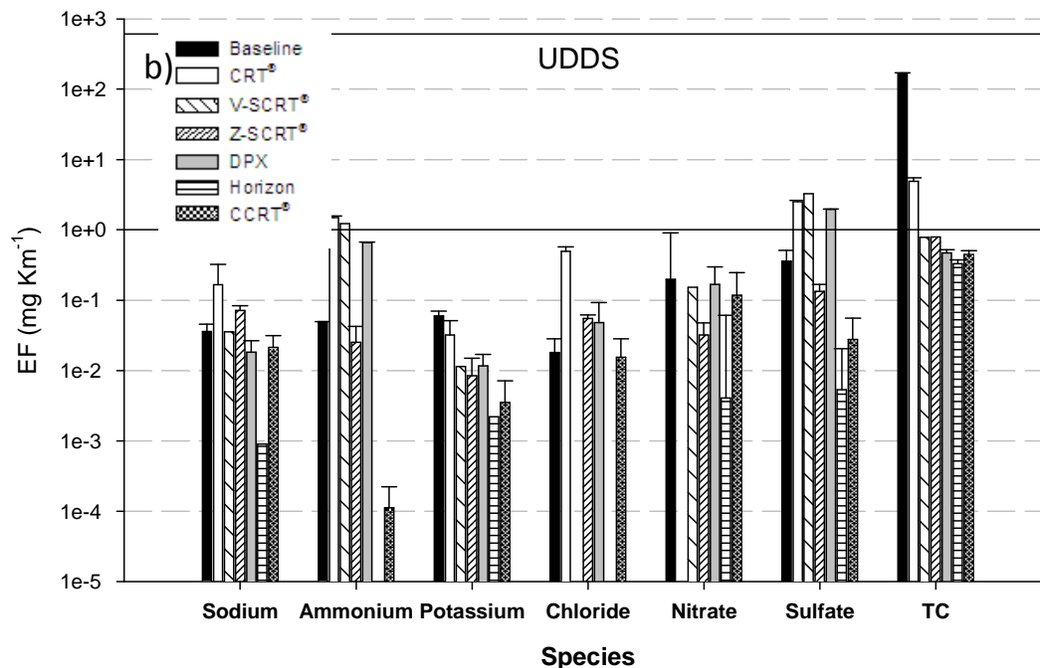
### 5.3 Results

#### **Emission Factors of Inorganic Ions and Total Carbon (TC)**

Figure 5.1 shows the EFs (expressed in mg of PM species per kilometer driven) of major chemical species integrated over the MOUDI-NanoMOUDI stages from 0.01- 2.5 μm. The low emission vehicles pose a challenge for PM chemical analysis due to very low mass loadings on individual substrates. The analytical uncertainties are shown by the error bars.

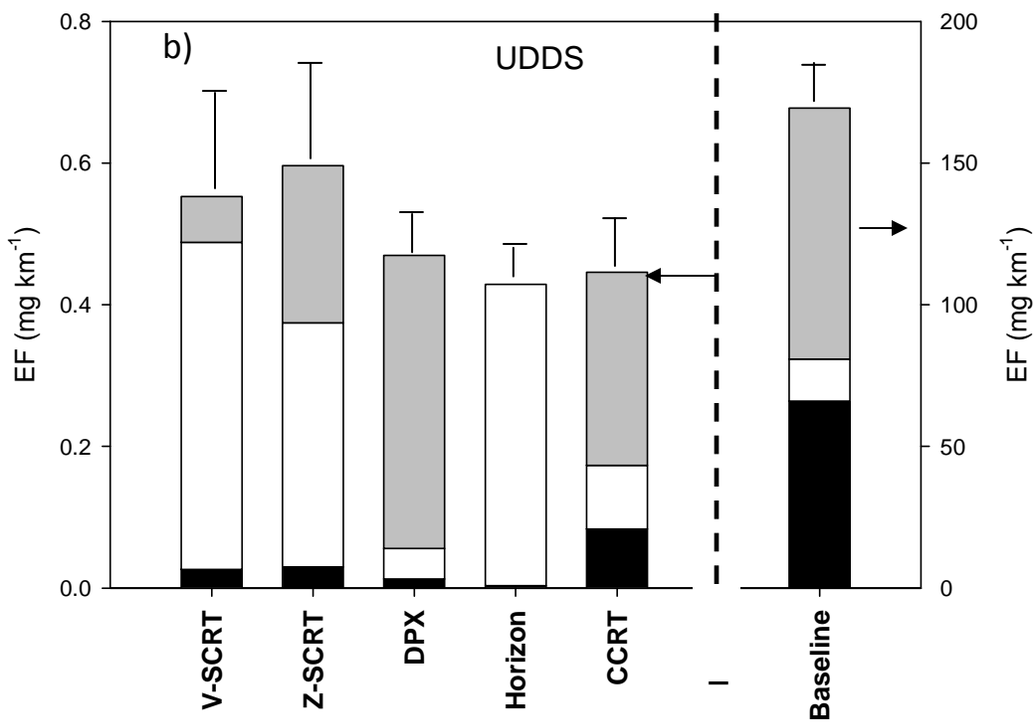
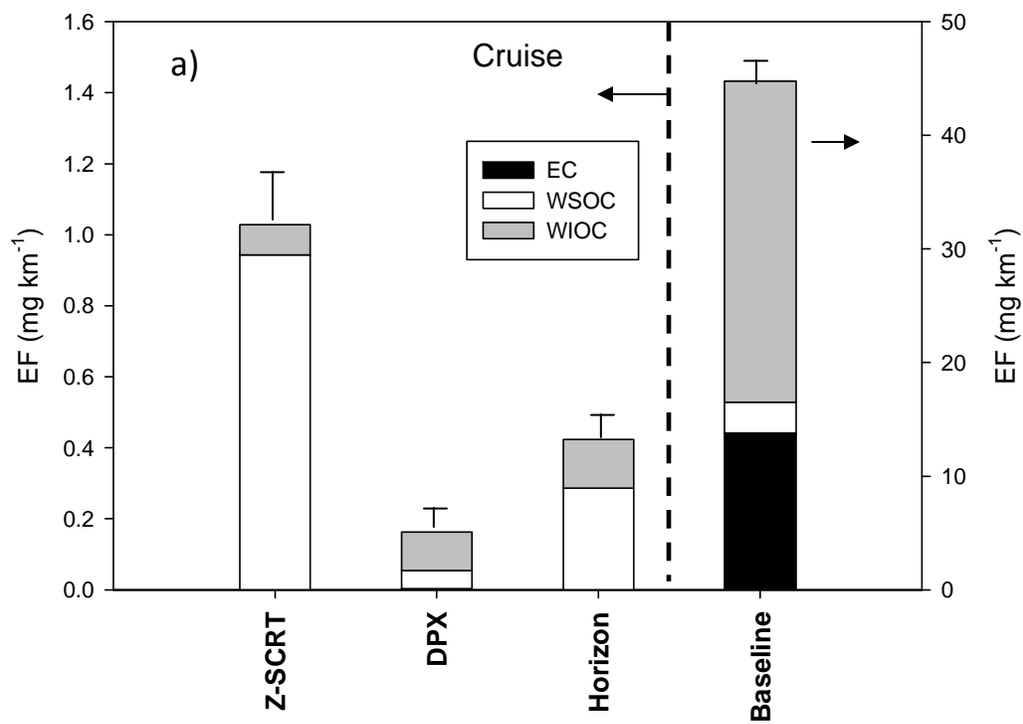
At both cruise and UDDS cycles, significant reductions in the emission of TC were observed for the retrofitted vehicles compared to the baseline vehicle. However, elevated emission rates of sulfate and ammonium were observed for vehicles and/or driving cycles that have a substantial fraction of nucleation mode particles, i.e. CRT, V-SCRT (measurements only available for UDDS cycle), Z-SCRT-cruise and DPX, compared to the baseline vehicle. The school bus retrofitted with an electric particulate filter (Horizon) was consistently the cleanest vehicle with the lowest emissions of inorganic ions and TC. This is also in agreement with the emission rates of PM based on gravimetric mass measurement (Section 3).

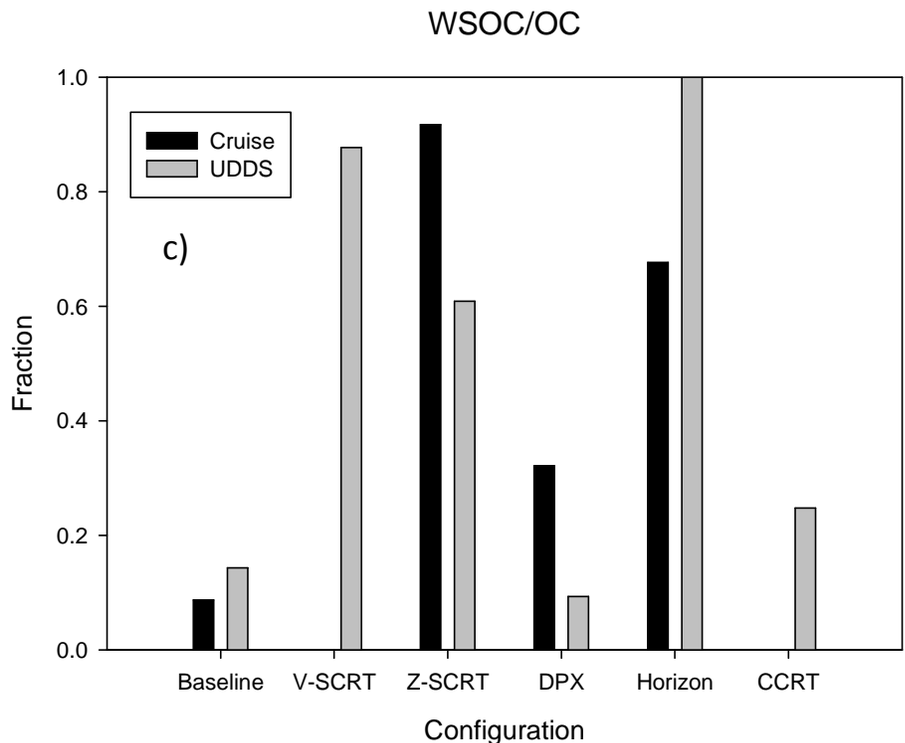




**Figure 5-1 Emission factors of integrated chemical species measured by MOUDI-NanoMOUDI. a). Cruise 50 mph b). Transient UDDS.**

The emission factors of TC were further split into EC, water-soluble and water-insoluble OC (WSOC and WIOC, respectively) and shown in Figure 5.2a, b. The baseline vehicle (shown on secondary axis) not only emits considerable amount of EC (Cruise: 13.8 mg km<sup>-1</sup>, UDDS: 65.9 mg km<sup>-1</sup>) but also elevated levels of WIOC (Cruise: 28.2 mg km<sup>-1</sup>; UDDS: 88 mg km<sup>-1</sup>) and WSOC (Cruise: 2.7 mg km<sup>-1</sup>; UDDS: 14 mg km<sup>-1</sup>). The low EFs of WIOCs (~0.1-0.4 mg km<sup>-1</sup>) and WSOC (0.05-0.95 mg km<sup>-1</sup>) for the retrofitted vehicles suggest that these control devices are also effective (65-99%) in reducing the major fraction of organic carbon in addition to EC. However, It is important to note here that only the baseline, V-SCRT, Z-SCRT and CRT configurations are directly comparable (same vehicle) and caution should be exercised while comparing with other vehicle configurations (DPX, Horizon, CCRT).





**Figure 5-2 a, b EC, WSOC and WIOC emission rates at a) Cruise 50mph and b) Transient UDDS c) Proportion of WSOC to OC.**

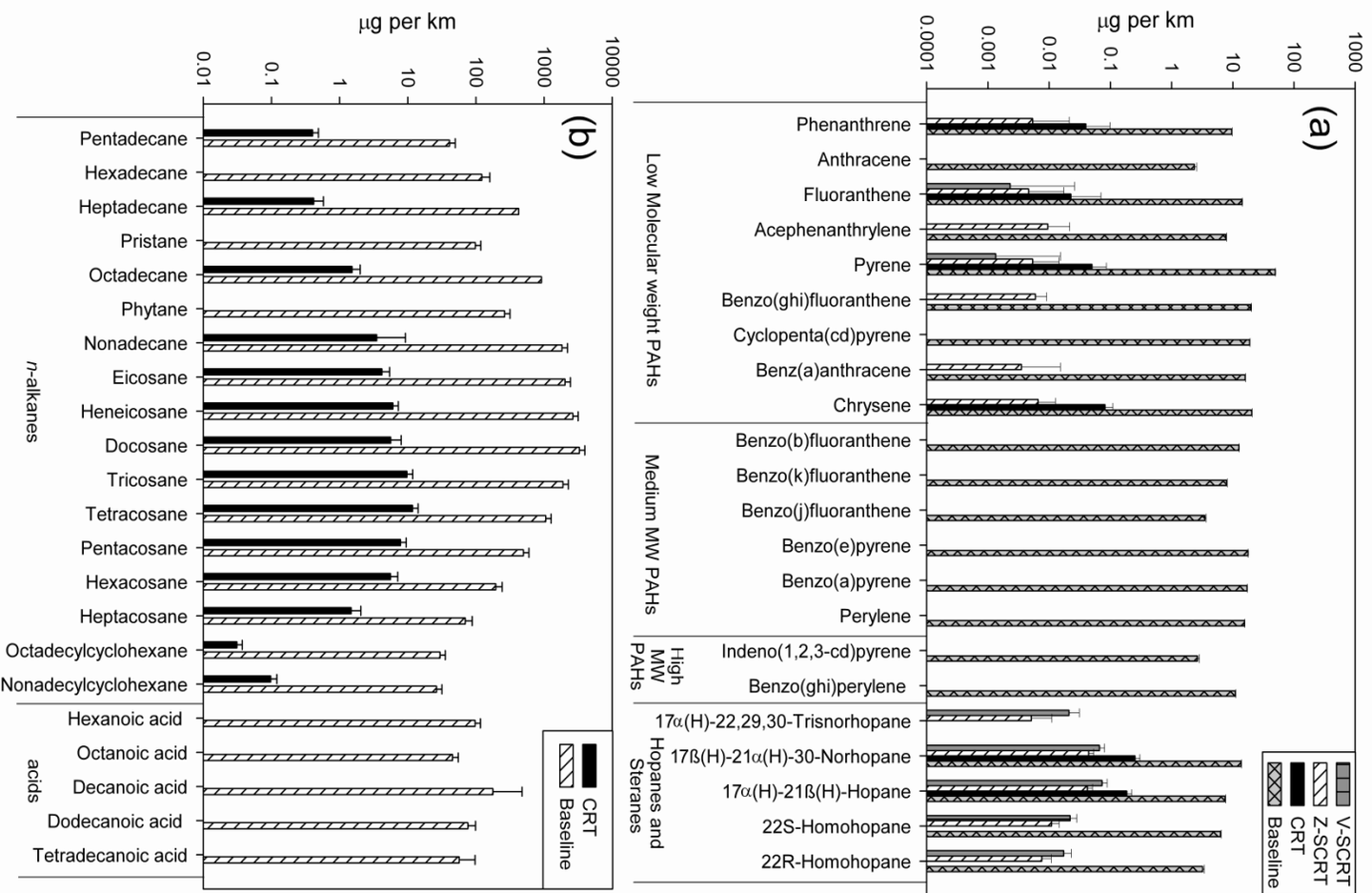
**Note: V-SCRT cruise and CCRT-Cruise cycles are not available**

In addition to the overall reduction of OC, the proportion of WSOC to OC, illustrated in Figure 5.2c varies with vehicle configurations and running cycles. The baseline vehicle shows the lowest OC solubility ( $WSOC/OC = 0.08-0.13$ ) during both cruise and UDDS cycles, suggesting freshly emitted primary OC. By comparison, OC emitted from Horizon and SCRTs consists primarily of WSOC. DPX and CCRT WSOC contents are intermediate between the baseline and other retrofitted vehicles. An earlier study by Hameri et al., (2001) indicated higher water solubility of OC in the nucleation/UF mode than the accumulation mode. This may explain, to a certain extent, the higher WSOC/OC ratio of SCRTs and Horizon, where significant fraction of OC is observed in the nucleation mode. In contrast, OC dominates in accumulation mode for the baseline vehicle. In addition to size distribution of OC, control technologies seem to have major influences on the percentage of WSOC emitted by vehicles. The DPF of Horizon and SCRTs (higher WSOC/OC) are uncatalyzed, while DPX and CCRT (lower WSOC/OC) have catalyzed filters. Although SCRTs and CCRT both have DOCs upstream of the DPF, the results suggest that DOCs, in comparison to DPF types (catalyzed or uncatalyzed) have a minimal effect on the WSOC/OC ratio. This conclusion, however, does not imply that the DOCs have not contributed in reducing OC from the total vehicle emissions.

### ***Emission Factors of Speciated Organic Compounds***

Figures 5.3a and 5.3b present the PM organic compound mass emission factors (EF) for the UDDS driving cycle of the baseline vehicle as well as the same vehicle equipped with the different after-treatment technologies noted earlier (CRT, V-SCRT and Z-SCRT respectively). The emission factors of PM bound PAHs, hopanes and steranes in UDDS range from 2.4 µg/km to 49.4 µg/km for the baseline vehicle, with an average of 14.5 µg/km and 7.6 µg/km for PAHs and hopanes and steranes, respectively. The emission factor of *n*-alkanes and organic acids range from 26.3 µg/km to 3287.2 µg/km with an average of 745.0 µg/km and 90.5 µg/km for *n*-alkanes and organic acids respectively. As shown in Figure 5.3a, low molecular weight PAHs with three to four rings are the largest contributor to the total PAH emissions. These light PAHs account for 47.8% of the measured total PAHs mass while the contribution of medium and high molecular weight PAHs are 33.6% and 18.6% respectively. This is consistent with other studies, which showed that three- and four-ring PAHs are present in HDDV exhaust in very high levels (Liu et al., 2008; Marr et al., 1999; Riddle et al., 2007). These PAHs are semi-volatile and can be found in both gas phase and particle form.

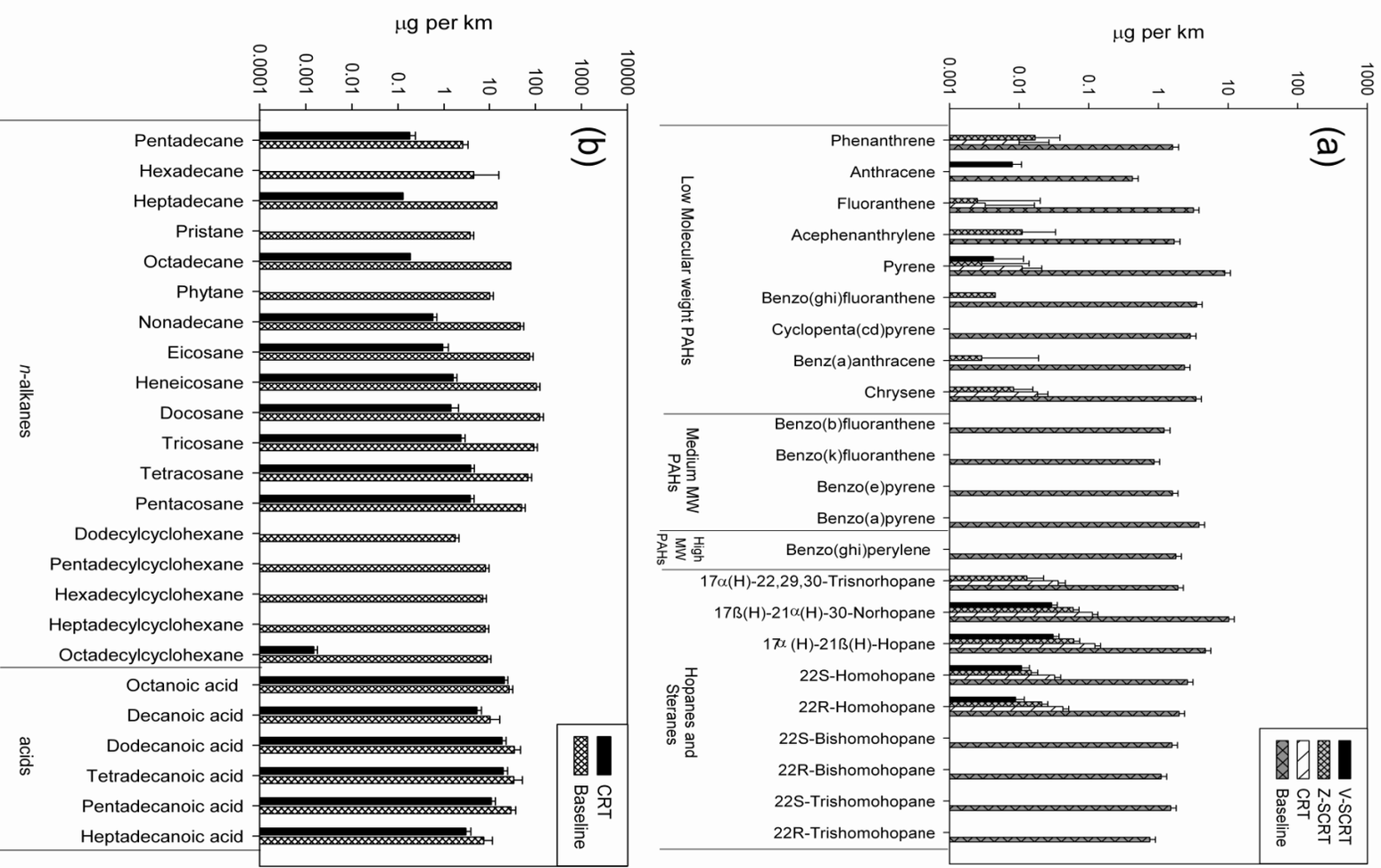
As shown in Figure 5.3a and 5.3b, both CRT and SCRT technologies effectively reduced the particle bound PAH, hopanes and steranes emissions compared with the baseline configuration by more than 99%. For medium and high molecular weight PAHs, the emission levels were below the detection limit of the analytical method with the introduction of both CRT and SCRT technologies. These higher molecular weight PAHs are often more carcinogenic according to the International Agency for Research on Cancer (IARC). In contrast to the non-detectable heavier PAHs, the presence of particle phase light PAHs (even in small amounts) in the exhaust of vehicles equipped with CRT and SCRT is attributed to the high volatility of these organic species. Light PAHs have lower vapor pressure and are most likely in the vapor phase in the hot engine exhaust. After passing through the control devices, these semi-volatile species may nucleate to form fresh particles, or condense onto existing particles under favorable dilution and temperature conditions (Kittelson et al., 2006; Vaaraslahti et al., 2004). These compounds may also condense or absorb on organic particles that have already nucleated. The occurrence of this PM formation mechanism has also been reported by Wehner et al., (2002) and Yu, (2001). In our studies, this observation is further supported by the substantial increase in number concentrations of nucleation mode particles for the vehicles equipped with control devices compared to the baseline truck, as reported in Section 3. For *n*-alkanes, there was a higher than 99% reduction in the emission factors after the introduction of CRT compared to baseline, whereas both V-SCRT and Z-SCRT devices effectively reduced the *n*-alkanes emission factors to below the detection limits. All three devices effectively reduced the organic acid content of the exhaust to under the detection limits.



**Figure 5-3 (a) Emission factors of PAHs, hopanes and steranes of baseline and control device-equipped vehicles in UDDS cycle. (b) Emission factors of n-alkanes and acids of baseline and control device-equipped vehicles in UDDS cycle.**

Figure 5.4a and 5.4b present the emission factors (EF) of particle bound PAHs, hopanes and steranes, *n*-alkanes and organic acids for the cruise driving cycle. The EF of PAHs, hopanes and steranes range from 0.42 µg/km to 10.17 µg/km with an average of 2.66 µg/km and 2.92 µg/km respectively for the baseline vehicle. The average emission factors of PAHs, hopanes and steranes at UDDS are 5.5 and 2.6 times higher than those at cruise cycle, respectively. For *n*-alkanes and organic acids at the cruise cycle, the emission factors range from 1.74 µg/km to 122.78 µg/km with an average of 36.6 µg/km and 23.9 µg/km for *n*-alkanes and organic acids respectively. The emission factors are also significantly lower than their corresponding values at the UDDS cycle. The reduction of these organic compounds may be attributed to possible adsorption of these species on the large surface area of the catalyst, followed by oxidation (Heck and Farrauto, 1995; Liu et al., 2008). PAH reduction by oxidation may be more efficient in the cruise compared to the UDDS cycle because the minimum activation temperature of the catalyst (for both V-SCRT and Z-SCRT) is reached much faster in the cruise cycle than the UDDS (Polidori et al., 2008). As the engine speed and load increase to a stable condition in cruise cycle, the exhaust temperature also increases to a level that enables full function of the catalyst.

All three control devices reduced the emissions of light molecular weight PAHs by more than 99%, while the emission factors of medium and high molecular weight PAHs after the devices were under the detection limits for the cruise cycle. The reduction of hopanes and steranes were equally high, i.e., 97.6%, 98.8% and 99.4% for CRT, Z-SCRT and V-SCRT, respectively. The CRT technology reduced *n*-alkanes and acids EF by 96% and 44% respectively, while both SCRT technologies showed a better performance by reducing *n*-alkanes and acids to levels below detection limits. Similar results were also observed for UDDS cycles. The improved performance of the SCRT catalyst may be due to their ability to break heavy hydrocarbons into smaller molecules (Ueno et al., 1998.).



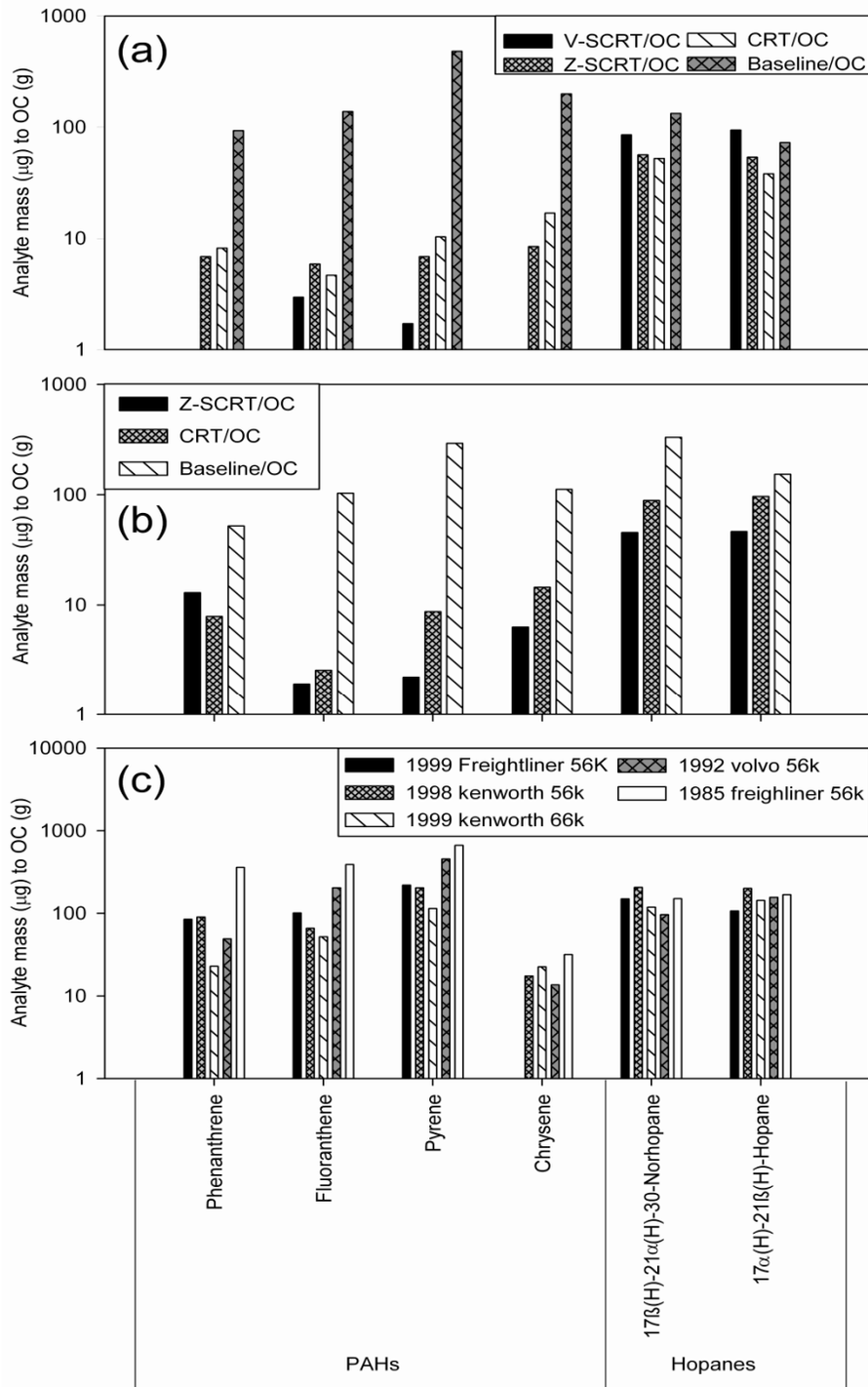
**Figure 5-4 (a) Emission factors of PAHs, hopanes and steranes of baseline and control device-equipped vehicles in cruise cycle. (b) Emission factors of n-alkanes and acids of baseline and control device-equipped vehicles in cruise cycle.**

Figures 5.5a, b and c show the concentration ratios of selected PAHs, hopanes and steranes to organic carbon of PM emissions from the baseline vehicle operating with and without control devices, running on UDDS (4.5a), and cruise (4.5b) cycles, and similar ratios of recently evaluated HDDV as reported by Riddle et al., (2007) (4.5c). As shown in these figures, the concentration ratios of the baseline vehicle displayed very similar profiles for these compounds between UDDS and cruise cycles, and in very similar magnitude as those observed by Riddle et al., (2007), which confirms the consistency in the sources of these PM organic tracers in both studies.

As shown in Figure 5.5a, PAHs-to-OC ratios are greatly reduced in vehicles equipped with control devices compared to baseline vehicle for the UDDS cycle. The average analyte mass ( $\mu\text{g}$ ) to OC mass (g) for the PAHs is 227.50 for baseline vehicle, and 10.0, 7.0, 2.3 for CRT, Z-SCRT, V-SCRT respectively, thus indicating a significant reduction of PAHs contributions to OC in the emissions with the control devices. The reduction in PAHs is very important from the perspective of public exposure because of their carcinogenicity (Harvey, 1985; Hewstone, 1994). The significant reduction of the relative contribution of PAHs to OC suggests that these compounds are either not formed as readily in combustion, or they are reduced very effectively in after-treatment devices using oxidation catalysts (Liu et al., 2008).

In contrast to PAHs, the ratios of analyte mass ( $\mu\text{g}$ ) to OC mass (g) for the hopanes and steranes are in the same order of magnitude for baseline vehicle (102.7) and controlled vehicles (45.1, 54.9 and 89.6 for CRT, Z-SCRT and V-SCRT respectively). Hopanes and steranes are exclusively found in lubricating oil. The similar ratios between retrofitted and baseline vehicles imply that even with the after-treatment devices, the relative contribution of the oil on the hopanes and steranes emissions are about the same order of magnitude as the baseline vehicle emissions. It should be noted that the ratio of hopanes and steranes to OC for the controlled vehicles is lower only by a factor of 2 or less (13%, 53% and 56% for CRT, Z-SCRT and V-SCRT, respectively) compared to the baseline vehicle. This suggests that the relative contribution of lubricating oil to the OC emissions in the controlled vehicles is more than half of those in baseline vehicles, even with the reduction in OC emissions by almost two orders of magnitude (95.0%, 99.2%, 99.2% for CRT, V-SCRT, Z-SCRT, respectively). It is possible that hopanes and steranes are oxidized at a slightly higher rate than the rest of the lubricating oil. However, a reduction by a factor of  $<2$  is quite small in the context of the overall OC reduction.

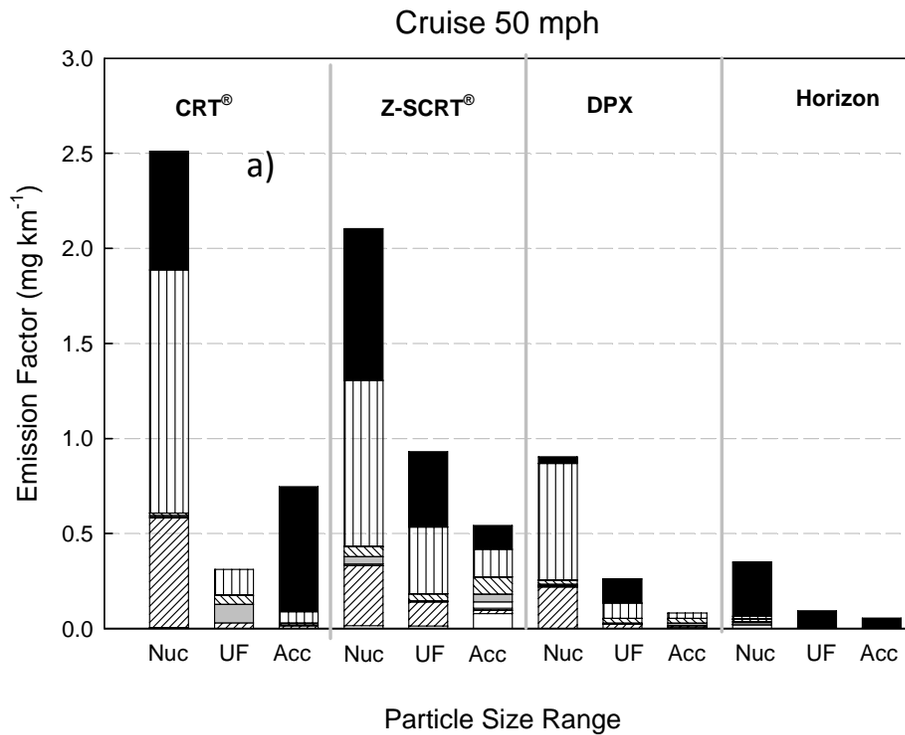
Similar results were also observed in the cruise driving cycle, as shown in Figure 5.5b. The average analyte mass ( $\mu\text{g}$ ) to OC mass (g) for the PAHs is 139.6 for baseline vehicle, 8.4 and 5.8 for CRT, Z-SCRT, respectively. The same ratio for the hopanes and steranes is 242, 92.4 and 45.9 for baseline, CRT and Z-SCRT, respectively. In general, the reduction efficiencies for high molecular weight PAHs, hopanes and steranes are consistently higher for the cruise cycle than the UDDS cycle. This is consistent with the fact that oxidation catalysts are more effectively activated to convert these organics by catalytic combustion in the cruise than the UDDS cycle, since the activation temperature is reached faster when the vehicles operate in the cruise cycle (Polidori et al., 2008). By comparison, low molecular weight PAHs, such as phenanthrene and fluoranthene, are largely in the gas-phase in the hot engine exhaust due to their high vapor pressure (Zielinska et al., 2004). Thus, their presence in the particle phase suggests that they are formed by nucleation or (more likely) by condensation onto pre-existing solid particles with the lowering of temperature as tailpipe emissions reach ambient conditions. Apparently, the effect of atmospheric dilution, which would favor the partitioning of these species in the gas phase, is outweighed by the drastic reduction in the exhaust temperature during cooling of the tailpipe (Kittelson et al., 2006; Vaaraslahti et al., 2004).

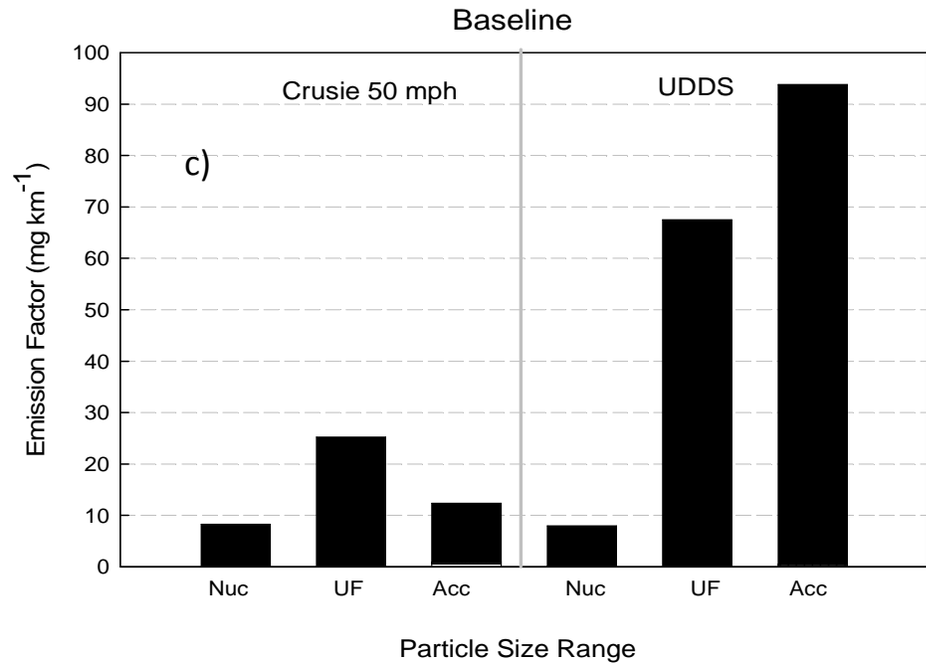
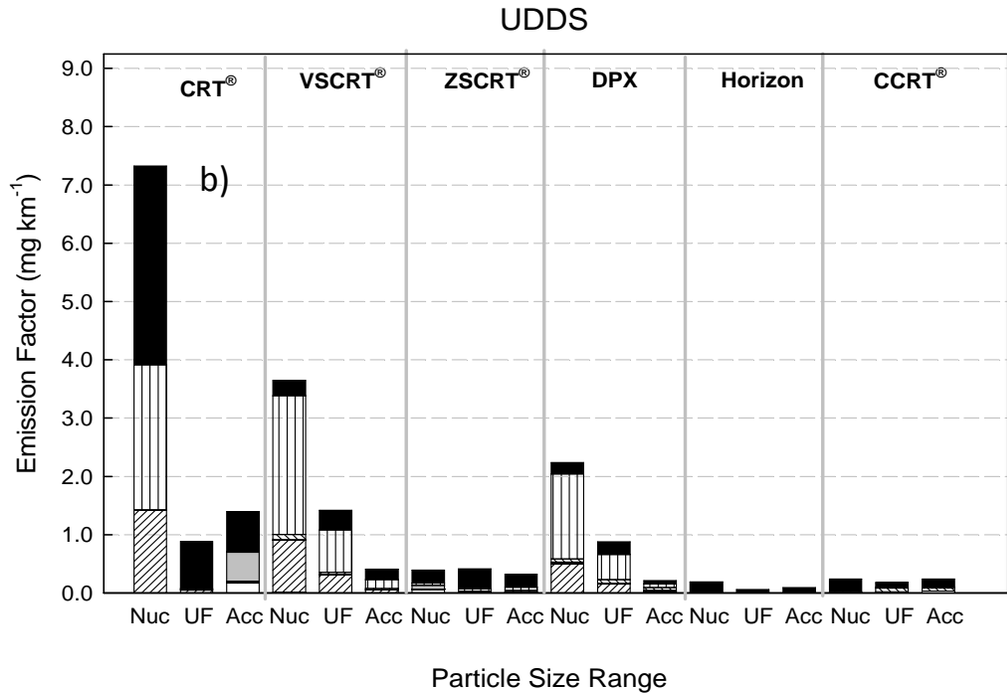


**Figure 5-5 Comparison of the ratios of PAHs, hopanes and steranes to organic carbon emission factors in μg/g from baseline and controlled vehicles. (a) running on UDDS cycle. (b) cruise cycle. (c) Ratios of PAHs, hopanes and steranes to organic carbon emission factors in μg/g as reported by Riddle et al., (2007)**

### Size-segregated Chemical Speciation

The ion and total carbon emission data are further segregated into three size ranges (Figure 5.6), i.e. 10-56nm, 56-180nm and 180-2500nm. The V-SCRT UDDS, Z-SCRT- cruise, CRT and DPX have displayed the highest content of sulfate in the nucleation mode. This finding is consistent with the conclusions of recent emission studies that nuclei mode particles from DPF equipped vehicles are predominantly sulfates (Grose et al., 2006). However, significant levels of TC were also present in these vehicles. During the analysis of the physical PM properties (Section 3), we observed that nucleation is suppressed in CCRT with un-aged catalyst and Horizon equipped with non-catalyst EPF, which have both a higher mass fraction of total carbon concentrations in all size ranges and less sulfate in nucleation mode PM. Unlike the SCRT cruise cycles, the UDDS cycles for Z-SCRT are unique in the sense that the emission rates of TC dominate those of sulfate. This may be again due to higher activation temperature of Zeolite catalysts, which may impede the conversion of SO<sub>2</sub> to sulfate during transient cycles, for which the exhaust temperature fluctuates continuously. Herner et al., (2007b) reported the critical temperatures (post after-treatment temperatures) required for Z-SCRT and V-SCRT to trigger nucleation were 373°C and 330 °C, respectively.





**Figure 5-6 Emission rates of size-resolved PM chemical species. a) Cruise 50 mph, b) Transient UDDS c) Baseline vehicle.**

**Note: Nucleation (Nuc, 10-56 nm), UF: 56-180 nm and Acc: 180 nm – 2.5 $\mu$ m.**

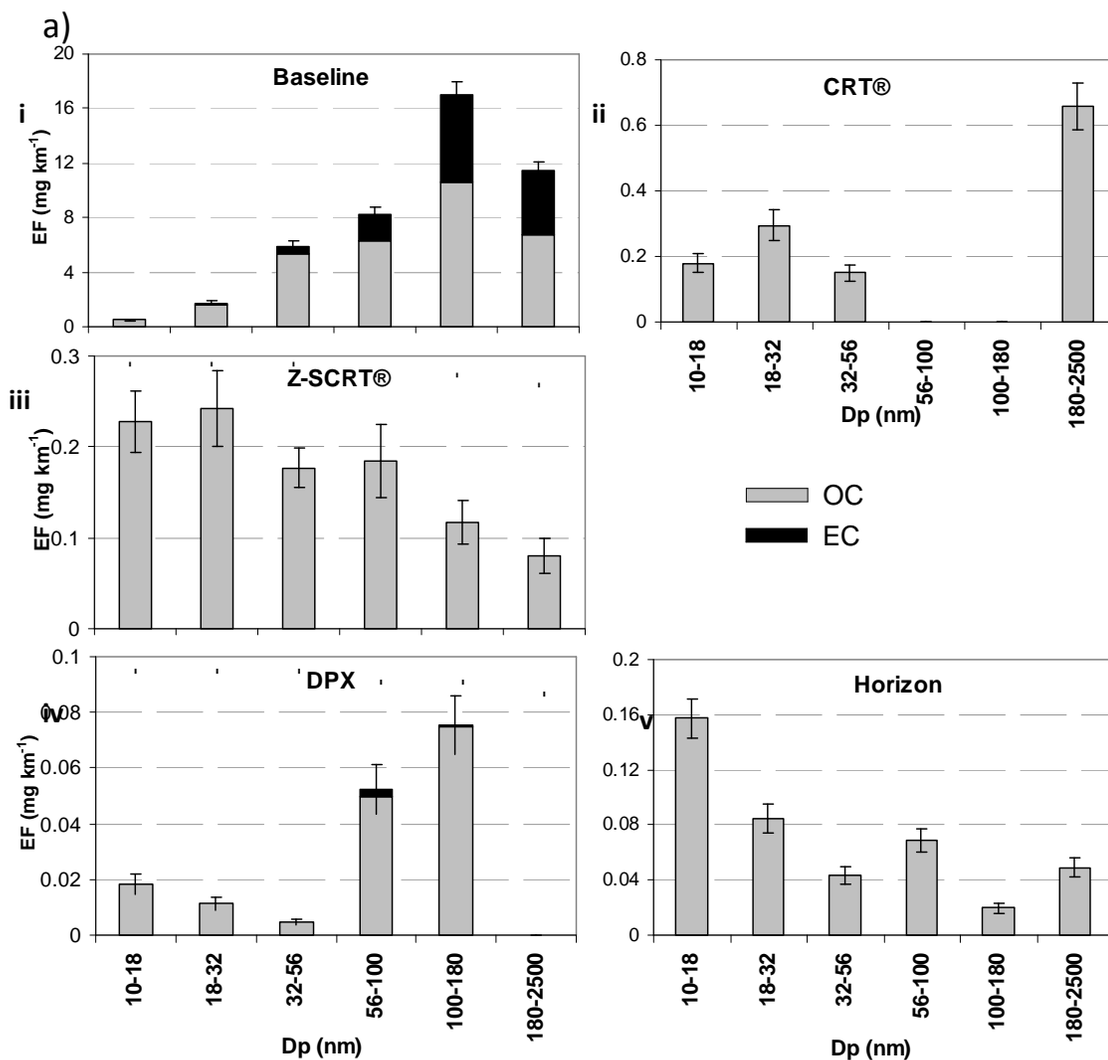
Ammonium is present exclusively in ultrafine PM, and especially in the nucleation mode particles. The formation of this species may be related to secondary mechanisms rather than being a constituent of the vehicle exhaust. Heeb et al., (2006) demonstrated the presence of gas-phase ammonia and particle phase ammonium in the exhaust of vehicles equipped with three way catalysts. They attributed this formation to catalytic reduction of NO in the exhaust, coupled with the low catalyst's selectivity for ammonia at the high temperature conditions of the exhaust.

It is interesting to note that ammonium was mainly observed for vehicles equipped with either a diesel oxidation catalyst (DOC) or a catalyzed filter, i.e. CRT, SCRTs, DPX, whereas vehicles operating without a DOC upstream of the DPF filter, such the Horizon, showed negligible emissions of ammonium. The CCRT is a notable exception to this observation. The lower ammonium level for CCRT or Horizon may also be due to their lower engine volume (5.9L), compared to SCRTs (11 L) and DPX (7.6 L). The measured ammonium is likely in the form of ammonium sulfate produced by gas-to-particle phase conversion processes catalyzed by the after-treatment devices. Grose and co-workers demonstrated that the nano-particles downstream of a CRT – equipped vehicle had a volatility profile similar to ammonium sulfate and concluded that the sulfates should be neutralized to some extent (Grose et al., 2006). Based on the experimental observations and discussion above, the formation of fresh nano-particles, especially for 'nucleating' vehicle configurations, may be attributed to a ternary nucleation process involving sulfuric acid, water vapor and ammonia. Meyer and Ristovski, (2007) demonstrated ternary nucleation pathway to be an important formation mechanism for diesel nanoparticles at high engine loads.

Figure 5.6c shows the mass size distribution of the chemical components of PM emissions from the baseline vehicle. Total carbon concentration outweighs any other species for the baseline vehicle (>99% of total PM mass concentration). Further, the UDDS cycle of Baseline vehicle emits almost 3-fold higher TC than the cruise cycle. This can be explained by the shift of TC peaks during cruise cycle in the 100-180nm (25mg km<sup>-1</sup>) size range to the even larger accumulation mode (180-2500nm, 90mg km<sup>-1</sup>) during UDDS cycle. The higher emission of TC during the transient cycles, however, was not observed for Z-SCRT and Horizon (Figure 5.6 b). The Z-SCRT-UDDS cycle emitted the least amount of PM mass among all the retrofitted cycles of Kenworth truck (CRT; SCRT cycles). This particular cycle also lacked a pronounced nucleation mode which was observed in CRT and other SCRT cycles. Higher activation temperatures for the Zeolite catalysts can inhibit the nucleation process (as mentioned before) especially during the fluctuating exhaust temperature of the UDDS cycle. Since a considerable fraction of TC (mostly OC for retrofitted vehicles) is associated with the nucleation mode, it is likely that the lack of nucleation can retard condensation of organics on these particles. Emissions of TC from both the Horizon cycles are low and comparable to the analytical limits of detection; therefore caution should be exercised in making inferences based on these data.

The size resolved total carbon (TC) is further segregated (in six size classifications of the MOUDI-nano MOUDI) in EC and OC and is shown in Figure 5.7 a, b. Here we should again emphasize that the EC-OC data are used to display trends, not necessarily precise values, due to the limitation of the analytical method as we discussed earlier, related to the use of Al foils for these measurements. OC clearly dominates EC emissions in vehicles retrofitted with diesel particulate filters, whereas by contrast, EC (30-40%) is comparable with OC (60-70 %) for the baseline vehicle. The EC is found mostly in the 100-180nm and 180-2500nm size ranges for the baseline vehicle. This is consistent with previous studies,

which also demonstrated that EC peaks in the higher size ranges of UF (> 80nm) as well as in the accumulation mode (Kleeman et al., 1999; Rose et al., 2006). The dominance of elemental carbon for the baseline vehicle was also inferred from our volatility analysis (Section 3), of which negligible loss of PM mass and number during heating process was observed.



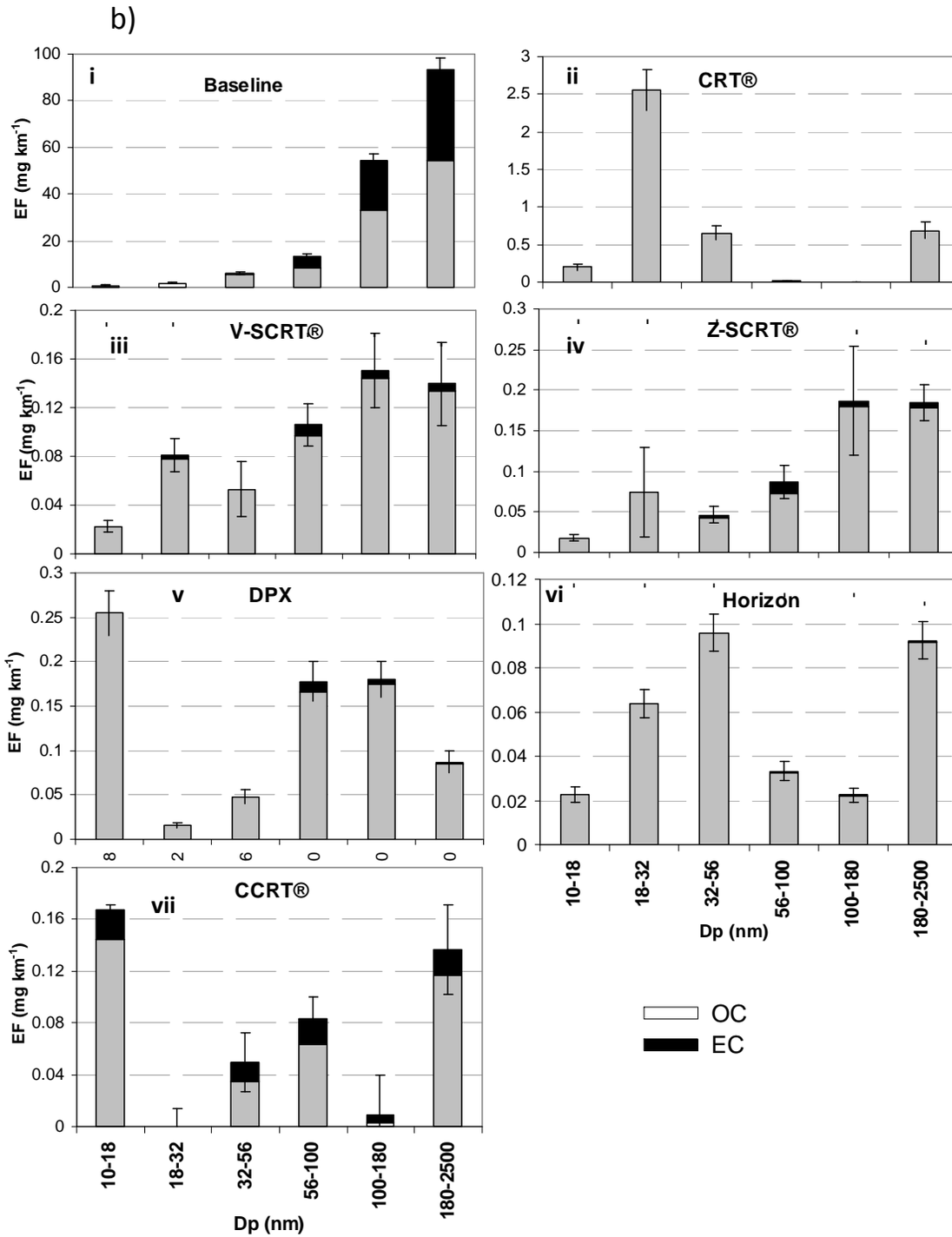


Figure 5-7 a, b: Size-resolved EC/OC mass distribution at a) Cruise mode b) UDDS mode.

For the vehicles equipped with DPF, soot particles are effectively captured, resulting in significant reduction of EC, hence a reduction in the particle surface area available for the condensation of semi-volatile organic vapors. On the other hand, the presence of catalysts will oxidize hydrocarbon (HC) and reduce the level of condensable semi-volatile OC. This will lead to the overall reduction in OC concentrations in the exhaust. The elimination of larger accumulation mode soot particles favors the formation and growth of nucleation mode particles by condensation of these organic species onto either partially or fully neutralized sulfate nano-particles. This is consistent with our findings discussed in the previous section regarding the increased sulfate content of nucleation mode PM in the DPF- equipped vehicles. During the UDDS cycle, a generally bimodal distribution of EC-OC was observed. Although insignificant in quantity, particles from UDDS cycles of DPF equipped vehicles have traces of EC, especially for the CCRT vehicle. The relatively higher EC fraction (22%) in CCRT might be associated with the higher accumulation mode observed in its number size distribution (Section 3). The transient cycles of diesel vehicles are generally known to emit more elemental carbon compared to cruise cycles (*Shah et al., 2004b*).

## 5.4 Conclusions

While comparing with a baseline vehicle, significant reduction (>90%) in EC, OC, WSOC and various organic compounds (PAHs, hopanes, steranes, n-alkanes and organic acids) emissions is achieved for vehicles with retrofit devices. The vehicles with significant nucleation (CRT, V-SCRT, Z-SCRT and DPX) mode particles produced considerable amount of sulfates especially during steady state operations. On the contrary, the non-nucleating configurations (Horizon, CCRT, and Z-SCRT-UDDS) were associated with higher amount of total carbon in the form of OC. In general, the transient cycles were associated with higher EC and OC. Soluble fraction of OC was highest for Horizon followed by SCRTs, DPX and baseline.

Individual ratios of speciated organic compounds to OC mass from the exhaust of baseline vehicle suggest that PAHs can form in combustion processes or originate from diesel fuel, whereas hopanes and steranes come from lubricating oils. With the introduction of control devices, the analyte to OC ratios reduced significantly for PAHs, while the reduction was more modest for hopanes and steranes, implying that fuel and lubricating oil have substantially different contributions to the OC emitted by vehicles operating with control devices compared to the baseline vehicle.

## 6. Toxicological Characteristics of Semi-volatile and Non-volatile Fraction of DEPs from Retrofitted Vehicles

### 6.1 Introduction

This section investigates the toxicological characteristics of PM from tested vehicles in comparison to the baseline vehicle. The oxidative activity was measured by two different assays – DTT assay and macrophage ROS assay. The DTT activity was determined for PM samples collected at both ambient and elevated temperature (thermally denuded of semi-volatile species), while ROS activity was measured only for undenuded samples. Further, to assess the role of transition metals in the measured ROS activity, undenuded samples were chelated using a Chelex® complexation method, followed by subsequent ROS analysis.

The study demonstrates that despite an increase in the intrinsic activity (both DTT and ROS, per mass basis) of exhaust PM with use of most control technologies, the overall activity (expressed per km or per hr) was substantially reduced for retrofitted configurations compared to the baseline vehicle. Significant reduction in DTT activity (by 50-100%) was observed for thermally-denuded PM from vehicles with retrofitted technologies (PM with significant semi-volatile fraction). On the other hand, Chelex treatment of undenuded PM samples removed a substantial ( $\geq 70\%$ ) fraction of the ROS activity.

Correlation analysis performed between measured activity and the chemical constituents showed that DTT activity is strongly associated ( $R=0.94$ ) with the water soluble organic carbon (WSOC), while the ROS activity was mostly driven by Fe content of the PM samples.

Exposure to DEPs has been linked with adverse health effects, including airway inflammation (Saldiva et al., 2002), pulmonary diseases such as asthma and bronchitis (Rusznak et al., 1994), mitochondrial dysfunction (Hiura et al., 2000; Seagrave et al., 2007), and lung cancer (Castranova et al., 2001; Garshick et al., 2004). As discussed in previous sections, that although PM mass emissions from retrofitted vehicles are significantly reduced ( $>90\%$ ), the number-based particles emissions increase under certain conditions. The chemical composition of particles emitted from these retrofits has also been shown to be substantially different than that of the untreated exhaust (Section 3). These changes in physico-chemical characteristics of PM emitted from retrofitted vehicles should also result in their diverse toxicity profiles, both on per mass and per distance traveled basis, compared to PM from unregulated vehicles, but this hypothesis remains to be examined.

The prime focus of this section is to investigate the oxidative activity of semi-volatile and non-volatile PM fractions from heavy-duty vehicles operating with and without emissions control technologies. The oxidative activity of the collected particles was measured by two independent assays: 1) the DTT assay, and 2) the macrophage ROS assay. In addition to assess the toxic potential of semi-volatile species, the role of transition

metals in inducing the cellular oxidative stress has also been examined by chelating the PM samples. For a subset of samples, we also used the dihydroxybenzoate (DHBA) assay to measure redox activity, and determine the ability of aqueous suspensions of PM from to generate hydroxyl radicals from hydrogen peroxide. Details of the DHBA assay are given by Cho et al (2005)

The results reported will be useful in assessing the efficacy of these control technologies and in establishing the future needs for additional measures to control these PM species.

## 6.2 Sample Analysis

The oxidative activity of the collected particles has been measured by two independent assays: 1) the DTT (Dithiothreitol) assay, and 2) the macrophage ROS (Reactive oxygen species) assay. As we discuss above, we also performed the DHBA assay on a limited subset of samples. For that assay, PM samples were collected in Biosamplers from the concentrated stream of versatile aerosol concentration enrichment system (VACES). The PM mass collected in Biosamplers were estimated by a parallel filter measurement from the same sampling line. Each cycle was repeated until sufficient PM mass (i.e., 3-5 mg) was collected for the analysis. Subsequently, PM collected in Biosamplers was analyzed for DHBA formation. Given the small number of samples and the different methodology used to collect them compared to the hi vol samples used for the DTT and ROS assays, these results are preliminary and a more detailed and comprehensive study should be conducted in the future, if there is sufficient interest in this assay. Some additional DHBA analysis was performed on left-over portions of the hi vol samples from selected vehicles and driving cycles (depending on the available PM mass left on the substrates), but as we noted earlier, these results are far too limited to be considered comprehensive and additional analysis in a separate study should be considered if deemed necessary.

The DTT assays were performed for denuded and undenuded PM collected on 47 mm Teflon filters and undenuded PM on HiVol (Teflon-coated glass fiber) filters. Particles deposited on these filters are first extracted into an aqueous suspension of Milli-Q water by sonication (Ultrasonic Cleaner, Model 5510R-MTH) for 20 minutes. The particle suspension is then incubated for a period between 0-30 minutes with 100mM DTT in 0.1 M phosphate buffer. An aliquot of mixture was added with 0.4 M TrisHCl (quench reaction), EDTA (Ethylenediaminetetraacetic acid) and 10mM DTNB (Beta dystrobrein) solution. The consumption rate of DTT was quantified by the formation of mercaptobenzoate at different times during the incubation. The assay is calibrated with each experiment using a standard diesel exhaust sample, whose overall rate varies by less than 10% among the experiments performed. The methodological details of this assay are provided by Cho et al., (2005).

Sub-sections of the HiVol PM samples were further analyzed for ROS activity by macrophage ROS assay. Samples were extracted with 15 mL of high purity (18 mega-ohm) water. Extractions were performed in 20 mL acid-leached, capped, polypropylene tubes, with continuous agitation for 16 hours at room temperature in the dark. Extracts were filtered through acid-leached 0.22  $\mu\text{m}$  polypropylene syringe filters. The ROS-activity of the PM extracts was measured by in vitro exposure to rat alveolar macrophage (NR8383, ATCC# CRL-2192) cells using 2',7'-dichlorofluorescein diacetate (DCFH-DA) as the fluorescent probe. DCFH-DA, a membrane permeable compound, is de-acetylated by cellular esterases generating 2',7'-dichlorodihydrofluorescein (DCFH). The oxidation of DCFH by ROS within the cell yields 2',7'-dichlorofluorescein (DCH), which is highly fluorescent and is monitored using a plate reader method. All samples, as well as positive and negative controls were analyzed in triplicate (3-wells). A minimum of six dilutions (each of them in triplicate) of every sample extract was run to ensure that a linear dose-response region could be identified. Methodological procedures of this assay are described in Landreman et al., (2008). Uncertainty in ROS method was estimated by propagating the standard deviation of triplicate ROS measurements with standard deviation of the applied method blank. These values ranged from 5 to 78% (median = 20%, n=24). Sub-samples of the primary water extracts (samples and controls) were further processed by chelation with the immobilized ligand iminodiacetate (Chelex chromatography) to remove the metal ions. For the Chelex treatment, mini-columns (1 mL polypropylene with Teflon frits) of Chelex were prepared with 0.2 g of 18 mega-ohm water-slurried Na-Chelex. Columns were rinsed with high purity water and buffered with 0.1 M sodium acetate. PM

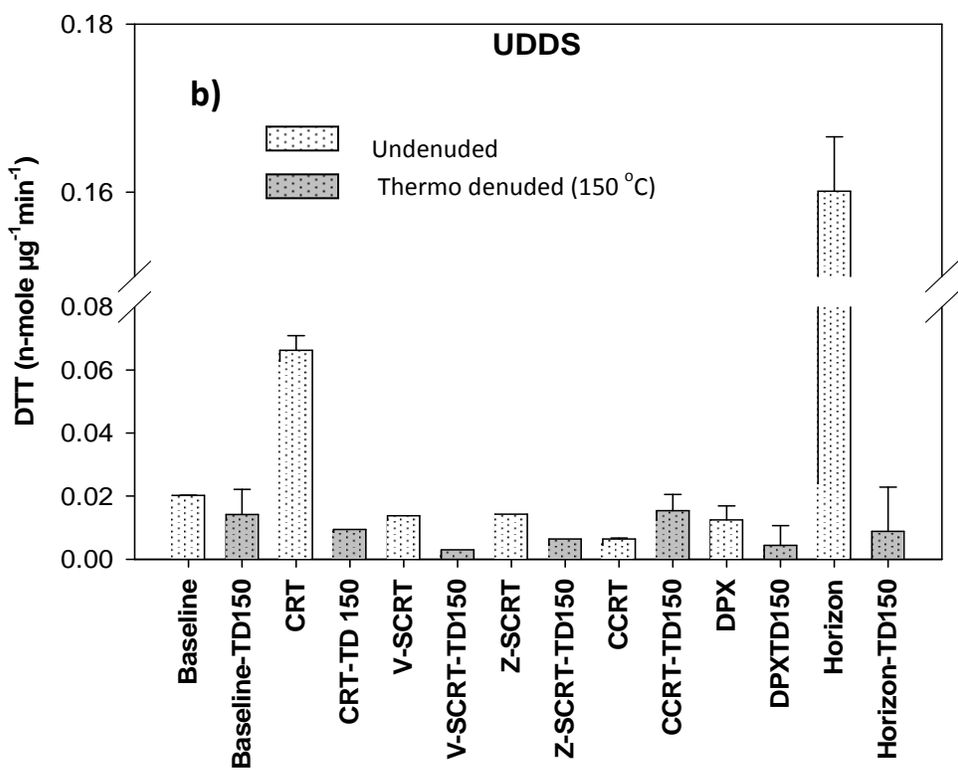
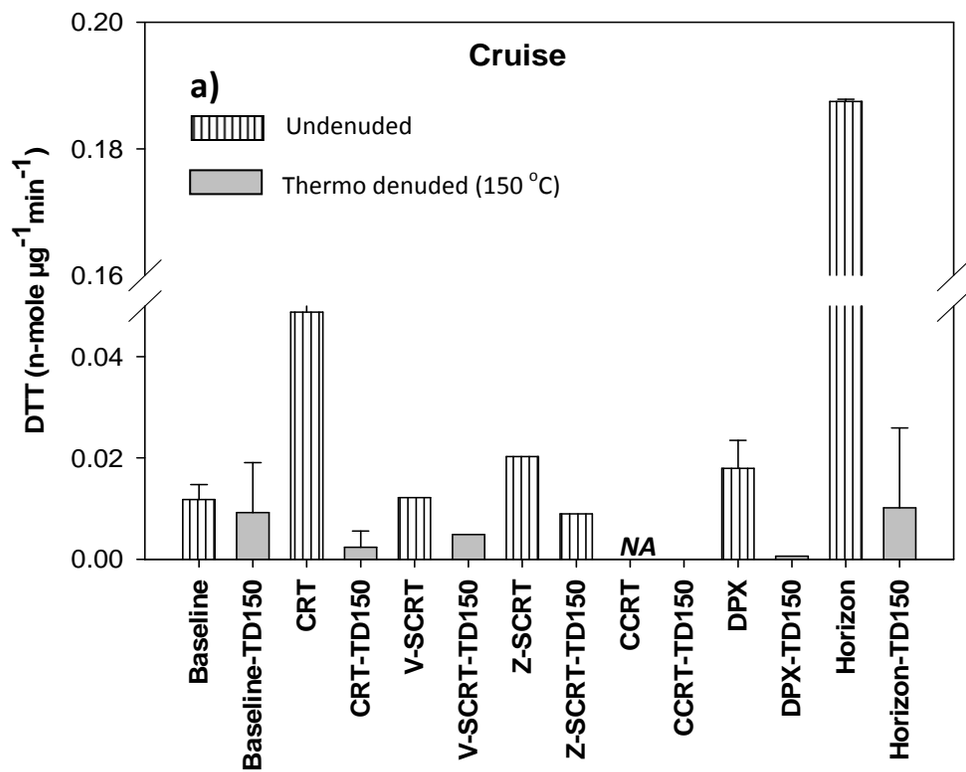
extracts (~1.75 mL) were processed through the columns, under gravity flow, in a solid phase extraction manifold, at a flow rate of ~1.0 mL/min. The Chelex-processed extract was collected and immediately assayed for ROS-activity and subsequent elemental analysis by magnetic sector inductively-coupled plasma mass spectrometry (SF-ICPMS). Control samples were processed through the Chelex columns to ensure that the treatment did not produce ROS active species or inhibit the activity of Zymosan (a  $\beta$ -1,3 polysaccharide of D-glucose). Zymosan was used as a positive control as it is recognized by TLR-2 (Toll-like receptors) on macrophage cells, activating a strong immuno-chemical and ROS response (Ciapetti et al., 1998). Blank-corrected ROS fluorescence data were normalized to the response of a standardized unit of Zymosan to correct for minor variations in method sensitivity between assay batches. ROS activity is therefore reported in terms of Zymosan units to facilitate the comparison of various data sets. None of the Chelex method blanks exhibited detectable ROS activity, and Chelex column blanks (high-purity water eluants) produced no measurable (<5%) suppression of ROS activity in Zymosan positive controls.

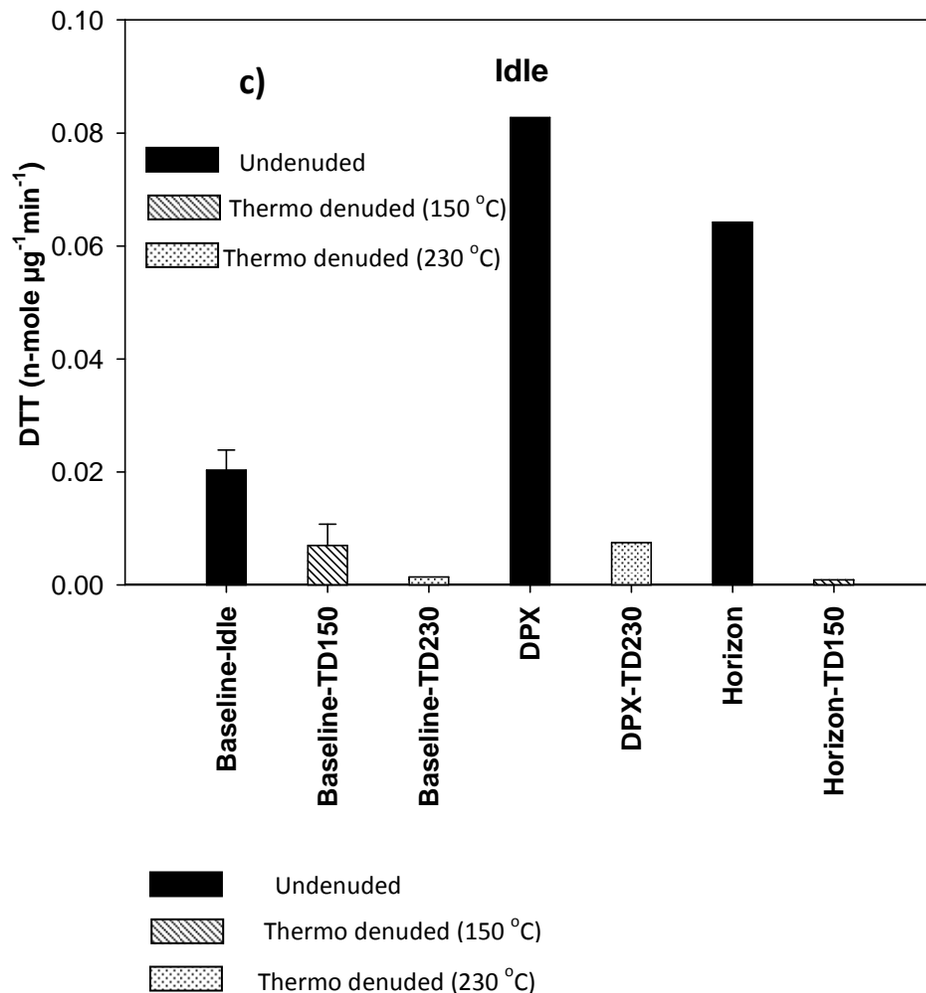
## 6.3 Results

### ***DTT Activity***

Blank-subtracted (filter blank  $\text{DTT} \sim 0\text{-}0.0005$  n-mole  $\mu\text{g}^{-1} \text{min}^{-1}$ ) average DTT consumption rates, normalized per unit mass of PM are reported in Figure 6.1 a,b,c for various vehicle configurations. The vehicle with the Horizon trap (School Bus) had the highest per mass DTT activity (0.16-0.19 n-mole  $\mu\text{g}^{-1} \text{min}^{-1}$  for cruise and UDDS) irrespective of driving conditions. The DTT consumption rates from both Vanadium and Zeolite-based SCRTs are on the same order of magnitude (0.01-0.02 n-mole  $\mu\text{g}^{-1} \text{min}^{-1}$ ). It is interesting to note that when the selective catalytic reduction (SCR) section from SCRT is removed from the exhaust stream and the vehicle is operated with only CRT (i.e., the DOC+ uncatalyzed filter), the DTT activity increased by a factor of almost 3, both for cruise and UDDS cycles. The CCRT (Catalyzed filter + DOC) is the most efficient among the test fleet with DTT rates as low as 0.006 n-mole  $\mu\text{g}^{-1} \text{min}^{-1}$ . Of particular note is the elevated level of DTT activity for the DPX and Horizon vehicles. The baseline vehicle emitted PM with similar DTT activity to those of newer vehicles with control technologies.

Figure 6.1 also reports the DTT values for the thermo denuded filters. There is a significant reduction (50-100%) in DTT activity as particles are heated to 150°C and their semi-volatile component is removed. The baseline vehicle, however, did not show any alteration in DTT response between denuded and undenuded exhaust stream, except while idling. This is because of the highly refractory nature (mostly soot) of these particles (Section 4).





**Figure 6-1 a, b, c: DTT consumption in n-moles  $\text{min}^{-1} \mu\text{g}^{-1}$  of PM of thermo-denuded and undenuded PM.** (DTT data are normalized to thermo-denuded PM and undenuded PM, respectively)

The increased DTT activity of the semi-volatile PM fraction is further highlighted by comparing with their corresponding semi-volatile PM mass and number fractions (Figure 6.2). For vehicles with (soot removing) control devices, the semi-volatile fraction contributed roughly 70-100% of the net oxidative activity compared to 20-30% for baseline vehicle. The sole exception is the CCRT vehicle, which hardly produced any DTT activity from its semi-volatile fraction, despite emitting 45% of semi-volatile PM by mass and 37% by number. For this vehicle (CCRT), the relatively low level of activity has originated from its residual non-volatile fractions. In general, particle number-based volatility, which is predominantly driven by the evaporation of the sub-50 nm, so-called nucleation mode particles (Kittelson, 1998), followed the activity trends better than the PM mass volatility.

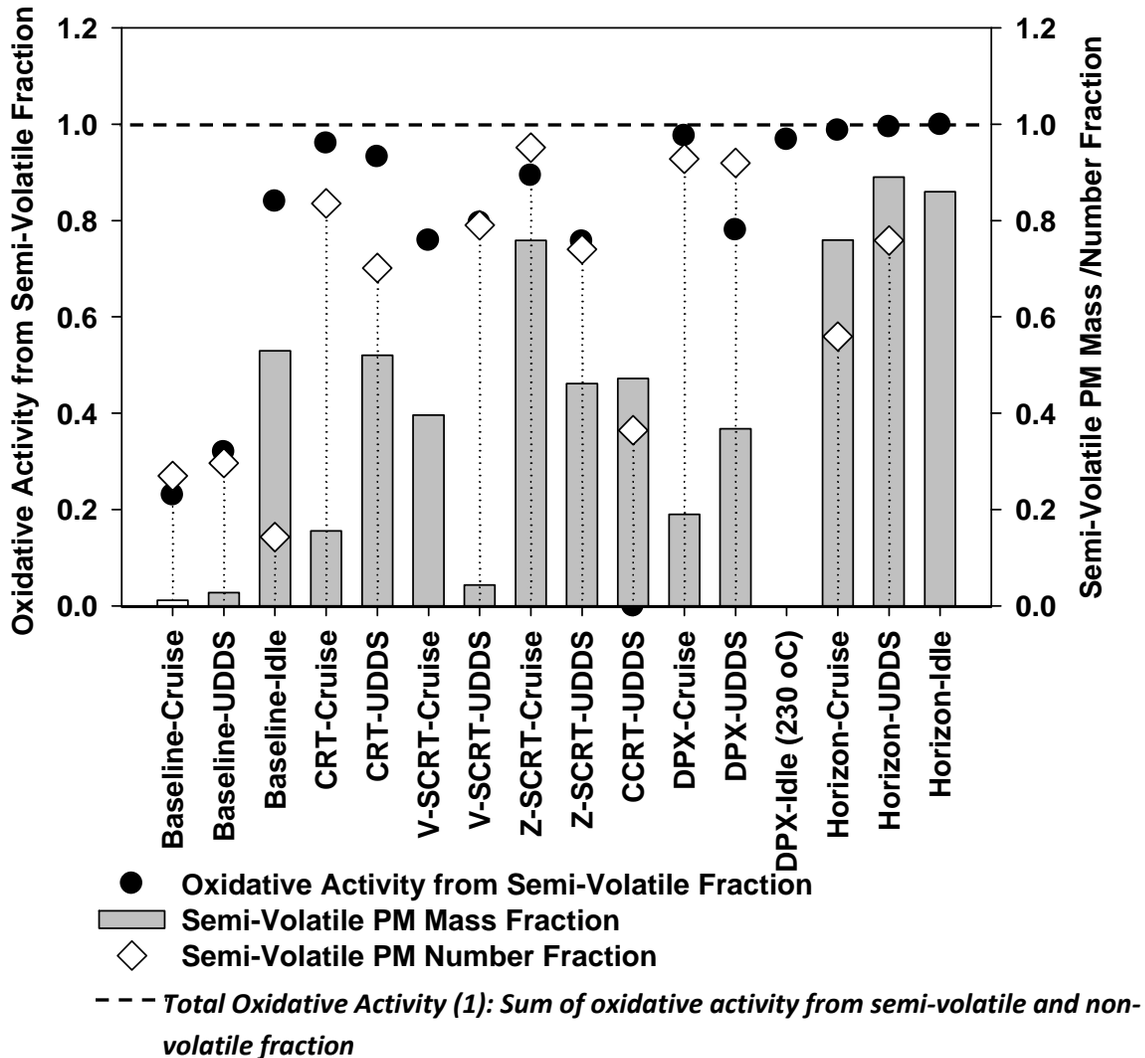


Figure 6-2 Relationship between oxidative activity and semi-volatile PM fraction at 150°C.

[Showing good correlation of DTT (filled circles) with particle number (hollow diamonds) and poor correlation with PM mass (bars)]

Note:

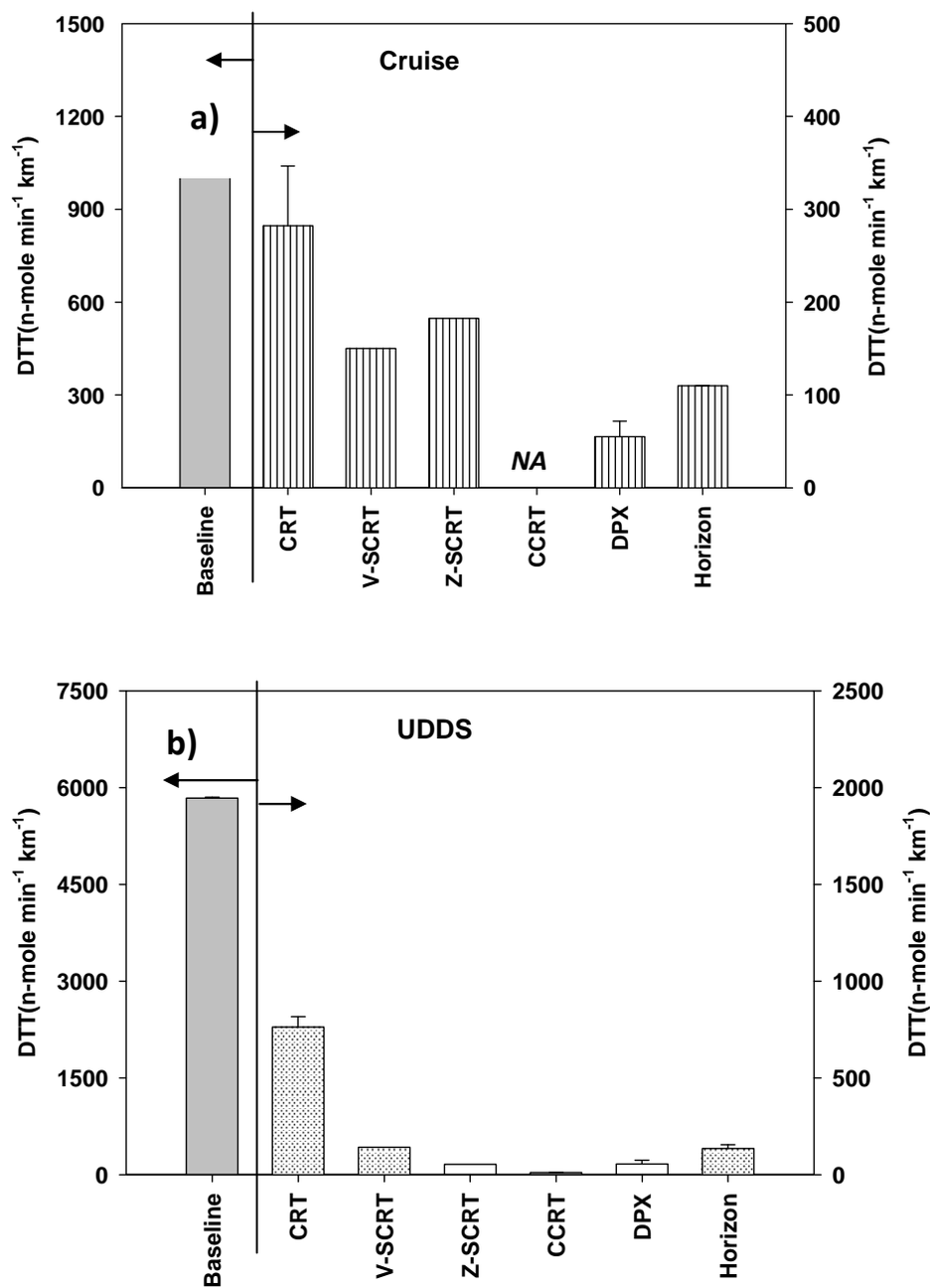
a) DTT activity from semi-volatile fraction =  $\frac{[(\text{DTT, undeneded, n-mole km}^{-1}\text{min}^{-1}) - (\text{DTT at } 150^{\circ}\text{C, n-mole km}^{-1}\text{min}^{-1})]}{[\text{DTT, Undeneded, n-mole km}^{-1}\text{min}^{-1}]}$

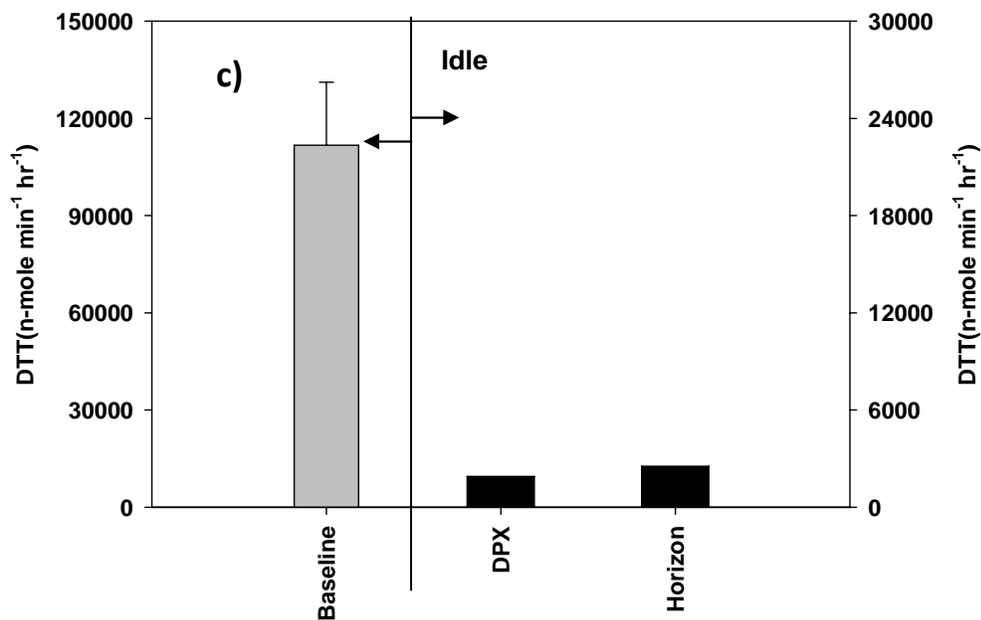
b) Semi-volatile PM mass/number fraction =  $\frac{[(\text{PM undeneded, mg or number km}^{-1}) - (\text{PM at } 150^{\circ}\text{C, mg or number km}^{-1})]}{[(\text{PM undeneded, mg or number km}^{-1})]}$

DPX Idle: DTT thermo denuded at 230°C; V-SCRT-Cruise, Horizon-Idle: Number volatility data not available

The DTT results expressed per unit vehicle distance traveled (Figure 6.3 a,b,c) provide a better idea of the total oxidative load imparted on the environment by these vehicles. They also provide a quantitative

assessment of the effectiveness of these after treatment devices. While we observed a comparable or lower oxidative response of the baseline vehicle per unit mass of PM, its net DTT consumption (per km) is substantially higher (>3 times) than retrofitted vehicles. This is because the baseline truck emits more than one order of magnitude higher particulate mass than other vehicles (Section 3).





**Figure 6-3a,b,c: DTT consumption per unit distance ( in n-moles min<sup>-1</sup> km<sup>-1</sup>) traveled by vehicles for Cruise and UDDS cycles and per hour( in n-moles min<sup>-1</sup> hr<sup>-1</sup>) for Idle.**

### ***Macrophage ROS activity***

Figure 6.4 shows the results of macrophage ROS assay conducted on the water extracts of exhaust PM samples from test vehicles. The results have been expressed both on a per mass of PM emitted basis ( $\mu\text{g ROS activity/mg of PM}$ ) (Figure 6.4a) as well as ROS activity per km (or per hr for idle) of vehicle driven (Figure 6.4b). The ROS activity per mass of particles emitted from all retrofitted configurations is higher than the baseline truck, with a sole exception of the hybrid vehicle. Again, the highest activity is observed for the school bus, which has been shown to be one of the most efficient vehicles in controlling both mass and number emissions (>99 %; Section 3). It is interesting to note that the hybrid vehicle, with similar particle mass and number emissions as that of the school bus, yields the lowest ROS activity. The DPX vehicle, which is also quite efficient in reducing PM mass emissions [up to 95 % (Section 3)], generates particles with high ROS activity (up to  $\sim 4600 \mu\text{g}$  of Zymosan units/mg of PM). Of particular note are also the elevated ROS levels observed for both vanadium and Zeolite based SCRTs compared to the baseline vehicle. Thus, although SCRTs effectively remove the targeted pollutants for which they were designed (i.e., hydrocarbons and NO<sub>x</sub>) in the exhaust emissions (Herner et al., 2009), their in-built catalysts appear to increase the fraction of certain redox active species in DEPs.

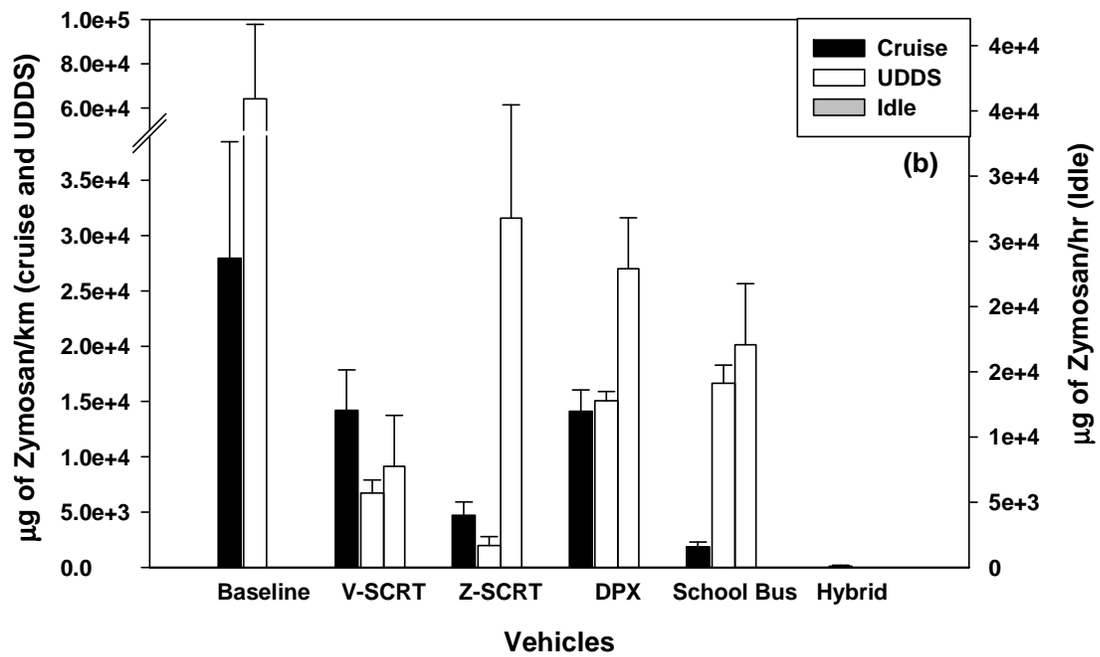
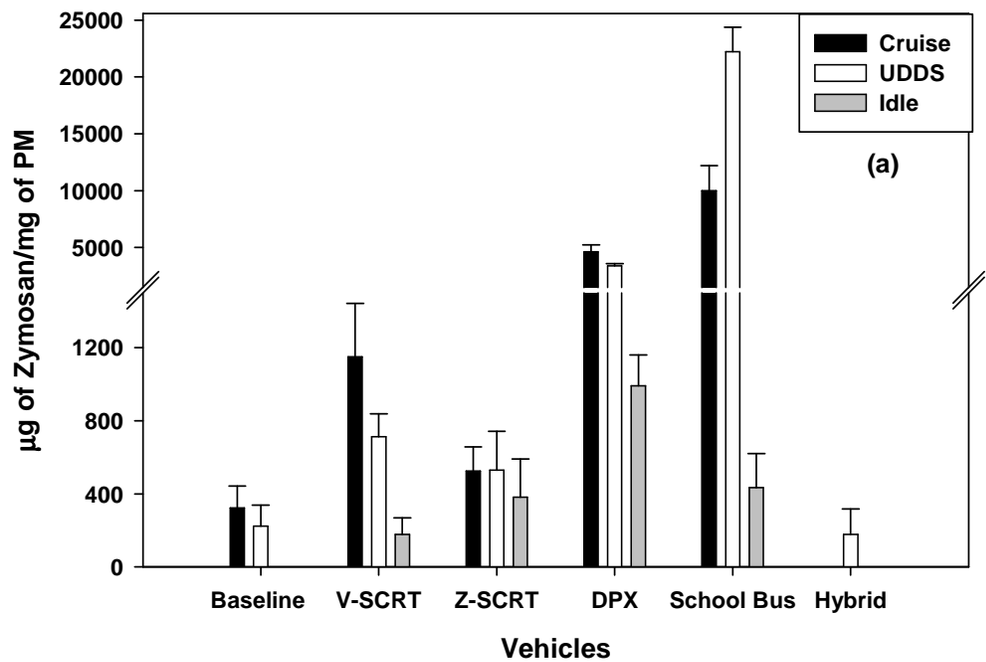
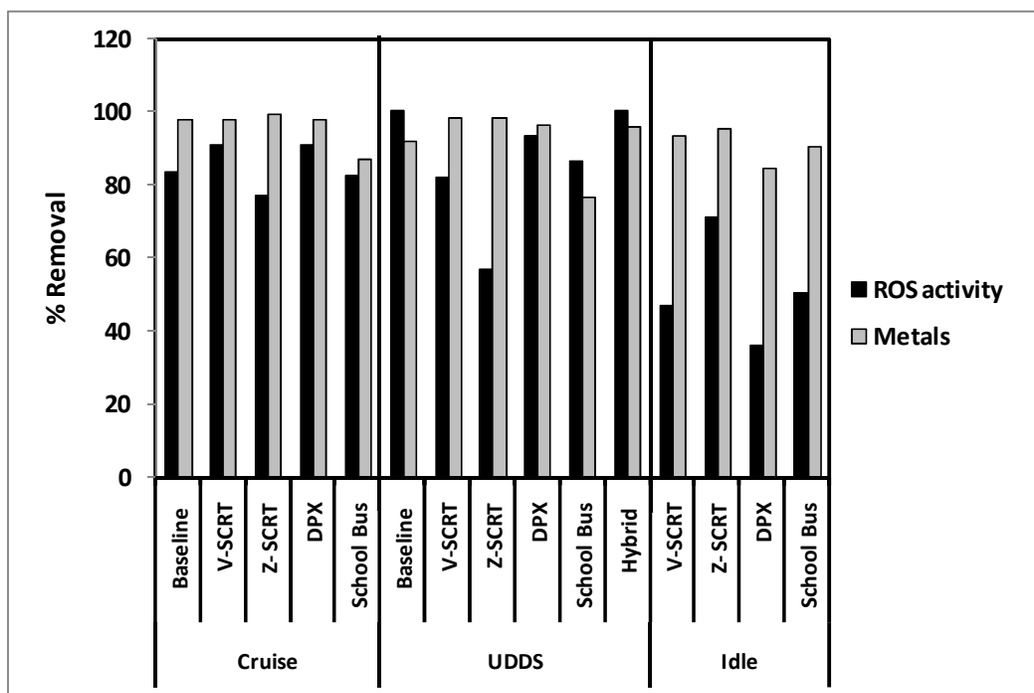


Figure 6-4 Reactive Oxygen Species (ROS) activity of PM from the tested vehicles, expressed as (a) per mass of PM, and (b) per km (or hr for idle) of vehicle driven.

It is of particular note that, although per PM mass activity is increased by most of the control technologies, all of the retrofitted configurations are efficient in reducing the per km (or per hr) ROS activity, compared to the baseline vehicle. Again, this reduction is a result of the lower particulate mass emissions (by more than one order of magnitude) from retrofitted configurations compared to the baseline truck (Section 3). The decrease in ROS activity is, however, not linear with the overall PM mass reduction in retrofitted vehicles. For example, an approximate 10-times reduction in PM mass emission achieved by Z-SCRT (cruise) lowers the ROS activity (per km) by only 3-4 times. The contrast between distance and mass based activity is consistent with the measurements using DTT assay. These findings have potential implications from the perspective of human exposure and risk assessment, and underscore the importance of both the intrinsic toxicity of PM, and its mass emission rate, while evaluating the net environmental impacts of an after-treatment technology.

Figure 6.5 shows the percent removal of ROS activity after Chelex treatment of the exhaust PM extracts from test vehicles under different driving cycles. To emphasize the correspondence between water-soluble metals and ROS activity, the percent removal of the sum of major metals from respective PM samples is also shown (Figure 6.5). Chelex treatment of DEPs water extracts removed a very large and fairly consistent fraction of ROS activity across most of the vehicles. The reduction in ROS activity averages at 77 ( $\pm 20$ ) % (range: 36% for DPX-idle to 100 % for baseline-UDDS and hybrid vehicles). The substantial reduction in ROS activity after metals chelation and removal highlights the dominant contribution of water soluble metals to ROS activity of DEPs.



**Figure 6-5 Percent removal of ROS activity in relation to that of aggregate water soluble metals after Chelex treatment of the exhaust PM samples from test vehicle-configurations under different driving cycles.**

### **Association between Chemical Species and Oxidative Activity**

To assess the contribution of various chemical species to the PM oxidative potential, regression analysis was conducted between the activity measured by two assays (DTT and macrophage ROS assay) and major chemical constituents. The Pearson correlation coefficient (R) and the associated significance levels (p-value) have been calculated and shown in Table 6.1 for the correlation of different chemical species with DTT values. The DTT was significantly correlated with WSOC (R=0.94, p<0.01) and organic acids (sum, R=0.91; p<0.01) and moderately correlated with organic carbon (R=0.76, P=0.02). All other species, including inorganic ions, PAHs, EC, and alkanes either have low or negative correlation.

WSOC is a complex mixture of diverse group of species, including dicarboxylic acids, poly acidic compounds, polyethers etc. (Decesari et al., 2000). Because of their hygroscopic nature, WSOC acts as surrogate for cloud condensation nuclei (Hori et al., 2003). Besides the well-documented implications of WSOC on global radiative balance, our work here indicates its relevance in health impacts, at least in the context of PM oxidative properties. More research however, is needed to identify the organic compounds present in water soluble components in DEPs. Only a few organic acids, constituting a small fraction (2-5% in most cases) of WSOC, were measured. Thus, it is difficult to conclude what specific species contribute to the overall WSOC and the related DTT consumption.

**Table 6-1 Correlation coefficient (R) and significance level (P) for DTT activity and selected chemical species**

| Species                       | Correlation with DTT |       |
|-------------------------------|----------------------|-------|
|                               | R                    | P     |
| EC                            | -0.35                | 0.37  |
| OC                            | 0.76                 | 0.02  |
| NO <sub>3</sub> <sup>-</sup>  | -0.09                | 0.77  |
| SO <sub>4</sub> <sup>2-</sup> | -0.32                | 0.27  |
| NH <sub>4</sub> <sup>-</sup>  | -0.25                | 0.26  |
| K <sup>+</sup>                | 0.43                 | 0.20  |
| Cl <sup>-</sup>               | 0.34                 | 0.15  |
| WSOC                          | 0.94                 | <0.01 |
| Alkanes (Alk.)                | 0.03                 | 0.54  |
| PAHs                          | -0.26                | 0.75  |
| Organic Acids (OA)            | 0.91                 | <0.01 |

On the other hand, ROS activity was found to be reasonably correlated with several transition metals such as Fe, Cr, Mn and Co (R>0.60; p<0.05) as shown in Table 6.2. The highest correlation was observed

for Fe (R=0.93; p=0.00), followed by Cr (R=0.67; p=0.00), Mn (R=0.62, p=0.01) and Co (R=0.61; p=0.04). Other metals (Mg, Al, Ca, V, Ni and Ba), semi-metals (As) and non-metals (P and S) are not significantly correlated with ROS activity.

**Table 6-2 Correlation coefficients and associated levels of significance for the regression between ROS activity and major water-soluble metals of PM**

| Elements | Correlation with ROS |          |
|----------|----------------------|----------|
|          | <i>R</i>             | <i>P</i> |
| Mg       | -0.19                | 0.36     |
| Al       | 0.38                 | 0.07     |
| P        | 0.27                 | 0.21     |
| S        | 0.13                 | 0.53     |
| Ca       | -0.01                | 0.95     |
| V        | -0.02                | 0.92     |
| Cr       | 0.67                 | 0.00     |
| Mn       | 0.62                 | 0.01     |
| Fe       | 0.93                 | 0.00     |
| Co       | 0.61                 | 0.04     |
| Ni       | 0.22                 | 0.31     |
| Cu       | 0.47                 | 0.02     |
| Zn       | 0.46                 | 0.02     |
| As       | 0.03                 | 0.89     |
| Cd       | 0.50                 | 0.01     |
| Ba       | 0.31                 | 0.15     |
| Pb       | 0.47                 | 0.02     |

Fe is among the most abundant transition metals in DEPs and is generally present in two oxidation states, Fe (II) and Fe (III). The solubility of Fe (II) is much higher than that of Fe (III); aqueous equilibrium concentrations of amorphous ferrous and ferric hydroxide [am-Fe(OH)<sub>2</sub> and am-Fe(OH)<sub>3</sub>] are nearly 10<sup>-5</sup> M and 10<sup>-10</sup> M, respectively (Nico et al., 2009). The redox cycling of Fe, generating an array of free radicals (e.g. OH\*, O<sub>2</sub>\*-, and HO<sub>2</sub>\*) through Fenton reaction, is well documented (Chen and Lippmann, 2009; Rose and Waite, 2005; Valko et al., 2005; Welch et al., 2002). Although, the exact mechanism of

the reaction is still a matter of debate, Fe (II) catalyzed reduction of molecular oxygen to release hydroxyl radicals [Haber Weiss reaction; (Winterbourn, 1987)] is considered to be largely responsible in Fe mediated toxicity. The comparatively higher water solubility of Fe (II) further makes it more bioavailable to probably participate in the reactions yielding ROS activity of DEP extracts.

## 6.4 Conclusions

This investigation explores the toxicological characteristics of the DEPs, emitted from retrofitted vehicles, in comparison to the baseline vehicle (without any control technology). The oxidative activity of the collected particles has been measured by two independent assays: 1) the DTT assay, and 2) the *macrophage* ROS assay. The effect of semi-volatile species on DTT assay was determined by comparing the DTT activity for PM samples collected at both ambient and elevated temperature. While, the contribution of transition metals in ROS activity was assessed by chelating the PM samples using a Chelex<sup>®</sup> complexation method.

The study demonstrates that despite generally similar reductions in PM mass emissions from diesel vehicles by various control technologies, the intrinsic oxidative activity (both DTT and ROS) of the emitted particles may vary dramatically with retrofit types. Although, mass based levels ( $\mu\text{g}/\text{mg}$  of PM) increased with the application of after-treatment devices, a significant reduction was observed in the overall oxidative activity (per km for cruise and UDDS and per hr for idle) for retrofitted configurations, compared to the baseline vehicle.

A substantial fraction of DTT activity of DEPs was associated with the semi-volatile fraction of the particles as demonstrated by a significant reduction in the activity (by 50-100%) observed for thermally-denuded PM. On the other hand, non-volatile transition metals drive the response of ROS assay as indicated by a substantial removal ( $\geq 70\%$ ) of the ROS activity after Chelex treatment of the PM samples. A univariate regression analysis further supported that DTT activity is strongly associated ( $R=0.94$ ) with the water soluble organic carbon (WSOC), while Fe is responsible for most of the variability ( $R=0.93$ ) in ROS levels.

## 7. List of Publications from this Study

1. Biswas, S., Hu, S.H., Verma, V., Herner, J.D., Robertson, W.H., Ayala, A., Sioutas, C., 2008. Physical properties of particulate matter (PM) from late model heavy-duty diesel vehicles operating with advanced PM and NO<sub>x</sub> emission control technologies. *Atmospheric Environment* 42 (22), 5622-5634.
2. Biswas, S., Verma, V., Schauer, J.J., Sioutas, C., 2009. Chemical speciation of PM emissions from heavy-duty diesel vehicles equipped with diesel particulate filter (DPF) and selective catalytic reduction (SCR) retrofits. *Atmospheric Environment* 43 (11), 1917-1925.
3. Pakbin, P., Ning, Z., Schauer J. J., Sioutas, C. 2009. Characterization of particle bound organic carbon from diesel vehicles equipped with advanced emission control technologies, *Environmental Science and Technology* 43 (13), 4679–4686.
4. Biswas, S., Verma, V., Schauer, J.J., Cassee, F.R., Cho, A.K., Sioutas, C., 2009. Oxidative potential of semi-volatile and non volatile particulate matter (PM) from heavy-duty vehicles retrofitted with emission control technologies. *Environmental Science & Technology* 43 (10), 3905-3912.
5. Verma, V., Shafer, M.M., Schauer, J.J. and Sioutas, C., 2010. Contribution of transition metals in the reactive oxygen species activity of PM emissions from retrofitted heavy-duty vehicles. *Atmospheric Environment*, 44 (39), 5165-5173.

## 8. Major Conclusions

This study has led to the following general conclusions.

- The newer diesel engines with emission control devices are very efficient in reducing the mass emission of particulate matter. However, enhanced formation of nucleation mode particles is observed in the exhausts of some retrofitted configurations. The vast majority of these nucleation mode particles are semi-volatile in nature.
- The total emissions (per distance of vehicle traveled) of the major chemical species (e.g. elemental and organic carbon) are substantially reduced in the retrofitted vehicles compared to the baseline vehicle. However, sulfate emissions increase for the configurations with enhanced nucleation mode particles in their exhaust. Although the retrofitted vehicles emit less water soluble organic carbon (WSOC) per mile of vehicle driven, the per PM mass water soluble fraction of the organic carbon (WSOC/OC) is increased for most configurations.
- With the introduction of retrofit control devices, the individual ratios of speciated organic compounds to OC are reduced significantly for PAHs, while the reduction was more modest for hopanes and steranes, implying that fuel and lubricating oil have substantially different contributions to the OC emitted by vehicles operating with control devices compared to the baseline vehicle. We hypothesize that PAHs can form in combustion processes and/or originate from diesel fuel, whereas hopanes and steranes come from lubricating oils.
- Despite an increase in the intrinsic oxidative activity (both DTT and ROS, per mass basis) of exhaust PM with the use of control technologies for most configurations, the overall activity (expressed per km or per hr) was substantially reduced for retrofitted configurations compared to the baseline vehicle.
- The semi-volatile fraction of the exhaust particles was observed to be highly oxidative in nature as demonstrated by a significant reduction in DTT activity (by 50-100%) observed for thermally-denuded PM. However, non-volatile species - particularly transition metals, are also responsible in cellular oxidative stress, as indicated by a substantial removal ( $\geq 70\%$ ) of the ROS activity after Chelex treatment of the PM samples.
- The correlation analysis showed that DTT activity is strongly associated ( $R=0.94$ ) with the water soluble organic carbon (WSOC), while Fe is responsible for most of the variability ( $R=0.93$ ) in ROS levels.
- The DTT and ROS assays, used in this study are two independent and intrinsically different types of analyses, and probably, are based on the generation of different oxidizing species by chemical PM

constituents. However, both of these assays provided important information for elucidating the health risks related to DEPs exposure from heavy-duty vehicles.

- An important caveat of the toxicological findings of this study is that they are all based on molecular or cellular assays that examine the toxicity of the PM suspension collected from a given vehicle and driving configuration based on PM mass. By their nature and design, these investigations did not take into account important parameters determining the toxicity and overall health effects attributable to the inhalation of an aerosol , such as particle size. The substantial reduction in the overall particle size distribution of newer vehicles creates an aerosol with a much higher lung deposition fraction than the baseline vehicle, and with considerably different toxicokinetics inside the human body once inhaled. Such important investigations can only be addressed by in vivo inhalation exposure studies to these aerosols, whether using animal models or human volunteers (or both), and are greatly needed in order to provide a more complete perspective to the results of this study.

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- University of Wisconsin Madison
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- University of California Los Angeles
  - John Froines, Arthur Cho ,Debra A Schmitz, Emma Di Stefano

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