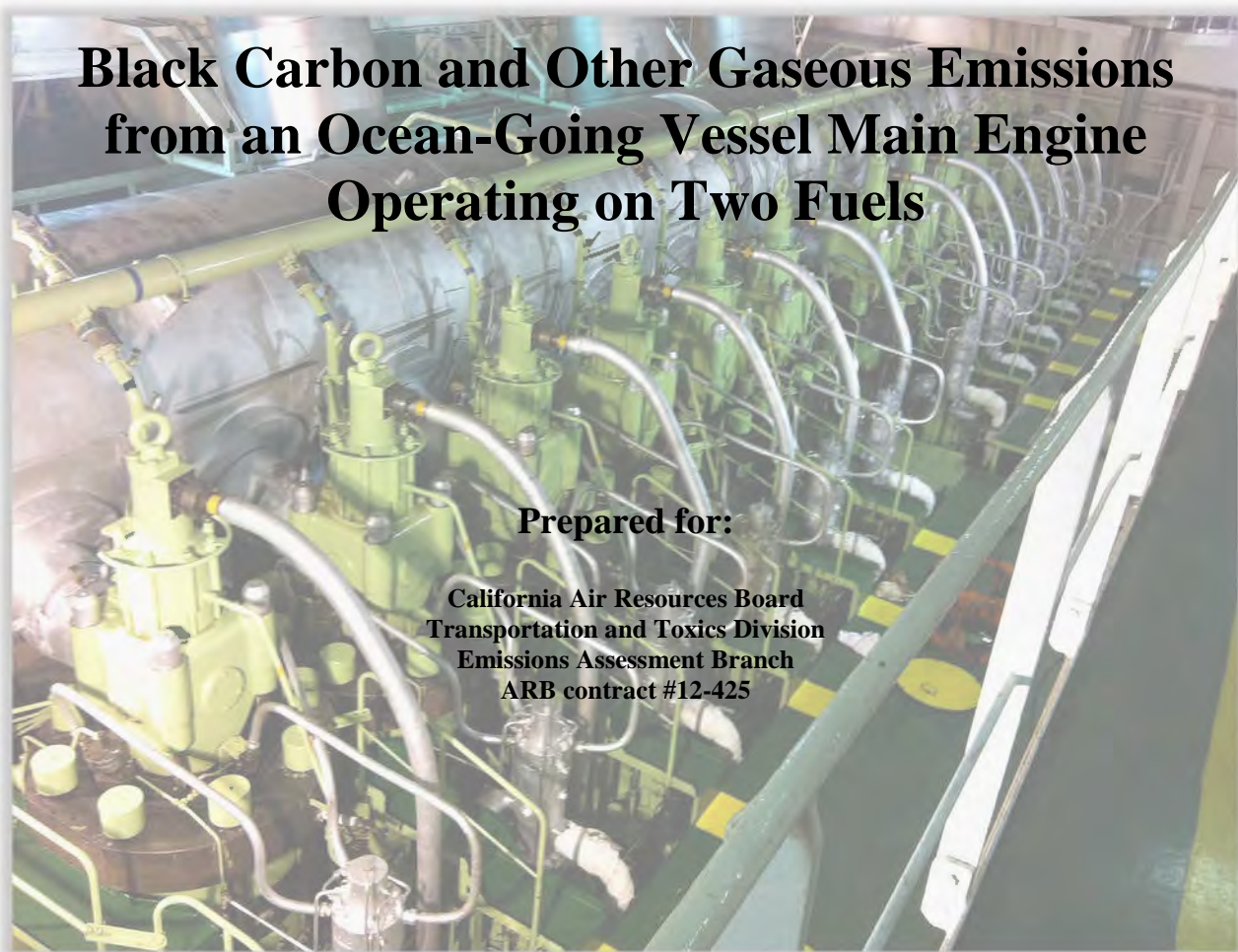


Final Report

Black Carbon and Other Gaseous Emissions from an Ocean-Going Vessel Main Engine Operating on Two Fuels



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Abstract

Movement of raw materials and finished goods by ocean going vessels represents a large emission source of smog and soot precursors. Accordingly there have been several efforts on the part of regulatory agencies, the ports and others to implement programs to reduce the emissions and mitigate the adverse impacts on near-by port communities and regional air quality. In addition to these actions, efforts are being devoted to better understand the impact of shipping on both the regional and global environment. The International Maritime Organization (IMO), has played a key role in bringing the shipping community and governments from around the world together to look at the impact of ship emissions including emerging issues associated with the release of black carbon (BC) from ships and the subsequent deposition on arctic ice. Interest is also growing in California as efforts are expended to identify ways to reduce emissions of short lived climate pollutants such as BC to help meet California's climate goals.

Recent measures enacted to reduce emissions from ocean-going vessels have required the use of cleaner low sulfur fuels. These measures do not specifically target BC emissions. Rather, the focus is on reducing sulfur oxides (SO_x) and total particulate matter (PM) which, for ships, includes sulfate, organics, elemental carbon, and metals. Ocean going vessels are a dominate source of sulfate PM due to the high sulfur levels in the fuel thus, many controls that target total PM reduction also reduce sulfate and organics well. It is unknown what impacts the control strategies that target PM have on BC. Control and quantification of BC emissions from ocean going vessels presents a challenging task since access to these vessels is difficult, limited, and there is no source of data available in the literature.

The purpose of this research was to quantify the black carbon (BC) and other emissions from a marine main engine operating on marine gas oil (MGO) and a heavy fuel oil (HFO) on an ocean voyage from Oakland to San Pedro, CA. Emissions measurements included Total Hydrocarbons (THCs), Nitrogen Oxides (NO_x), Carbon Dioxide (CO_2), and PM, including $\text{PM}_{2.5}$ mass (2.5 μm fine particles only), elemental, organic and BC. Several real time or semi-real time BC measurement techniques based on the principles of gravimetric net weight change, flame ionization detection (FID), photoacoustic (PA) measurements, and aerosol light scattering were employed.

The measured bsBC emissions by the Micro Soot Sensor (MSS) 483 and the Photoacoustic Extinctionmeter (PAX) were essentially the same for MGO fuel and for the HFO fuel. The correlation coefficients for these BC measurements versus EC measurements by the IMPROVE and NIOSH methods all exceed 0.96 indicating a significant correlation. The slopes of the correlations vary from 0.75 to 1.6 depending upon the sensor and the fuel.

The bs NO_x emissions for MGO fuel were slightly lower than those for the HFO fuel, averaging 21.9 for the MGO fuel and 24.6 for the HFO fuel for the 20 to 60% loads. Brake specific SO_2 (bs SO_2) emissions were lower for the MGO fuel compared to the HFO fuel at the same load, consistent with the lower sulfur content of the MGO fuel. Emissions on a brake specific basis were generally lower at the lowest load point, which was 5% for this study.

Acronyms and Abbreviations

ARB	Air Resources Board
bs	brake specific
BC	black carbon
CARB	California Air Resources Board
CE-CERT	College of Engineering-Center for Environmental Research and Technology (University of California, Riverside)
CFR	Code of Federal Regulations
CO	carbon monoxide
COV	coefficient of variation
CO ₂	carbon dioxide
Dp	particle diameter
DPF	diesel particulate filter
DR	dilution ratio
EC	elemental carbon
FID	flame ionization detector
g/hp-h	grams per brake horsepower hour
HFO	heavy fuel oil
lpm	liters per minute
MDL	minimum detection limit
MFC	mass flow controller
MGO	marine gas oil
nm	nanometers
NO _x	nitrogen oxides
OC	organic carbon
OEM	original equipment manufacturer
PA	photo acoustic
PEMS	portable emissions measurement systems
PM	particulate matter
PM _{2.5}	ultra-fine particulate matter less than 2.5 μm
RPM	revolutions per minute
scfm	standard cubic feet per minute
SEE	standard error estimate
SOF	soluble organic fraction
SO _x	Sulfur Oxides
THC	total hydrocarbons
UCR	University of California at Riverside

Executive Summary

Black Carbon (BC) is a climate forcing agent formed through incomplete combustion of fuels. BC has direct and indirect climate effects. Black carbonaceous material can directly affect climate via the absorption of visible solar radiation or indirectly effect climate via interactions with warm and cold clouds. BC emissions are the second strongest contributor to global warming, trailing behind carbon dioxide. BC stays in the atmosphere for weeks, whereas carbon dioxide has an atmospheric lifetime of over a century. Identifying the most effective way to test BC emissions on ocean-going vessels and to improve the understanding of the effectiveness of different ocean-going vessel emission control options is important for informing efforts to reduce emissions from ocean-going vessels that contribute to climate change.

The objective of this work is to evaluate emissions from a marine main engine operating on marine gas oil (MGO) and a heavy fuel oil (HFO) were measured on an ocean voyage from Oakland, CA to San Pedro, CA. Emissions measurements included SO_x, THCs, NO_x, CO₂, and PM, including PM_{2.5} mass (2.5 μm fine particles only), elemental, organic and BC. Several real time or semi-real time BC measurement techniques based on the principles of gravimetric net weight change, flame ionization detection, photoacoustic measurements, and aerosol light scattering were employed.

The main findings from this work can be summarized as:

- Brake specific NO_x (bsNO_x) emissions were relatively similar for the different loads for test loads higher than 20%. The bsNO_x emissions for MGO fuel were slightly lower than those for the HFO fuel, averaging 21.9 for the MGO fuel and 24.6 for the HFO fuel.
- Brake specific CO₂ (bsCO₂) emissions varied from ~600 g/kWhr to ~690 g/kWhr from high (62%) to low (20%) load. But when the test load was around 5%, brake specific CO₂ (bsCO₂) emissions were ~1000 g/kWh. These values are consistent with expected bsCO₂ values and agree with typical published data, where higher brake specific fuel consumption exists at lighter loads. Additionally, the bsCO₂ emissions were similar for the MGO and HFO fuel, suggesting the two test conditions were similar.
- BC measurements and the existing ISO 8178 sampling procedures work well for in-situ Photoacoustic type measurement methods.
- For the same fuel at the same engine load, the two photoacoustic meters give essentially the same results.
- At the 5% engine load point both photoacoustic meters showed higher BC emissions for the HFO fuel while at all other load points the BC emissions are statistically the same for both fuels.
- EC as determined by the NIOSH and IMPROVE method correlated well with the BC measurements for both fuels. The R-squared correlation coefficients are between 0.9619 and 0.9937 for correlations of bsBC with bsEC. The slopes of the equations vary between 0.7504 and 1.576 depending upon the BC and EC method and the fuel.
- The differences in the bsEC between the NIOSH and IMPROVE method is related to the method and not the sample since the samples were taken from the same quartz filter.

1 Introduction

Movement of raw materials and finished goods by large ocean going vessels results in significant emissions of smog and soot precursors. Accordingly, there are several efforts on the part of regulatory agencies, the ports and others to implement both regulatory and voluntary programs to reduce the emissions and mitigate the adverse impacts on near-by port communities and regional air quality. These efforts, such as requiring ocean-going vessels to use cleaner fuels, to slow down when approaching ports and to connect to shore power when at-berth have resulted in dramatic reductions in emissions of sulfur oxides (SO_x), particulate matter (PM), and to a lesser extent nitrogen oxides (NO_x). While some information is available regarding the impacts of these measures on PM emissions, very little data is available on the impacts of these programs on black carbon (BC).

Black carbon (BC) is known to strongly absorb visible light. It is often formed from the incomplete combustion of fuels and is thus considered an anthropogenic emission. BC has an average atmospheric life time of a few weeks, but it can significantly modify the earth's energy balance during that short time. Hence BC is a short lived climate forcing agent. Thus, the reduction of atmospheric BC emissions is being considered as a near-term mitigation strategy for climate impacts.

BC has both direct and indirect climatic effects. BC is a dominant absorber of solar radiation in the atmosphere. Furthermore, BC is transported over long distances and can mix with other aerosols to form transcontinental plumes of brown clouds. Anthropogenic sources of BC are concentrated in the tropics where high solar irradiance occurs. BC's high absorption properties, regional distribution aligned with high solar irradiance, and the capacity to mix and form widespread brown clouds make the emissions of BC the second strongest contribution to global warming, trailing behind only carbon dioxide. Furthermore, the deposition of BC darkens snow and ice surfaces, contributing to accelerating melting in Arctic sea ice.

Recently, the International Maritime Organization's (IMO) Marine Environment Protection Committee (MEPC) agreed to develop a work plan to address the impact of carbon emissions from ships and instructed the Sub-Committee on Bulk Liquids and Gases (BLG) to develop a definition for black carbon emissions from international shipping. The group is to consider measurement methods for BC and identify the most appropriate method for measuring black carbon emissions from international shipping; investigate appropriate control measures to reduce the impacts of black carbon emissions from international shipping in the Arctic and submit a final report to MEPC 65.

The work presented in this report describes the evaluation of the ocean going vessel APL England's main engine on a voyage from Oakland to San Pedro. One of the main focuses of this effort was the evaluation of several BC measurement methods to evaluate their performance limitations for marine applications. Impacts on other pollutants are also evaluated.

2 Objectives

The objectives for this work are to quantify the black carbon (BC) and other emissions from a marine main engine operating on marine gas oil (MGO) and a heavy fuel oil (HFO) on an ocean voyage from Oakland to San Pedro, CA. Emissions measurements included SO_x, THC_s, NO_x, CO₂, and PM, including PM_{2.5} mass (2.5 μm fine particles only), elemental, organic and BC. The real time or semi-real time BC measurement techniques evaluated in this program are based on flame ionization detection, photoacoustic measurements, and aerosol absorption.

3 Approach and Procedures

This section describes the test article (the main marine engine), measurement approach, and the real time instruments selected. This testing was conducted on a voyage from Oakland, CA to San Pedro, CA.

3.1 Test article and matrix

The engine tested was a 74640 BHP x 94 RPM Samsung-MAN B&W, model ML-0241 main propulsion engine; see Table 3-1 below. This is the main engine of the APL England, as shown in Figure 3-1. Appendix A provides additional information about the ship's particulars and engine specifications.

Table 3-1 Summary of selected main engine specifications

Description	Value	Units
Eng Disp	19467	liters
Max Power	55,659	kW
Safe Power	30222	kW



Figure 3-1 Samsung-MAN B&W main propulsion engine tested under voyage

Two fuels were used for this study, heavy fuel oil (HFO) and marine gas oil (MGO). The properties of these fuels are provided in Table 3-2 and in more detail in Appendix A. The test matrix points are provided in Table 3-3. A total of four load points were tested on each of the test fuels, including loads at 60%, 40%, 20%, and 5% (representing vessel speed reduction VSR). A log of the load points and tests conducted throughout the voyage is provided in Appendix B.

Table 3-2 Selected properties of tested fuel

Parameters	HFO	MGO	Units	Specification Limits
Density @ 15C	990.9	867.9	kg/m ³	991.0 Max
Viscosity @ 50C	130.2		cSt	380.0 Max
Sulfur	0.95	0.3	% (mass)	1.00 Max
Net Specific Energy	40.85		MJ/kg	

Table 3-3 Engine load and related information for the test setup

Mode #	Fuel	Speed knots	Power ³ kW	BSFC g/kWhr	Meas ¹ RPM	Load ⁴		Pres bar	Temp C	Exh flow ² Nm ³ /min
						% max	% recmd			
1-MGO	MGO	22.4	34299	149.7	79.7	62%	113%	1.6	41.0	3296
2-MGO	MGO	19.8	23082	171.0	72.0	41%	76%	1.1	37.0	2520
3-MGO	MGO	16.3	12293	212.4	59.5	22%	41%	0.5	35.0	1531
VSR-MGO	MGO	11.6	3043	310.9	40.7	5%	10%	0.2	37.0	869
1-HFO	HFO	21.7	33773	154.8	79.5	61%	112%	1.6	42.8	3270
2-HFO	HFO	19.5	24546	172.1	71.9	44%	81%	1.1	38.3	2449
3-HFO	HFO	16.6	11719	212.9	59.4	21%	39%	0.5	41.0	1498
VSR-HFO	HFO	11.6	2870	315.0	40.0	5%	9%	0.2	37.0	853

¹ According to the engine manual (report # ENG-00160) Samsung - MAN B&W the engine RPM ranges from 25 rpm (idle) to 94 RPM max. Rated power at 80, 72, and 60 RPM is slightly lower than measured by 10%

² Engine rated at max load of 74,640 BHP (55,659 BKW) at 94 RPM. The displacement is based on the diam. cylinder = 900 mm with a 2,550 mm stroke and 12 cylinders. Thus, the total displacement is $\pi \cdot 0.900^2 / 4 \cdot 2.550 \cdot 12 \cdot 1000 = 19466.8788$

The engine is boosted where intake P and T are the intake pressure and temperature for each point. VR is the volume recirculated which varies by RPM.

³ Engine load was recorded from real time on-line instruments in the engine room. These values were in slightly higher than published load data from the engine manual (report # ENG-00160 Samsung - MAN B&W) in the control room.

⁴ Percent engine load is based on maximum available power of 55,659 BKW as reported in the manuals and recommended safe operating power of 30222 kW (as recommended).

3.2 Sample system

The sampling approach follows UCR’s standard practice for on-vessel emissions testing seen in the schematic in Figure 3-2. The sampling approach is consistent with ISO 8178-1¹ and ISO 8178-2², which specify the measurement and evaluation methods for gaseous and particulate exhaust emissions when combined with combinations of engine load and speed provided in ISO 8178- Part 4: *Test cycles for different engine applications*. IMO ship pollution rules and measurement methods are contained in the “International Convention on the Prevention of

¹ International Standards Organization, ISO 8178-1, *Reciprocating internal combustion engines - Exhaust emission measurement -Part 1: Test-bed measurement of gaseous particulate exhaust emissions*, First edition 1996-08-15

² International Standards Organization, ISO 8178-2, *Reciprocating internal combustion engines - Exhaust emission measurement -Part 2: Measurement of gaseous and particulate exhaust emissions at site*, First edition 1996-08-15

Pollution from Ships”, known as MARPOL 73/78³, and sets limits on NO_x and SO_x emissions from ship exhausts. The intent of the UCR protocol is to conform as closely as practical to both the ISO and IMO standards.

UCR uses a close-coupled sample system where the transfer line has been eliminated to prevent PM losses during in-use ship testing. UCR has found significant PM losses for moderate length (5-10 feet) metallic heated transfer lines during ship testing. As such, UCR continues to approach all ship testing with direct sampling from the ship’s exhaust. The drawback with this approach is a longer setup time and more difficult testing layout since the equipment needs to be at the sample location. More details are provided on the transfer line in Appendix C.

A properly designed sampling system is essential to accurately collect a representative sample from the exhaust. ISO points out that PM must be collected in either a full flow or partial flow dilution system. UCR uses a partial flow dilution system with single venturi, as shown in Figure 3-2

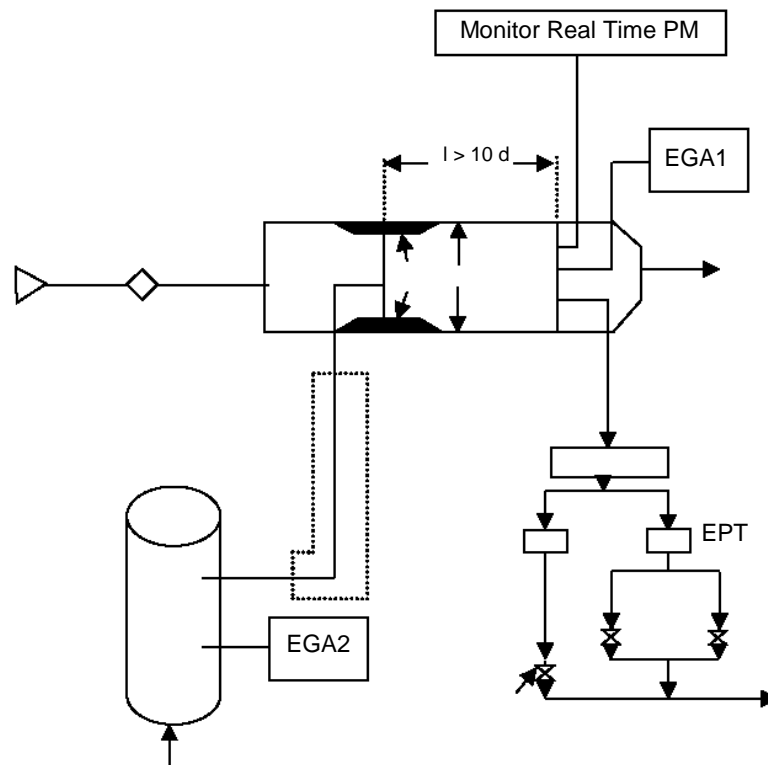


Figure 3-2 UCR’s sample system schematic (ISO method)

UCR’s sample system collects raw exhaust gas from the exhaust pipe (EP) through a sampling probe (SP) and the transfer tube (TT) to a dilution tunnel (DT) due to the negative pressure created by the venturi (VN) in the DT. The gas flow rate through TT depends on the momentum

³ International Maritime Organization, *Annex VI of MARPOL 73/78 “Regulations for the Prevention of Air Pollution from Ships and NOx Technical Code”*.

exchange at the venturi zone and is therefore affected by the absolute temperature of the gas at the exit of TT. Consequently, the exhaust split for a given tunnel flow rate is not constant, and the dilution ratio at low load is slightly lower than at high load. More detail on the key components is provided in Appendix C.

3.3 Gaseous measurements

Measurement of the concentration of the main gaseous constituents is one of the key activities in measuring emission factors. UCR utilized reference methods for the measurements of the gaseous species. These include the following:

- Heated flame ionization detector (HFID) for the measurement of hydrocarbons;
- Non-dispersive infrared analyzer (NDIR) for the measurement of carbon monoxide and carbon dioxide;
- Heated chemiluminescent detector (HCLD) or equivalent for measurement of nitrogen oxides;
- Paramagnetic detector (PMD) or equivalent for measurement of oxygen.

Table 3-4 Measurement species, detection method, and instrument reactions.

Component	Detector	Ranges
Nitrogen Oxides (NO_x)	Heated Chemiluminescence Detector (HCLD)	0-25, 50, 100, 250, 500, 1000, & 2500 ppmv
Carbon Monoxide (CO)	Non dispersive Infrared Absorption (NDIR)	0-200, 500, 1000, 2000, & 5000 ppmv
Carbon Dioxide (CO₂)	Non dispersive Infrared Absorption (NDIR)	0-5, 10, & 20 vol%
Sulfur Dioxide (SO₂)	Non dispersive Infrared Absorption (NDIR)	0-200, 500, 1000, & 3000 ppmv
Oxygen	Zirconium oxide sensor	0-5, 10, & 25 vol%

For SO₂, ISO recommends and UCR concurs that the concentration of SO₂ is calculated based on the fact that 95+% of the fuel sulfur is converted to SO₂. As such SO₂ is not measured, but will be reported as a calculation from the sulfur in the fuel.

3.4 PM measurements

Table 3-5 lists the PM measurements and instrumentation used for this program. These include batch samples for total PM_{2.5} from Whatman Teflo filters and PM composition utilizing Quartz fiber filters and the NIOSH and IMPROVE analysis method. Pall Gelman (Ann Arbor, MI) 47 mm Tissuquartz™ fiber filters were used to collect PM for EC and OC analysis. Each quartz filter was pretreated in a furnace at 600°C for 5 h. Each filter was stored in a separate sealed

petri dish at 10°C prior to and after sampling. A Sunset Labs (Forest Grove, OR) Thermal/Optical Carbon Aerosol Analyzer analyzed a 1.5 cm² quartz filter punch following the temperature program outlined in the NIOSH 5040 method (NIOSH, 1996)⁴. A second 1.5 cm² from the same filter was taken for analysis by the IMPROVE method. EC/OC was collected at 15 LPM onto precleaned (600 °C, 8 hours) 47mm diameter QAT Tissuquartz quartz fiber filters (Pall-Gelman, Ann Arbor, MI, USA). OC analysis was performed using an Thermal/Optical Carbon Aerosol Analyzer (Sunset Laboratory, Forest Grove, OR, USA) in the thermal-optical transmittance (TOT) mode following the National Institute for Occupational Safety and Health (NIOSH) Method 5040 protocol (Birch and Cary, 1996, NIOSH method 5040).

EC/OC was also analyzed using thermal-optical reflectance (TOR) following the DRI IMPROVE protocol on the Sunset Laboratories EC/OC analyzer modified to perform TOR correction and following the temperature program prescribed by the IMPROVE protocol (Chow, Watson, & Chen, 2015).

Table 3-5 PM Instruments and their measurement principles

Instrument	Model	Principle	Output	Wavelength
Micro Balance	UMX2	Gravimetric net weight change	Total PM _{2.5} measurement	n/a
Sunset Laboratory Carbon analyzer	Lab OC-EC	Flame Ionization Detection (FID)	Operationally defined organic and elemental carbon via transmittance	n/a
Micro Soot Sensor (MSS)	MSS 483	Photoacoustic (PA)	BC mass from real time in-situ signal (mg/m ³)	808 nm
Photoacoustic Extinctionmeter (PAX)	PAX ¹	Photoacoustic + scattering	Measure of light absorption and scattering from real time in-situ signal (ug/m ³ and mg/m ³)	870 nm

3.5 PM measurements: Black Carbon

There are several methods to measure black carbonaceous material and the reported terminology can be mixed. For example, the terms “black carbon, soot, elemental carbon, equivalent black carbon and refractory black carbon” refer to the light-absorbing component of particles, but the underlying definitions for these terms and associated measurement methods are different.⁵ Hence, BC measurements using different techniques are required to understand the nature and quantity of Black Carbon from emission sources. Published articles are available in the literature that describes the different BC measurement methods in detail.⁶

⁴ NIOSH (1996) NIOSH Manual of Analytical Methods. National Institute of Occupational Safety and Health, Cincinnati, OH.

⁵ Petzold, A., Ogren, J. A., Fiebig, M., Laj, P., Li, S.-M., Baltensperger, U., Holzer-Popp, T., Kinne, S., Pappalardo, G., Sugimoto, N., Wehrli, C., Wiedensohler, A., and Zhang, X.-Y. (2013) Recommendations for reporting "black carbon" measurements, Atmos. Chem. Phys., vol. 13, pp. 8365-8379, doi:10.5194/acp-13-8365-2013.

⁶ Moosmuller, H, Chakrabarty R, Arnott, W., (2009) Aerosol light absorption and its measurement: A review, Journal of Quantitative Spectroscopy and Radiative Transfer, vol. 110, pp 844-878.

Popular BC instrumentation methods use optical, thermal, or incandescence methods to estimate BC mass concentrations. BC emission inventories are mainly based on emission factors derived from thermal-optical methods that detect the carbon evolving from a heated filter sample, while data from atmospheric monitoring stations are mostly derived from optical absorption.⁷

For this project, real-time PM instruments were selected to characterize the black carbon emissions. These include the MSS 483 photoacoustic micro-soot sensor (BC_{PA-MSS}) manufactured by AVL and the photoacoustic extinction meter (BC_{PA-PAX}) by Droplet Measurement Technologies meter. Photoacoustic measurements use the intensity of a sound wave generated by the contraction and expansion of gas molecules when the BC particle is pulsed by a laser beam. The PA method has a large PM measurement range of the real time instruments and can measure up to $50,000 \mu\text{g}/\text{m}^3$, see Table 3-5. Other instruments require significant dilution of ship emissions to reach their desired BC concentration range (less than $100 \mu\text{g}/\text{m}^3$). The PA-soot method uses a conversion factor calibrated from gravimetric methods to derive the soot content. The PAX uses a photoacoustic meter in combination with light scattering. Thermo Scientific Multi-Angle Absorption Photometer (MAAP), and Magee Aethalometer (AE33) are other continuous filter based BC instruments that were not used for the testing on this voyage, but were used for a companion study on an auxiliary engine for the APL England.

In addition to mass concentration the PAX reports the single scattering albedo which is important to understand the absorption of black carbon. Scattering albedo is proportional to the absorption coefficient divided by the sum of the absorption coefficient and the scattering coefficient. These results will be presented in this report separately from the mass concentration.

⁷ Petzold, A., Ogren, J. A., Fiebig, M., Laj, P., Li, S.-M., Baltensperger, U., Holzer-Popp, T., Kinne, S., Pappalardo, G., Sugimoto, N., Wehrli, C., Wiedensohler, A., and Zhang, X.-Y. (2013) Recommendations for reporting "black carbon" measurements, *Atmos. Chem. Phys.*, vol. 13, pp. 8365-8379, doi:10.5194/acp-13-8365-2013.

4 Quality Control

This section describes UCR standard practices for calibrations, verifications, and control checks performed before, during and after testing. This section also describes checks performed to validate the data provided in the report. Additional instrument accuracy, precision, and standard verifications is provided in Appendix C.

4.1 Pre-test calibrations

Prior to departing from UCR all systems were cleaned and verified for the testing campaign. The MSS system included cleaning the internal pollution window and performing a span calibration using an internal pollution window. All systems were found to be within specifications and the systems were prepared for testing.

4.2 On-site calibrations

Pre- and post-test calibrations were performed on the gaseous analyzer using NIST traceable calibration bottles. Post-test dilution ratio was verified by removing the probe from the dilution tunnel and sampling from the raw exhaust. This method has been used in addition to operating two gas analyzers and has been shown to be reliable. Hourly zero checks were performed with each of the real time PM instruments. Leak checks were performed for the total PM_{2.5} system prior to each sample point.

4.3 Post-test and data validation

Post-test evaluation includes verifying consistent dilution ratios between points and verifying brake specific fuel consumption (BSFC) with reported manufacturer numbers. Typically, this involves corresponding with the engine manufacturer to discuss the results for emissions of interest. The brake specific fuel consumption results were with-in reason and thus suggest the load and mass of emissions measured are reasonable and representative. Thus, this suggests the data collected for the APL England main engine out emissions are accurate and representative of a properly functioning system.

5 Results

This section covers the emission results for the tests on the main engines on the APL England container vessel on the MGO and HFO fuels. The results are organized into four sections, 1) Gaseous and PM emissions, 2) PM_{2.5} and OC emissions, and 3) Black Carbon measurement techniques.

5.1 Real-time emissions

Figure 5-1, Figure 5-2, and Figure 5-3 show the real time PM-soot, NO_x and CO₂ emissions as a function of time for the main engine tested. The results of full voyage condition are presented by Figure 5-1. For the full voyage, the dilute NO_x concentration varied from 50 to 200 ppm and the dilute CO₂ concentration varied from 0.06% to 1.0% during dilute sampling. The PA-soot concentration varied from 0 mg/m³ to 0.8 mg/m³. The PAX PA-soot concentration also varied from 0 mg/m³ to 0.8 mg/m³, but it was slightly higher than MSS PA-soot concentration. The green stars represent when filter batch samples were collected for the Teflon and Quartz filter media. Typically three samples are collected, but due to limited time and vessel crew demands, some tests only utilized two filter samples. The large spikes in the gaseous emissions are a result of the dilution ratio quantification. The dilution ratio was checked after each batch PM filter sample by moving the dilute sample probe to the stack for raw sampling. The raw CO₂ concentrations reached 6% which resulted in spikes from less than 1% to ~6% in a short amount of time.

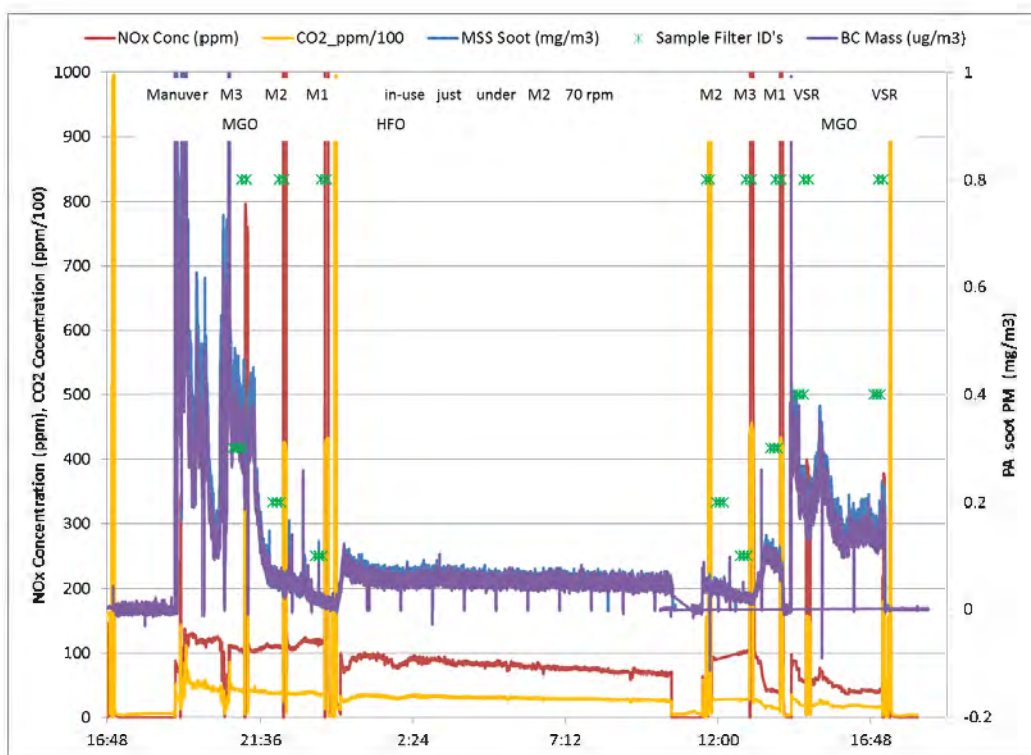


Figure 5-1 Real time emissions measurement during the full voyage

Figure 5-2 and Figure 5-3 show the time representing the departure from Oakland and arrival at San Pedro, respectively. BC emissions based on PA-soot and MSS soot concentrations for these specific parts of the voyage fluctuated a lot and were relatively higher than other time periods for the voyage. When the ship departed Oakland, BC showed a maximum concentration of 0.8 mg/m^3 , which was higher than for the remainder of the voyage. For gaseous emissions, NO_x and CO_2 concentrations, showed a series of peaks when the ship departed Oakland and arrived at San Pedro.

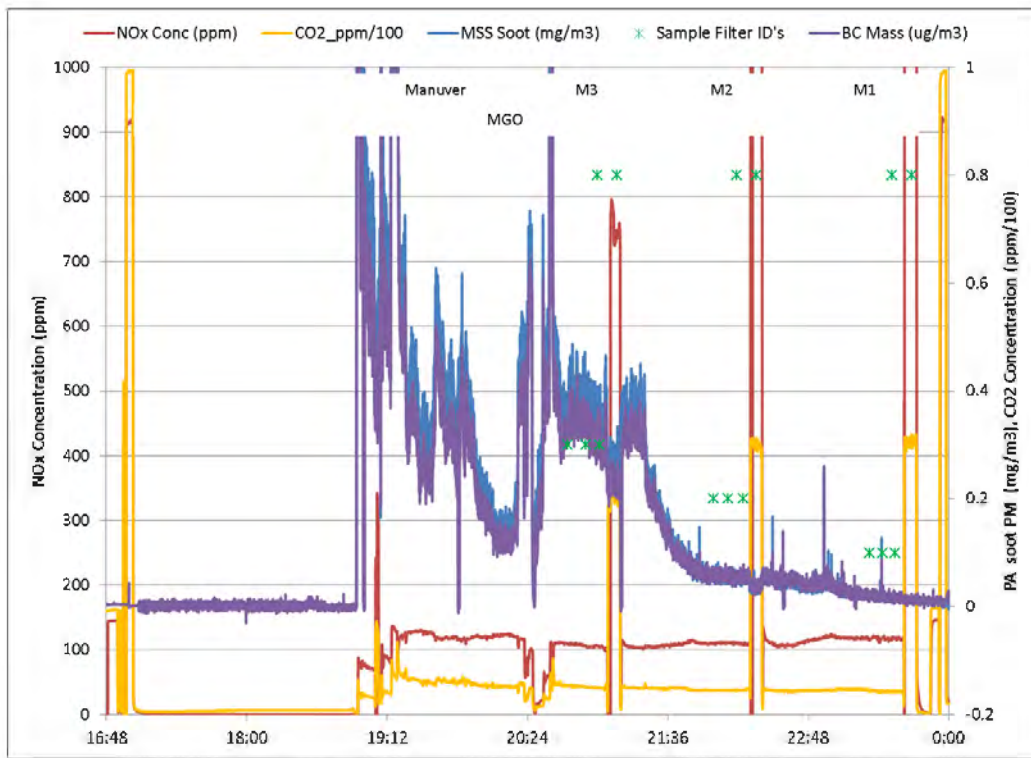


Figure 5-2 Real time emissions measurement when departing Oakland

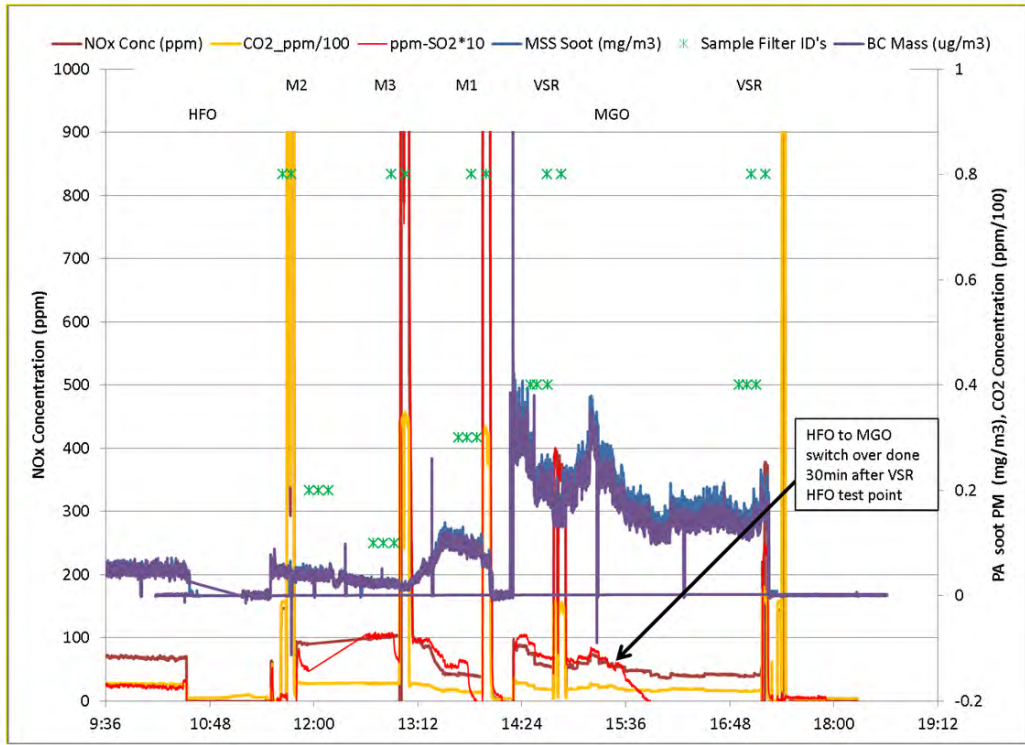


Figure 5-3 Real time emissions measurement when arriving San Pedro.

5.2 Gaseous emission factors

Table 5-1, Table 5-2, and Table 5-3 list the brake specific (bs), time specific, and fuel specific regulated emissions results, respectively. The load on the engine varied from just under 62% to approximately 5% for the MGO and HFO fuel test points. When the test load was higher than 20%, the brake specific NOx (bsNOx) emissions were relatively similar for the different loads. The bsNOx emissions for MGO fuel were slightly lower than those for the HFO fuel, as shown in Table 5-1, averaging 21.9 for the MGO fuel and 24.6 for the HFO fuel. The brake specific CO₂ (bsCO₂) emissions varied from 600 g/kWhr to 690 g/kWhr from high (62%) to low (20%) load. But when the test load was around 5%, brake specific CO₂ (bsCO₂) emissions were higher (~1000 g/kWhr) compared to the higher load test points. These values are consistent with expected bsCO₂ values and agree with typical published data, where higher brake specific fuel consumption exists at lighter loads. Additionally, the bsCO₂ emissions were similar for the MGO and HFO fuel, suggesting the two test conditions were similar. See Appendix D for a presentation of the measured concentration for all the reported test points.

Table 5-1 Brake specific gas-phase emission results (g/kWhr basis)

Mode #	Power kW	Load %	NOx g/kWh	CO g/kWh	CO ₂ g/kWh	DR n/a
1-MGO	34299	61.6%	21.65	0.29	607.9	15.9
2-MGO	23082	41.5%	19.20	0.30	611.5	13.4
3-MGO	12293	22.1%	18.86	1.01	693.1	12.0
VSR-MGO	3043	5.5%	27.89	1.74	1006.3	20.8
1-HFO	33773	60.7%	23.59	0.51	595.9	20.0
2-HFO	24546	44.1%	21.94	0.57	606.6	21.2
3-HFO	11719	21.1%	20.71	1.89	671.7	33.0
VSR-HFO	2870	5.2%	32.22	2.06	992.5	16.0

¹Engine load was recorded from real time on-line instruments in the engine room. These values were in good agreement with published load

Table 5-2 Time specific gas-phase emission results (g/hr basis)

Mode #	Power kW	Load %	NOx kg/hr	CO kg/hr	CO ₂ kg/hr	DR n/a
1-MGO	34299	61.6%	742.55	9.95	20851	15.9
2-MGO	23082	41.5%	443.16	7.01	14115	13.4
3-MGO	12293	22.1%	231.86	12.45	8520	12.0
VSR-MGO	3043	5.5%	84.87	5.29	3062	20.8
1-HFO	33773	60.7%	796.87	17.09	20127	20.0
2-HFO	24546	44.1%	538.62	14.09	14891	21.2
3-HFO	11719	21.1%	242.72	22.17	7872	33.0
VSR-HFO	2870	5.2%	92.48	5.93	2848	16.0

Table 5-3 Fuel specific gas-phase emission results (g/kg fuel basis)

Mode	Power	Load	NOx	CO	CO2	DR
#	kW	%	g/kg fuel	g/kg fuel	g/kg fuel	n/a
1-MGO	34299	61.6%	116.04	1.55	3258.4	15.9
2-MGO	23082	41.5%	100.14	1.58	3189.5	13.4
3-MGO	12293	22.1%	89.77	4.82	3298.8	12.0
VSR-MGO	3043	5.5%	91.41	5.70	3297.7	20.8
1-HFO	33773	60.7%	126.37	2.71	3191.8	20.0
2-HFO	24546	44.1%	115.01	3.01	3179.5	21.2
3-HFO	11719	21.1%	98.28	8.98	3187.5	33.0
VSR-HFO	2870	5.2%	105.48	6.76	3249.0	16.0

5.2.1 CO₂

Figure 5-4 plots the bsCO₂ emissions versus load for the MGO and the HFO. The HFO appears to use slightly less fuel as would be expected due to the higher energy content of HFO compared to MGO. The overlap of the CO₂ standard deviations for MGO and HFO at each load, indicate there is no statistically significant difference in the bsCO₂ emissions for MGO and HFO at any of the tested engine load points.

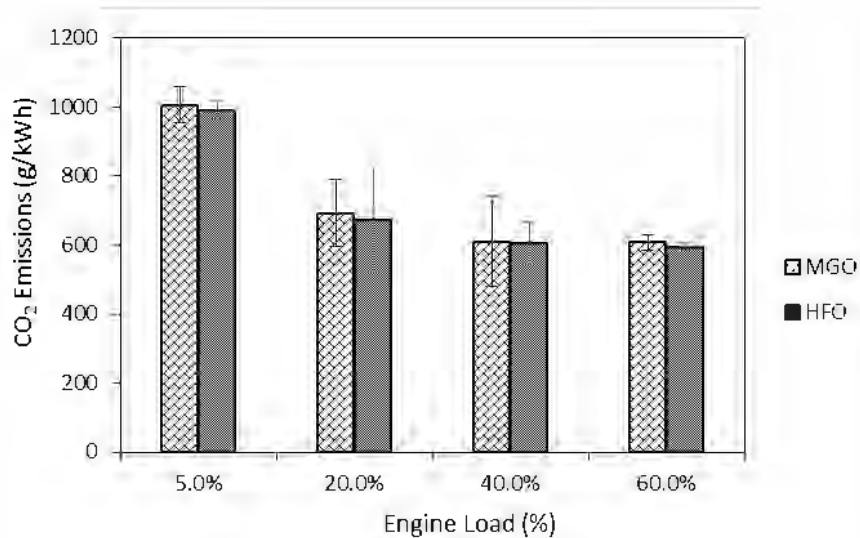


Figure 5-4 Brake Specific CO₂ Emissions versus Engine Load

5.2.2 NO_x

Figure 5-5 plots the bsNO_x emissions versus load for the MGO and the HFO. The NO_x emissions are higher from the HFO than from the MGO at all engine load points, but only the differences at the 5% and 40% engine load points are statistically different.

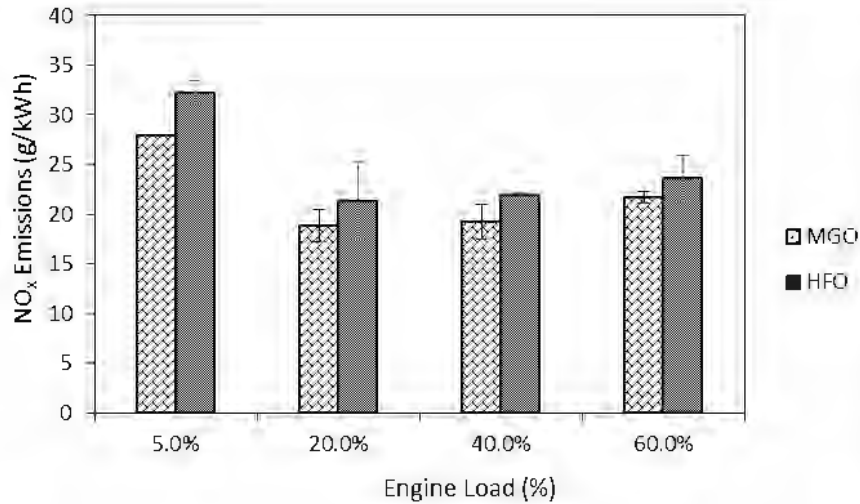


Figure 5-5 Brake Specific NO_x Emissions versus Engine Load

5.2.3 CO

Figure 5-6 plots the bsCO emissions versus load for the MGO and the HFO. As was the case for the bsNO_x the bsCO emissions are higher from the HFO than from the MGO. The differences are statistically different at the 5% engine load point and may be marginally statistically different at the 60% engine load point.

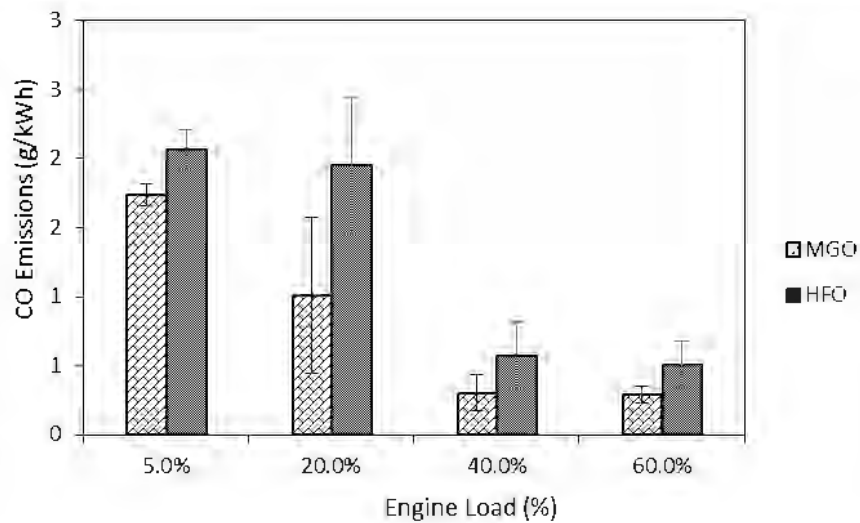


Figure 5-6 Brake Specific CO Emissions versus Engine Load

5.3 PM emission factors

Table 5-4 shows the brake specific emissions of PM_{2.5}, BC_{PA-MSS}, BC_{PA-PAX}, and EC/OC by NIOSH and IMPROVE for both the MGO and HFO fuels. The engine out bsPM_{2.5} ranged from 207 mg/kWhr to 501 mg/kWhr over the range of loads and test fuels tested. The higher bsPM_{2.5} emissions at lighter loads for the HFO agrees with typical diesel engine emission rates. However,

the lowest bsPM_{2.5} at the lightest load for the MGO is atypical of diesel engine emission rates. The bsPM_{2.5} emissions were lower for the MGO fuel than the HFO fuel for all test points, except the 20% load. The HFO fuel showed increases of 56%, 42%, -24%, and 143% compared to the MGO fuel for the 60%, 40%, 20%, and 5% loads, respectively.

The organic PM from the MGO was essentially the same for the NIOSH and the IMPROVE method except at the lightest load. The IMPROVE OC was 5%, -8%, -2%, and 12% higher than the NIOSH OC for the MGO at the engine load points of 60%, 40%, 20% and 5%, respectively, see Table 5-4. The IMPROVE OC was 144%, 30%, 11%, and -4% higher than the NIOSH OC for the HFO at the engine load points of 60%, 40%, 20% and 5%, respectively, see Table 5-4. The IMPROVE EC emissions were higher than the NIOSH EC emissions at all load points for both fuels, ranging from 25% to 165% higher for the MGO and from 71% to 1140% from HFO.

Marine engines running on high sulfur fuels tend to have a PM composition dominated by OC and Sulfate. The EC+OC masses were approximately 50% of the total PM_{2.5}. Although sulfate PM was not measured, one can infer the sulfate PM from the measurement of SO₂ in the exhaust. It is expected that the difference in the PM_{2.5} and the sum of OC and EC (including mass factors) is approximately the sulfate PM mass. The total PM composition is thus, approximately 45% sulfate and 45% OC, with a small fraction of EC (10%).

Table 5-4 Total PM, BC, and EC/OC brake specific PM emissions

Mode #	Power kW	Load %	PM _{2.5} mg/kWh	BC _{PA-MSS} ¹ mg/kWh	BC _{PAPAX} ² mg/kWh	NIOSH		IMPROVE	
						EC mg/kWh	OC mg/kWh	EC mg/kWh	OC mg/kWh
VSR-MGO	3043	5.5%	206.5	53.0	46.5	39.81	290.5	61.47	325.2
3-MGO	12293	22.1%	424.9	32.5	29.2	32.26	258.2	40.20	253.6
2-MGO	23082	41.5%	277.9	4.7	4.9	2.56	143.2	5.59	131.8
1-MGO	34299	61.6%	251.4	1.4	1.7	1.21	171.1	3.21	179.8
VSR-HFO	2870	5.2%	501.3	64.6	61.3	39.64	255.0	67.72	243.9
3-HFO	11719	21.1%	323.1	24.1	23.1	17.45	156.9	33.30	173.5
2-HFO	24546	44.1%	393.9	5.6	5.0	1.10	145.5	13.69	188.9
1-HFO	33773	60.7%	392.8	3.1	2.6	2.47	148.9	8.13	363.1

¹ **PA MSS** is the AVL 483 micro soot sensor that measures the acoustic properties of PM or black carbon. The laser wavelength was xx nm. PA MSS showed a zero offset of -0.016 mg/m³ during 1-MGO (tested after 1 and 2). Adjustment provided in post processing with a sliding scale from 22:30 to 24:00. All other PA MSS data was corrected with on-site zero adjustments. ² **PA PAX** is the PAX photo acoustic instrument for black carbon in units of mg/m³. The laser wavelength was xx nm. PA PAX instrument designed for frequent zero adjustments (1 per hour). Hourly adjustments recommended for the PA type instruments. ³ **ScatPAX** is the PAX single absorption scattering coefficient. This is a unit less ratio of absorption over scattering+absorption ⁴ **ISO-8178-1** 2006 Chapter 12.4 "Adjustment of the dilution ratio": Specification is minimum of 4 to 1.

5.3.1 PM_{2.5}

Figure 5-7 plots the bsPM_{2.5} emissions from both fuels. The differences between the bsPM_{2.5} noted above are statistically significant based on the non-overlap of the standard deviations for each fuel at each load point.

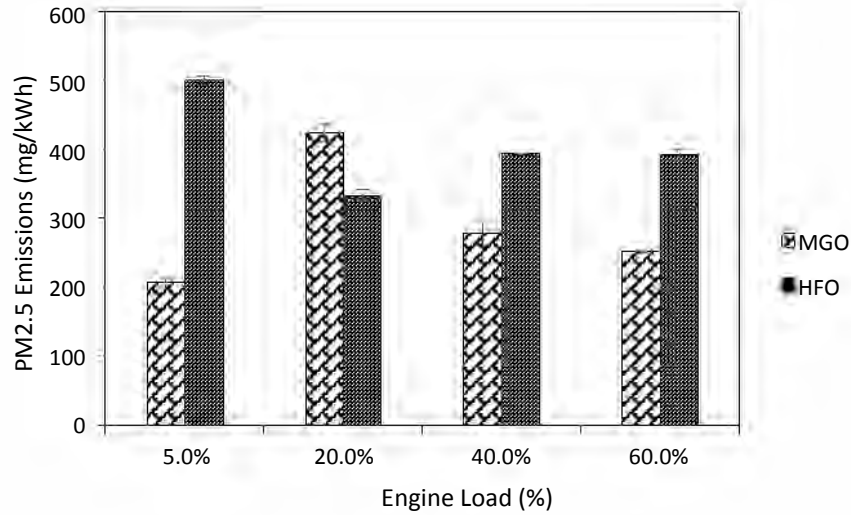


Figure 5-7 PM2.5 as a Function of Engine Load for MGO and HFO

5.3.2 BC

Figure 5-8 plots BC emissions for both fuels and two test methods for all engine load points. Both methods give essentially the same bsBC at all engine load points for the same fuel. At the 5% engine load point both methods show higher bsBC emissions for the HFO fuel while at all other load points the bsBC are statistically the same for both fuels.

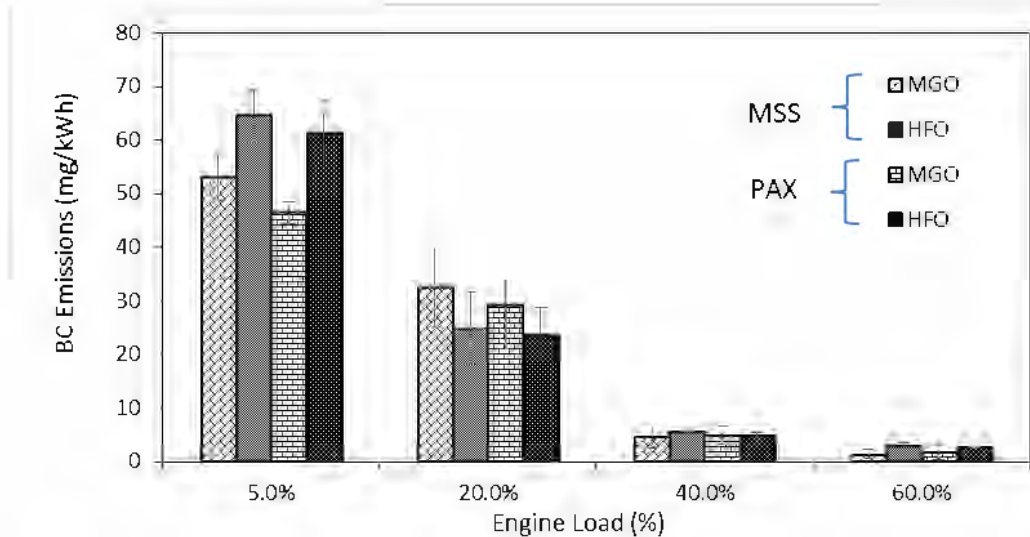


Figure 5-8 BC by two methods as a Function of Engine Load for MGO and HFO

5.3.3 EC

Figure 5-9 plots the bsEC emissions for both fuels by the IMPROVE and the NIOSH methods. The results are mixed. At the 5% load the NIOSH method gives much lower results for both fuels and only the IMPROVE method gives higher emissions for the HFO versus the MGO. At the 20% load the NIOSH method gives lower results for both fuels but only much lower for the

HFO. Both the IMPROVE and the NIOSH show higher emissions for the MGO versus the HFO at the 20% load. At the 40% load the NIOSH shows lower emissions than the IMPROVE for both fuels but for MGO the differences are not statistically different. The IMPROVE has statistically higher emissions for the HFO versus the MGO at the 40% load while the NIOSH has higher emissions from the MGO than from the HFO but the difference is not statistically significant. At the 60% load the NIOSH method results are lower than the IMPROVE results for both fuels and both methods have higher emissions for the HFO versus the MGO.

Since the samples for the EC/OC analysis were taken from the same filter these results indicate that the calculated μg of EC on the filter is dependent upon the method used to measure it.

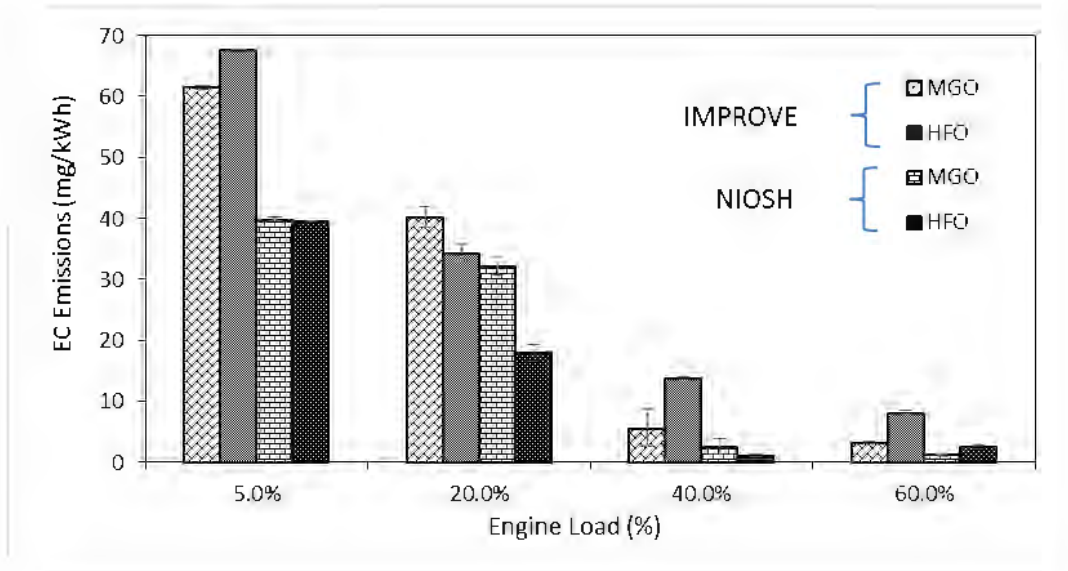


Figure 5-9 EC by two methods as a Function of Engine Load for MGO and HFO

Figure 5-10 plots the $\text{bsBC}_{\text{PA-MSS}}$ versus the IMPROVE bsEC for both fuels and there is a very high correlation between the methods for both fuels. However, while there is nearly a 1 to 1 correlation for the HFO fuel the $\text{bsBC}_{\text{PA-MSS}}$ results are ~13% lower than the bsEC . As is seen in Table 5-5, while there is variation in the slopes and intercepts for all other possible correlations, they all have very high correlation coefficients.

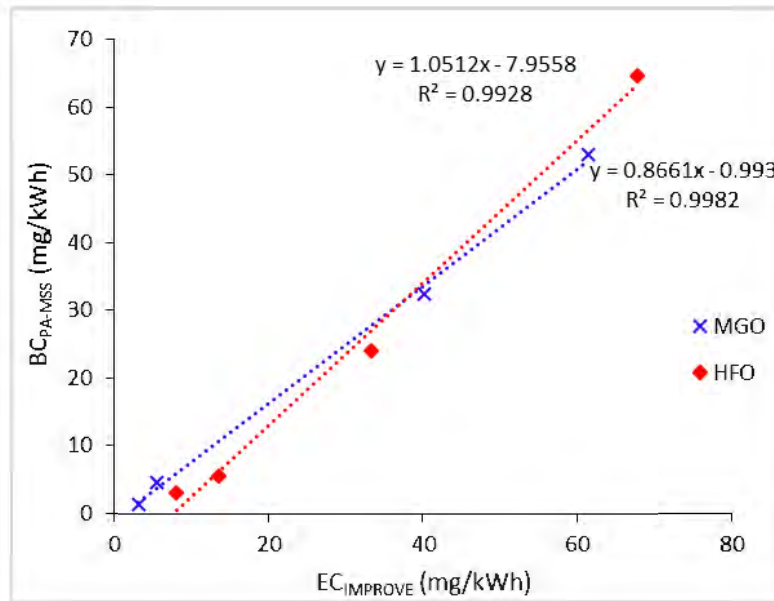


Figure 5-10 Black Carbon by PA-MSS versus EC by IMPROVE

Table 5-5 BC Photoacoustic Correlation Equations with EC by Improve and NIOSH Methods

BC Method	EC Method	Fuel	y =	R ²
PA-MSS	IMPROVE	MGO	0.8661x - 0.993	0.9982
		HFO	1.0512x - 7.9558	0.9928
PA-MSS	NIOSH	MGO	1.2008x + 0.1575	0.9619
		HFO	1.576x + 0.4264	0.9874
PA-PAX	IMPROVE	MGO	0.7504x - 0.1462	0.9985
		HFO	1.0033x - 7.8064	0.9937
PA-PAX	NIOSH	MGO	1.0419x + 0.8228	0.9649
		HFO	1.50456x + 0.189	0.9887

5.3.4 OC

Figure 5-11 plots the OC emissions versus engine load for both fuels by the IMPROVE and NIOSH methods. Because OC depends on the dilution ratio, which wasn't constant for all tests, (see chapter 5), quantitative comparisons of these emissions are not valid.

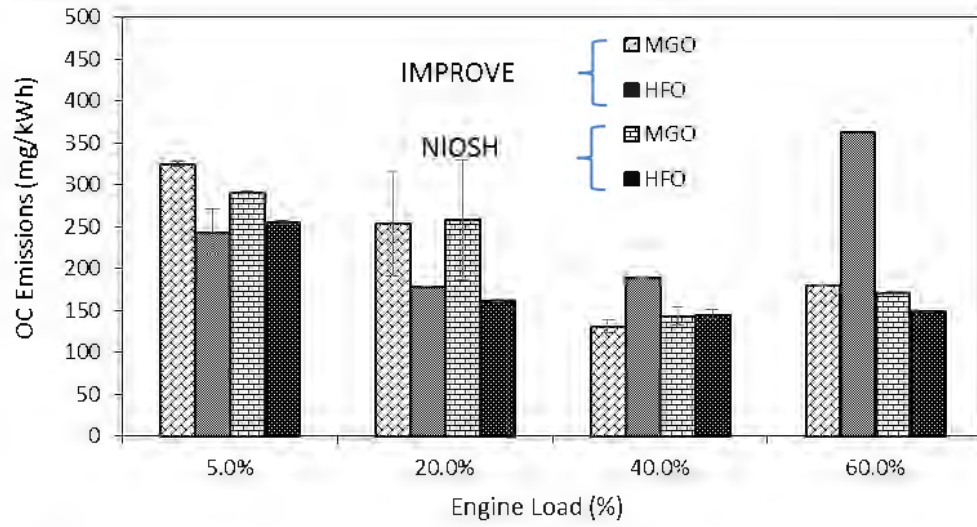


Figure 5-11 OC by two methods as a Function of Engine Load for MGO and HFO

6 Data qualification

The primary purpose of the research was real time measurement of black carbon emissions. During previous testing with real time BC measurement systems data was lost because the concentration exceeded the range of the instruments. To prevent signal range issues in this testing program, a smaller sample orifice tube for the UCR ISO dilution system was used to increase the sample dilution (see Appendix C for details). During testing the dilution ratio (DR) increased from sample #1 through sample #6 where the system was identified as needing cleaning, see Figure 6-1. After cleaning the sample probe, the dilution factor was reduced from 33 to about 15. Figure 6-2 shows the dilution factor for the fuels and test points. With the exception of the 5% load point (VSR), the MGO was tested at a dilution factor of ~15 while the HFO fuel was tested at a dilution factor of ~20 and a dilution factor of ~34 at the 20% load.

Others have shown that changes in dilution factor can have an impact on PM formation for volatile organics PM but not necessarily black carbon or elemental carbon PM⁸. These studies, though, were with low sulfur on-road fuels and medium speed diesel 4-stroke heavy duty engines so direct comparisons may not be accurate, but the trends should be similar. Some marine HFO fuel DR studies have been performed but there is speculation with their results and are thus not referenced here. In general experience suggests maintaining a similar dilution ratio between pre and post-tests A/B comparisons is important and recommended during all vessel testing. The fact that the DR changed was unfortunate, and it is unclear if the change in dilution ratio would have affected the black carbon measurements, which is not expected. It is expected there would be an impact in the total and organic PM emission results.

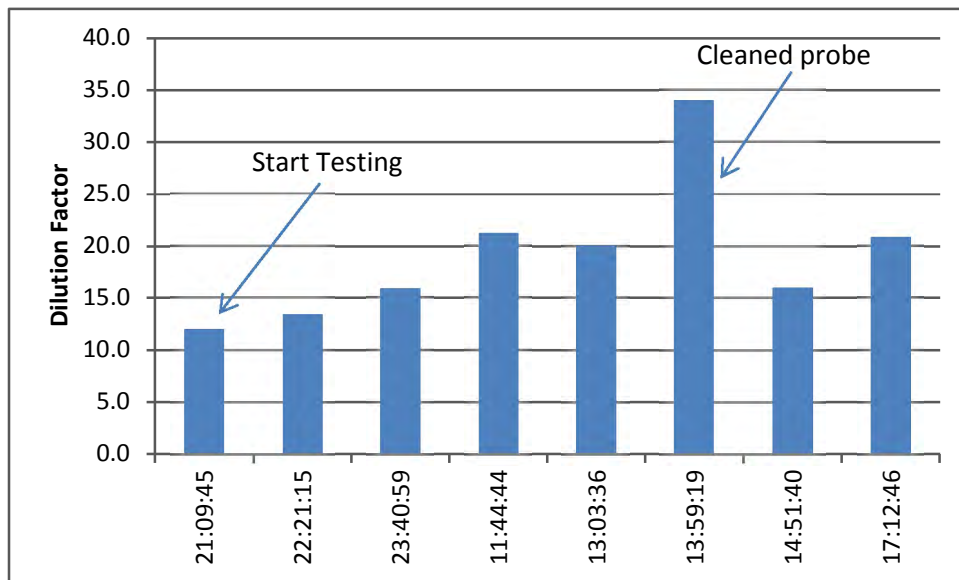


Figure 6-1 Dilution ratio organized chronologically for the APL main engine testing

⁸ Khalek, I., "Diesel Particulate Measurement Research. 2007, SwRI. p. Project E-66-Phase 1, 2, and 3," 2007.

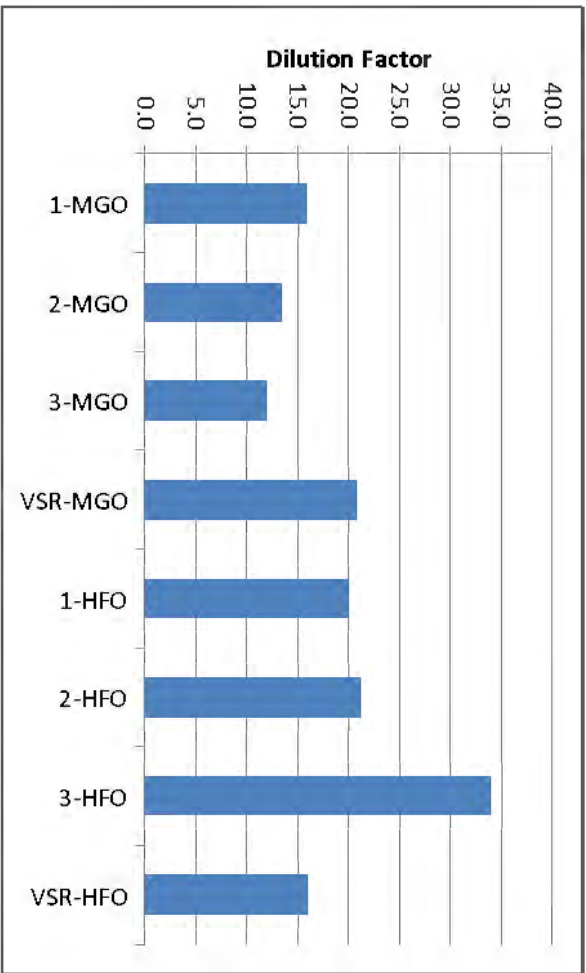


Figure 6-2 Dilution ratio organized by test mode for the APL main engine testing

7 Summary and Conclusions

Emissions from a marine main engine operating on marine gas oil (MGO) and a heavy fuel oil (HFO) were measured on an ocean voyage from Oakland, CA to San Pedro, CA. Emissions measurements included SO_x, THC_s, NO_x, CO₂, and PM, including PM_{2.5} mass (2.5 μm fine particles only), elemental, organic and BC. BC was measured by two photoacoustic instruments and by extraction from a quartz filter followed by FID determination by the NIOSH and IMPROVE methods.

The overall results and conclusions from this work can be summarized in the following highlights:

- Brake specific NO_x (bsNO_x) emissions were relatively similar for the different loads for test loads higher than 20%. The bsNO_x emissions for MGO fuel were slightly lower than those for the HFO fuel, averaging 21.9 for the MGO fuel and 24.6 for the HFO fuel.
- Brake specific CO₂ (bsCO₂) emissions varied from ~600 g/kWhr to ~690 g/kWhr from high (62%) to low (20%) load. But when the test load was around 5%, brake specific CO₂ (bsCO₂) emissions were ~1000 g/kWh. These values are consistent with expected bsCO₂ values and agree with typical published data, where higher brake specific fuel consumption exists at lighter loads. Additionally, the bsCO₂ emissions were similar for the MGO and HFO fuel, suggesting the two test conditions were similar.
- BC measurements and the existing ISO 8178 sampling procedures work well for in-situ Photoacoustic type meters.
- For the same fuel at the same engine load, the two photoacoustic meters give essentially the same results.
- At the 5% engine load point both photoacoustic meters showed higher BC emissions for the HFO fuel while at all other load points the BC emissions are statistically the same for both fuels.
- EC as determined by the NIOSH and IMPROVE method correlated well with the BC measurements for both fuels. The R-squared correlation coefficients are between 0.9619 and 0.9937 for correlations of bsBC with bsEC. The slopes of the equations vary between 0.7504 and 1.576 depending upon the BC and EC method and the fuel.
- The differences in the bsEC between the NIOSH and IMPROVE method is related to the method and not the sample since the samples were taken from the same quartz filter.

Appendix A. Ship and Fuel Specifications

This Appendix contains records utilized during the testing of the ocean going vessel. These records were either obtained from the engine control room manuals (engine related details) or from the captain's fuel records during bunkering. Selected items were greyed out for each vessel manufacturer. Only the pertinent information relating to the testing was provided.

```

Barge : FDH 35-3
Sample Grade : RMG380LS
Sample Seal No : 0495018 - Sealed
Bunker Quantity : 453.016 MT
Bunker Density @15C : 990.2 kg/m3
Bunker Viscosity @50C : 127.0 cSt
Sulphur Content : 0.92 %
Water Content : 0.20 %
Source of the sample : MANIFOLD
Sampling Method : DRIP
    
```

Figure A-1 HFO analysis reference

120.000 Tonnes Marine Diesel Oil / Gas Oil

We undertake to supply you with the above grade(s) of bunkers. Some basic characteristics of the bunkers are as follows.

Product Name	Product was blended on board in advance? (Yes / No)	Kinematic Viscosity @ 40° or 50° C, mm ² /s ISO 3104	Density @15° C, kg/m ³ ISO 3675 ISO 12185	Water Content % V/V ISO 3733	Flash Point °C ISO 2719	Sulphur Content %, m/m ISO 14596 ISO 8754
MGO	No	3.5	0.8679	TRACE	76	0.300

We will supply MGO first, followed by . The approximate delivery temperature is °C.

The rated pumping capacity of our bunker tanker is 30-100 tonnes per hour.

Figure A-2 MGO analysis reference

Port of Registry	: Singapore
Official Number	: 389193
IMO Number	: 9218650
Call Sign	: 9VDD2
Satcom B Telex & Phone	: Tlx:356372270; Ph:356372250 (Bridge) ;356372251 (Master)
Satcom B FAX	: 356372260
Satcom C	: 456372240
MMSI	: 563722000
Owner	: APL (Bermuda) Ltd
Operator	: APL Co.Pte Ltd
Manager	: Neptune Shipmanagement Services Pte.Ltd. Singapore
P & I Club	: Britannia Steamship Insurance Association Ltd.
Date of Build	: 04 Sept 2000
Delivery Date	: 20 Feb 2001
Hull No	: 1335
Bulder	: SAMSUNG HEAVY INDUSTRIES - KOREA
Class	: DNV +A1 Container Carrier EO LCS(SI) Nauticus
LSA Capacity	: 30 Persons
LOA/LDP	: 277.255m/ 263.00m
Bridge to Bow/Stern	: Bridge to stem=188.84m, Bridge to stern=88.16m
Breadth Moulded	: 40.00m
Height	: 59.99 (From Keel to Highest Point)
Moulded Depth	: 24.30m
Summer Freeboard	: 3639mm
Summer Draft (Fb)	: 14.026m
Design Draft	: 12.50m
Summer Displacement	: 92439.2mt
Summer TPC	: 91.25mt/cm
GRT / NRT	: 65792 / 35494
Suez GRT/NRT	: 69673.55 / 57339.77
Lightship / FWA	: 24,452.6 , FWA=256
Displacement (Summer)	: 92439.2 mt
Deadweight (Summer)	: 67986.6 mt
Container Capacity (6Tier)	: Below Deck - 2602 TEU, On Deck - 2904 TEU Total - 5506
Reefer Capacity	: 650 units @ 440V
Ballast Capacity	: 14693.6 m ³
Heavy Fuel Oil Capacity	: 9030.0 m ³
Diesel Oil Capacity	: 449.0 m ³
Fresh Water Capacity	: 526.4 m ³
Main Engine	: SAMSUNG B&W 12K 90MC
Main Engine Rating MCR	: 55659kW / 74640 BHP @ 94 RPM
Main Engine Rating NCR	: 67180 BHP @ 90.8RPM
Service Speed	: 25.00 Kts.
Generators	: 3 x 2900 kW (3625 kVA) - B&W 7L32/40 x 720 rpm
Shaft Generator	: 2100KW ABT 80-104% of ME rpm
Emergency Generator	: 1 x 150kW (188 kVA)
Bow Thrusters	: 2000kW
Propeller	: 1 x Right Handed 6 Bladed 8450mm Diameter

Updated 07th March 2007

Figure A-3 Ship particulars and specifications

2012/10/15

PARTICULARS OF ENGINE & EQUIPMENT

ENGINE TYPE	SAMSUNG – MAN B&W
ENGINE NO.	ML – 0241
NUMBER OF CYLINDER	12
DIAMETER OF CYLINDERS	900 mm
STROKE	2,550 mm
OUTPUT (M.C.R)	74,640 BHP x 94 RPM
FIRING ORDER	
ROTATING DIRECTION	
CYLINDER CONSTANT	3.6761

PARTICULAR OF DYNAMOMETER

MAKER AND TYPE	
COEFFICIENT	
MAXIMUM CAPACITY	

SPECIFICATION OF OIL USED AT SHP TRIAL

	FUEL OIL	SYSTEM OIL	CYL. OIL	T/C OIL	CAM OIL
KIND OF OIL	BUNKER-A				
SPEC. GRAVITY 15/4 °C	0.9012				
FLASH POINT °C	82				
VISCOSITY(CST) 40 °C	15				
RE. CARBON(WT%)	2.6				
ASH (WT%)	0.002				
WATER (WT%)	0.22				
SULPHUR (WT%)	0.4				
NET CAL. VALUE Kcal/kg	10,031				

Figure A-4 Engine information

SUMMARY DATA OF LOAD												
ENGINE TYPE	12K90MC	ENG.NO	ML-0241	DATE			Oct.27.2000		WEATHER		FINE	
DATA SHEET NO			01	02	03	04	05	06	07	08	09	
LOAD (%)			25%	50%	75%	90%	100%(1)	100%(2)	110%			
TIME			10:50	11:55	13:10	14:10	14:50	15:15	15:55			
SPEED(RPM)			59.2	74.6	85.4	90.8	94.0	94.0	97.0			
BHP(PS)			18660	37320	55980	67180	74640	74640	82104			
IHP(PS)												
MECH.EFF. (%)												
PI (bar)												
FUEL	MEASURED	135.92	129.79	127.27	128.29	131.23	131.56	135.24				
CONSUMP	CORRECTED											
g / BHP.H												
Pmax.(bar)												
Pcomp.(bar)												
PUMP INDEX												
V.I.T INDEX												
EXH.GAS	CYL. OUT	290.6	281.7	287.9	309.2	331.7	334.2	364.2				
TEMP	BEF. T / C	350.0	360.0	375.0	402.5	435.0	440.0	480.0				
	AFT. T / C	298.8	265.0	242.5	243.8	255.0	258.8	277.5				
T/C	NO.1											
SPEED	NO.2											
	(rpm)	NO.3										
	NO.4											
AUX. BLOWER (on / off)			ON	OFF	OFF	OFF	OFF	OFF	OFF			
SCAV. AIR TEMP.(°C)			23	24	32	36	40	41	43			
TEST	°C	21.5	23.5	25.0	24.8	25.5	25.5	26.0				
ROOM	mmHg	767.3	767.3	767.3	767.3	767.3	767.3	768.0				

NOTE : THE FUEL OIL CONSUMPTION IS CORRECTED TO ISO CONDITION
F.O. LOWER CALORIFIC VALUE : 10031kcal/kg

Figure A-4 Engine specifications relevant to emissions testing

Appendix B. Test logs

This Appendix summarizes the details of the test logs utilized for the analysis. Table B-1 shows the engine details measured from the vessels engine control room. Table B-2 shows the sample log for the Teflon and Quartz filters sampled as part of the PM measurement systems. The footnotes provide context for special observations with each log record.

Table B-1 Log samples of engine load, fuel rate, BSFC, and exhaust flow parameters

Time	Fuel Rate kg/hr	Torque kN*m	BSFC g/kWhr	Engine		Fixed ¹ RPM	Meas Spd knots	Fuel	Operation
				Intake P_bar	Intake T_C				
19:52	3047	2390	198	0.5	34	62.1		MGO	Leave Port
19:54	3223	2402	197.8	0.5	34	63.7		MGO	Leave Port
19:58	3374	2409	196.9	0.7	34	65		MGO	Leave Port
20:09	3573	2427	195.9	0.82	34	64		MGO	Leave Port
20:18	2260	2374	200	0.4	34	57		MGO	Leave Port
20:21	2702	2378	220	0.41	34	57.7		MGO	Leave Port
20:22	1155	2294	313	0.2	34	42		MGO	Leave Port
20:26	1100	2200	300	0.20	34	45		MGO	Leave Port
20:27	100	2200	450	0.10	34	28		MGO	Leave Port
20:34	800	200	350	0.01	34	34		MGO	Leave Port
20:36	1100	2200	300	0.20	34	45		MGO	Leave Port
20:36	2260	2374	200	0.4	34	58		MGO	Leave Port
20:37	2260	2374	200	0.4	34	58		MGO	set RPM
20:40				0.4	35	60	16.3	MGO	mode 3
20:44	2712	2375	220	0.4	35	59.6		MGO	mode 3
20:46	2580	2370	210	0.4	35	59.5		MGO	mode 3
20:48	2618	2364	210	0.4	35	59.9		MGO	mode 3
20:50	2550	2371	209	0.4	35	59.4		MGO	mode 3
20:53	2600	2366	210	0.4	35	59.4		MGO	mode 3
20:55	2630	2379	210	0.4	35	59		MGO	mode 3
20:56	2611	2370	220	0.4	35	58.9		MGO	mode 3
20:56			210	0.4	35	60		MGO	mode 3
21:23	2611	2370	210	0.4	35	60		MGO	set new RPM
21:52	3944	2426	171	1	37	72	19.8	MGO	mode 2
22:09	3967	2423	171	1	37	72		MGO	mode 2
22:41	3930		171	1	37	72		MGO	mode 2
22:41	3930	2423	171	1	37	72		MGO	set new RPM
23:13	5120	2471	150	1.6	41	80	22.4	MGO	mode 1
23:34	5100	2475	149.5	1.6	41	79.1		MGO	mode 1
23:50	5180	2470	149.5	1.6	41	79.9		MGO	mode 1
23:51	5180	2470	149.5	1.6	41	79.9		MGO	set fuel and RPM
0:20				0.8	36	70		HFO	in-use
8:32	3550		181	0.8	37	69		HFO	
11:32	3325		190.5	0.71	37	67		HFO	set new RPM
11:46	3941	2418	180.4	0.92	38	70.4		HFO	setting new RPM
11:50	4313	2441	172	1.05	38	71.9	19.1	HFO	mode 2
12:02	4054	2436	172.1	1.02	38	72	19.6	HFO	mode 2
12:15	4306	2443	172.2	1.08	39	71.8	19.9	HFO	set new RPM
12:40	5298	2473	155.9	1.5	42	79.4	21.7	HFO	mode 1
12:47	5256	2474	156.6	1.5	43	79.2	21.7	HFO	mode 1
12:56		2483	156.1	1.55	43	79.6		HFO	mode 1
13:04	5365	2479	150.4	1.57	43	79.6	21.6	HFO	set new RPM
13:36	2510	2360	212.9	0.4	41	59.5	16.2	HFO	mode 3
13:42	2454	2362	212.8	0.5	41	59.3		HFO	mode 3
13:47	2478	2370	212.7	0.5	41	59.5	16.8	HFO	mode 3
13:59	2539	2364	213.3	0.5	41	59.3	16.8	HFO	set VSR speed
14:21	904	2287	315	0.15	40	40	11.6	HFO	set VSR speed
17:08	946	2294	310.9	0.15	37	40.7	11.6	MGO	mode VSR

¹ According to the engine manual (report # ENG-00160) Samsung - MAN B&W the engine RPM ranges from 25 rpm (idle) to 94 RPM max

Table B2 Filter log and summary weights

Table Filter media and flows for each test point sampled							PM total mg/filter	Anatlical NIOSH		IMPROVE		¹ Teflo V m3	¹ Quartz V m3	PM total mg/m3	NIOSH		IMPROVE		
Test Day	Mode	Fuel	Start	End	Teflon ID	Quartz ID		PM EC mg/filter	PM OC mg/filter	PM EC mg/filter	PM OC mg/filter				PM EC mg/m3	PM OC mg/m3	PM EC mg/m3	PM OC mg/m3	
1	15	3	MGO	20:44:30	20:47:30	WT130272	QU201310_01	0.300	0.024	0.182	0.028	0.167	0.049	0.045	6.13	0.53	4.07	0.63	3.75
2	15	3	MGO	20:54:00	20:57:00	WT130271	QU201310_02	0.222	0.014	0.113	0.018	0.114	0.049	0.045	4.55	0.32	2.53	0.40	2.55
3	15	3	MGO	21:01:00	21:04:00	WT130274	QU201310_03	0.221	0.015	0.123	0.019	0.129	0.049	0.045	4.53	0.33	2.75	0.43	2.89
	15	PreDR																	
	15	ChkDR																	
4	15	2	MGO	21:59:30	22:02:30	WT130276	QU201310_04	0.183	0.002	0.140	0.006	0.116	0.049	0.045	3.74	0.05	3.13	0.13	2.60
5	15	2	MGO	22:07:00	22:10:00	WT130277	QU201310_05	0.155	0.000	0.000	0.000	0.000	0.049	0.045	3.18	0.00	0.00	0.00	0.00
6	15	2	MGO	22:14:30	22:17:30	WT130278	QU201310_06	0.159	0.002	0.097	0.003	0.102	0.049	0.045	3.24	0.04	2.17	0.08	2.28
	15	PreDR																	
	15	DR																	
7	15	1	MGO	23:19:30	23:22:30	WT130279	QU201310_07	0.145	0.001	0.092	0.002	0.098	0.049	0.045	2.96	0.02	2.06	0.04	2.18
8	15	1	MGO	23:26:00	23:29:00	WT130280	QU201310_08	0.144	0.001	0.087	0.001	0.088	0.049	0.045	2.95	0.01	1.95	0.03	1.98
9	15	1	MGO	23:32:30	23:35:30	WT130282	QU201310_09	0.140	0.001	0.091	0.002	0.098	0.049	0.045	2.87	0.01	2.04	0.04	2.20
	16	PreDR																	
	16	DR																	
10	16	2	HFO	11:57:00	12:00:00	WT130283	QU201310_11	0.162	0.000	0.064	0.006	0.075	0.049	0.045	3.31	0.01	1.43	0.14	1.67
11	16	2	HFO	12:03:30	12:06:30	WT130284	QU201310_12	0.170	0.001	0.052	0.004	0.067	0.049	0.045	3.47	0.02	1.17	0.09	1.49
12	16	2	HFO	12:10:00	12:13:00	WT130286	QU201310_13	0.155		0.050	0.005	0.075	0.049	0.045	3.16	0.00	1.12	0.12	1.67
	16	PreDR																	
	16	DR																	
13	16	1	HFO	12:41:30	12:44:30	WT130287	QU201310_14	0.188	0.001	0.065	0.003	0.074	0.049	0.045	3.85	0.02	1.45	0.07	1.65
14	16	1	HFO	12:48:30	12:51:30	WT130288	QU201310_15	0.171	0.001	0.060	0.003	0.073	0.049	0.045	3.50	0.03	1.34	0.07	1.64
15	16	1	HFO	12:55:30	12:58:30	WT130289	QU201310_16	0.170	0.001	0.061	0.004	0.306	0.049	0.045	3.48	0.02	1.37	0.09	6.85
	16	PreDR																	
	16	DR																	
16	16	3	HFO	13:40:00	13:43:00	WT130292	QU201310_17	0.074	0.005	0.036	0.007	0.038	0.049	0.045	1.52	0.12	0.81	0.16	0.85
17	16	3	HFO	13:46:00	13:49:00	WT130290	QU201310_18	0.058	0.002	0.026	0.007	0.033	0.049	0.045	1.19	0.05	0.58	0.16	0.74
18	16	3	HFO	13:53:00	13:56:00	WT130291	QU201310_19	0.068	0.002	0.028	0.005	0.029	0.049	0.045	1.39	0.05	0.63	0.11	0.64
	16	PreDR																	
	16	DR																	
19	16	VSR	HFO	14:30:00	14:33:00	WT130293	QU201310_20	0.123	Filters not in lab			0.049	0.045	2.51					
20	16	VSR	HFO?	14:34:30	14:37:30	WT130294	QU201310_21	0.085	Filters not in lab			0.049	0.045	1.73					
21	16	VSR	HFO?	14:42:00	14:45:00	WT130295	QU201310_22	0.069	0.007	0.043	0.012	0.041	0.049	0.045	1.40	0.15	0.97	0.26	0.93
	16	PreDR																	
	16	DR																	
22	16	VSR	MGO	16:54:30	16:57:30	WT130296	QU201310_25	0.050	0.005	0.037	0.009	0.041	0.049	0.045	1.02	0.10	0.82	0.19	0.92
23	16	VSR	MGO	17:00:00	17:03:00	WT130297	QU201310_26	0.005	0.006	0.042	0.007	0.043	0.049	0.045	0.10	0.14	0.95	0.16	0.95
24	16	VSR	MGO	17:06:30	17:09:30	WT130298	QU201310_27	0.036	-0.296	105.730	0.009	0.049	0.049	0.045	0.74	-6.63	2365.32	0.21	1.10
	16	PreDR																	
	16	DR																	

¹ Note: Red highlighted samples were outliers and were removed from the analysis. Blue are filters that were damaged and not weighed.

Appendix C. **Sampling system description**

ISO 8178-1⁹ and ISO 8178-2¹⁰ specify the measurement and evaluation methods for gaseous and particulate exhaust emissions when combined with combinations of engine load and speed provided in ISO 8178- *Part 4: Test cycles for different engine applications*. The emission results represent the mass rate of emissions per unit of work accomplished. Specific emission factors are based on brake power measured at the crankshaft, the engine being equipped only with the standard auxiliaries necessary for its operation. Per ISO, auxiliary losses are <5 % of the maximum observed power.

IMO ship pollution rules and measurement methods are contained in the “International Convention on the Prevention of Pollution from Ships”, known as MARPOL 73/78¹¹, and sets limits on NO_x and SO_x emissions from ship exhausts. The intent of this protocol was to conform as closely as practical to both the ISO and IMO standards.

Gaseous and Particulate Emissions

A properly designed sampling system is essential to accurate collection of a representative sample from the exhaust and subsequent analysis. ISO points out that particulate must be collected in either a full flow or partial flow dilution system and UCR chose the partial flow dilution system with single venturi as shown in Figure C-1.

⁹ International Standards Organization, ISO 8178-1, *Reciprocating internal combustion engines - Exhaust emission measurement -Part 1: Test-bed measurement of gaseous particulate exhaust emissions*, First edition 1996-08-15

¹⁰ International Standards Organization, ISO 8178-2, *Reciprocating internal combustion engines - Exhaust emission measurement -Part 2: Measurement of gaseous and particulate exhaust emissions at site*, First edition 1996-08-15

¹¹ International Maritime Organization, *Annex VI of MARPOL 73/78 “Regulations for the Prevention of Air Pollution from Ships and NO_x Technical Code”*.

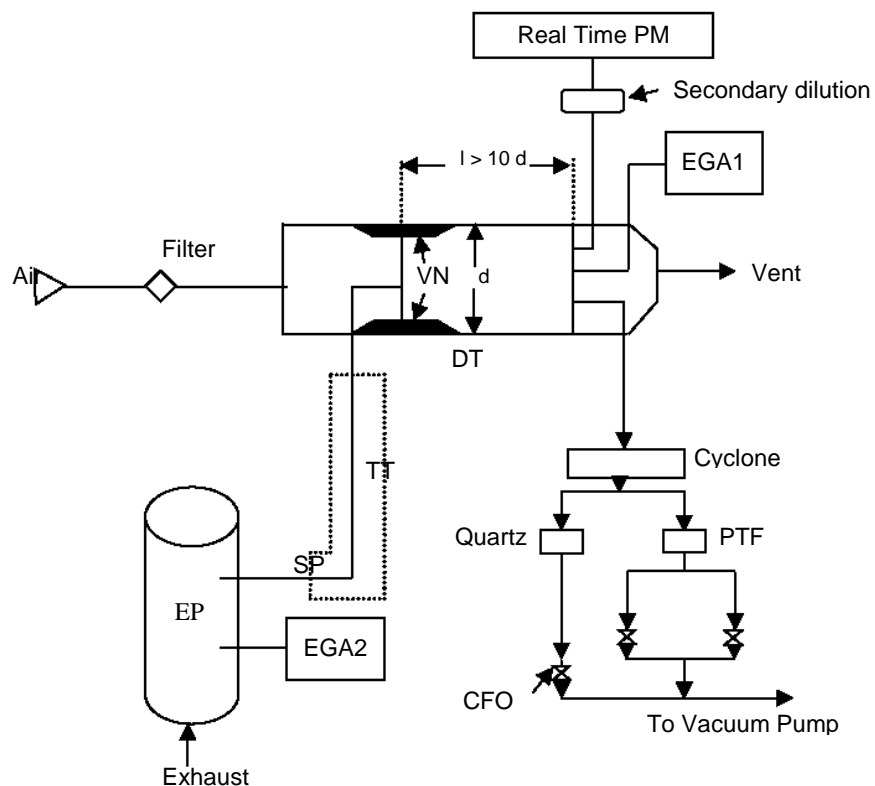


Figure C-1 Partial Flow Dilution System with Single Venturi

A partial flow dilution system was selected based on cost and the impossibility of a full flow dilution for “medium and large” engine testing on the test bed and at site. The flow in the dilution system eliminates water condensation in the dilution and sampling systems and maintains the temperature of the diluted exhaust gas at $<52^{\circ}\text{C}$ before the filters. ISO cautions the advantages of partial flow dilution systems can be lost to potential problems such as: losing particulates in the transfer tube, failing to take a representative sample from the engine exhaust and inaccurately determining the dilution ratio.

An overview of UCR’s partial dilution system in Figure C-1 shows that raw exhaust gas is transferred from the exhaust pipe (EP) through a sampling probe (SP) and the transfer tube (TT) to a dilution tunnel (DT) due to the negative pressure created by the venturi (VN) in DT. The gas flow rate through TT depends on the momentum exchange at the venturi zone and is therefore affected by the absolute temperature of the gas at the exit of TT. Consequently, the exhaust split for a given tunnel flow rate is not constant, and the dilution ratio at low load is slightly lower than at high load. More detail on the key components is provided in Table C-1.



Figure C-2 Dilution system and measurement layout on the auxiliary engine exhaust stack

Dilution Air System

A partial flow dilution system requires dilution air and UCR uses compressed air in the field as it is readily available. ISO recommends the dilution air be at $25 \pm 5^{\circ}\text{C}$, filtered and charcoal scrubbed to eliminate background hydrocarbons. The dilution air may be dehumidified. To ensure the compressed air is of a high quality UCR processes any supplied air through a field processing unit that reduces the pressure to about 30psig as that level allows a dilution ratio of about 5/1 in the geometry of our system. The next stages, in sequence, include: a liquid knock-out vessel, desiccant to remove moisture with silica gel containing an indicator, hydrocarbon removal with activated charcoal and a HEPA filter for the fine aerosols that might be present in the supply air. The silica gel and activated carbon are changed for each field campaign. Figure C-3 shows the field processing unit in its transport case. In the field the case is used as a framework for supporting the unit



Figure C-3 Field Processing Unit for Purifying Dilution Air in Carrying Case

Table C-1 Components of a Sampling System: ISO/IMO Criteria & UCR Design

Section	Selected ISO and IMO Criteria	UCR Design
Exhaust Pipe (EP)	In the sampling section, the gas velocity is > 10 m/s, except at idle, and bends are minimized to reduce inertial deposition of PM. Sample position is 6 pipe diameters of straight pipe upstream and 3 pipe diameters downstream of the probe.	UCR follows the ISO recommendation, as closely as practical.
Sampling Probe (SP) -	The minimum inside diameter is 4 mm and the probe is an open tube facing upstream on the exhaust pipe centerline. No IMO code.	UCR uses a stainless steel tube with diameter of 8mm placed near the center line.
Transfer Tube (TT)	<ul style="list-style-type: none"> • As short as possible and < 5 m in length; • Equal to/greater than probe diameter & < 25 mm diameter; • TTs insulated. For TTs > 1m, heat wall temperature to a minimum of 250°C or set for < 5% thermophoretic losses of PM. 	UCR no longer uses a transfer tube.
Dilution Tunnel (DT)	<ul style="list-style-type: none"> • shall be of a sufficient length to cause complete mixing of the exhaust and dilution air under turbulent flow conditions; • shall be at least 75 mm inside diameter (ID) for the fractional sampling type, constructed of stainless steel with a thickness of > 1.5 mm. 	UCR uses fractional sampling; stainless steel tunnel has an ID of 50mm and thickness of 1.5mm.
Venturi (VN) --	The pressure drop across the venturi in the DT creates suction at the exit of the transfer tube TT and gas flow rate through TT is basically proportional to the flow rate of the dilution air and pressure drop.	Venturi proprietary design provided by MAN B&W; provides turbulent mixing.
Exhaust Gas Analyzers (EGA)	One or several analyzers may be used to determine the concentrations. Calibration and accuracy for the analyzers are like those for measuring the gaseous emissions.	UCR uses a 5-gas analyzer meeting IMO/ISO specs

Calculating the Dilution Ratio

According to ISO 8178, “it is essential that the dilution ratio be determined very accurately” for a partial flow dilution system such as what UCR uses. The dilution ratio is simply calculated from measured gas concentrations of CO₂ and/or NO_x in the raw exhaust gas, the diluted exhaust gas and the dilution air. UCR has found it useful to independently determine the dilution ratio from both CO₂ and NO_x and compare the values to ensure that they are within ±10%. UCR’s experience indicates the independently determined dilution ratios are usually within 5%. At systematic deviations within this range, the measured dilution ratio can be corrected, using the calculated dilution ratio. According to ISO, dilution air is set to obtain a maximum filter face temperature of <52°C and the dilution ratio shall be > 4.

Dilution System Integrity Check

ISO describes the necessity of measuring all flows accurately with traceable methods and provides a path and metric to quantifying the leakage in the analyzer circuits. UCR has adopted the leakage test and its metrics as a check for the dilution system. According to ISO the maximum allowable leakage rate on the vacuum side shall be 0.5 % of the in-use flow rate for the portion of the system being checked. Such a low leakage rate allows confidence in the integrity of the partial flow system and its dilution tunnel. Experience has taught UCR that the flow rate selected should be the lowest rate in the system under test.

Measuring the Gaseous Emissions: CO, CO₂, HC, NO_x, O₂, SO₂

Measurement of the concentration of the main gaseous constituents is one of the key activities in measuring emission factors. This section covers the ISO/IMO protocols and that used by UCR. For SO₂, ISO recommends and UCR concurs that the concentration of SO₂ is calculated based on the fact that 95+% of the fuel sulfur is converted to SO₂.

Measuring Gaseous Emissions: ISO & IMO Criteria

ISO specifies that either one or two sampling probes located in close proximity in the raw gas can be used and the sample split for different analyzers. However, in no case can condensation of exhaust components, including water and sulfuric acid, occur at any point of the analytical system. ISO specifies the analytical instruments for determining the gaseous concentration in either raw or diluted exhaust gases.

- Heated flame ionization detector (HFID) for the measurement of hydrocarbons;
- Non-dispersive infrared analyzer (NDIR) for the measurement of carbon monoxide and carbon dioxide;
- Heated chemiluminescent detector (HCLD) or equivalent for measurement of nitrogen oxides;
- Paramagnetic detector (PMD) or equivalent for measurement of oxygen.

ISO states the range of the analyzers shall accurately cover the anticipated concentration of the gases and recorded values between 15% and 100% of full scale. A calibration curve with five points is specified. However, with modern electronic recording devices, like a computer, ISO allows the range to be expanded with additional calibrations. ISO details instructions for establishing a calibration curve below 15%. In general, calibration curves must be < ±2 % of each calibration point and by < ±1 % of full scale zero.

ISO outlines their verification method. Each operating range is checked prior to analysis by using a zero gas and a span gas whose nominal value is more than 80 % of full scale of the measuring range. If, for the two points considered, the value found does not differ by more than ± 4 % of full scale from the declared reference value, the adjustment parameters may be modified. If $>4\%$, a new calibration curve is needed.

ISO & IMO specify the operation of the HCLD. The efficiency of the converter used for the conversion of NO_2 into NO is tested prior to each calibration of the NO_x analyzer. The efficiency of the converter shall be > 90 %, and >95 % is strongly recommended.

ISO requires measurement of the effects from exhaust gases on the measured values of CO , CO_2 , NO_x , and O_2 . Interference can either be positive or negative. Positive interference occurs in NDIR and PMD instruments where the interfering gas gives rise to the same effect as the gas being measured, but to a lesser degree. Negative interference occurs in NDIR instruments due to the interfering gas broadening the absorption band of the measured gas, and in HCLD instruments due to the interfering gas quenching the radiation. Interference checks are recommended prior to an analyzer's initial use and after major service intervals.

Measuring Gaseous Emissions: UCR Design

The concentrations of CO , CO_2 , NO_x and O_2 in the raw exhaust and in the dilution tunnel are measured with a Horiba PG-250 portable multi-gas analyzer. The PG-250 simultaneously measures five separate gas components with methods recommended by the ISO/IMO and USEPA. The signal output of the instrument is connected to a laptop computer through an RS-232C interface to continuously record measured values. Major features include a built-in sample conditioning system with sample pump, filters, and a thermoelectric cooler. The performance of the PG-250 was tested and verified under the U.S. EPA ETV program.



Figure C-4 Setup Showing Gas Analyzer with Computer for Continuous Data Logging

Details of the gases and the ranges for the Horiba instrument are shown in Table C-2. Note that the Horiba instrument measured sulfur oxides (SO_2); however, the UCR follows the protocol in ISO and calculates the SO_2 level from the sulfur content of the fuel as the direct measurement for SO_2 is less precise than calculation.

Table C-2 Detector Method and Concentration Ranges for Monitor

Component	Detector	Ranges
Nitrogen Oxides (NO_x)	Heated Chemiluminescence Detector (HCLD)	0-25, 50, 100, 250, 500, 1000, & 2500 ppmv
Carbon Monoxide (CO)	Non dispersive Infrared Absorption (NDIR)	0-200, 500, 1000, 2000, & 5000 ppmv
Carbon Dioxide (CO₂)	Non dispersive Infrared Absorption (NDIR)	0-5, 10, & 20 vol%
Sulfur Dioxide (SO₂)	Non dispersive Infrared Absorption (NDIR)	0-200, 500, 1000, & 3000 ppmv
Oxygen	Zirconium oxide sensor	0-5, 10, & 25 vol%

For quality control, UCR carries out analyzer checks with calibration gases both before and after each test to check for drift. Because the instrument measures the concentration of five gases, the calibration gases are a blend of several gases (super-blend) made to within 1% specifications. Experience has shown that the drift is within manufacturer specifications of ±1% full scale per day shown in Table E-3. The PG-250 meets the analyzer specifications in ISO 8178-1 Section 7.4 for repeatability, accuracy, noise, span drift, zero drift and gas drying.

Table C-3 Quality Specifications for the Horiba PG-250

Repeatability	±0.5% F.S. (NO _x : ≤/ 100ppm range CO: ≤/ 1,000ppm range) ±1.0% F. S.
Linearity	±2.0% F.S.
Drift	±1.0% F. S./day (SO ₂ : ±2.0% F.S./day)

Measuring the Particulate Matter (PM) Emissions

ISO 8178-1 defines particulates as any material collected on a specified filter medium after diluting exhaust gases with clean, filtered air at a temperature of $\leq 52^{\circ}\text{C}$, as measured at a point immediately upstream of the primary filter. The particulate consists of primarily carbon, condensed hydrocarbons and sulfates, and associated water. Measuring particulates requires a dilution system and UCR selected a partial flow dilution system. The dilution system design completely eliminates water condensation in the dilution/sampling systems and maintains the temperature of the diluted exhaust gas at $< 52^{\circ}\text{C}$ immediately upstream of the filter holders. IMO does not offer a protocol for measuring PM. A comparison of the ISO and UCR practices for sampling PM is shown in Table C-4.

Table C-4 Measuring Particulate by ISO and UCR Methods

	ISO	UCR
Dilution tunnel	Either full or partial flow	Partial flow
Tunnel & sampling system	Electrically conductive	Same
Pretreatment	None	Cyclone, removes $>2.5\mu\text{m}$
Filter material	Fluorocarbon based	Teflon (TFE)
Filter size, mm	47 (37mm stain diameter)	Same
Number of filters in series	Two	One
Number of filters in parallel	Only single filter	Two; 1 TFE & 1 Quartz
Number of filters per mode	Single or multiple	Multiple
Filter face temp. $^{\circ}\text{C}$	< 52	Same
Filter face velocity, cm/sec	35 to 80.	~ 33
Pressure drop, kPa	For test < 25	Same
Filter loading, μg	> 500	500-1,000 + water w/sulfate
Weighing chamber	$22 \pm 3^{\circ}\text{C}$ & $\text{RH} = 45\% \pm 8$	Same
Analytical balance, LDL μg	10	0.5
Flow measurement	Traceable method	Same
Flow calibration, months	< 3 months	Every campaign

Sulfur content. According to ISO, particulates measured using ISO 8178 are “conclusively proven” to be effective for fuel sulfur levels up to 0.8%. UCR is often faced with measuring PM for fuels with sulfur content exceeding 0.8% and has extended this method to those fuels as no other method is prescribed for fuels with a higher sulfur content.

Added Comments about UCR’s Measurement of PM

In the field UCR uses a raw particulate sampling probe fitted close to and upstream of the raw gaseous sample probe and directs the PM sample to the dilution tunnel. There are two gas streams leaving the dilution tunnel; the major flow vented outside the tunnel and the minor flow directed to a cyclone separator, sized to remove particles $>2.5\mu\text{m}$. The line leaving the cyclone separator is split into two lines; each line has a 47 Gelman filter holder. One holder collects PM on a Teflon filter and the other collects PM on a quartz filter. UCR simultaneously collects PM on Teflon and quartz filters at each operating mode and analyzes them according to standard procedures.

Briefly, total PM was collected on Pall Gelman (Ann Arbor, MI) 47 mm Teflo filters and weighed using a Metler Toledo UMX2 microbalance with a 0.1 ug resolution. Before and after collection, the filters were conditioned for 24 hours in an environmentally controlled room (RH = 40%, T= 25 °C) and weighed daily until two consecutive weight measurements were within 3 µg or 2%. It is important to note that the simultaneous collection of PM on quartz and Teflon filters provides a comparative check of PM mass measured by two independent methods and serves as an important Quality Check for measuring PM mass.

Measuring Non-Regulated Gaseous Emissions

Neither ISO nor IMO provide a protocol for sampling and analyzing non-regulated emissions. UCR uses peer reviewed methods adapted to their PM dilution tunnel. The methods rely on added media to selectively collect hydrocarbons and PM fractions during the sampling process for subsequent off-line analysis. A secondary dilution is constructed to capture real time PM.

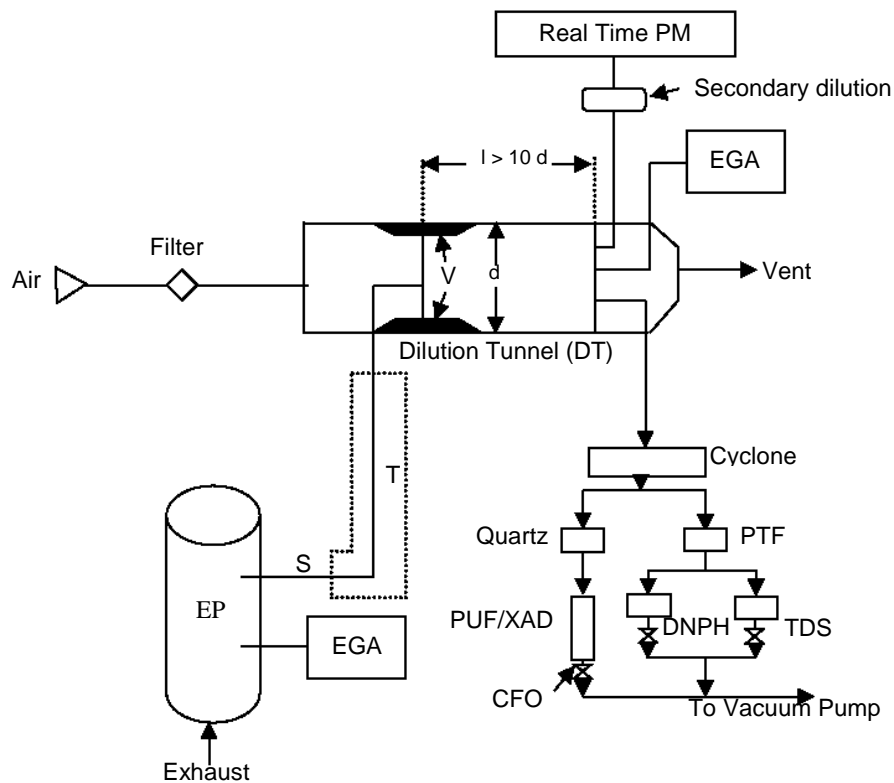


Figure C-5 Extended setup of the PFDS for non-regulated emissions

Appendix D. Additional Emissions Results

This appendix contains additional information collected on the APL main engine. Table D1 shows the measured concentrations during testing and the dilution ratio, D2 shows the PM concentration for the real time instruments and PM and EC/OC samples, and D3 and D4 show the percent differences between MGO and HFO for the different PM measurement systems.

Table D1 Average measured concentration and dilution ratio

Mode	NOx	CO	CO2	O2	SO2	DR ⁴
1-MGO	122.7	2.7	0.36%	21%	2.2	16.0
2-MGO	114.0	3.0	0.38%	20%	3.6	14.0
3-MGO	109.6	9.7	0.42%	20%	3.1	12.0
VSR-MGO	40.8	4.2	0.15%	21%	-3.2	18.0
1-HFO	105.7	3.7	0.28%	21%	9.3	20.0
2-HFO	90.0	3.9	0.26%	21%	4.8	22.0
3-HFO	42.7	6.4	0.14%	21%	2.7	34.0
VSR-HFO	58.9	6.2	0.19%	21%	7.6	14.0

Table D2 Average measured concentration and dilution ratio

Mode	#	BC ^{PA-MSS} ₁		BC ^{PA-PAX} ₂		ScatPAX ₃		DR ⁴		PM ^{2.5}		EC		OC	
		mg/m ³	mg/m ³	mg/m ³	mg/m ³	n/a	n/a	mg/m ³	mg/m ³	mg/m ³	mg/m ³	mg/m ³	mg/m ³	mg/m ³	mg/m ³
1-MGO	0.015	0.018	0.687	15.9	2.73	0.013	1.855	0.035	1.949	0.063	1.497	0.063	1.497	0.063	1.497
2-MGO	0.054	0.056	0.494	13.4	3.16	0.029	1.626	0.063	1.497	0.063	1.497	0.063	1.497	0.063	1.497
3-MGO	0.362	0.324	0.324	12.0	4.72	0.358	2.869	0.447	2.817	0.447	2.817	0.447	2.817	0.447	2.817
VSR-MGO	0.148	0.130	0.312	20.8	0.58	0.111	0.813	0.172	0.910	0.172	0.910	0.172	0.910	0.172	0.910
1-HFO	0.026	0.023	0.436	20.0	3.36	0.021	1.275	0.070	3.109	0.070	3.109	0.070	3.109	0.070	3.109
2-HFO	0.044	0.039	0.330	21.2	3.09	0.009	1.141	0.107	1.481	0.107	1.481	0.107	1.481	0.107	1.481
3-HFO	0.095	0.091	0.295	33.0	1.27	0.069	0.618	0.131	0.684	0.131	0.684	0.131	0.684	0.131	0.684
VSR-HFO	0.226	0.214	0.303	16.0	1.75	0.139	0.891	0.237	0.852	0.237	0.852	0.237	0.852	0.237	0.852

¹ PA-MSS is the AVL 483 micro soot sensor that measures the acoustic properties of PM or black carbon. The laser wavelength was xx nm.

PA MSS showed a zero offset of -0.016 mg/m³ during 1-MGO (tested after 1 and 2). Adjustment provided in post processing with a sliding scale from 22:30 to 24:00. All other PA MSS data was corrected with onsite zero adjustments. ² **PA PAX** is the PAX photo acoustic instrument for black carbon in units of mg/m³. The laser wavelength was xx nm. PA PAX instrument designed for frequent zero adjustments (1 per hour). Hourly adjustments recommended for the PA type instruments. ³ **ScatPAX** is the PAX single absorption scattering coefficient. This is a unit less ratio of absorption over scattering+absorption
⁴ ISO-8178-1 2006 Chapter 12.4 "Adjustment of the dilution ratio": Specification is minimum of 4 to 1.

Table D3 Percent gaseous emissions change from MGO on a work specific basis

Nominal Load	Power kW	Load %	SO ₂ %	PM _{2.5} %	EC %	OC %	NO _x %	CO %	CO ₂ %
1	45643	61%	-81%	-36%	-51%	16%	-8%	-42%	3%
2	31935	43%	-49%	-29%	134%	-1%	-12%	-47%	1%
3	16100	22%	-61%	27%	79%	60%	-12%	-48%	0%
VSR	3964	5%	-152%	-59%	-1%	13%	-14%	-17%	0%

Table D4 Percent particle emissions change from MGO on a work specific basis

Engine Conditions			Real Time			Grav.	NIOSH		IMPROVE	
Nominal Load	Power kW	Load %	BC _{PA-MSS} ¹ %	BC _{PA-PAX} ² %	ScatPAX ³ %	PM _{2.5} %	EC %	OC %	EC %	OC %
1	45643	61%	-54%	-36%	58%	-36%	-51%	15%	-61%	-50%
2	31935	43%	-15%	-1%	50%	-29%	132%	-2%	-59%	-30%
3	16100	22%	35%	26%	10%	31%	85%	65%	21%	46%
VSR	3964	5%	-18%	-24%	3%	-59%	0%	14%	-9%	33%

¹ MSS is the AVL 483 micro soot sensor that measures the acoustic properties of PM or black carbon

² MAAP is a mass aerosol particle. The MAAP over ranged at 50 ug/m³ (concentration ranged from 200 to 10,000 ug/m³)

³ Aeth is the aethalometer which uses the light scattering principle to measure PM concentration. Aeth over ranged also.

⁴ Mode 3 post scrubber was not stable for PM. See real time MSS data. Filter samples showed PM increase, not MSS